

#### Pipeline for Haplotype Frequencies Estimation from Pooled Targeted Sequencing in Maize

Minguella Raphaël, Aurélie Canaguier, Delphine Madur, Aurélie Bérard, Isabelle Le Clainche, Agustin O. Galaretto, Damien Hinsinger, Stephane Nicolas, Patricia Faivre Rampant

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Minguella Raphaël<sup>1</sup>, Canaguier Aurélie<sup>1</sup>, Madur Delphine<sup>2</sup>, Berard Aurélie<sup>1</sup>, Le Clainche Isabelle<sup>1</sup>, Galaretto Agustin-Oscar<sup>2</sup>, Hinsinger Damien D.<sup>1</sup>, Nicolas Stéphane D.<sup>2</sup>, Faivre Rampant Patricia<sup>1</sup>

<sup>2</sup> INRAE, GQE, Gif-sur-Yvettes, <sup>1</sup> INRAE, EPGV, Evry, France France

#### 1. Background

Haplotypes are useful markers in population genetics due to a tighter link to populations history than SNP and are therefore considered more informative for populations structuration analyses. However, capturing populations diversity can be challenging because it requires to genotype many individuals which can be very expensive. An usual solution is to genotype populations in **pool**, meaning that several individuals of a same population are mixed and their DNA extraction is done in pool. Unfortunately, information about haplotypes is lost during the process because the DNA is fragmented and mixed. Several approaches have been proposed to rebuild haplotypes with reads overlap. Here implemented one of these algorithm in a pipeline to estimates short haplotypes frequencies from targeted genotyping by sequencing (tGBS) technology, which reduces costs by sequencing only desired genomic regions.

We assessed the accuracy of our pipeline using control pools with known haplotypes frequencies and we show that our pipeline gives correct haplotypic frequencies estimations when frequencies are higher than 5%.

