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Maritime pine (*Pinus pinaster*) de novo transcriptome assembly



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Rationale

1

Fast increase of world population implies increased wood demand (heating, pulp & paper, construction).
 Developing the wood resource calls for cultivated forest species such as maritime pine.

2

Major steps of a typical sequence processing pipeline include sequence cleaning, fragment assembly and clustering.

3

The sequence cleaning step usually includes filtering of duplicated reads, short reads, low quality reads, contaminations, and reads containing ambiguous bases (N) above a certain threshold.

Starting Point

Pinus pinaster contigs database: TGICL assembly mainly based on Roche 454 GS-FLX sequences

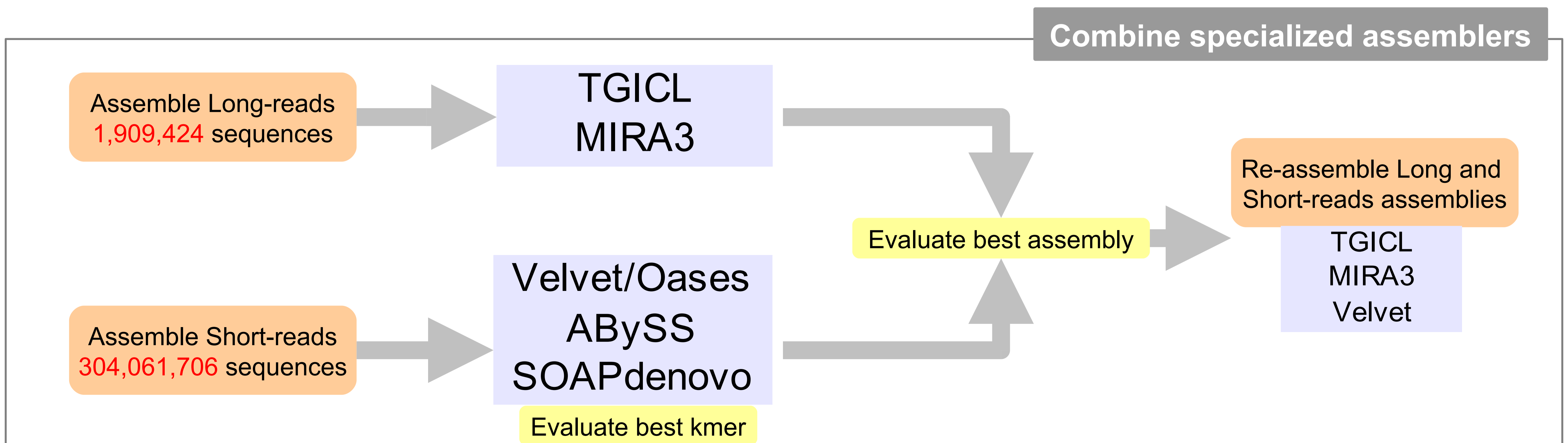
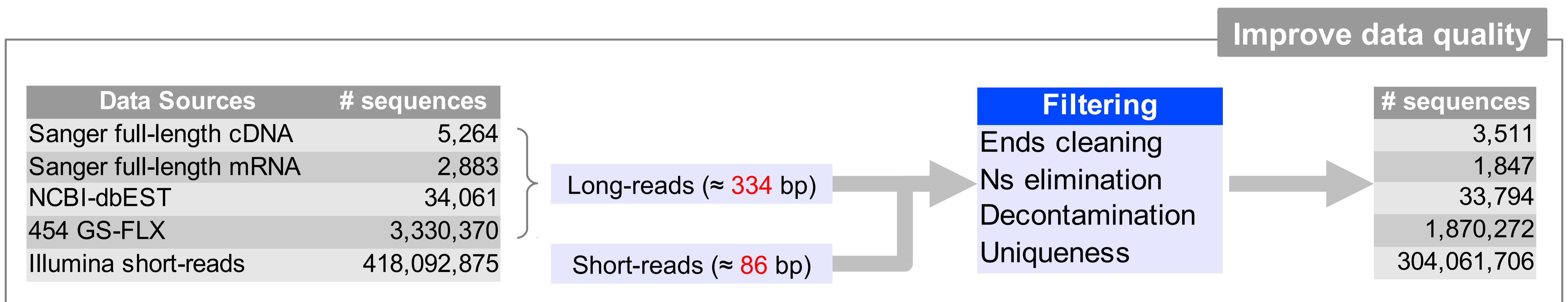


Assembly:	lpp2, July 2010
Number of base pairs:	102,376,712
Number of sequences:	2,016,588
Total number of contigs:	198,425
Number of singlets:	124,542
Polymorphism:	July 2010
Number of putative SNPs:	30,803
Number of contigs including SNP:	11,936

http://genotoul-contigbrowser.toulouse.inra.fr:9092/Pinus_pinaster2/index.html



Present Strategy



StatSeq'2011

14-15 April, Toulouse. The third StatSeq workshop on
 "Statistical challenges on the 1000€ genome sequences in plants"

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