From sample to fastq

Outline

• Brief overview of library preparation procedure

Sequencing costs

• Estimate cost for your own experiment

Requirements for library preparation protocol

- To prepare libraries for hundreds of samples, we need a protocol that is
 - Cheap
 - Efficient
 - Reliable

• Sometimes robustness to sample degradation is also important

One example of a library preparation technique



RESEARCH ARTICLE

Inexpensive Multiplexed Library Preparation for Megabase-Sized Genomes

Michael Baym¹°, Sergey Kryazhimskiy^{2,3}°, Tami D. Lieberman¹°, Hattie Chung¹°, Michael M. Desai^{2,3,4}*, Roy Kishony^{1,5}*

- 1 Department of Systems Biology, Harvard Medical School, Boston, Massachusetts, United States of America, 2 Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, Massachusetts, United States of America, 3 FAS Center for Systems Biology, Harvard University, Cambridge, Massachusetts, United States of America, 4 Department of Physics, Harvard University, Cambridge, Massachusetts, United States of America, 5 Faculty of Biology and Department of Computer Science, Technion-Israel Institute of Technology, Haifa, Israel
- These authors contributed equally to this work.
- * mmdesai@fas.harvard.edu (MB); roy_kishony@hms.harvard.edu (RK)



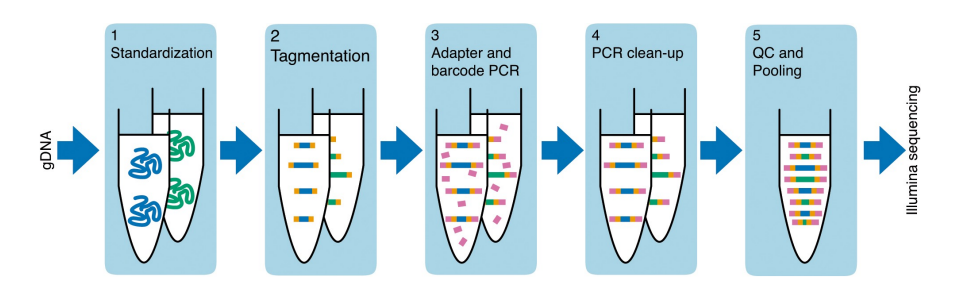
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Abstract

Whole-genome sequencing has become an indispensible tool of modern biology. However, the cost of sample preparation relative to the cost of sequencing remains high, especially for small genomes where the former is dominant. Here we present a protocol for rapid and inexpensive preparation of hundreds of multiplexed genomic libraries for Illumina sequencing. By carrying out the Nextera tagmentation reaction in small volumes, replacing costly re-

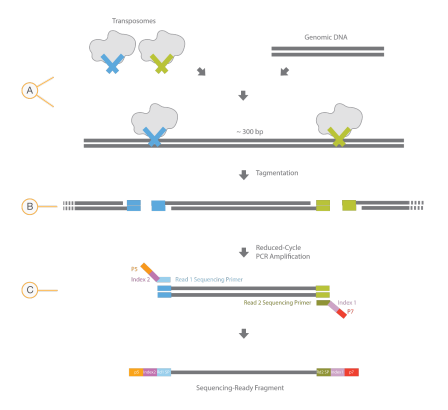
Library preparation protocol



Transposome with adapters combined with template DNA

Tagmentation to fragment and add adapters

Limited-cycle PCR to add index adapter sequences



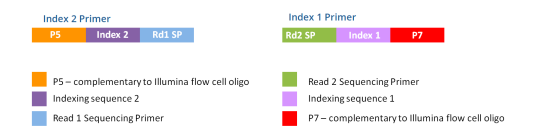
Transposome with adapters combined with template DNA

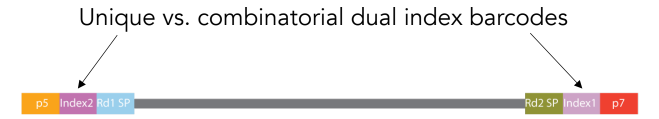
Tagmentation to fragment and add adapters

Limited-cycle PCR to add index adapter sequences

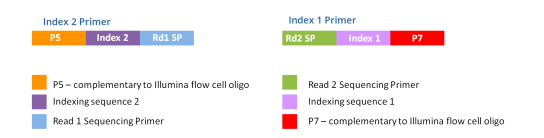
Other great library preparation methods work by adapter ligation (rather than tagmentation)



