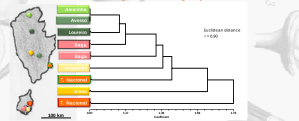
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Populational analysis of Saccharomyces cerevisiae strains from different appellations of origin and grape varieties by microsatellite analysis L. Pereira^{1*}, R. Franco-Duarte, P. Ramos^{2*}, F. Alemão^{2*}, P. Gomes^{3*}, S. Sousa^{3*} M. Santos^{3,4}, F. L. Duarte², M. Casal¹, D. Schuller¹ * contributed equally Centro de Biologia Molecular e Ambiental, Universidade do Minho, Braga Estação Vitivinícola Nacional, Quinta da Almoínha, Dois Portos BIOCANT, Centro de Inovação em Biotecnologia, Cantanhede 🙆 biocant Departamento de Biologia, Universidade de Aveiro, Aveiro erials and Methods Molecular identification Computer assisted data analysis Samples dura. Palmela and Alen Fermented beverages and foods have a longstanding cultural and historical tradition in most societies and the economical interest of fermonitation technologies is constantly increasing. Saccharomyces cerevisiae is one of the world's premier fermenting microorganisms. This yeast is predominantly found in association with human activities, particularly the production of alcoholic beverages. Whether the vineyard is the natural environment of S. cerevisae, is still a matter of controversy, but recent studies have shown that specialized strains for the production of alcoholic controversy. een Saccharomyces and eliminary discrimination bet Groups of strains with unique micro The sampling plan included 20 vineyards, and 9 grape varieties in several Portuguese appellations origin. In each region, appellation-specific recommended grape varieties were collectied in th vineyards comprising several grape varieties or single grape varieties, as shown below. Crape samp were obtained from six sampling points in each vineyard, and the yeast flord 20 randomly select isolate) form pointenously fereneuting grape juice (300 ml was analysed when the must weight w reduced by 70 g/l, corresponding to the consumption of about 2/3 of the sugar content. non-Saccharomyces yeast was based on the inability to grow in YNB medium containing L-lysine [4]. S. profiles were considered as population: corresponding to each vineyard. The pattern and degree of temporal and spatial divergence in the nuclear microsatellites among subpopulations was estimated by F_{gr} cerevisiae strains were further characterised by analysis of 6-10 S. cerevisiae specific microsatellite loci [5, 6]. beverages have been derived from natural populations unassociated with alcoholic beverage production, rather than the opposite [1]. The grape's yeast flora depends on a large variety of factors such as climatic conditions including temperature and rainfalls, the geographic localization of the vineyard, antifungal applications, the harvest technique, grape variety, the vineyard's age as well as determination over all loci by AMOVA analysis >Vinho Verde solated strains were analysed by mitochondrial DNA restriction patterns (miDNA RFLP) [7]. Strains with identical miDNA RFLP patterns were grouped and one representative strain was further characterised by >Bairrada >Estremadura >Palm-(Arlequin software (91), A similarity matrix of allelic frequencies was computed by the program NTSYSpc 2.0 [10], based on Euclidean the soil type. Several ecological surveys report a large diversity of Saccharomyces sp. strains among the enological fermentative flora. Some strains seem to be widely distributed in a given viticultural Alentejo (Évora) age linkage (UPGMA). the enological termentative flora. Some strains seem to be widely distributed in a given viticultural region, can be found in several consecutive years and are also predominant in the termenting flora, hypothesizing the occurrence of specific native strains that can be associated to a terroir [2-3]. The objective of the present study was to evaluate populational relationships among 5 cervisiae Brains isolated from some of the Portuguese most important grapevine varieties in different appendiations of origin, using polymorphic microsatellites. ce and av Isolated strains were analysed by PCR-amplification of interdelta sequences [8] as preliminary screen, followed by microsatellite analysis for in-depth characterization. The equivalent discriminatory power of mtDi interdelta analysis has been previously repor er of mtDNA RFLP and Interdelta Strains collected Allelic frequencies Distribution of the most frequent alleles among vineyards Distribution of the most frequent (🔚) microsatellite alleles of the Vinho Verde Region in consecutive years 8 S. cerevisiae strains involved in spontaneous fermentation 000 The ten markers revealed a high degree of genetic variability, being ScAAT1, C5 and YOR267 the most polymorphic markers with 43, 26 and 26 lleles, respectively. Most frequent alleles among S. cerevisiae strains from grape varieties of the Vinho Verde and Bairrada Regions 10 Alvarinho (30 strain oureiro (27 strains) Avesso (12 strains) Arinto (10 strains) Nacional (Minho) S. cerevisiae populations isolated from the Alvarinho, Avesso ıda and Loureiro grape varieties share the most frequent alelles, _____, that are equally distributed among populations isolated Jul The State ation by non-Sacchai 🐞 Fen omyces species Total: 501 S. cerevisiae strains from consecutive sampling years. Less frequent alleles that were found in S. cerevisiae Baga 18 strains Baga 14 strains From a total of 300 grape samples 4470 isolates were obtained; 192 samples (2850 ٠ Aragonès 13 stra T. Nacional populations from the three vineyards showed a differential isolates) were collected in the five wine regions during the grape harvest of 2006. distribution The fermentative yeast flora from the Vinho Verde and Bairrada Regions were composed by S. cerevisiae strains, whereas spontaneous fermentation of grapes Vineyard- pecific alleles that ocurred in one or two locations were also detected from the other regions were mainly conducted by non-Saccharomyces species. A total of 501 S. cerevisiae strains was obtained. Q. Less frequent alleles showed more variations in their distribution in consecutive years The number of S, cerevisiae strains is indicated for each spontaneou. fermentation, that corresponds to a sqare in each column related to the sampling The colored circles mark the three most frequent alleles for S. cerevisiae isolates from different grape varieties in the Vinho Verde and Bairrada regions year and grape variety. The composition of S. cerevisiae strains was very variable, randing from 1 – 22 among 30 randomly collected isolates. . Five alleles were shared by more S. cerevisiae populations, whereas alleles 20(ScAAT4), 20(C4), 21(C4), 22(C4) and 10(C5) can be considered as characteristic of the vineyards from where these isolates were obtained. ø The allelic distribution of S. cerevisae strains strains from Baga and Touriga Nacional grapes shows that the genetic constitution of the strains is rather vineyard-specific than grape variety- specific Similarity of S. cerevisiae populations Conclusions from distinct wine regions and grape varieties References [1]