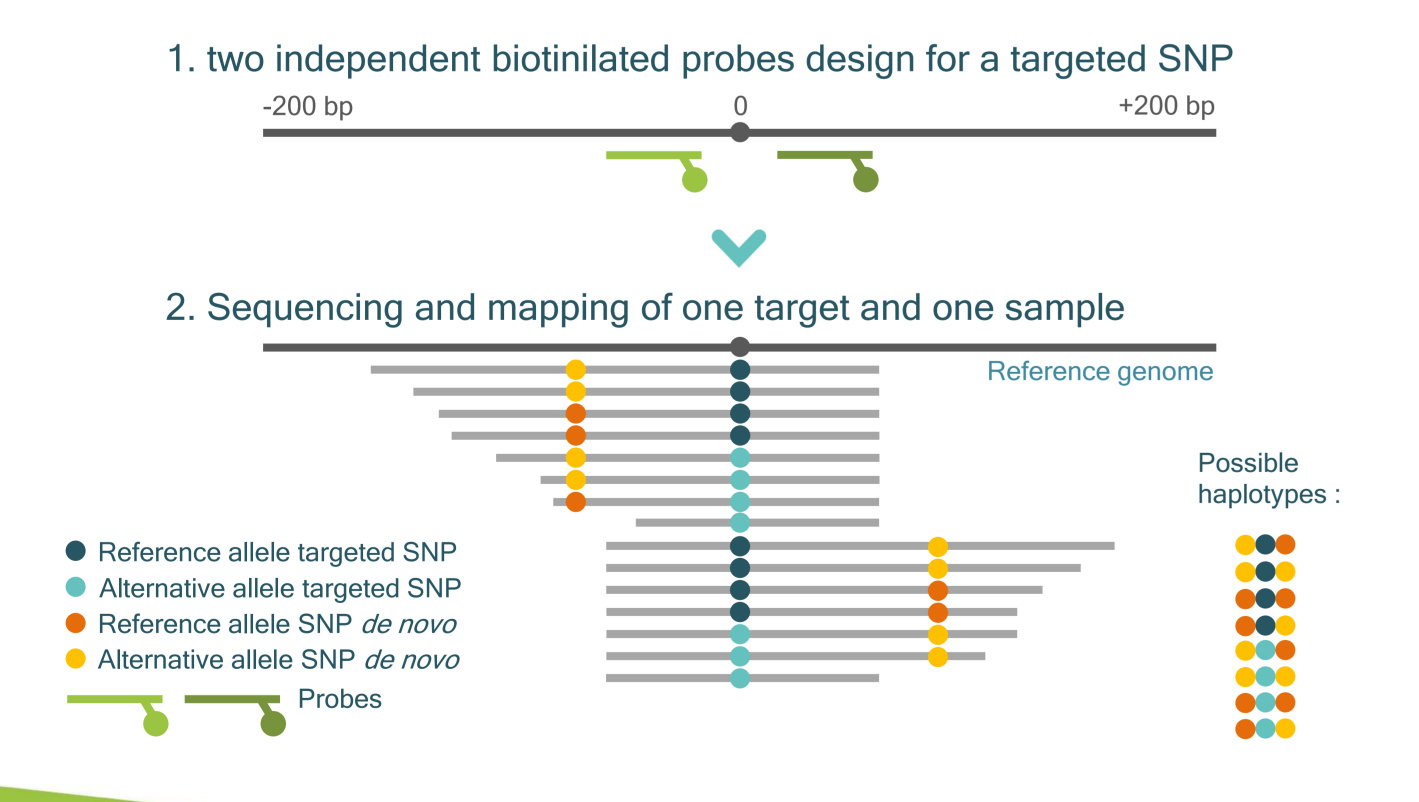
#### 2. Control pools design

To evaluate haplotypic frequencies estimation quality we designed control pools that are mixes of DNA from homozugous inbred lines in known proportions (or known F1 hybrids). Therefore, we can calculate expected haplotype frequencies for each control pool from the haplotyping of inbred lines that is assumed to be correct, and then, compare it to the observed frequencies in each control pools. We used 30 control pools: 5 F1 hybrids and 25 mixes of 3 inbred lines including different genetic groups.



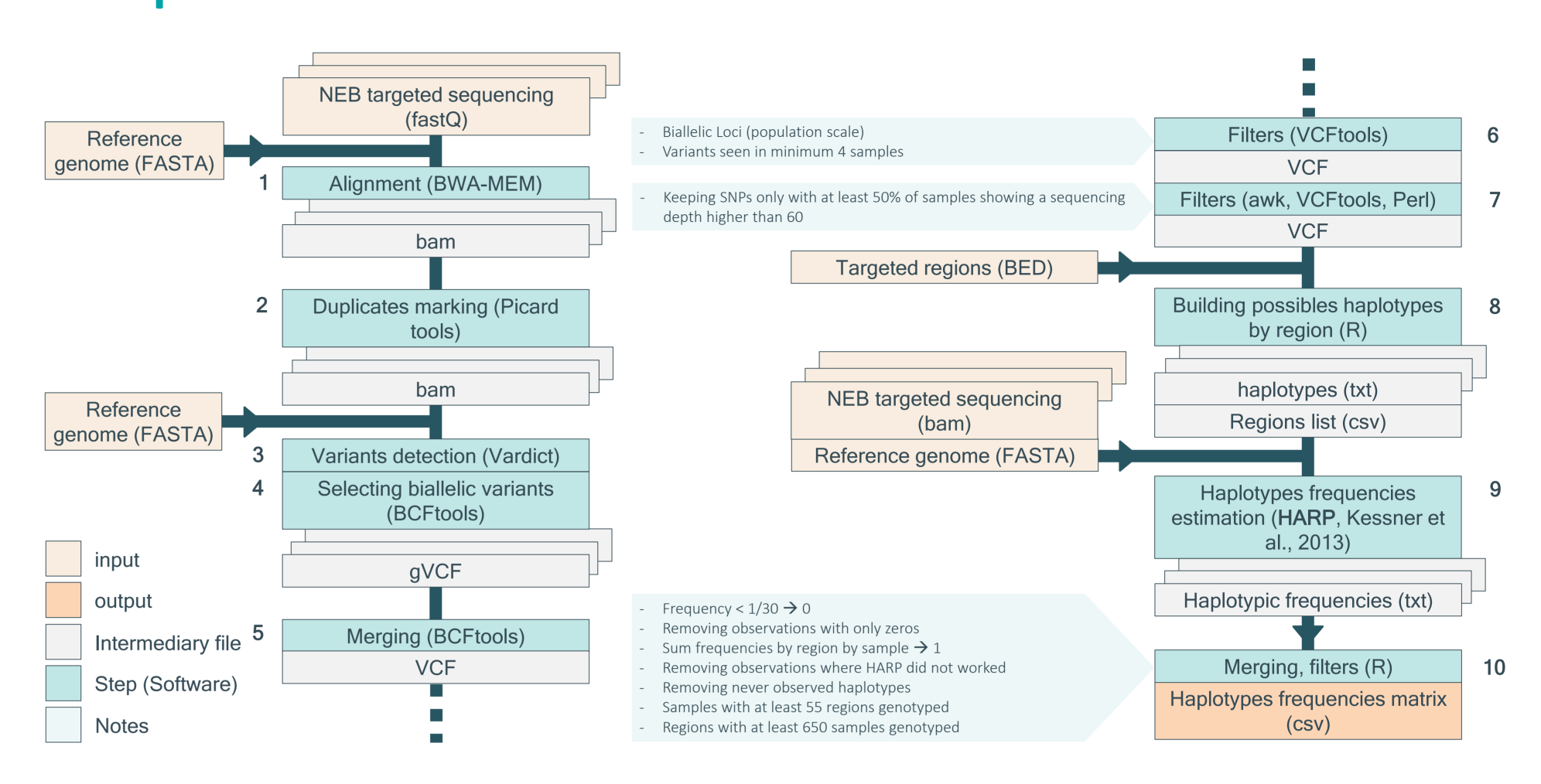
Pooled *Illumina®* sequencing of inbred lines and control pools

