

Genome assembly post-processing

Nadège Guiguelmoni

Read pre-processing

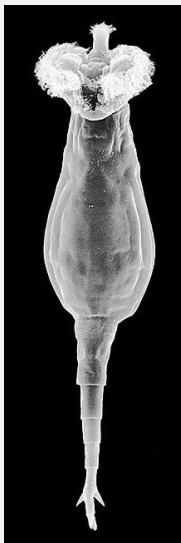
- ▶ **Adapter trimming**
- ▶ **Read filtering:** select longest/highest quality: Filtlong
- ▶ **Read correction:** reduce error rate of long reads
 - ▷ **self correction:** long reads only
 - ▷ **hybrid correction:** long reads & short reads

Assembly post-processing

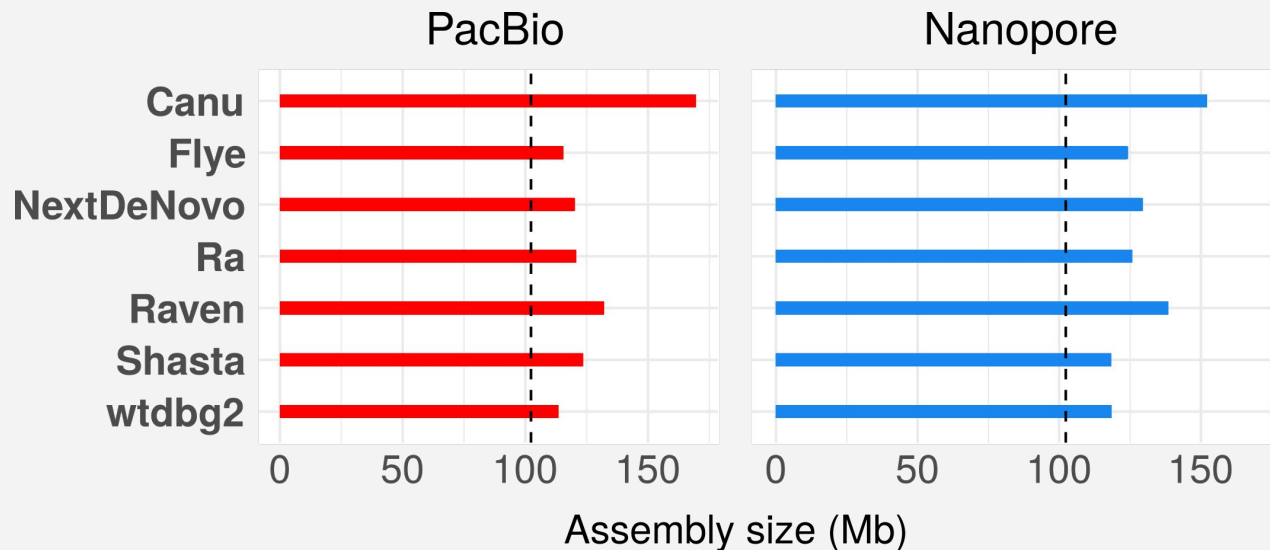
- ▶ **Polishing:** reduce errors
- ▶ **Haplotig purging:** remove uncollapsed haplotypes
- ▶ **Scaffolding:** increase contiguity
- ▶ **Gap filling:** find missing sequences

Haplotig purging

Adineta vaga



Expected haploid size 102 Mb



Who Needs Sex (or Males) Anyway?
Liza Gross, PLoS Biology, 2007

Haplotig purging

Haplotype 1 ATTACCAGTCTCAAT**TGGATGGCTACTCTTTGACGATAGCT**

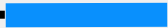
Haplotype 2 ATTACCAGTCTCAA**AGGCTGCTAGTGTTTGACGATAGCT**

Assembly process




Assembly output

Good haploid assemblies

contig 1 ———  ——— ✓

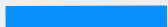
OR


contig 1 ———  ——— ✓

Problematic assembly

contig 1 ———

contig 2 ———

contig 3  X

contig 4 

Haplotig purging

HaploMerger2

HaploMerger2: rebuilding both haploid sub-assemblies from high-heterozygosity diploid genome assembly

Shengfeng Huang*, Mingjing Kang and Anlong Xu

Identifying and removing haplotypic duplication in primary genome assemblies

Dengfeng Guan^{1,2}, Shane A. McCarthy ², Jonathan Wood³, Kerstin Howe ³,
Yadong Wang^{1,*} and Richard Durbin ^{2,3,*}

