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the Nacional cocoa variety

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5 Guittard, United-States

Montpellier, France December 5-7





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Project Leader: C. Lanaud/R. Boulanger

Total Budget total: 427K€

- MUSE
- Valrhona











International Symposium or

Cocoa Research







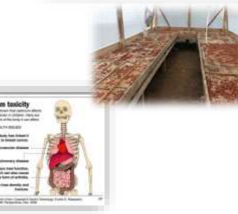
- History of past domestication of aromatic fine cocoa trees using paleogenomics
- Rescue and exploitation of new genetic resources related to fine cocoa
- > Study of genetic and biochemical determinants of fine cocoa aromas
- Genetic basis of cadmium accumulation
- > Study of dynamics of fine cocoa markets

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Location of cultivation sites in Ecuador



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The plant material were composed of a collection from Ecuador :

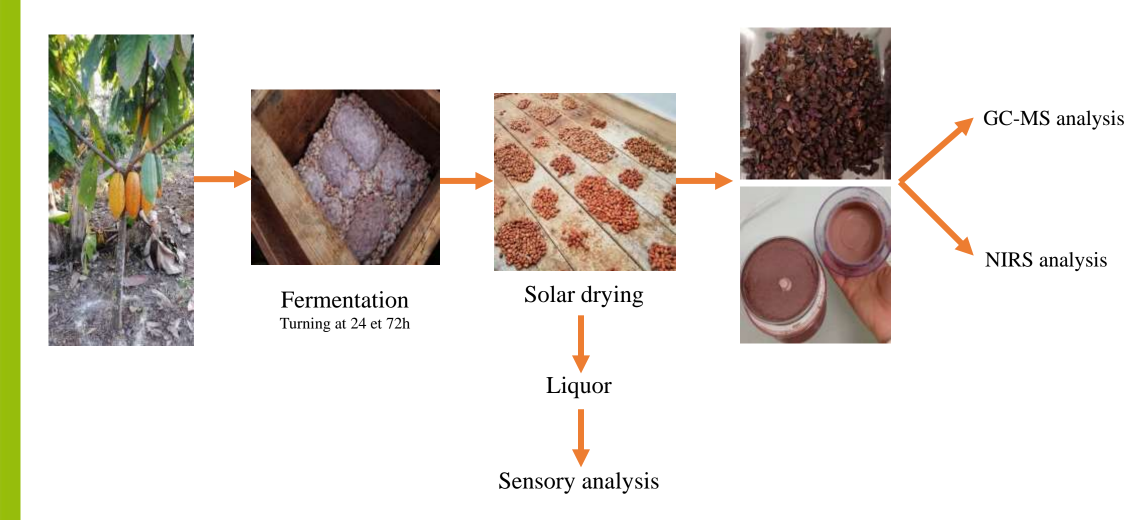
- Pichilingue experimental station of INIAP
- Coleccion de Cacao de Aroma Tenguel (CCAT) of Tenguel

This population (152) represents the Nacional variety currently grown in Ecuador

A second population of native cocoa trees (202) in the south of the Ecuadorian Amazon in the area of origin of Nacional



Phénotypes for Genome Wide Association Study (GWAS)