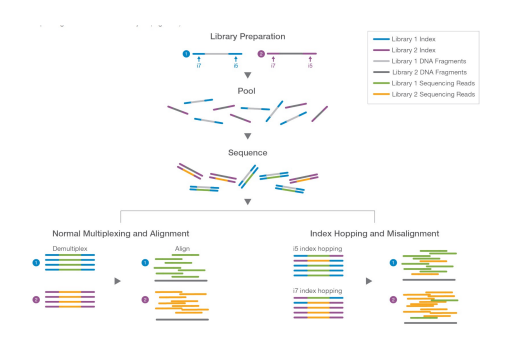




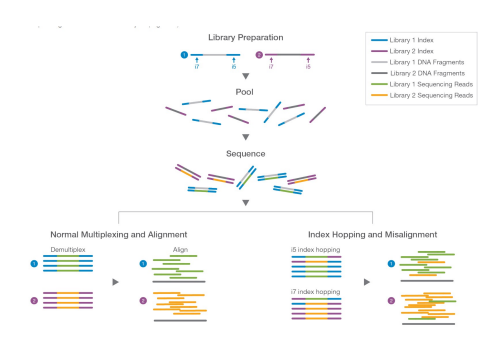
Sequencing-Ready Fragment



Beware that index hopping can cause misassigned sequence reads when using combinatorial index barcodes

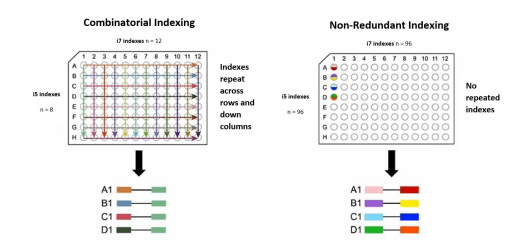


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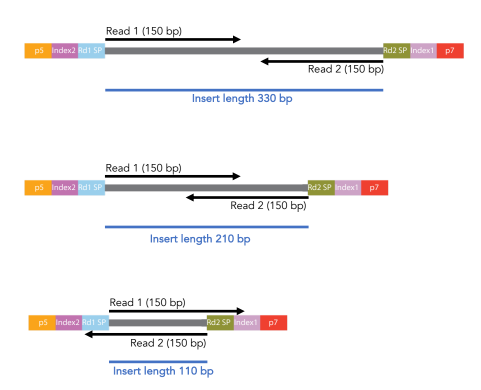


Index hopping often affects 0.1-2% of reads!

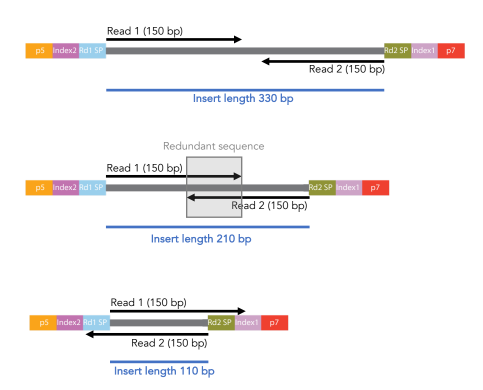
Unique dual index recommended even though they are more expensive than combinatorial dual index adapters



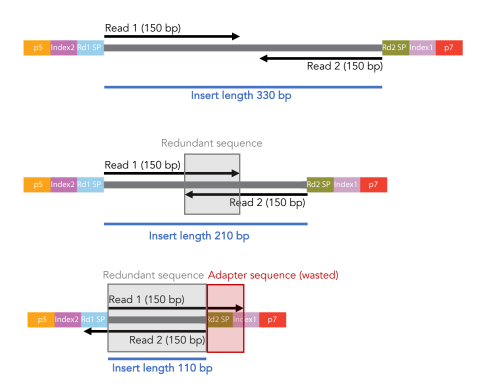
Insert length relative to read length



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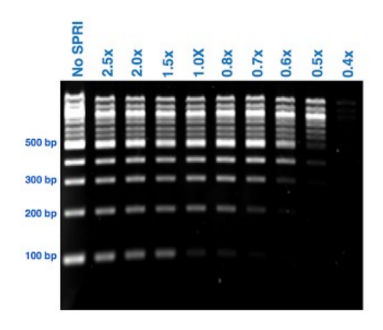


Insert length relative to read length

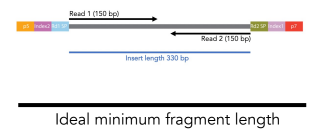


Size selection with Ampure beads

Tune the size distribution of your library fragments to minimize "waste" of sequence due to paired-end overlap and adapter read-through

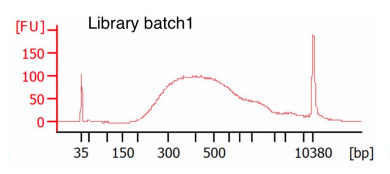


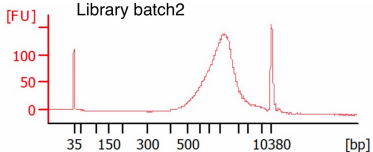
Ideally, we want all library fragments to be greater than the adapter length plus 2 x the read length (for PE)



