

Long-read, whole-genome shotgun sequence data for five model organisms

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Abstract

Single molecule, real-time (SMRT) sequencing from Pacific Biosciences is increasingly used in many areas of biological research including *de novo* genome assembly, structural-variant identification, haplotype phasing, mRNA isoform discovery, and base-modification analyses. High-quality, public datasets of SMRT sequences can spur development of analytic tools that can accommodate unique characteristics of SMRT data (long read lengths, lack of GC or amplification bias, and a random error profile leading to high consensus accuracy). In this paper, we describe eight high-coverage SMRT sequence datasets from five organisms (*Escherichia coli*, *Saccharomyces cerevisiae*, *Neurospora crassa*, *Arabidopsis thaliana*, and *Drosophila melanogaster*) that have been publicly released to the general scientific community (NCBI Sequence Read Archive ID SRP040522). Data were generated using two sequencing chemistries (P4-C2 and P5-C3) on the PacBio RS II instrument. The datasets reported here can be used without restriction by the research community to generate whole-genome assemblies, test new algorithms, investigate genome structure and evolution, and identify base modifications in some of the most widely-studied model systems in biological research.

Background and Summary

Single-molecule, real-time (SMRT®) DNA sequencing occurs by optically detecting a fluorescent signal when a nucleotide is being incorporated by a DNA polymerase [1-4]. This relatively new technology enables detection of DNA sequences that have unique characteristics, such as long read lengths, lack of CG bias, and random error profiles, and can yield highly accurate consensus sequences [5]. Kinetic information such as pulse width and interpulse duration are also recorded and can be used to detect base modifications [6-8].

Since its introduction, investigators have published on a range of applications using SMRT sequencing. For example, the developers of GATK (Genome Analysis Toolkit) demonstrated that single nucleotide polymorphisms (SNPs) could be detected using SMRT sequences [9, 10] due to their lack of context-specific bias and systematic error [5, 10]. Likewise, the developers of PBcR (PacBio error correction) [11, 12] showed that complete bacterial genome assemblies using SMRT sequence data had greater than Q60 base quality [12]. PBcR was later incorporated as the “pre-assembly” step in the HGAP (hierarchical genome assembly process) system [13], followed by consensus polishing using the Quiver algorithm [13] to produce a complete assembly pipeline for SMRT sequence data. In addition, other third-party tools now support long reads for various applications such as mapping [14, 15], scaffolding [16], structural-variation discovery [17], and genome assembly [11, 18]. Other applications such as 16S rRNA sequencing [19], characterization of entire transcriptomes [20, 21], genome-editing studies [22], base-modification studies [7, 8, 23-25], and validation of CRISPR targets [26] have also been published.

To encourage interest in further applications and tool development for SMRT sequence data, we report here the release of whole-genome shotgun-sequence datasets from five model organisms (*E. coli*, *S. cerevisiae*, *N. crassa*, *A. thaliana*, and *D. melanogaster*). These organisms have among the most complete and well-annotated reference genome sequences, due to continual refinement by dedicated teams of scientists. Despite continued improvement of these genome sequences with new technologies, few are completely finished with fully contiguous assemblies of all chromosomes. The gaps remaining arise from complex structures such as transposable elements, repeats, segmental duplications, or other dynamic regions of the genome that cannot be easily assembled. Structural differences in these regions can account for variability in millions of nucleotides within every genome, and mounting evidence suggest that such mutations are important for human diversity and disease susceptibility in many complex traits including autism and schizophrenia [27-29]. SMRT sequencing data can therefore play an important role in the completion of these and other reference genomes, providing a platform for new insights into genome biology.

Methods

We generated eight whole-genome shotgun-sequence datasets from five model organisms using the P4C2 or P5C3 polymerase and chemistry combinations, totaling nearly 1000 gigabytes (GB) of raw data (See Data Records section). Genomic DNA was either purchased from commercial sources or generously provided by collaborators.