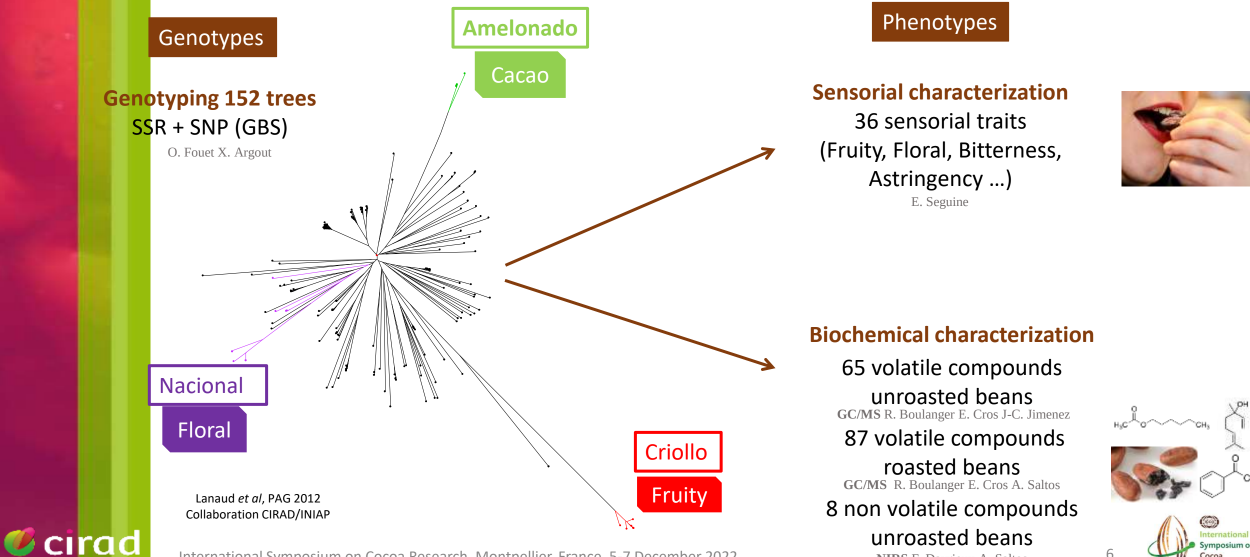
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Genome Wide Association Study (GWAS)