purge_dups

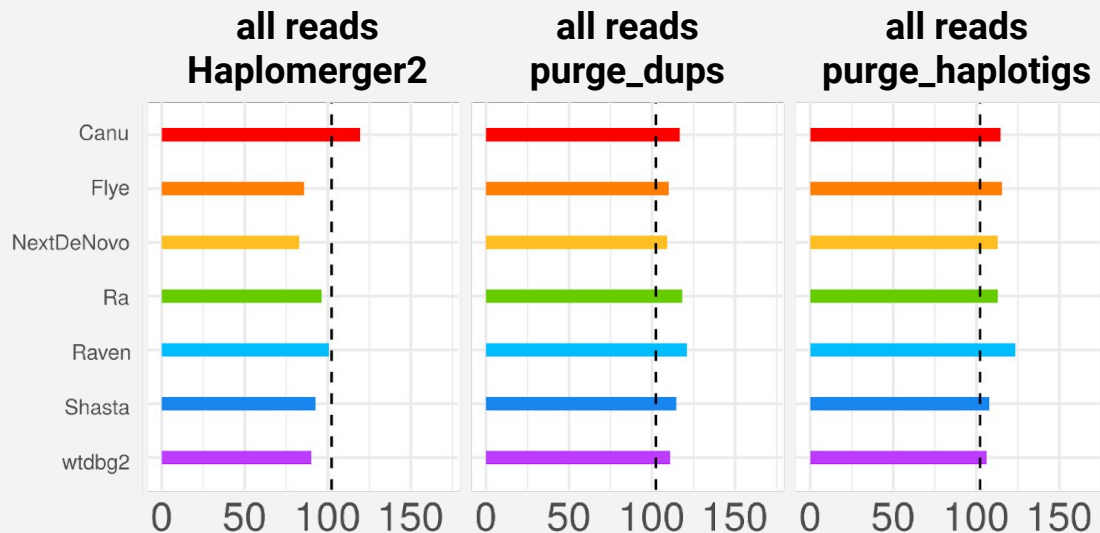
Purge Haplotigs

Purge Haplotigs: allelic contig reassignment for third-gen diploid genome assemblies

