Genomic history of the origin and domestication of common bean unveils its closest sister species

Additional file 1.

Figure S1. Geographic origin of wild *P. vulgaris* accessions.
**Figure S2.** Phylogenetic trees produced using the ML method and SH-aLRT branch support implemented in PhyML, considering non-unique SNPs from chromosomes 1 to 4 according to the pseudoassembly of *P. vulgaris* cv. BAT93 against *P. vulgaris* G19833. Branch support: aLRT>0.95, circles; aLRT [0.75-0.95], squares.
Figure S3. Phylogenetic trees produced using the ML method and SH-aLRT branch support implemented in PhyML, considering non-unique SNPs from chromosomes 5 to 8 according to the pseudoassembly of *P. vulgaris* cv. BAT93 against *P. vulgaris* G19833. Branch support: aLRT>0.95, circles; aLRT [0.75-0.95], squares.
**Figure S4.** Phylogenetic trees produced using the ML method and SH-aLRT branch support implemented in PhyML, considering non-unique SNPs from chromosomes 9 to 11 according to the pseudoassembly of *P. vulgaris* cv. BAT93 against *P. vulgaris* G19833. Branch support: aLRT>0.95, circles; aLRT [0.75-0.95], squares.
**Figure S11.** Metabolic profile of *Phaseolus* accessions considering the top 100 most abundant metabolites.  
a. Hierarchical clustering/bootstrap tree of *Phaseolus* accessions; AU (Approximately Unbiased) and bootstrap probabilities are highlighted in red and green, respectively. Coloured boxed enclose the independent clades of *P. coccineus* (red), AH - *P. pseudovulgaris* accessions (green) and *P. vulgaris* (blue).  
b. Metabolic heatmap and clustering of the accessions.
Figure S12. Principal component analysis of *Phaseolus* accessions considered in the metabolomics fingerprinting. a. PCA using the 30 most informative metabolites obtained following a machine learning approach. b. PCA using the 100 most abundant metabolites in the screening.
Figure S13. Introgression signal between wild *P. vulgaris* subpopulations. $f_d$ values in 5kb non-overlapping windows of each chromosome are represented in Manhattan plots on left panels; the red threshold lines show the top 5% $f_d$ outliers in each comparison, and strong signals of introgression ($f_d+d_{XY}$) are highlighted in green. The directionality of genomic flux corresponds to: North towards West (a), South towards North (b), South towards West (c), North towards Center (d). Absolute genetic divergence ($d_{XY}$) calculated at the genome-wide scale (left), in $f_d$ 5% outliers (middle) and regions with introgression signal ($f_d+d_{XY}$, right) are shown in boxplots on the right panel of each comparison.
**Figure S14.** Introgression signal between wild and domesticated (DMA) *P. vulgaris* subpopulations. \( f_d \) values in 5kb non-overlapping windows of each chromosome are represented in Manhattan plots on left panels; the red threshold lines show the top 5\% \( f_d \) outliers in each comparison, and strong signals of introgression \((f_d + d_{XY})\) are highlighted in green. The directionality of genomic flux corresponds to: North towards DMA (a), South towards DMA (b), Center towards DMA (c), West towards DMA (d). Absolute genetic divergence \( (d_{XY}) \) calculated at the genome-wide scale (left), in \( f_d \) 5\% outliers (middle) and regions with introgression signal \((f_d + d_{XY}, \text{right})\) are shown in boxplots on the right panel of each comparison.
Figure S15. Introgression signal between domesticated (DMA) and wild *P. vulgaris* subpopulations. *f*<sub>d</sub> values in 5kb non-overlapping windows of each chromosome are represented in Manhattan plots on left panels; the red threshold lines show the top 5% *f*<sub>d</sub> outliers in each comparison, and strong signals of introgression (*f*<sub>d</sub> +<em>d<sub>XY</sub></em>) are highlighted in green. The directionality of genomic flux corresponds to: DMA towards Center (a), DMA towards West (b), DMA towards North (c), DMA towards South (d). Absolute genetic divergence (<em>d<sub>XY</sub></em>) calculated at the genome-wide scale (left), in *f*<sub>d</sub> 5% outliers (middle) and regions with introgression signal (*f*<sub>d</sub> +<em>d<sub>XY</sub></em>, right) are shown in boxplots on the right panel of each comparison.
Figure S16. Introgressed PCG. The number of genes encoded in each introgressed block is represented in scatter plots. Coloured lines: linear (red) and local (blue) regressions.
Figure S17. Introgression signal between *P. pseudovulgaris* and *P. vulgaris* subpopulations. $f_d$ values in 5kb non-overlapping windows of each chromosome are represented in Manhattan plots on left panels; the red threshold lines show the top 5% $f_d$ outliers in each comparison, and strong signals of introgression ($f_d+d_{XY}$) are highlighted in green. The directionality of genomic flux corresponds to: AH towards DMA (a), AH towards WMA (b), AH towards AND (c), AN towards AH (d). Absolute genetic divergence ($d_{XY}$) calculated at the genome-wide scale (left), in $f_d$ 5% outliers (middle) and regions with introgression signal ($f_d + d_{XY}$, right) are shown in boxplots on the right panel of each comparison.
Figure S18. Genome-wide nucleotide diversity (π) across *P. vulgaris* subpopulations.
Figure S19. Introgression signal between *Phaseolus* spp. and *P. vulgaris* / *P. pseudovulgaris* subpopulations. *f*<sub>d</sub> values in 5kb non-overlapping windows of each chromosome are represented in Manhattan plots on left panels; the red threshold lines show the top 5% *f*d outliers in each comparison, and strong signals of introgression (*f*d +*d*<sub>XY</sub>) are highlighted in green. The directionality of genomic flux corresponds to: *P. dumosus*/*P. costaricensis* towards AH (a), *P. dumosus*/*P. costaricensis* towards WMA (b), *P. coccineus* towards AH (c), *P. coccineus* towards WMA (d). Absolute genetic divergence (*d*<sub>XY</sub>) calculated at the genome-wide scale (left), in *f*<sub>d</sub> 5% outliers (middle) and regions with introgression signal (*f*<sub>d</sub> +*d*<sub>XY</sub>, right) are shown in boxplots on the right panel of each comparison.
Figure S20. Enriched categories among PCGs introgressed from wild MA subpopulations into domesticated genotypes.
Figure S21. Enriched categories among PCGs introgressed between wild MA subpopulations.