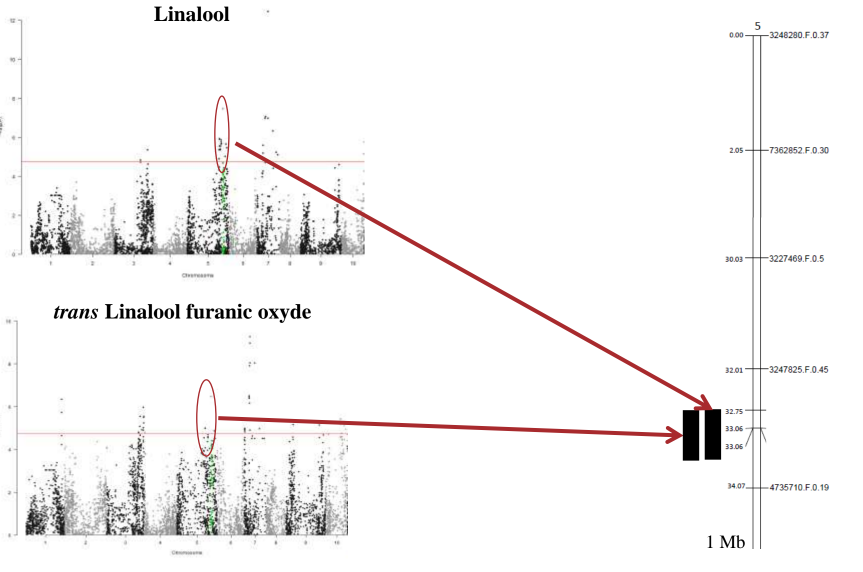


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Research

NIRS F. Davrieux A. Saltos

Association genetics (GWAS): example of results for aroma compounds

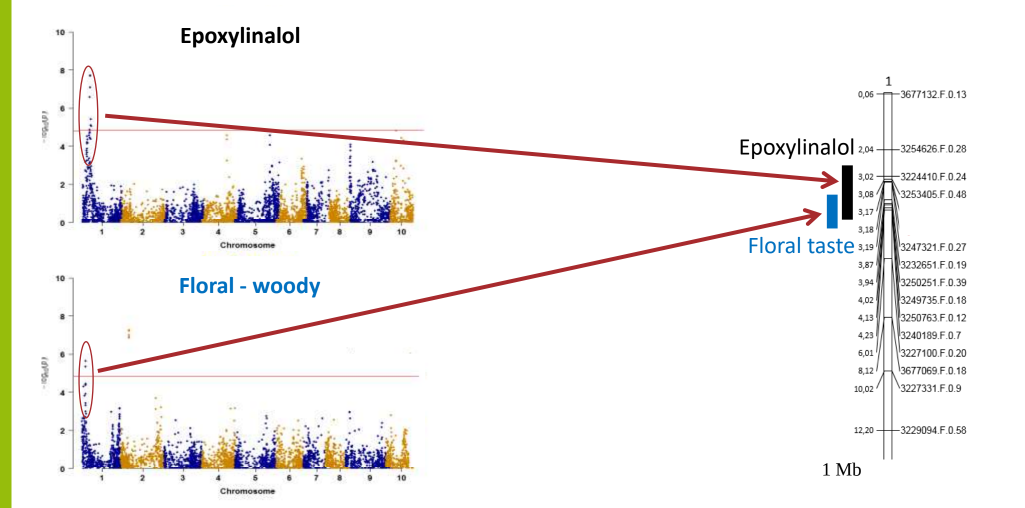


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Association genetics (GWAS): example of results for floral notes



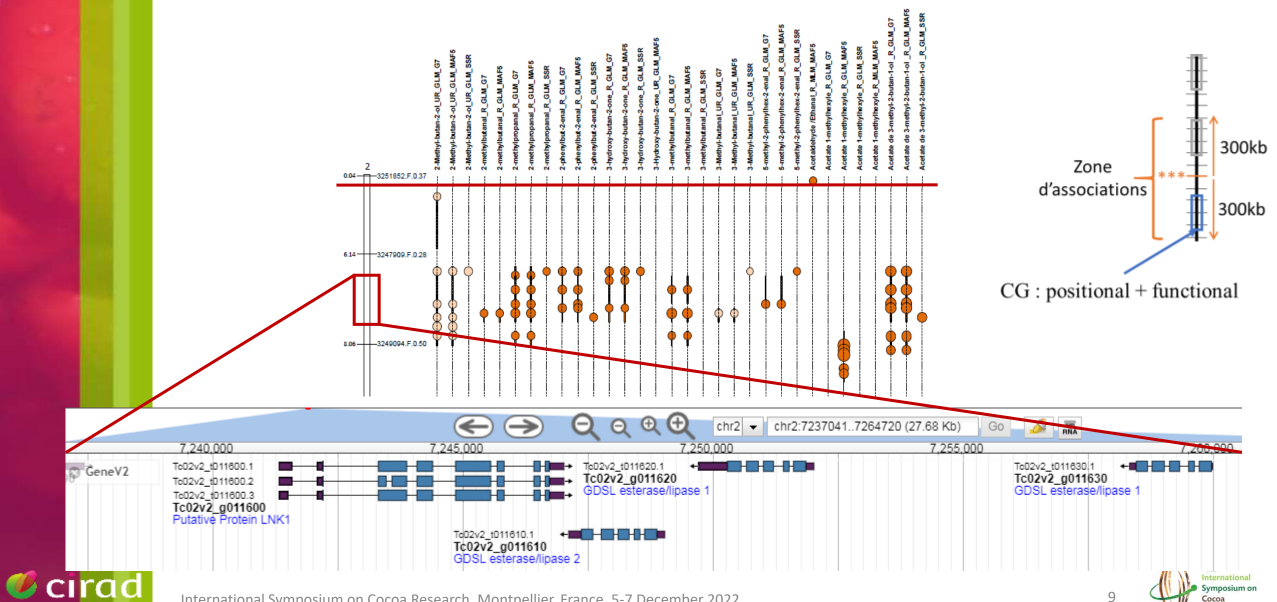


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Candidates genes research in association areas



Research

Determining the floral and fruity aromas of modern Nacional

Floral notes - 16 sensory descriptors, 27 VOCs

48 Association areas (9 sens.; 39 VOCs)

2 Main biosynthetic pathways:

- Monoterpene biosynthetic pathway (9 Candidate Genes)
- L-phenylalanine degradation pathway (19 Candidate Genes)