DNA from the reference K12 strain of *E. coli* was purchased from Lofstrand Labs Limited (K12 MG1655 *E. coli*, cat# L3-4001SP2). DNA from the reference OR74A strain of *N. crassa* was purchased from the Fungal Genetics Stock Center (FGSC). A standard Ler-0 strain of *A. thaliana* plants was grown from seeds purchased from Lehle seeds (WT-04-19-02) and DNA was extracted at Pacific Biosciences. The protocol is available on Sample Net [30] and summarized in the organism-specific methods section of this paper. DNA from the 9464 strain of *S. cerevisiae* was provided by J. Li at University of California San Francisco. The 9464 strain is a daughter of the reference WG303 strain. DNA from the T1 strain of *N. crassa* was obtained from D. Catcheside at Flinders University who has an interest in polymorphic genes regulating recombination. The T1 strain is an A mating type strain which, like OR74A, was derived from a cross between the Em a 5297 and Em A 5256 strains. DNA from the ISO1 strain [31] of *D. melanogaster* was obtained from S. Celniker at Lawrence Berkeley National Laboratory. This is the reference strain of *D. melanogaster* that was originally chosen to be the first large genome to be sequenced and assembled using a whole-genome shotgun approach [32]. It continues to serve as the reference strain in subsequent releases and numerous annotations of the *D. melanogaster* genome.

DNA extraction methods were species-specific and optimized for each organism (See organism-specific methods below). In general, the steps are: (1) remove debris and particulate material, (2) lyse cells, (3) remove membrane lipids, proteins and RNA, (4) DNA purification.

SMRTbell™ libraries for sequencing [9] were prepared using either 10 kb [33, 34] or 20 kb [35] preparation protocols to optimize for the most high-quality and longest reads. The main steps for library preparation are: (1) Shearing (2) DNA damage repair, (3) blunt end-ligation with hairpin adapters supplied in the DNA Template Prep Kit 2.0 (Pacific Biosciences), (4) size selection, and (5) binding to polymerase using the DNA Sequencing Kit 3.0 (Pacific Biosciences).

Table 1: Summary of DNA Samples. The NCBI sample ID associated with each dataset is provided. DNA was extracted in a species-specific manner, yielding genomic DNA of various sizes. All DNA was size selected using the Blue Pippin system (Sage Sciences), and select samples were sheared with g-TUBEs (Covaris).

Dataset Name	Sample ID	DNA extraction	gDNA size (kb)	Shearing	Size selection
<i>E. coli</i> MG1655 P4C2	SAMN02951645	ammonium acetate or SDS, proteinase K, phenol-chloroform	10	none	Blue Pippin (7kb)
<i>E. coli</i> MG1655 P5C3	SAMN02743420	ammonium acetate or SDS, proteinase K, phenol-chloroform	10	none	Blue Pippin (7kb)
<i>S. cerevisiae</i> 9464 P4C2	SAMN02731377	contact J. Li at UCSF	>40	g-TUBE	Blue Pippin (17kb)
<i>N. crassa</i> OR74A P4C2	SAMN02724975	BashingBeads, Zymo Research kit	6	none	Blue Pippin (4kb)
<i>N. crassa</i> T1 P4C2	SAMN02724976	SDS, proteinase K, phenol-chloroform, RNAase, isopropanol	15	none	Blue Pippin (7kb)
<i>A. thaliana</i> Ler-0 P5C3	SAMN02724977	CTAB, chloroform:isoamyl, isopropanol precip.	>40	g-TUBE	Blue Pippin (15kb)
<i>A. thaliana</i> Ler-0 P4C2	SAMN02731378	CTAB, chloroform:isoamyl, isopropanol precip.	>40	g-TUBE	Blue Pippin (7kb)
<i>D. melanogaster</i> ISO1 P5C3	SAMN02614627	SDS, phenol-chloroform, CsCl banding, ethanol precip.	>40	g-TUBE	Blue Pippin (17kb)

E. coli collection, DNA Extraction, and SMRTbell Library Preparation

Both P4C2 and P5C3 samples were prepared in the same way. *E. coli* K12 genomic DNA was ordered and purified by Lofstrand Labs Limited (K12MG1655 *E. coli*, cat# L3-4001SP2). Field Inversion Gel Electrophoresis (FIGE) was run to ensure presence of high-molecular-weight gDNA. Ten micrograms of gDNA was sheared using g-TUBE devices (Covaris, Inc) spun at 5500 rpm for 1 minute. Three microliters of elution buffer (EB) was added to rinse the upper chamber, spun at 6000 rpm, and spun again at 5500 rpm after inverting the g-TUBE device. SMRTbell libraries were created using the Procedure & Checklist – 20 kb Template Preparation using BluePippin™ Size Selection protocol[35]. Briefly, the library was run on a BluePippin system (Sage Science, Inc., Beverly, MA, USA) to select for SMRTbell templates greater than 10 kb. The resulting average insert size was 17 kb based on 2100 Bioanalyzer instrument (Agilent Technologies Genomics, Santa Clara, CA., USA). Sequencing primers were annealed to the hairpins of the SMRTbell templates followed by binding with the P5 sequencing polymerase and MagBeads (Pacific Biosciences, Menlo Park, CA, USA). One SMRT Cell was run on the PacBio® RS II system with an on-plate concentration of 150 pM using P5-C3 chemistry and a 180-minute data-collection mode.