Michael J. Roach*, Simon A. Schmidt and Anthony R. Borneman

Haplotig purging

PacBio assemblies

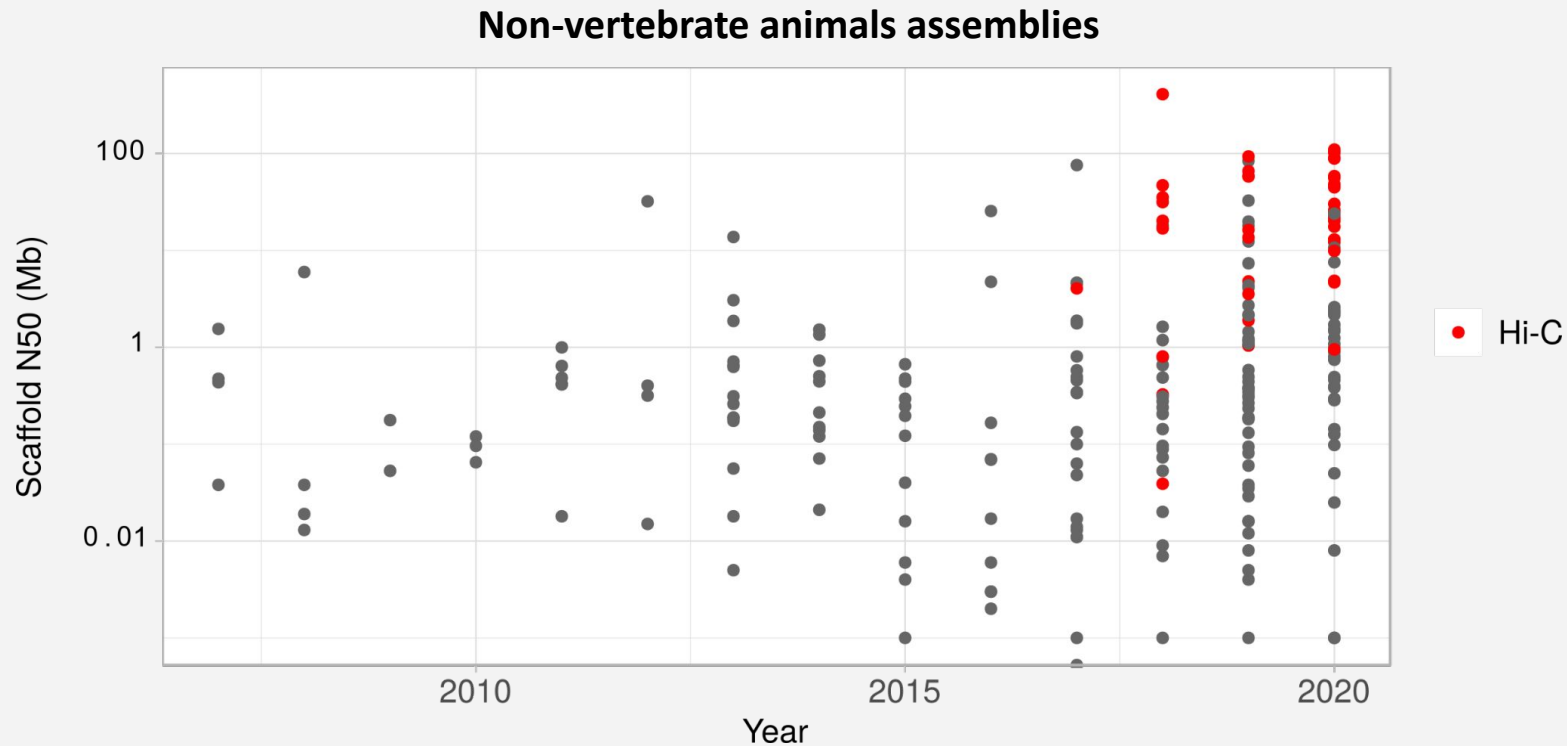


Scaffolding approaches

Scaffolding: grouping and orienting contigs to build chromosome-level scaffolds

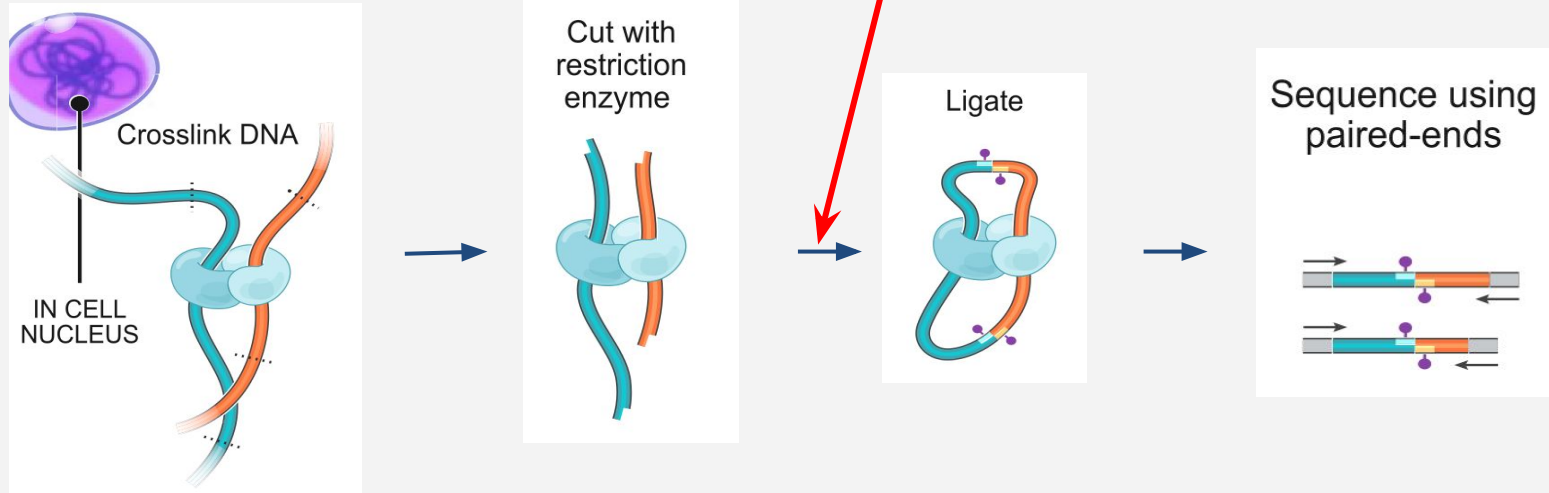
- ▶ **Long reads**
- ▶ **Linked reads:** barcoded short reads
- ▶ **Hi-C**

Scaffolding approaches: Hi-C scaffolding



Scaffolding approaches: Hi-C scaffolding

Hi-C



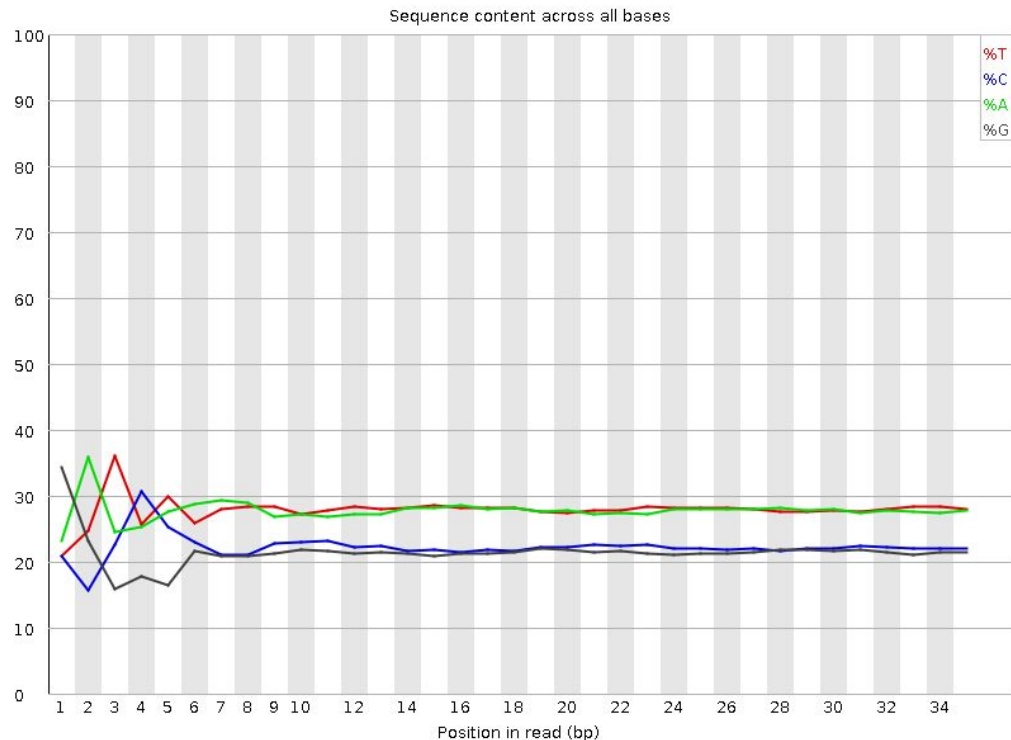
A 3D map of the human genome at kilobase resolution reveals principles of chromatin looping. Rao et al., 2014

