

Supplementary Material

1 Supplementary Tables

Table S1. Kendall's correlation in the clustering of datasets.

	GBS-B73	GBS-Mock
SNP-array	0.99**	0.96**
GBS-B73	-	0.97**

SNP-array: Affymetrix® Axiom Maize Genotyping array; GBS-B73: genotyping-by-sequencing with SNP calling using B73 as reference genome; GBS-Mock: genotyping-by-sequencing with SNP calling using the mock reference built with all parental lines.

** Empirical significance level from permutations.

Fable S2 Variance components and	l genomic heritabilit	y of traits from SNP	datasets.
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		GY	РН	EH
	SNP-array	0.14	34.27	31.55
$\sigma^{2}a$	GBS-B73	0.15	30.36	31.34
	GBS-Mock	0.14	35.82	32.49
	SNP-array	0.07	5.83	1.95
σ^2 d	GBS-B73	0.09	7.88	3.08
	GBS-Mock	0.08	3.62	1.01
	SNP-array	0.17	17.92	6.70
σ^2 r	GBS-B73	0.16	18.02	7.03
	GBS-Mock	0.16	20.55	7.61
	SNP-array	0.56	0.69	0.83
H ²	GBS-B73	0.60	0.68	0.83
	GBS-Mock	0.57	0.66	0.81
h²	SNP-array	0.36	0.59	0.78
	GBS-B73	0.37	0.54	0.76
	GBS-Mock	0.37	0.60	0.79

SNP-array: Affymetrix® Axiom Maize Genotyping array; GBS-B73: genotyping-by-sequencing with SNP calling using B73 as reference genome; GBS-Mock: genotyping-by-sequencing with SNP calling using the mock reference built with all parental lines.

GY: grain yield; PH: plant height: EH: ear.

 $\hat{\sigma}_a^2$, $\hat{\sigma}_d^2$ and $\hat{\sigma}_r^2$: Additive, dominance and residual variances, respectively: H²: broad-sense heritability; h²: narrow-sense heritability.

Table S3 Coincidence in the division of the parental lines into heterotic groups among datasets.

		GBS-B73	GBS-Mock
GY	SNP-array	0.94**	1.00**
	GBS-B73	-	0.94**
РН	SNP-array	0.94**	0.93**
	GBS-B73	-	0.87**
EH	SNP-array	0.94**	0.93**
	GBS-B73	-	0.87**

SNP-array: Affymetrix® Axiom Maize Genotyping array; GBS-B73: genotyping-by-sequencing with SNP calling using B73 as reference genome; GBS-Mock: genotyping-by-sequencing with SNP calling using the mock reference built with

all parental lines.

** Significant at the 0.01 probability level by the t-test.

Fable S4 SCA's and GC	A's correlation of the	lines between the	SNP datasets.
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		GBS-B73	GBS-Mock
			SCA
GY	SNP-array	0.97**	0.96**
	GBS-B73	-	0.97**
РН	SNP-array	0.97**	0.96**
	GBS-B73	-	0.97**
EH	SNP-array	0.97**	0.96**
	GBS-B73	-	0.97**
			GCA
GY	SNP-array	0.99**	1.00**
	GBS-B73	-	0.99**
РН	SNP-array	1.00**	1.00**
	GBS-B73	-	1.00**
EH	SNP-array	1.00**	1.00**
	GBS-B73		1.00**

SNP-array: Affymetrix® Axiom Maize Genotyping array; GBS-B73: genotyping-by-sequencing with SNP calling using B73 as reference genome; GBS-Mock: genotyping-by-sequencing with SNP calling using the mock reference built with all parental lines.

** Significant at the 0.01 probability level by the t-test.

2 Supplementary Figures



Supplementary Figure 1. Optimal number of clusters formed among the 330 parental lines for all SNP datasets by Total Within Sum of Square (WSS) method. A SNP-array; **B** GBS-B73; **C** GBS-Mock.



Supplementary Figure 2. A Variance explained by the principal components (PCA) from SNParray, GBS-B73 e GBS-Mock SNP datasets for 330 tropical parental lines; **B** Cumulative explained variance estimated by principal components from SNP-array, GBS-B73 e GBS-Mock SNP datasets for 360 tropical parental lines.











Supplementary Figure 5. Heatmaps of the Rogers genetic distance matrices estimated from **A** SNP-array, **B** GBS-B73 and **C** GBS-Mock-All SNP datasets for 330 tropical parental lines. Lines and columns of each plot were clustered according to the Euclidian distance performed in the Roger genetic distance matrix from the SNP-array dataset.



Supplementary Figure 6. Heatmaps of the (A, B and C) additive genomic relationship (*Ga*), and (D, E and F) dominance genomic relationship (*Gd*) matrices estimated from (A and D) SNP-array, (B and E) GBS-B73, and (C and F) GBS-Mock SNP datasets for 751 tropical maize single crosses. Lines and columns of each plot were clustered according to the Euclidian distance performed in the genomic relationship matrices from the SNP-array dataset.