Fruity notes - 13 sensory notes and 35 VOCs

233 Association areas (39 sens.; 194 VOCs)

5 Biosynthetic pathways:

- Monoterpene biosynthetic pathway
- Pathway of L-phenylalanine
- Degradation of proteins, sugars and fatty acids

(514 Candidate Genes)

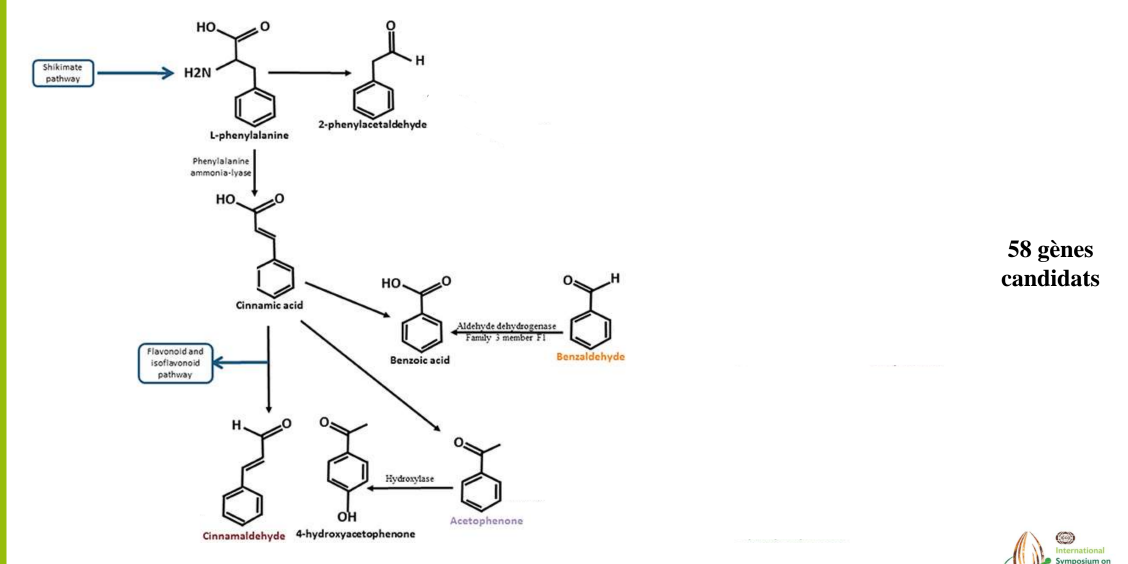




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Colonges et al, 2021 (Front. Plant Sci.) Colonges et al, 2022 (Plant Physiology and Biochemistry)

Example for candidate genes, floral and fruity notes L-phenylalanine pathway



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Determining the floral and fruity aromas of National ancestral

A second population of native cocoa trees resulting from surveys carried out in the south of the Ecuadorian Amazon in the area of origin of Nacional

Floral notes – 16 sensory descriptors, 27 COVs

393 Zones d'associations (38 sens.; 355 COVs)

• monoterpene biosynthetic pathway (114 Candidate Genes)

• L-phenylalanine degradation pathway (41 Candidate Genes)

Fruity notes - 13 sensory descriptors and 35 COVs

502 Zones d'associations (22 sens.; 480 COVs)

- monoterpene biosynthetic pathway
- pathways of L-phenylalanine

(1108 Candidate Genes)

) 55 () 56 ()

• degradation of proteins, sugars and fatty acids

54 Common genes between the two populations



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Colonges et al, 2022 (PlantsPeoplePlanet)



Determining the floral and fruity aromas of modern Nacional

- > 5 biosynthetic pathways highlighted:
- 2 for the floral notes
- 5 for the fruity notes 🛛 🍯 🧼 🚴 🙈

> Characterization of genome zones in relation to aroma types

 \succ Selection of areas of the genome favourable to the desired notes

Nest step

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Expression analyses at different stages





Thanks for your attention



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