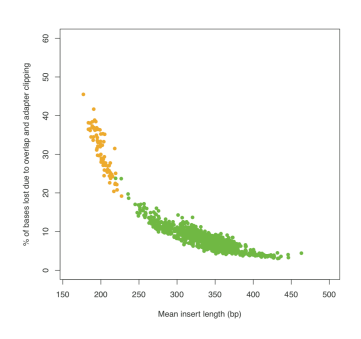
http://enseqlopedia.com/2012/04/how-do-spri-beads-work/

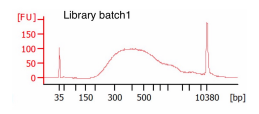
Two examples of our library pools

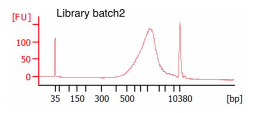




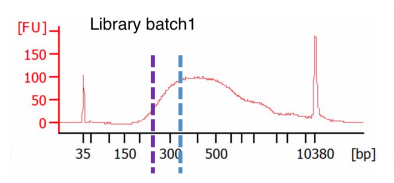
The library fragment size distribution can substantially influence the amount of data lost in data QC steps

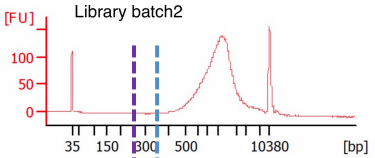






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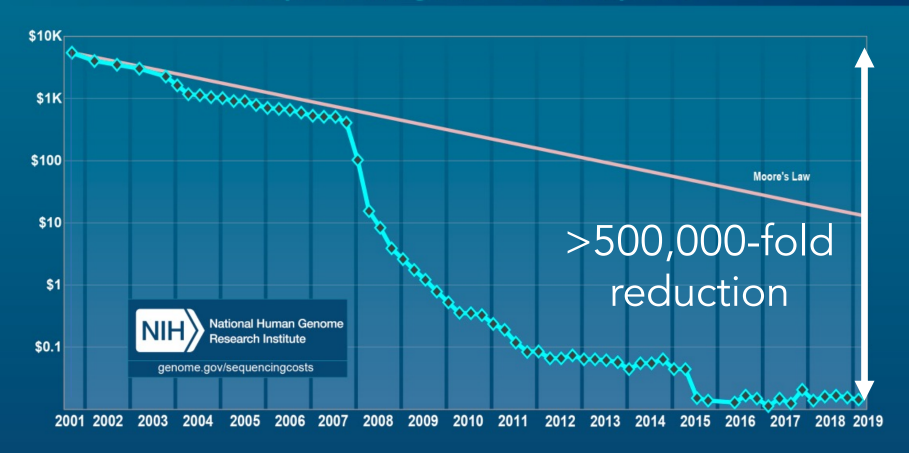
The length of Nextera adapters is 138 bp and libraries were sequenced with 2*125bp reads

- → Minimum fragment length to avoid overlap
- Minimum fragment length to avoid adapter read-through

388bp

250bp

Cost per Raw Megabase of DNA Sequence



What is the current price for 2x sequencing of an Atlantic silverside (including library preparation)?

Genome size ~650 Mb

\$12	
\$25	
\$48	
\$72	
\$213	



1 USD ≈ 1 EURO



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Example costs for other genome sizes Incl. library preparation and sequencing to 2x genome coverage*

	Cost per san	nple (USD) ^a		
Genome size (Gb)	1× coverage	2× coverage	Example organisms	
0.2	11 (3)	13 (5)	Fruit fly, honeybee, arabidopsis	
0.65	16 (8)	25 (17)	Atlantic silverside, stickleback, eastern oyster	
1	21 (13)	34 (26)	Zebra finch, chicken, purple sea urchin	
3	47 (39)	86 (78)	Human, Atlantic salmon, African clawed frog	

^{*}Cost estimates do not include labor and assume sequencing costs ~13 USD per Gb in shared S4 lanes on an Illumina NovaSeq and 8 USD per sample for library preparation

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Compare to:

\$30 per sample for RADseq \$15 per sample for RADcapture

Meek and Larson, 2019, Mol Ecol Res

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Exercise – how much will your experiment cost?

- Assumed costs:
 - Library preparation: \$8 per sample
 - Sequencing: \$13 per Gb
 - Target coverage per sample: Expect to lose at least 30-50% of your data in filtering

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- Assumed costs:
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- Example: I would like to have 1x coverage for downstream analysis for 40 individuals from each of 5 populations (200 individuals total) of my favorite animal with a genome size of ~800 Mb
- Calculation: I will target 2x coverage raw sequencing. This means
 - 2 * 800 Mb/individual * 200 individuals = 320,000 Mb (320 Gb)

My total cost is thus (320 Gb * \$13/Gb) + (200 libraries * \$8 per library) = **\$5,760**