## 3. tGBS: NEBnext® direct genotyping solution



## Pipeline for Haplotype Frequencies Estimation from Pooled Targeted Sequencing in Maize

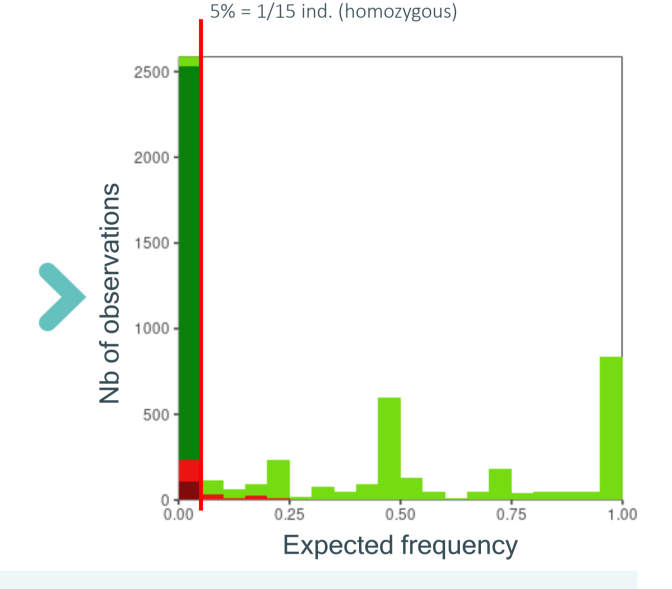
#### 4. Pipeline



#### 5. Performances

Haplotypes presence or absence To estimate the quality of haplotypes detection, we assessed the qualitative detection of this haplotype (right) for each haplotypes of each region and each pool. Unexpected and not observed Then, we calculated for each expected frequency the proportion of each category Unexpected but observed that we plotted (right). When haplotype frequency >5%: - Very few false positives (=unexpected

Good haplotypes detection if found in at least 1/15 ind. in the pool



Nb of expected haplotypes

but observed haplotypes)

