

Microevolutionary changes of commercial Saccharomyces cerevisiae strains recovered from vineyard environments identified by comparative genome hybridization on array

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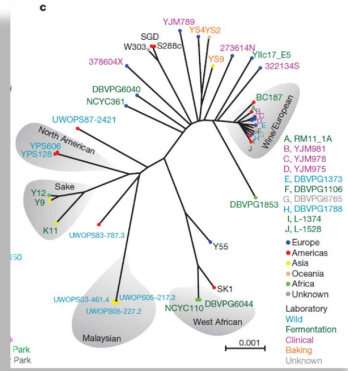






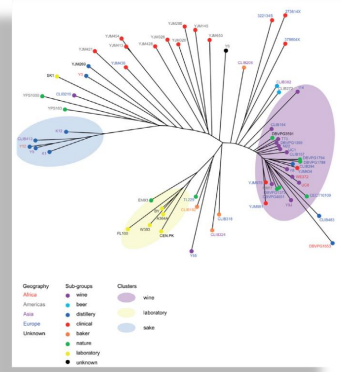
## **INTRODUCTION**

#### The population structure of Saccharomyces cerevisiae



#### Liti et al., Nature, 2009

#### 235,127 SNPs 14,051 nucleotide insertions or deletions



#### Schacherer et al., Nature, 2009

1.89 x 10<sup>6</sup> SNP (30,097 SNPs per strain) 3,985 deletios (200 bp length)

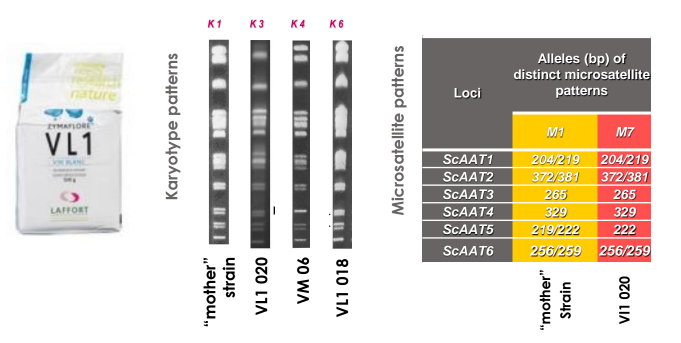
- low coverage genome sequencing
- high density arrays
- few well-defined, geographically isolated lineages
- many different mosaics of these lineages (wine, laboratory and saké strains)

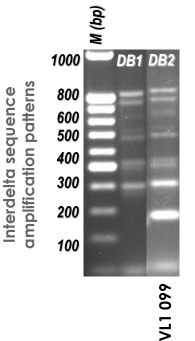
# INTRODUCTION

#### S. cerevisiae commercial winemaking strains



- ✓ Extensive use of commercial S. cerevisiae wine strains
- Such strains are disseminated from the winery and can be recovered from locations in close proximity (10-200m)
  Valero et al., 2005
- Re-isolation of 100 isolates of the commercial strain VL1 from vineyards close to the winery where this strain has been used during many years Schuller and Casal, 2007