S. cerevisiae collection, DNA Extraction, and SMRTbell Library Preparation

Please contact J. Li at University of California, San Francisco to obtain the protocol.

A. thaliana collection, DNA Extraction, and SMRTbell Library Preparation

Plants were grown from seeds provided by Lehle seeds (WT-04-19-02). Shoots and leaves were harvested at three weeks and ground in liquid nitrogen using a mortar and pestle. The complete protocol is described in the document “Preparing *Arabidopsis* Genomic DNA for Size-Selected ~20 kb SMRTbell™ Libraries” [36]. This protocol can be used to prepare purified *Arabidopsis* genomic DNA for size-selected SMRTbell templates with average insert sizes of 10 to 20 kb. We recommend

starting with 20-40 grams of three-week-old *Arabidopsis* whole plants, which can generate >100 µg of purified genomic DNA. SMRTbell libraries were created using the document “Procedure & Checklist – 20 kb Template Preparation using BluePippin™ Size Selection protocol” [35]. Eighty-five SMRT Cells were run on the PacBio RS II system using P4-C2 chemistry and a 180-minute data-collection mode. Forty-six SMRT Cells were run on the PacBio RS II system using P5-C3 chemistry and a 180-minute data-collection mode.

***N. crassa* OR74A, collection, DNA Extraction, and SMRTbell Library Preparation**

The T1 strain of N. crassa, is an A mating type strain derived by DG Catcheside from a cross between the Em A 5297 and Em A 5256 strains he obtained from Stirling Emerson in 1955. The fungus was grown in shake culture for 72 hr at 25°C in 500 ml Vogel’s [37] minimal medium containing 2% sucrose. Mycelium was harvested by filtration, ground in liquid nitrogen, resuspended in 10 ml of a buffer containing 0.15 M NaCl, 0.1 M EDTA, 2% SDS at pH 9.5, and incubated overnight at 37°C with 1 mg protease K. Debris was precipitated by centrifugation and 10 ml distilled water was added to the supernatant, which was extracted once with an equal volume of water saturated phenol and once with chloroform. Nucleic acids were precipitated from the aqueous phase with 0.6 volumes of isopropanol. Following centrifugation, the pellet was dried and dissolved in 1 ml TE buffer (TRIS 10 mM, 1 mM EDTA pH 8.0). RNA and protein were digested by overnight incubation at 37°C with RNAase (50 µg) followed by addition of protease K (50 µg) and further incubation for 2 hr. The digest was extracted once with water-saturated phenol and once with chloroform. DNA was collected by precipitation with 0.6 volumes of isopropanol and, following centrifugation, the pellet was dried, dissolved in 500 µl TE buffer and stored at 4°C. Field Inversion Gel Electrophoresis (FIGE) was run to ensure presence of high-molecular-weight gDNA. The genomic DNA was approximately 25 kb and was not sheared. SMRTbell libraries were created using the document “Procedure and Checklist – 10 kb Template Preparation and Sequencing (with Low-Input DNA)” [33]. Two SMRT Cells were run on the PacBio RS II system using P4C2 chemistry and a 180-minute data collection mode.

