

# Genetic Evidence of Intra-cultivar Variability within Iberian Olive Cultivars

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**Abstract.** A collection of 130 olive samples, originating from diverse areas in Europe and corresponding to 67 different cultivars denominations, was genotyped at 14 microsatellite loci. In total, 135 alleles with a mean number of 9.6 alleles per locus were detected. All but 30 accessions showed unique genotypes. Several cases of synonymy listed in the FAO database of olive germplasm could not be confirmed, as different allelic profiles were obtained from putatively synonymous cultivars. The existence of homonyms or mislabeled samples in olive germplasm collections was evidenced by allele differences of up to 60% between samples of the same denomination. An allele-sharing phenogram of the analyzed genotypes revealed several cultivars with high levels of intra-varietal polymorphism, as well as cultivar families consisting of closely related cultivars with similar denominations. Our work shows that the current designations of olive cultivars fall short of describing the genetic variability among economically important plant material. A thorough investigation of the existing variability will prove of major importance for both management and economic production of olive trees.

Olive (*Olea europaea* L.) is one of the most ancient cultivated trees and its domestication probably occurred in the Near-East (Zohary and Hopf, 1994). Olive cultivation, which is widespread throughout the Mediterranean basin, is important both to the rural economy and to the environmental balance of the producing regions.

The great number of olive cultivars grown all over the world, estimated as about 1200 with over 3000 synonyms (Rugini and

Lavee, 1992) and the increasing demand for olive oil (Besnard et al., 2001a; [http://www.internationaloliveoil.org/downloads/consumation1\\_ang.PDF](http://www.internationaloliveoil.org/downloads/consumation1_ang.PDF)) combined, increase the importance of the correct identification of the different varieties.

The large amount of cultivars causes a huge problem in the management of germplasm collections and genomic traceability of olive oils, as there is a considerable uncertainty about the names of many cultivars and as olive cultivars are morphologically very similar (Cipriani et al., 2002). Moreover, it was shown that genetic variability could occur within the same cultivated populations (Gregoriou, 1996; Wiesman et al., 1998), which could justify heterogeneity in production and quality traits (Besnard et al., 2001a).

Clonal vegetative propagation has been practised in olives for several thousand years (García-Díaz et al., 2003), using cuttings that were planted directly, or more recently, grafted onto indigenous oleasters (Amane et al., 1999). The identification of olive tree cultivars has been traditionally carried out by morphological and agronomic traits, which are influenced by environmental or cultivation factors (Hernández et al., 2001), increasing the risk of accidental misidentification or mislabeling.

The recognition of the Protected Designation of Origin [Regulations (EEC) No. 2081/92 and (EEC) No. 2082/92] for olive oil increased

the need of a reliable verification of cultivars being grown, particularly to nurserymen and growers.

Microsatellite markers or simple-sequence repeats (SSRs) are a suitable tool for cultivar characterization and a number of loci have already been identified for olive (Carriero et al., 2002; Cipriani et al., 2002; Rallo et al., 2000; Sefc et al., 2000). The combination of data of several highly polymorphic microsatellite loci results in individual allelic profiles for different genotypes, while their codominant manner of inheritance allows parentage analysis (Sefc et al., 1997). The general reproducibility of SSR genotyping results between laboratories (Jones et al., 1998) supports the establishment of an olive worldwide genotyping database based on microsatellite data, as suggested for other species (Botta et al., 1995).

In the present study 130 individuals were typed with 14 SSR to evaluate the genetic variability in olive germplasm (between and within cultivars) mainly in the Iberian Peninsula.

## Material and Methods

Eighty two accessions corresponding to 36 Portuguese varieties of several geographical locations were chosen. Also several other Iberian and Mediterranean cultivars were selected for comparison, yielding a total of 130 samples (Table 1).

DNA was extracted from leaves following the protocol described by Fabri et al. (1995). The plants were genotyped for 14 SSR loci: ssrOeUA-DCA1, ssrOeUA-DCA3, ssrOeUA-DCA4, ssrOeUA-DCA5, ssrOeUA-DCA7, ssrOeUA-DCA8, ssrOeUA-DCA9, ssrOeUA-DCA11, ssrOeUA-DCA13, ssrOeUA-DCA14, ssrOeUA-DCA15, ssrOeUA-DCA16, ssrOeUA-DCA17, and ssrOeUA-DCA18 (Sefc et al., 2000).

Polymerase chain reactions (PCR) were carried out in a total volume of 20 µL containing 50 ng DNA, 1.5 mM MgCl<sub>2</sub>, 100 µM of each dNTP, 10 pmol of each primer and 0.5U *Taq* DNA polymerase in storage buffer B (Promega) in supplied reaction buffer. Reactions were performed in a UNO II Biometra thermocycler with the temperature regime described by Sefc et al. (2000). PCR products were analyzed in an automated sequencer (ABI Prism 310 genetic analyser, PE Applied Biosystems) and fragment lengths were determined with the help of internal size standards (Genescan 350 TAMRA size standard, PE Applied Biosystems).

Gene diversity ( $H_e$ ) (Nei, 1973), observed heterozygosity ( $H_o$ ) (Brookfield, 1996), probability of identity (PI) (Paetkau et al., 1995) and estimation of null allele frequency from the heterozygote deficiency ( $r$ ) (Brookfield, 1996) were calculated using IDENTITY (Wagner and Sefc, 1999). This program was also used to detect identical genotypes.