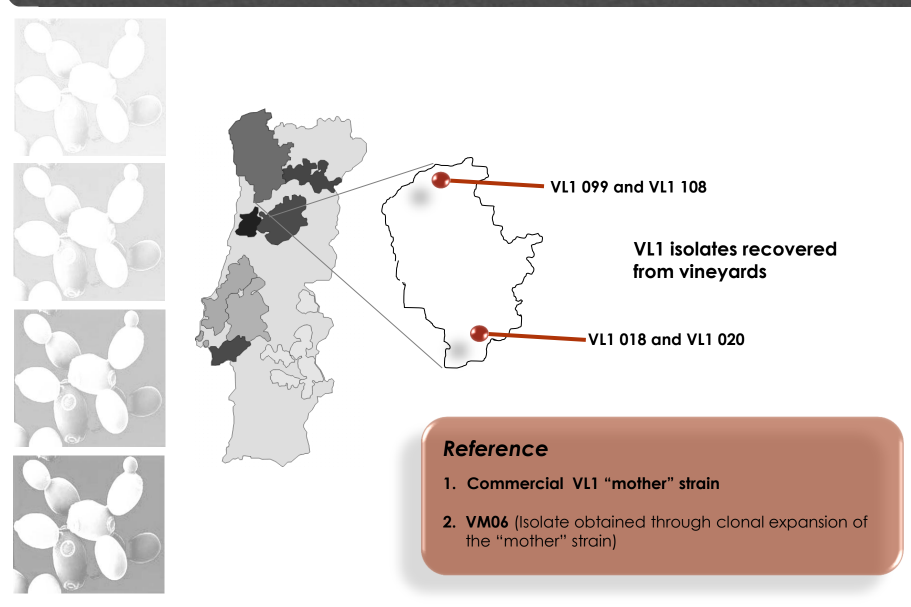


#### **Objectives**

- Evaluation of genome variations among isogenic isolates of the commercial strain Saccharomyces cerevisiae Zymaflore VL1 that were re-isolated from vineyards surrounding the wineries where this industrial strain was applied, using Comparative Genome Hybridization on array (aCGH);
- Conclude about adaptive mechanisms that occur during the strain's permanence in vineyard environments