Scaffolding approaches: Hi-C scaffolding

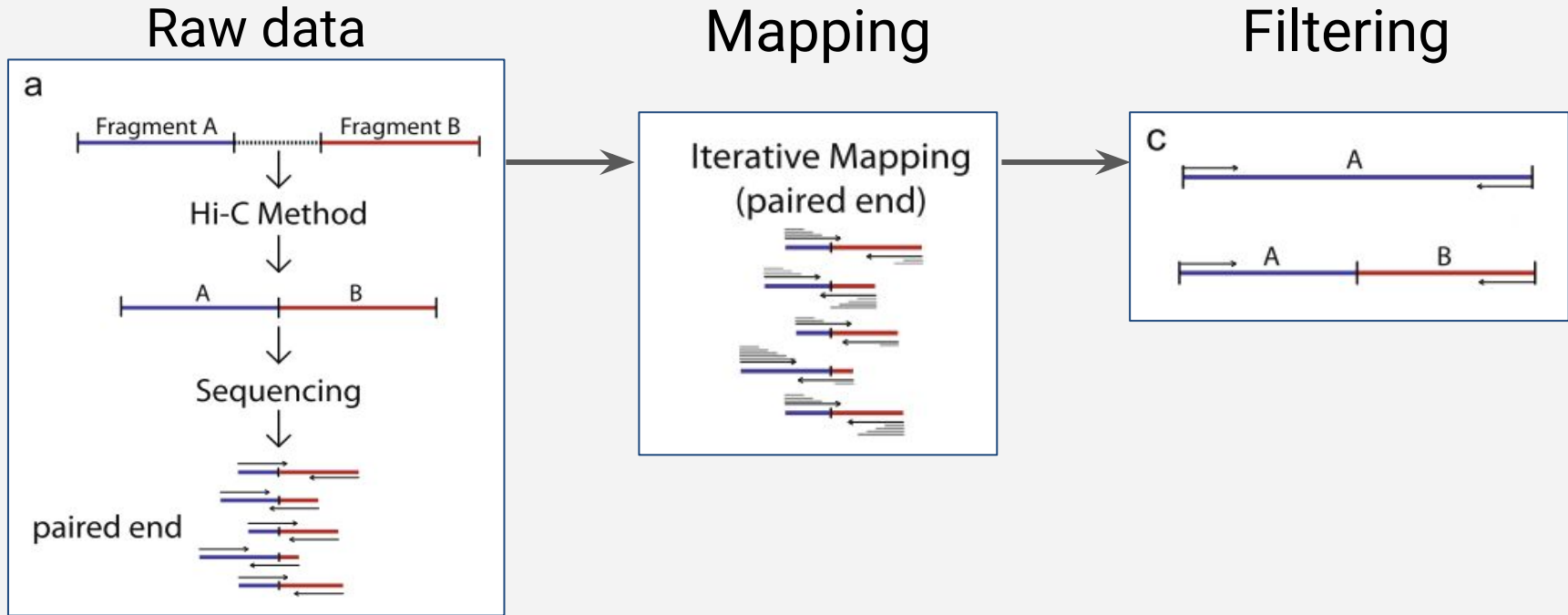
Summary

- ✓ [Basic Statistics](#)
- ✓ [Per base sequence quality](#)
- ✓ [Per tile sequence quality](#)
- ✓ [Per sequence quality scores](#)
- ⚠ [Per base sequence content](#)
- ✓ [Per sequence GC content](#)
- ✓ [Per base N content](#)
- ✓ [Sequence Length Distribution](#)
- ⚠ [Sequence Duplication Levels](#)
- ✓ [Overrepresented sequences](#)
- ✓ [Adapter Content](#)

