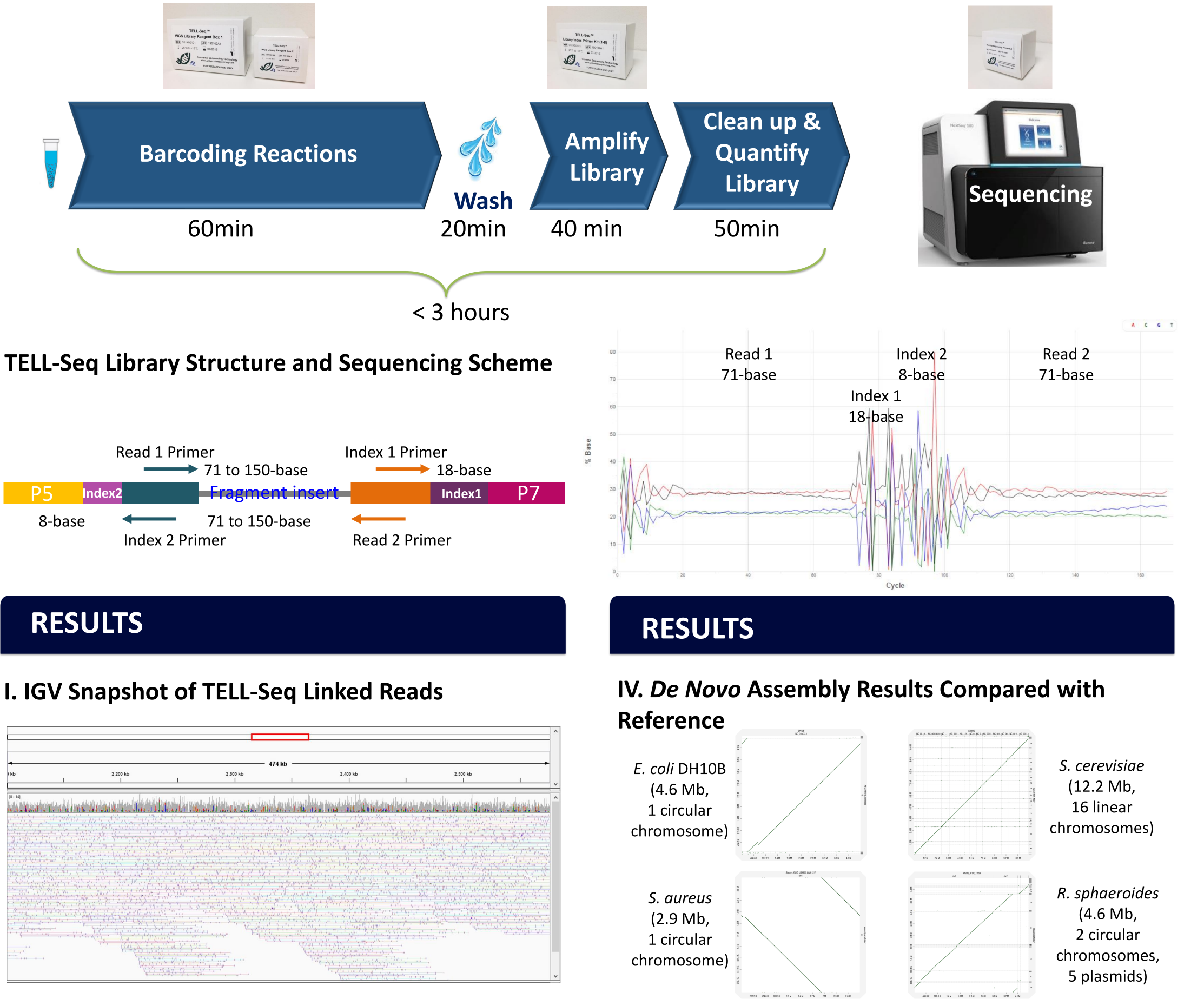
Simple and scalable genome analysis with TELL-SeqTM: from haplotype phasing to de novo assembly in a tube

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INTRODUCTION

<u>Transposase Enzyme Linked Long-read</u> <u>Sequencing</u> (TELL-SeqTM) is a simple and scalable NGS library technology to generate barcode linked-reads for genome scale haplotype phasing and/or *de novo* sequencing. In a PCR tube, under a standard NGS laboratory setting and without a need for any expensive protocol-specific instrument, TELL-Seq Whole Genome Sequencing (WGS) Library Prep Kit will generate an Illumina sequencing library in 3 hours. Multiple samples can be easily processed parallelly in a 96-well format when needed. The library protocol can be adjusted and used for various sizes of genomes ranging from bacteria to human. TELL-Seq library will replace both fragment library and mate-pair library altogether and become a new standard library method for WGS.

TELL-Seq Library Preparation and Sequencing Workflow



METHODS AND MATERIALS

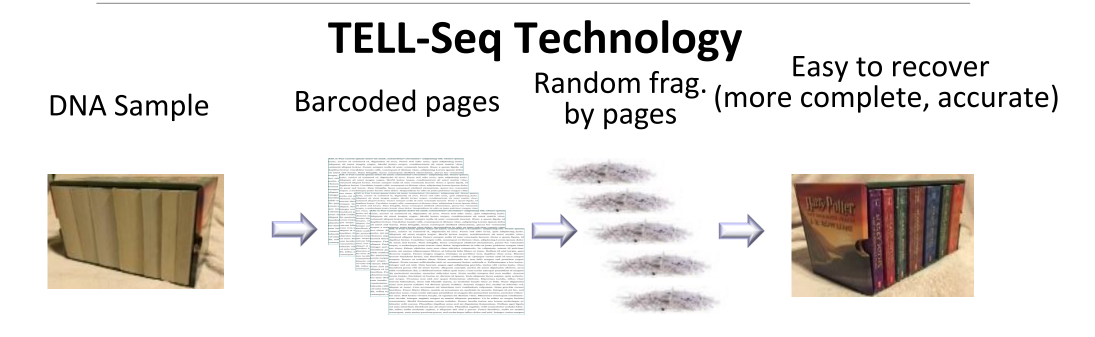
Standard NGS Sequencing

DNA Sample

Random fragmentation Difficult to recover (incomplete, error-prone) Shotgun sequencing