***N. crassa* T1 collection, DNA Extraction, and SMRTbell Library Preparation**

The T1 strain of N. crassa, is an A mating type strain derived by DG Catcheside from a cross between the Em A 5297 and Em A 5256 strains he obtained from Stirling Emerson in 1955. The fungus was grown in shake culture for 72 hr at 25°C in 500 ml Vogel’s N [37] minimal medium containing 2% sucrose. Mycelium was harvested by filtration, ground in liquid nitrogen, resuspended in 10 ml of a buffer containing 0.15 M NaCl, 0.1 M EDTA, 2% SDS at pH 9.5, and incubated overnight at 37°C with 1 mg protease K. Debris was precipitated by centrifugation and 10 ml distilled water was added to the supernatant, which was extracted once with an equal volume of water-saturated phenol and once with chloroform. Nucleic acids were precipitated from the aqueous phase with 0.6 volumes of isopropanol. Following centrifugation, the pellet was dried and dissolved in 1 ml TE buffer (TRIS 10 mM, 1 mM EDTA pH 8.0). RNA and protein were digested by overnight incubation at 37°C with RNAase (50 µg) followed by addition of protease K (50 µg) and further incubation for 2 hr. The digest was extracted once with water saturated phenol and once with chloroform. DNA was collected by precipitation with 0.6 volumes of isopropanol and, following centrifugation, the pellet was dried, dissolved in 500 µl TE buffer and stored at 4°C. Field Inversion Gel Electrophoresis (FIGE) was run to ensure presence of high-molecular-weight gDNA. The genomic DNA was approximately 25 kb and was not sheared. SMRTbell libraries were created using the document “Procedure and Checklist – 10 kb Template Preparation and Sequencing (with Low-Input DNA)” [33]. Eighteen SMRT Cells were run on the PacBio RS II system using P4-C2 chemistry and a 180-minute data-collection mode.

***D. melanogaster* collection, DNA Extraction, and SMRTbell Library Preparation**

A total of 1.2 g of adult male ISO1 flies corresponding to 1950 animals were collected, starved for 90-120 min and frozen. The flies ranged in age from 0-7 days based on four collections (1) 0-2 days old, 500 males, 0.33 g; (2) 0-4 days old, 500 males, 0.29 g; (3) 0-7 days old, 500 males, 0.29 g; (4) 0-2 days old, 450 males, 0.29 g. Flies were ground in liquid nitrogen to a fine powder and genomic DNA was purified by phenol-chloroform extraction and CsCl banding in the ultracentrifuge. Briefly, the pulverized fly extract was gently re-suspended in 5 ml of HB buffer (7 M Urea, 2% SDS, 50 mM Tris pH7.5, 10 mM EDTA and 0.35 M NaCl) and 5 ml of 1:1 phenol/chloroform. The mixture was shaken slowly for 30 minutes and then centrifuged at 18K rpm for 10 min at 20°C. The aqueous phase was re-extracted twice as above and then precipitated by adding two volumes of ethanol and centrifuging at 18K rpm for 10 min at 20°C. The pellet was re-suspended in 3 ml of TE (10 mM Tris 1 mM EDTA pH 8.0) by gentle

inversion. To the re-suspended DNA, 3 g CsCl and 0.3 ml of 10 mg/ml ethidium bromide (EtBr) were added and the mixture centrifuged at 45K rpm for 16 hrs at 15°C. The DNA band was collected and the EtBr removed by extraction with CsCl-saturated butanol. The DNA was diluted three-fold with TE, 1/10 vol, 5 M NaCl was added and the DNA precipitated with two volumes of ethanol. After centrifugation, the pellet was washed in 70% ethanol. The DNA was resuspended in 100 µl TE at a concentration of 1.4 µg/µl and quantified using a Nanodrop instrument. This protocol routinely yields at least 10 ng DNA per mg of flies with an estimated DNA size >100 kb.

Genomic DNA was sheared, using a g-TUBE device (Covaris), at 4800 RPM, 150 ng/µl and purified using 0.45x volume ratio of AMPure PB beads. SMRTbell libraries were created using the Procedure & Checklist – 20 kb Template Preparation using BluePippin™ Size Selection [35]. Libraries were ligated with excess adapters and an overnight incubation was performed to increase the yield of ligated fragments larger than 20 kb. Smaller fragments and adapter dimers were then removed by >15 kb size selection using the BluePippin DNA size selection system by Sage Science. Forty-two SMRT Cells were run on the PacBio RS II system. The first run was composed of four SMRT Cells, loaded at 75 pM, 150 pM, 300 pM, and 400 pM in order to determine the optimal loading concentration of the sample. The remaining 38 SMRT Cells were loaded at 400 pM.

Data Records

After DNA extraction, libraries were generated and sequenced at Pacific Biosciences of California, uploaded to Amazon Web Services' Simple Storage Service (S3), and then submitted to the Sequence Read Archive at NCBI under Project ID SRP040522. The corresponding accession numbers and file sizes are listed in Table 1. More detailed information including md5 checksums and links to download the original data from AWS S3 are provided in Supplementary Table S1.

Table 2: Summary of Datasets. Eight datasets from five organisms are described in this paper. Data can be accessed from the Sequence Read Archive (SRA) using the accession numbers provided.