⚠ Per base sequence content

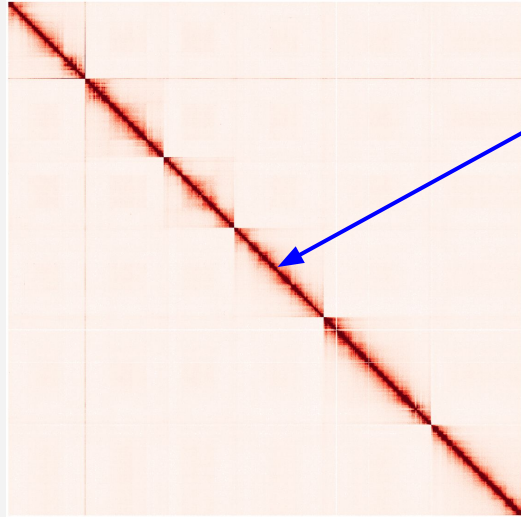


Scaffolding approaches: Hi-C scaffolding



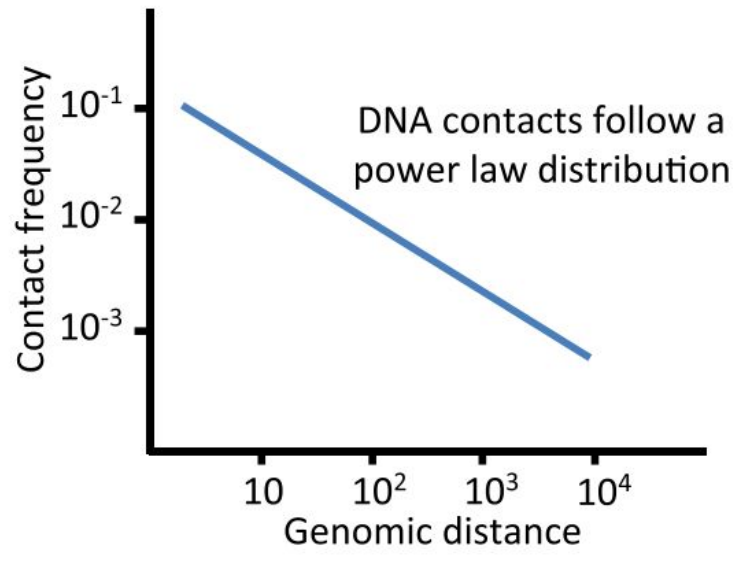
The Hitchhiker's guide to Hi-C analysis: Practical guidelines. Lajoie et al., 2015

Scaffolding approaches: Hi-C scaffolding



Contact map of
Caenorhabditis elegans

contact frequency = $f(\text{genomic distance})$



Contact genomics: scaffolding and phasing (meta)genomes using chromosome 3D physical signatures. Flot et al., 2015

Scaffolding approaches: Hi-C scaffolding

High-throughput genome scaffolding from *in vivo* DNA interaction frequency

Noam Kaplan ✉ & Job Dekker ✉

dnaTri

Lachesis

Chromosome-scale scaffolding of *de novo* genome assemblies based on chromatin interactions

Joshua N Burton ✉, Andrew Adey, Rupali P Patwardhan, Ruolan Qiu, Jacob O Kitzman & Jay Shendure

High-quality genome (re)assembly using chromosomal contact data

Hervé Marie-Nelly ✉, Martial Marbouty, Axel Cournac, Jean-François Flot, Gianni Liti, Dante Poggi Parodi, Sylvie Syan, Nancy Guillén, Antoine Margeot, Christophe Zimmer ✉ & Romain Koszul ✉

GRAAL

Scaffolding approaches: Hi-C scaffolding

De novo assembly of the *Aedes aegypti* genome using Hi-C yields chromosome-length scaffolds

Olga Dudchenko^{1,2,3,4}, Sanjit S. Batra^{1,2,3,*}, Arina D. Omer^{1,2,3,*}, Sarah K. Nyquist^{1,3}, Marie Hoeger^{1,3}, Neva C. Durand^{1,...}

3D-DNA

SALSA2

Integrating Hi-C links with assembly graphs for chromosome-scale assembly

Jay Ghurye, Arang Rhie, Brian P. Walenz, Anthony Schmitt, Siddarth Selvaraj, Mihai Pop, Adam M. Phillippy , Sergey Koren 

instaGRAAL: chromosome-level quality scaffolding of genomes using a proximity ligation-based scaffolder

[Lyam Baudry](#), [Nadège Guiguelmoni](#), [Hervé Marie-Nelly](#), [Alexandre Cormier](#), [Martial Marbouty](#), [Komlan Avia](#), [Yann Loe Mie](#), [Olivier Godfroy](#), [Lieven Sterck](#), [J. Mark Cock](#), [Christophe Zimmer](#), [Susana M. Coelho](#)  & [Romain Koszul](#) 

instaGRAAL

Scaffolding approaches: Hi-C scaffolding

And in 2021

EndHiC: assemble large contigs into chromosomal-level scaffolds using the Hi-C links from contig ends

Sen Wang, Hengchao Wang, Fan Jiang, Anqi Wang, Hangwei Liu, Hanbo Zhao, Boyuan Yang, Dong Xu, Yan Zhang, Wei Fan

Efficient iterative Hi-C scaffolder based on N-best neighbors

Dengfeng Guan^{1,2,4}, Shane A. McCarthy^{2,3}, Zemin Ning³, Guohua Wang^{1*}, Yadong Wang^{1*} and Richard Durbin^{2,3*}

YaHS: yet another Hi-C scaffolding tool

Chenxi Zhou¹, Shane A. McCarthy^{1,2}, Richard Durbin^{1,2}

Scaffolding approaches: Hi-C scaffolding



Cheetah



Yellow fever mosq...