I. IGV Snapshot of TELL-Seq Linked Reads



- Standard NGS laboratory setting without any protocol specific capital expenditure
- All barcoding reactions are done in a 0.2ml PCR tube
- 0.5ng 5ng genomic DNA input based on the genome size
- Works for both small and large genomes ranging from bacteria to human genome
- Highly scalable workflow for preparation of multiple samples in parallel

TELL-Seq Barcoding Reactions



Transposon

GC Content Escherichi (50% GC) SLF3 SLF2 0P 90 Chromosome >K+K+ aureus (38% GC) 1,200 Linked reads 1,000 800 600 Rhodobacter sphaeroides 400 -(69% GC) Distal 200 reads Distance (bp) to next alignment (\log_{10}

III. TELL-Seq Results Compared with Mate Pair Data

E. coli DH10B	Genomic coverage (%)	Average gap size (bp)	Number of gaps	Total gap length (bp)	Full genes⁺	Sample Processing
Illumina mate pair data 1*	96.72	3,576	43	153,750	4,181	Input DNA: 1-4µg
Illumina mate pair data 2*	96.59	3,191	50	159,572	4,164	Lib.Prep Time: 2 days
TELL-Seq data 1: 2x71bp	96.23	3,213	55	176,739	4158	Input DNA: 1ng
TELL-Seq data 2: 2x148bp	96.36	2,889	59	170,500	4166	Lib.Prep Time: 3 hrs

V. Whole Genome Haplotype Phasing Results for NA12878

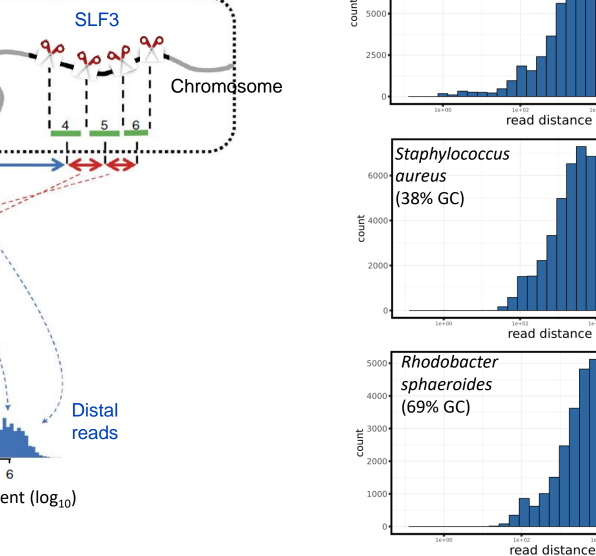
		7
Method	TELL-Seq	1e7 run190111nx:A500 DNA mass - SLFs size
Depth of genome coverage (Unique)	26X	4- sapiens
Largest Phased Block	27.6 Mb	SSE 3 - VI A A A A A A A A A A A A A A A A A A
N50 Phased Block	4.6 Mb	1-
Switch Error Rate	0.3%	0 50000 100000 150000 200000 250000 size of SLFs (>4k), mean=39080.10, median=32207.00

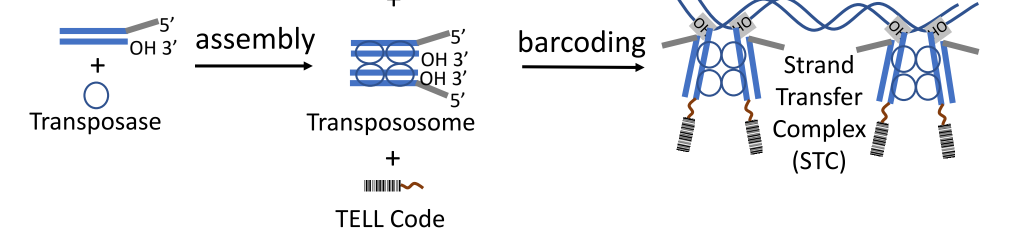
CONCLUSION

TELL-Seq library technology enables short read 2nd generation sequencing systems to generate long read like sequencing data with innovative molecular barcoding

250000 300

II. TELL-Seq Results for Three Bacteria with Different





TELL-Seq WGS Library Prep kit includes

- TELL-Seq[™] WGS Library Reagent Box 1
- TELL-Seq[™] WGS Library Reagent Box 2
- TELL-Seq[™] Library Index Primer Kit



TELL-Seq[™] Illumina Sequencing Primer Kit

⁺ The number of genes covered at 100%

* https://www.illumina.com/documents/products/appnotes/appnote-nextera-mate-pairbacteria.pdf

- library method. It is
- used for various sized genomes
- a single tube reaction without specialized instrument
- highly scalable for doing multiple samples in parallel
- one-stop solution for *de novo* sequencing and whole genome phasing
- automation friendly

TELL-Seq WGS library prep kit is available for early access soon. Visit <u>www.universalsequencing.com</u> for more information.



TELL-SeqTM: Tell you the true genome today!