## Materials and Methods

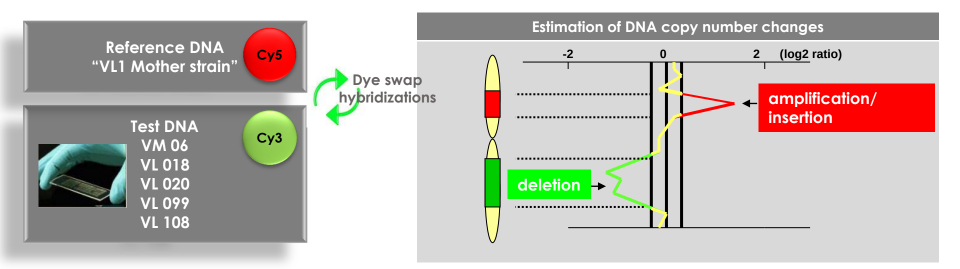
#### Saccharomyces cerevisiae isolates

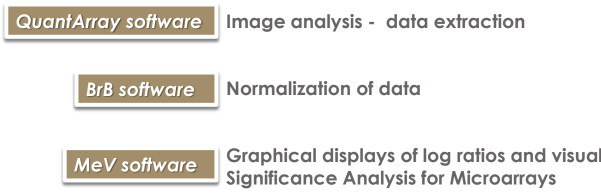


# Materials and Methods

#### Array Chromosome Genome Hybridization (aCGH)

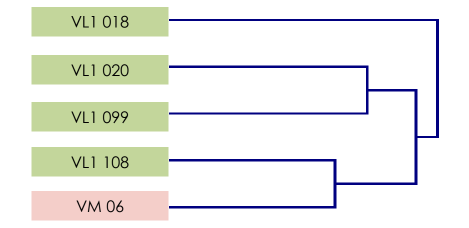
Spot





Graphical displays of log ratios and visual representation of data

#### Clustering of aCGH profiles

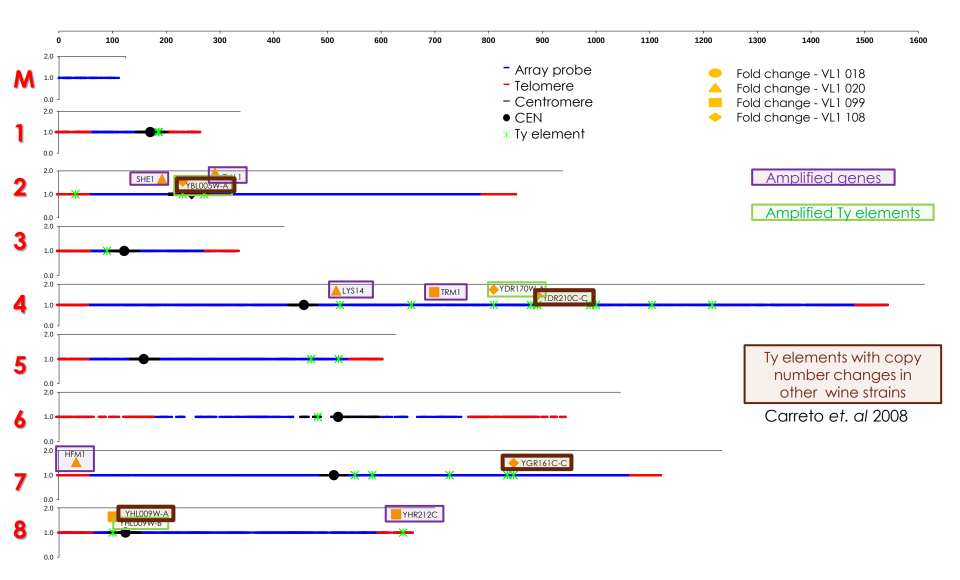


(Hierarchical clustering, Pearson correlation, average linkage)

No clear separation between VL1 isolates obtained from nature () and an isolate derived from the "mother" strain ()

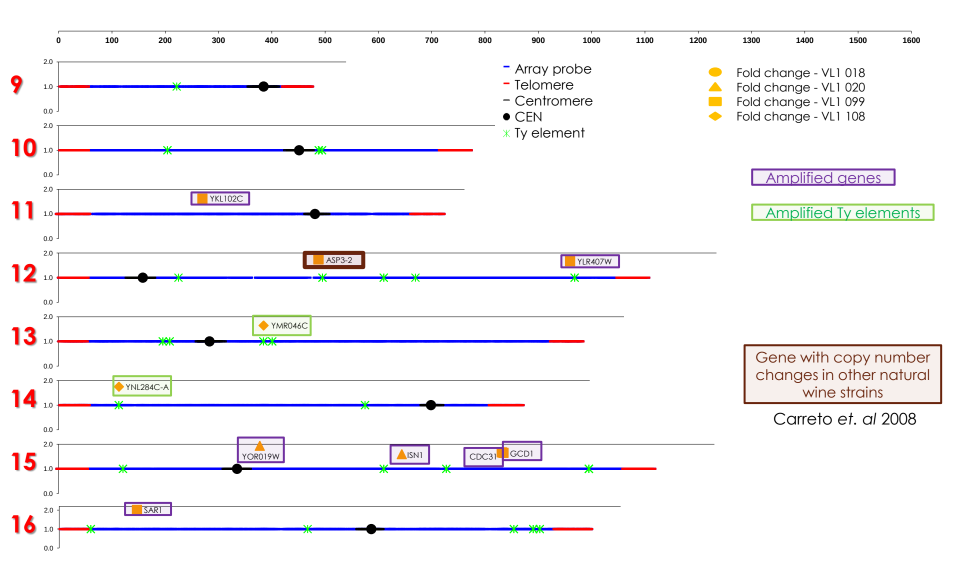
Results

#### Gene Copy number alterations – SAM analysis



Results

#### Gene Copy number alterations – SAM analysis



Results

#### Phenotypic characterization



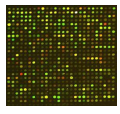
- Wine must + compound
- 30 °C
- 200 rpm
- quadruplicate