Hoary bat



Red panda



Allen's Swamp Mo...



American alligator



Chinese alligator



Asian small-clawed...



California sea hare



Golden eagle



Peanut



Hog deer



Bryde's whale



Ringtail



Cacomistle



www.dnazoo.org

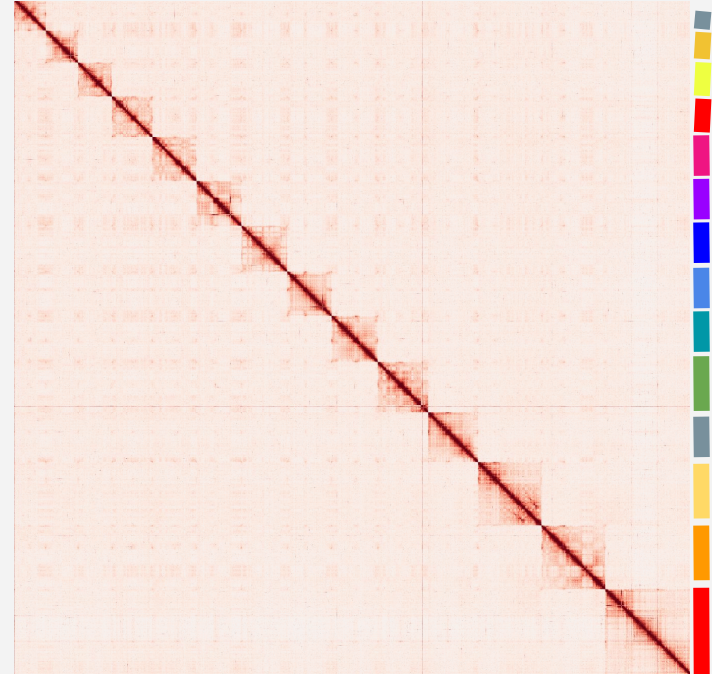
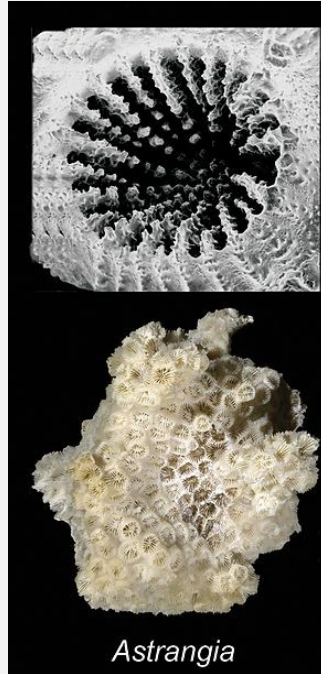
Scaffolding approaches: Hi-C scaffolding

Astrangia poculata

(coral)

14 scaffolds

455 Mb

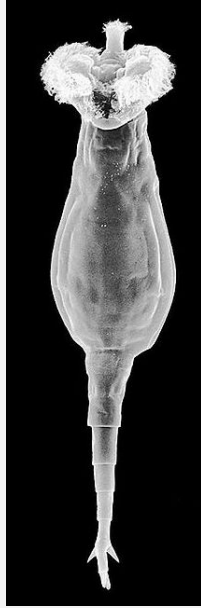


Hi-C contact map of *Astrangia poculata*

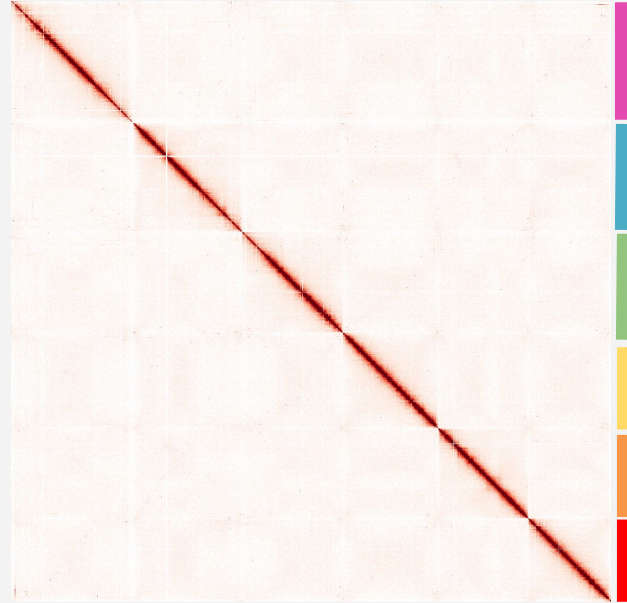
Scaffolding approaches: Hi-C scaffolding

Adineta vaga (rotifer)

6 scaffolds



Who Needs Sex (or Males) Anyway?
Liza Gross, PLoS Biology, 2007



Hi-C contact map of *Adineta vaga*

Scaffolding approaches: Hi-C scaffolding

“What coverage should I get?”