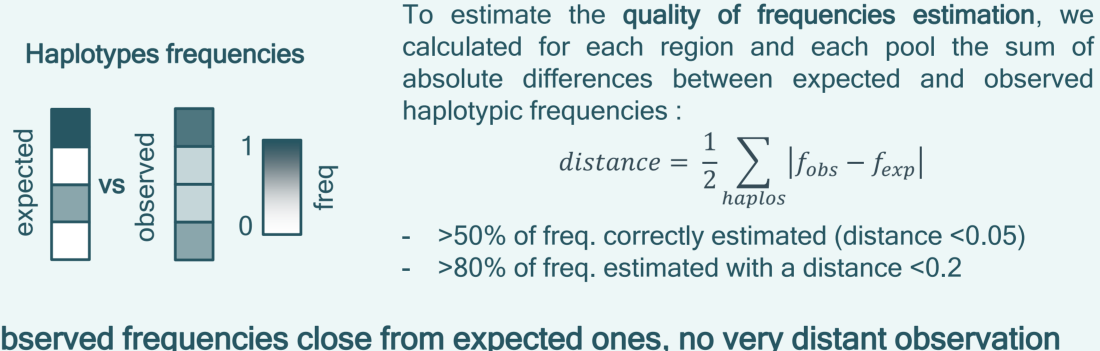
haplotypes)

- Very few false negatives (=undetected

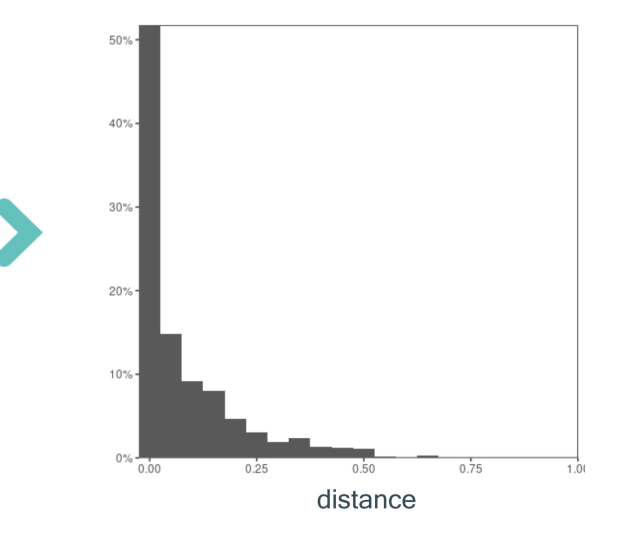
To estimate the ability of our pipeline to properly detect all haplotypes in a pool, we calculated the number of observed and expected haplotypes for each region of each control pool:

- Correct number of detected haplotypes (1-2 haplotypes in a pool) - When >2 haplotypes in the pool : a few undetected haplotypes
- >3 haplotypes observed due to residual heterozygosity in inbred lines
- Few cases with detected hap. > expected hap. (HARP errors)

The pipeline retrieves the expected number of haplotypes



Observed frequencies close from expected ones, no very distant observation



#### 6. Conclusions & perspectives

- Haplotypes are detected if there frequency is more than 5% (=1/15 individual in the pool) and frequencies are correctly estimated in pools
- Cheap short haplotypes sequencing approach, but improvement could be useful during probes design
- First results on actual data show that maize landraces are more diversified than maize inbred lines
- Maize landraces harbor haplotypes that are not in inbred lines

# Île-de-France – Versailles-Saclay

## Acknowlegment







## Bibliography

NEBnext® direct genotyping solution:

Emerman AB, Bowman SK, Barry A, Henig N, Patel KM, Gardner AF, Hendrickson CLJCPiMB: NEBNext Direct: A Novel, Rapid, Hybridization-Based Approach for the Capture and Library Conversion of Genomic Regions of Interest. 2017;119(1):7.30. 31-37.30. 24.

#### HARP:

Darren Kessner, Thomas L. Turner, John Novembre, Maximum Likelihood Estimation of Frequencies of Known Haplotypes from Pooled Sequence Data, Molecular Biology and Evolution, Volume 30, Issue 5, May 2013, Pages 1145- 1158, https://doi.org/10.1093/molbev/mst016



support-epgv@inrae.fr