	Phenotypic tests																			
Strain	30°C	18°C	40°C	рН 2	pH 8	KCI 0.75M	NaCI 1.5M	CuSO4 5mM	SDS 0.01%	Etanol 6%	Etanol 10%	Etanol 14%	Iprodion (0.05mg/mL)	lprodion (0.1mg/mL)	Procymidon (0.05mg/mL)	Procymidon (0.1mg/mL)	KHSO3 (150 mg/l)	KHSO <sub>3</sub> (300 mg/l)	Vinho + glucose 0.5%	Vinho + glucose 1%
VL1 018	3	1	3	0	2	2	1	0	0	3	2	1	3	3	3	3	3	1	1	1
VL1 020	3	1	3	0	2	3	1	0	0	3	2	1	3	3	3	3	3	1	1	1
VL1 099	3	1	3	0	2	2	1	0	0	3	2	1	3	3	3	3	3	2	0	0
VL1 108	3	1	3	0	2	2	0	0	0	3	2	1	3	3	3	3	3	2	0	0
VM06	3	1	3	0	2	2	1	0	0	3	2	1	3	3	3	3	3	2	1	1
"Mother" strain	3	0	3	0	2	2	1	1	1	3	2	1	3	3	3	3	3	2	0	1

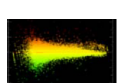
**0** – Abs<sub>640nm</sub> 0.1

 $1 - Abs_{640nm} 0.2-0.4$   $2 - Abs_{640nm} 0.5-1.2$   $3 - Abs_{640nm} \ge 1.3$ 

# SUMMARY AND CONCLUSIONS



- Isogenic isolates of the commercial wine yeast strain Zymaflore VL1 recovered from nature show genetic differences in comparison with the "mother" strain:
  - Gene amplifications
  - Ty element amplifications
  - Apparent stochastic distribution



• Generation of intra-strain phenotypic variability

The transition from nutrient-rich musts to nutritionally scarce natural environments is correlated with microevolutionary changes that may reflect adaptative responses



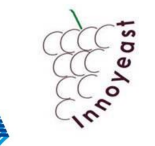
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- Nuno Fonseca
- Eugénia Vieira •

PROGRAMME







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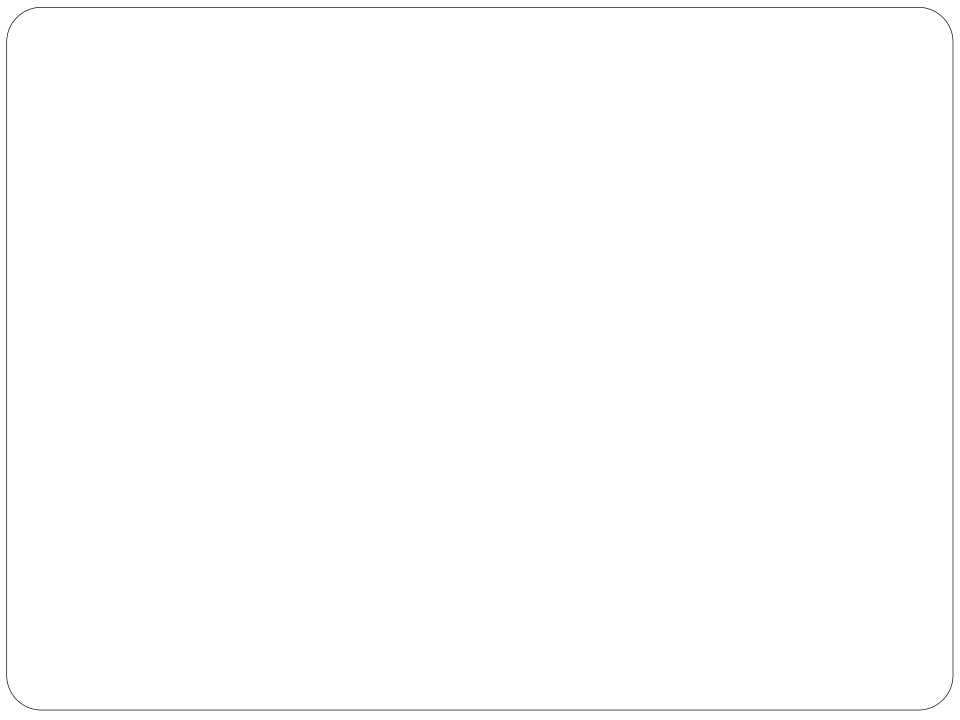
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GOVERNO DA REPÚBLICA PORTUGUE