Organism	Strain	Origin	Polymerase & Chemistry Library kits	SRA Accession	Size (GB)
<i>E. coli</i>	MG1655	Lofstrand Labs	P4C2	SRX669475	6.0
<i>E. coli</i>	MG1655	Lofstrand Labs	P5C3	SRX533603	3.8
<i>S. cerevisiae</i>	9464	J. Li	P4C2	SRX533604	38
<i>N. crassa</i>	OR74A	FGSC	P4C2	SRX533605	29
<i>N. crassa</i>	T1	D. Catcheside	P4C2	SRX533606	143
<i>A. thaliana</i>	Ler-0	Lehle Seeds	P4C2	SRX533608	263
<i>A. thaliana</i>	Ler-0	Lehle Seeds	P5C3	SRX533607	252
<i>D. melanogaster</i>	ISO1	S. Celniker	P5C3	SRX499318	187

Raw data was transferred from the instrument to a storage location and organized first by the run name, and then by the SMRT Cell directory. Each run contained one or more SMRT Cells. Each SMRT Cell produced a metadata.xml file that recorded the run conditions and barcodes of sequencing kits, three bax.h5 files that contained base call and quality information of actual sequenced data, and one bas.h5

file that acted as a pointer to consolidate the three bax.h5 files. The “h5” suffix denotes that these are Hierarchical Data format 5 (HDF5) files. The specific contents and structure of a PacBio bax.h5 file is described in more detail in online documentation [38].

Recall the “SMRT bell” structure that underwent sequencing was created by the library preparation process [9]. Sequenced SMRT Bells corresponded to raw reads that may pass around the same base multiple times. A raw read could therefore have a structure that is composed of left adapter → DNA insert → right adapter → reverse complement of DNA insert → left adapter → DNA insert → and so on. This raw read is typically processed downstream to remove adapters and create subreads composed of the DNA sequence of interest to the investigator. Typical filtering conditions for high-quality SMRT sequence data are read score > 0.8, read length > 100, subread length > 500. In addition, the ends of reads are trimmed if they are outside of high-quality (HQ) regions, and adapter sequences between subreads are removed.

The post-filter statistics of each dataset are listed in Table 3. While raw read lengths reflect the true sequencing capacity of the instrument; only subreads are summarized in Table 3 because it is used in downstream analysis algorithms such as *de novo* assemblers. Multiple subreads can be contained within one raw read, and subreads exclude adapters and low quality sequence. N50 is a statistic used to describe the length distribution of a collection of reads, contigs, or scaffolds, and is defined as the length where 50% of all bases are contained in sequences longer than that length. The N50 filtered subread lengths ranged from 7.6 kb to 10.5 kb for datasets generated with P4-C2 chemistry and ranged from 12.2 kb to 14.2 kb for datasets generated with P5-C3 chemistry. With the exception of *N. crassa* OR74A, all datasets were sequenced to high-coverage (>68X) and sufficient for *de novo* genome assembly applications.

Table 3: Summary statistics of filtered data. Results shown for each dataset are based on output of SMRT Portal analysis using the default filtering parameters (see text for details). Fold coverage is calculated relative to the estimated genome size.

Dataset Name	Number of filtered subreads	N50 filtered subread length (nt)	Maximum filtered subread length (nt)	Total filtered subread (nt)	Estimated genome size (Mb)	Fold coverage
<i>E. coli</i> MG1655 P4C2	61,019	7,586	22,609	331,516,965	5	66X
<i>E. coli</i> MG1655 P5C3	43,063	12,041	28,647	373,874,428	5	75X
<i>S. cerevisiae</i> 9464 P4C2	269,145	8,821	30,164	1,597,871,118	12	133X
<i>N. crassa</i> OR74A P4C2	175,926	7,617	30,845	981,884,113	40	25X
<i>N. crassa</i> T1 P4C2	210,480	10,462	36,227	11,497,185,440	40	287X
<i>A. thaliana</i> Ler-0 P4C2	1,338,320	8,769	41,753	8,129,670,483	120	68X
<i>A. thaliana</i> Ler-0 P5C3	2,067,212	12,188	47,445	17,714,447,516	120	148X
<i>D. melanogaster</i> ISO1 P5C3	1,561,929	14,214	44,766	15,194,174,294	160	95X

Technical Validation

DNA and Sample preparation

To assess the quality of genomic DNA received, we used Qbit (Life Technologies) and Nanodrop (Thermo Scientific) to measure the concentration of genomic DNA. Ideal samples had similar concentration estimates on both platforms, with $A_{230/260/230}$ ratios close to 1:1.8:1, corresponding to what is expected of pure DNA. All samples presented here passed this screening criterion.