→ Arima recommends 200 millions pairs per Gb

Species	Size	# fragments	# Hi-C pairs	Hi-C mapping
<i>Adineta vaga</i>	101 Mb	30	55 millions	83%
<i>Astrangia poculata</i>	455 Mb	2995	723 millions	67%
<i>Flaccisagitta enflata</i>	929 Mb	6612	489 millions	37%
<i>Mercenaria mercenaria</i>	1.86 Gb	5118	455 millions	55%

And then...

- ▶ **Gap filling:** TGS-GapCloser...
- ▶ **Polishing:** using high-accuracy reads, HyPo, Racon...

Gap filling & Polishing

	Scaffolds	After TGS-Gapcloser	After HyPo
<i>Flaccisagitta enflata</i>	9,239	3,694	1,476
<i>Norana najaformis</i>	860	748	632
<i>Lucinoma borealis</i>	24,786	5,093	2,135

Assembly pipeline

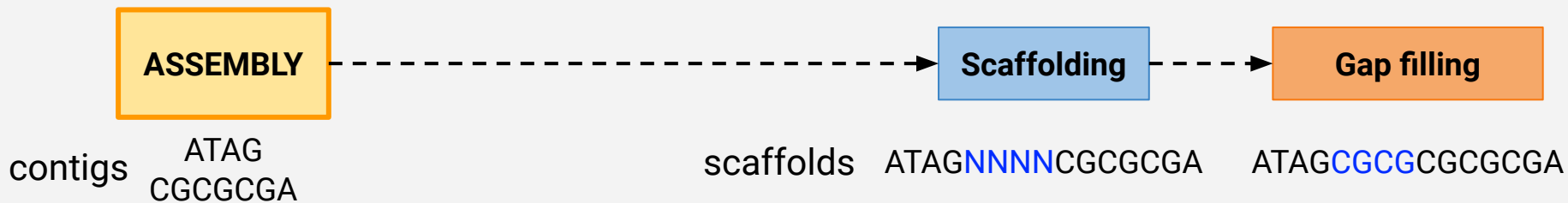
ASSEMBLY

reads	ATTTGTACG GTACGGACA GGACATAGTA
contig	ATTTGTACGGACATAGTA

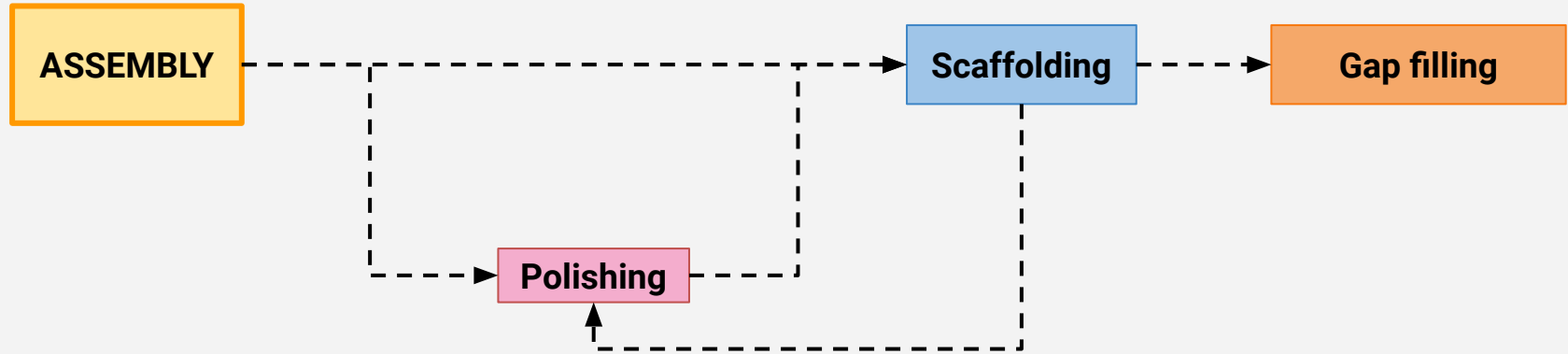
Assembly pipeline



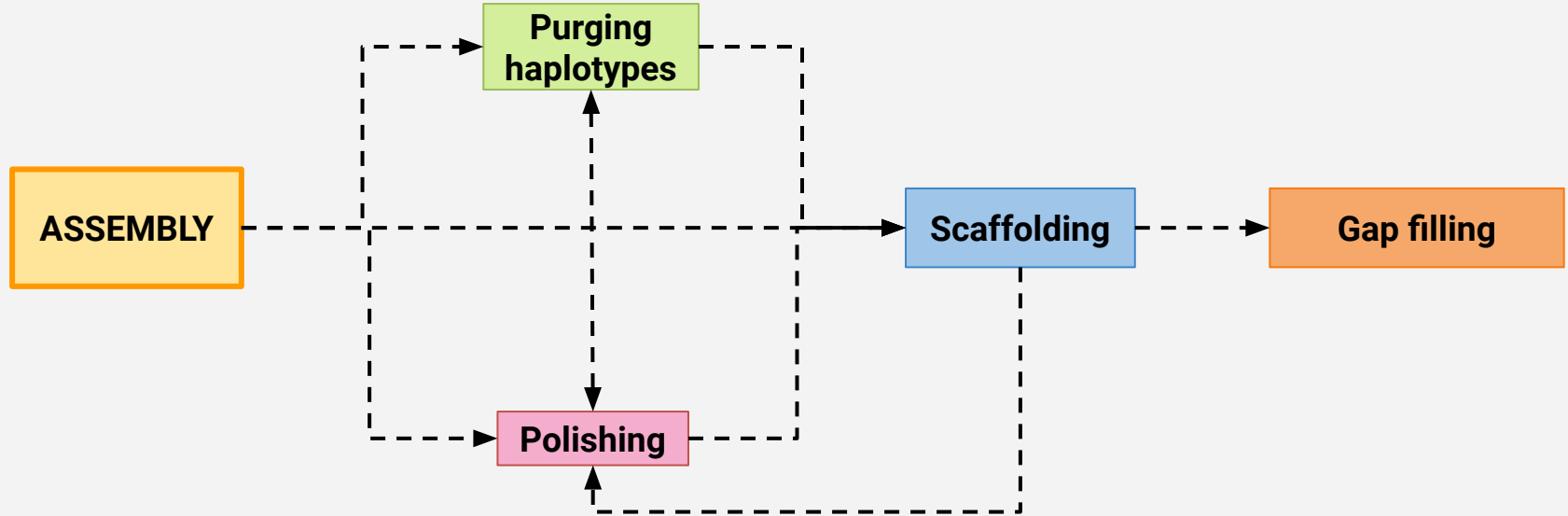
Assembly pipeline



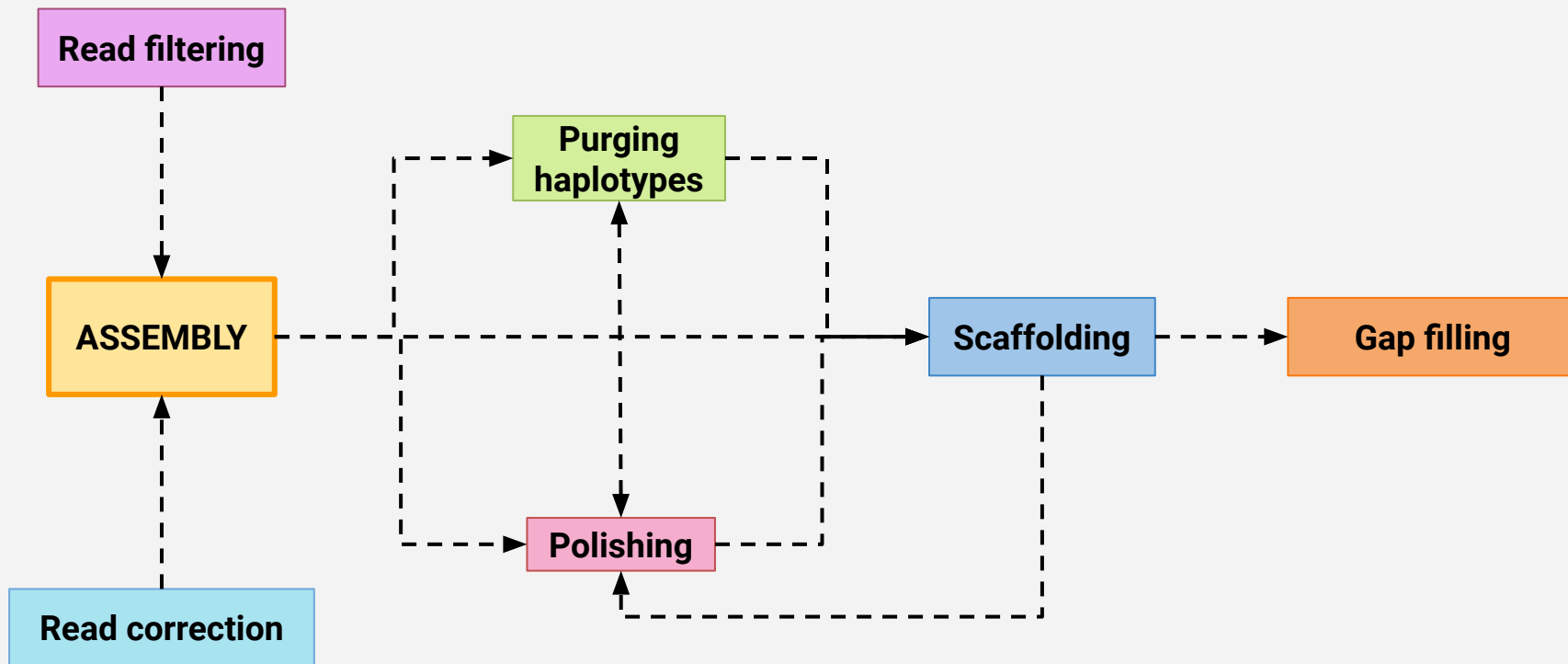
Assembly pipeline



Assembly pipeline



Assembly pipeline



Assembly pipeline