- ee of spatial divergence in ten nuclear microsatellite markers was determined by clus analysis (UPGMA) based on a Euclidean distance dissimilarity matrix of allelic frequencies.
- Vineyard-specific populational structures show that genetic proximity is not related with the geographic distance. The most similar populations were obtained from vineyards in the Vinho Verde Region (Alvarinho and Avesso, located at a distance of ca. 150 km). Contrarily, yeast populations of vineyards in closer locations (e.g. Avesso and Arinto, at a distance of ca. 20 km) were more distinctive.
- Genetic proximity is not related with the grape variety. Populations isolated from Baga and Touriga Nacional grapes in different locations are not closely related.

yeast flora of grapes from the Vinho Verde and Bairrad osed by S. cerevisiae, contrarily to the prevailing non-is yeast flora in southern Portuguese wine regions. Thes y be assigned to climatic or other ecological factors.

g of 501 S. cerevisiae strains over four years reveales a l versity of S. cerevisiae strains in vineyard environments

tical analysis of the distribution of 192 alleles from 10 mid

- lifferences among S. cerevisiae populations derive ic "vineyard-, specific alleles and the accumulati menov differences across ten microsatellite loci a
 - vineyard contains differentiated S. cerevisiae populations. Th orted by the finding of identical or similar S. cerevisiae strains

- [6] [7] [8] [9] [9]

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