Comparison of vineyard-associated Saccharomyces cerevisiae populations by microsatellite analysis

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Winery

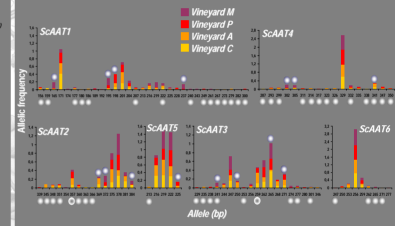


tions was estimated by Fst determination over al. supropulations was estimated by Fst determination over a loci by AMOVA analysis (computed by the Arlequin software [9]). A similarity matrix of allelic frequencies was computed by the program NTSYSpc 2.0 [10], based on the Euclidean

Winery	Year	Number of collected samples	Number of spontaneous fermentations	Number of Isolates	Number of Non-S. cerevisiae isolates	Number of genolypes	
	2001	12	3	90	0	11	1
Α	2002	6	6	180	0	34	0 1 0
	2003	12	6	180	0	41	• 1
С	2001	12	8	240	0	26	
	2002	3	1	30	0	1	
	2003	12	7	210	0	35	0 0 2
Р	2001	12	8	240	0	64	0 2 0
	2002	9	5	150	0	12	02
	2003	12	10	300	0	59	022
М	2001	12	11	330	129	51	0 0 0
	2002	12	12	360	359	1	
	2003	12	6	180	59	21	6

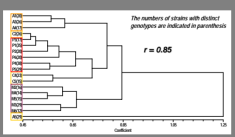
- patterns according their atletic distribution.

 The highest S. cerevisiae biodiversity was observed in winery M (323 isolates, 73 patterns) followed by es P (690 isolates, 135 patterns), A (450 isolates, 86 patterns) and C (480 isolates, 62 patter
- everal genotypes showed a wider temporal and geographical generalized pattern of sporadic prosence and reappearance across sampling sites, vineyards or years.
- Non-Saccharomyces strains belonging to the genus Kloeckera



- The six markers revealed a high degree of genetic variability, being ScAAT1 and ScAAT3 the most polymorphic markers with 31 and 19
- Besides the 41 ScAATi-ScAAT6 alleles previously described for 51 strains [3], 52 new alleles were identified in the present study.

 Some newly described alleles o occur with relative high frequency and may be used as indicative alleles for the Vinho Verde Win.
- The vast majority of alleles were evenly distributed among S. cerevisiae populations belonging to vineyards A, C and P and M, but differences are notorious for few alleles, which can be considered as vineyard(s) or Wine Region indicative



- Vineyard-specific populational substructure is shown by several clusters, comprising sampling sites of vineyards C, P, A and M. Populations within groups C and P are more closely related, while S. cerevisiae populations belonging to vineyard A are much more heterogeneous and also more distinct from C and P.
- The C2 population lies within the P-cluster, indicating that genetic differences do

Source o	AG	APWG	WP		P (r < 0)		
	2001		3.03	9.03	87.94	0.12	< 0.0001
	2002	A/P	6.38	13.28	80,33	0.20	0.0001
	2003		2.76	11.29	85.95	0.14	0.0001
	2001		-4.16	16.66	87.51	0.12	0.059
	2003	A/C	1.09	16.20	82.71	0.17	< 0.000
	2001	C/P	-1.21	8.31	92.89	0.07	0.0001
Among vineyards	2003	CIF	0.48	8.10	91.42	0.09	< 0.000
	2001	M/A	5.87	7.33	86.80	0.13	< 0.000
	2003	W/A	7.38	14.59	78.04	0.22	< 0.000
	2001		0.03	5.72	94.25	0.06	0.016
	2003	M/C	3.85	9.25	86.90	0.13	0.001
	2001	MP	2.75	5.44	91.80	0.08	< 0.000
	2003	IVE	3.48	4.54	91.98	0.08	< 0.000
	2001 / 2002	Α	-2.45	13.94	88.51	0.11	0.034
		P	0.79	9.94	89.27	0.11	0.0001
	2002 / 2003	Α	1.29	15.79	83.0	0.17	< 0.000
	2002 / 2003	P	1.68	7.73	90.59	0.09	0.052
Among years -		Α	-2.45	20.48	82.05	0.18	< 0.000
	2001 / 2002	С	-1.56	12.67	88.89	0.11	0.0001
	2001 / 2003	M	-0.25	5.63	94.61	0.05	0.07
		Р	0.37	6.30	93.33	0.07	0.0001

Pair wise association of populations from different vineyards showed that the closer vineyards A/P and A/C (30 - 50 km) are distant vineyards P/C (ca. 80 km).

F_{st} values for the pair wise association of S cerevisiae populations from France and Portugal (A/M, C/M and P/M) are similar to the values

observed among Portuguese populations.

Populations variation within a vineyard in consecutive years is similar to the variation observed between vineyards, being more variable in A (Fst = 0.11 - 0.18) compared to P

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Conclusions

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