Next we assessed the size of the genomic DNA received. For genomic DNA where the size range was less than 17kb, we used the Bioanalyzer 21000 (Agilent) to determine the actual size distribution. For genomic DNA where the size range was greater than 17kb, we opted for pulse field gel electrophoresis to better estimate the larger size distributions. The sizes of the genomic DNA for each sample are listed in Table 1.

To ensure that the library insert sizes were in the optimal size range, we sheared genomic DNA using gTubes if the apparent size was greater than 40 kb. Alternatively, if the size was less than 40kb, then the DNA was not sheared and carried straight through to library preparation. Extremely small fragments (<100bp) and adapter dimers are eliminated by Ampure Beads. Adapter Dimer (0-10bp) and small inserts (11-100bp) represented less than 0.01% of all the reads sequenced in all datasets. We additionally use the Blue Pippin (Sage Science) to select ensure that the libraries had a physical size of 10kb or greater. The size cutoffs used for each sample are listed in Table 1.

Analysis and Quality Filtering

To assess the quality of the libraries sequenced, we examined the percent of bases filtered by a standard QC procedure. Filtering conditions for high-quality SMRT sequence data are read score > 0.8, read length > 100, subread length > 500. In addition, the ends of reads are trimmed if they are outside of high-quality (HQ) regions, and adapter sequences between subreads are removed. All samples retained 71-97% of the bases after filtering.

To ensure that the sequences matched the model organism of interest, we examined the percent of post-filter bases that were mapped to the closest reference genome available. All samples had a mapping rate of 81-95%, with the exception of the *Neurospora* T1 sample that had a mapping rate of 62%. This sample may have some damaged DNA as it had been stored in a freezer for over 20 years. Nonetheless, preliminary unpublished results show that the sequence from the *Neurospora* T1 sample can be successfully assembled into a genome that is more contiguous than the existing reference genome for *Neurospora* [39].

Usage Notes

The datasets described in this paper were first released on DevNet [40], the PacBio Software Developer Community Network website, with brief descriptions on the PacBio blog. DevNet typically hosts open-source software; SampleNet [30], the PacBio Sample Preparation Community Network website, typically hosts protocols for DNA extraction and library preparation. These websites provide valuable data and documentation about the technology, but are not considered a part of the traditional academic record. This paper in *Nature Scientific Data* provides an opportunity to describe the methodology and characteristics of the eight datasets in more detail, creates a citable entity for the scientific community, and allows the data to be continually hosted and maintained by the Sequence Read Archive.

DNA sequencing instruments and chemistries change rapidly, and PacBio SMRT sequencing is no exception. The datasets presented here are from P4-C2 and P5-C3 polymerase-chemistry combinations, spanning release dates from late-2013 to early-2014. These datasets represent some of the longest read lengths to date for these chemistries, and can be used to benchmark and develop new algorithms and the state of the art as the technology evolves.

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NBACC, or Battelle National Biodefense Institute (BNBI) have any responsibility or liability for any use, misuse, inability to use, or reliance upon the information contained herein. The Department of Homeland Security does not endorse any products or commercial services mentioned in this publication. CMB was supported by Human Frontier Science Program Young Investigator grant RGY0093/2012.

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Author contributions

KEK prepared libraries, sequenced, and analyzed data for the *N. crassa* OR74A, *N. crassa* T1, and *D. melanogaster* samples. PP and DRR grew plants from seed, prepared libraries, and sequenced the *A. thaliana* P4C2 and P5C3 datasets. PB prepared libraries and sequenced the *E. coli* datasets. PJY and DE provided DNA for *N. crassa* T1. CY and SEC extracted DNA, and WWF collected male flies for the *D. melanogaster* dataset. NAR and JL extracted DNA and prepared libraries for and PP sequenced the *S. cerevisiae* 9464 sample. JML deposited data to the SRA. CSC, AP, CMB and JML analyzed the data and prepared the manuscript. CMB and JML coordinated the project.

Competing Financial Interests

The authors declare competing financial interests. KEK, PP, PB, CSC, NAR, DRR, and JML are employees of Pacific Biosciences of California, Inc., a company commercializing DNA sequencing technologies.

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