## Significant altered genes

Strain	Systematic Name	Classical Name	Description	Chromosome			
020	YBL031W	SHE1	Mitotic spindle protein that interacts with components of the Dam1 (DASH) complex, its effector Sli15p, and microtubule-associated protein Bim1p; also localizes to nuclear microtubules and to the bud neck in a ring-shaped structure				
	YOR019W	NA	Protein of unknown function that may interact with ribosomes, based on co-purification experiments	15			
	YGL251C	HFM1/MER3	Meiosis specific DNA helicase involved in the conversion of double-stranded breaks to later recombination intermediates and in crossover control; catalyzes the unwinding of Holliday junctions; has ssDNA and dsDNA stimulated ATPase activity	7			
	YOR155C	ISN1	Inosine 5'-monophosphate (IMP)-specific 5'-nucleotidase, catalyzes the breakdown of IMP to inosine, does not show similarity to known 5'-nucleotidases from other organisms				
	YDR034C	LYS14	Transcriptional activator involved in regulation of genes of the lysine biosynthesis pathway; requires 2-aminoadipate semialdehyde as co-inducer	4			
	YBR020W	GAL1	Galactokinase, phosphorylates alpha-D-galactose to alpha-D-galactose-1-phosphate in the first step of galactose catabolism; expression regulated by Gal4p	2			
	YDR120C	TRM1	tRNA methyltransferase; two forms of the protein are made by alternative translation starts; localizes to both the nucleus and mitochondrion to produce the modified base N2,N2-dimethylguanosine in tRNAs in both compartments	4			
	YLR407W	NA	Putative protein of unknown function; null mutant displays elongated buds and a large fraction of budded cells have only one nucleus	12			
099	YOR260W	GCD1/TRA3	Gamma subunit of the translation initiation factor eIF2B, the guanine-nucleotide exchange factor for eIF2; activity subsequently regulated by phosphorylated eIF2; first identified as a negative regulator of GCN4 expression	15			
	YKL102C	NA	Dubious open reading frame unlikely to encode a functional protein; deletion confers sensitivity to citric acid; predicted protein would include a thiol-disulfide oxidoreductase active site	11			
	YOR257W	CDC31/DSK1	Calcium-binding component of the spindle pole body (SPB) half-bridge, required for SPB duplication in mitosis and meiosis II; homolog of mammalian centrin; binds multiubiquitinated proteins and is involved in proteasomal protein degradatio	15			
	YHR212C	NA	Dubious open reading frame unlikely to encode a functional protein, based on available experimental and comparative sequence data	8			
	YLR157C	ASP3-2	Cell-wall L-asparaginase II involved in asparagine catabolism; expression induced during nitrogen starvation; ORF contains a short non- coding RNA that enhances expression of full-length gene; reference strain \$288C has four copies of ASP3	12			
	YPL218W	SAR1	GTPase, GTP-binding protein of the ARF family, component of COPII coat of vesicles; required for transport vesicle formation during ER to Golgi protein transport	16			

# **RESULTS**

Significant altered genes

#### Ty elements:

Strain	Systematic Name	Chromosome	Fold Change			
018	YMR046C	13	1.6474975			
099	YHL009W-A	8	1.5785116			
	YHL009W-B	8	1.646452			
	YGR161C-C	7	1.4855264			
108	YBL005W-A	2	1.5453535			
	YDR210C-C	4	1.4554093			
	YDR170W-A	4	1.7406232			
	YNL284C-A	14	1.7531929			
	YMR046C	13	1.7273986			