Deviations of observed heterozygosity values from Hardy-Weinberg expectations were analyzed using GENEPOP (Raymond and Rousset, 1995). Genetic distances between cultivars were calculated in MICROSAT (Minch, 1997) as 1-proportion of shared alleles and a phenogram was drawn using the UPGMA

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Table 1. Olive cultivars studied. The number in parentheses represents the number of individuals analyzed for each accession and the letters indicate the country of origin of the denominations (Fr = France; Gr = Greece; It = Italy; Mo = Morocco; Pt = Portugal; Sp = Spain; Tu = Tunisia; Tk = Turkey).

Alberquia (3) (Sp)	Lianolia (Gr)
Arbequina (Sp)	Maçanilha (Sp)
Ascolana (2) (It)	Maçanilha Carrasquenha de Almendralejo (3) (Pt)
Ayvalik (Tk)	Maçanilha de Elvas (Pt)
Azeiteira (5) (Pt)	Maçanilha de Jaen (Sp)
Bical Castelo Branco (Pt)	Maçanilha de Tavira (4) (Pt)
Bico de Corvo (Pt)	Madural (2) (Pt)
Blanqueta (3) (Sp)	Manzanilha (Sp)
Borrenta (2) (Pt)	Meski (Tu)
Carrasca de Trás-os-Montes (Pt)	Mora (Sp)
Carrasquenha (3) (Pt)	Negrinha (3) (Pt)
Carrasquenha de Elvas (Pt)	Negrinha (Pt)
Carrasquinha (Pt)	Negroa do Estanqueiro (Pt)
Cima di melfi (It)	Nocellara messinese (It)
Cipressino (It)	Nociara (It)
Cobrançosa (4) (Pt)	Pendolino (It)
Conserva das Barrancas (Pt)	Picholine (2) (Fr)
Conserva de Elvas (3) (Pt)	Picholine Marocaine (Mo)
Coratina (2) (It)	Pico Limon (Sp)
Cordovil (3) (Pt)	Picual (4) (Sp)
Cordovil de Castelo Branco (4) (Pt)	Planalto (Pt)
Cordovil de Elvas (4) (Pt)	Quinta do Portado (Pt)
Cordovil de Moura (Pt)	Redondil (4) (Pt)
Cordovil de Serpa (4) (Pt)	Sant'Agostino (It)
Cornezuelo (3) (Sp)	Santulhana (Sp)
Cornicabra (Sp)	Tentilheira (Pt)
Frantoio (3) (It)	Termite di Bitetto (It)
Galega (8) (Pt)	Uovo di Piccione (Tu)
Galega Grada de Serpa (Pt)	Verde Verdelho (3) (Pt)
Galego Évora (Pt)	Verdeal Alentejana (3) (Pt)
Golosinha (Pt)	Verdeal de Elvas (Pt)
Gordal (Sp)	Verdeal de Serpa (3) (Pt)
Hojiblanca (2) (Sp)	Verdeal Transmontana (Pt)
Leccino (2) (It)	

algorithm in PHILIP (Felsenstein, 1989) and the program TREEVIEW (Page, 1996).

## Results

*Characterization of 14 SSR markers in olive cultivars from different origins.* Among the 130 accessions comprising 67 different denominations, 100 different SSR profiles were obtained. A total of 135 alleles were detected, ranging from 4 to 16 alleles (Table 2), with a mean of 9.6 alleles per locus.

Probabilities of identity varied between loci due to uneven allele frequency distributions. The highest information content was provided by locus *ssrOeUA-DCA16*, while high PI val-

ues were estimated for loci *ssrOeUA-DCA14* and *ssrOeUA-DCA5*. In *ssrOeUA-DCA14* the two most common alleles account for 90% of the frequencies, while the remaining 10% were shared by three alleles. In *ssrOeUA-DCA5* the most common allele has a frequency of 75%, while the remaining five alleles have frequencies between 1% and 12%. The combined probability for identical genotypes across all loci is low with  $9.4 \times 10^{-13}$ .

Both expected heterozygosity and PI values indicate *ssrOeUA-DCA16* as the most informative and *ssrOeUA-DCA5* as the least informative marker.

At 10 of the 14 loci, the observed heterozygosity was higher than the expected values. In

contrast, it was lower than the expected values at loci *ssrOeUA-DCA4*, *ssrOeUA-DCA7*, *ssrOeUA-DCA11* and *ssrOeUA-DCA13*. In consequence the probability of occurrence of null alleles is positive at these loci (Table 2).

*Intra-varietal polymorphism, cultivar families, synonymous and homonymous cultivars.* Synonymous and homonymous cultivar designations indicated in the database of Olive Germplasm: Cultivars and World-Wide Collections (available from FAO's Seed and Plant Genetic Resources Service at [www.fao.org/ag/agp/agps/seed/olive.htm](http://www.fao.org/ag/agp/agps/seed/olive.htm)) were compared with the data obtained by the characterization of the 130 accessions at 14 SSR loci (data available at [www.angra.uac.pt/biotecagri/publicacoes/htm](http://www.angra.uac.pt/biotecagri/publicacoes/htm)). The phenogram in Fig. 1 illustrates the genetic similarities among the analyzed cultivars.

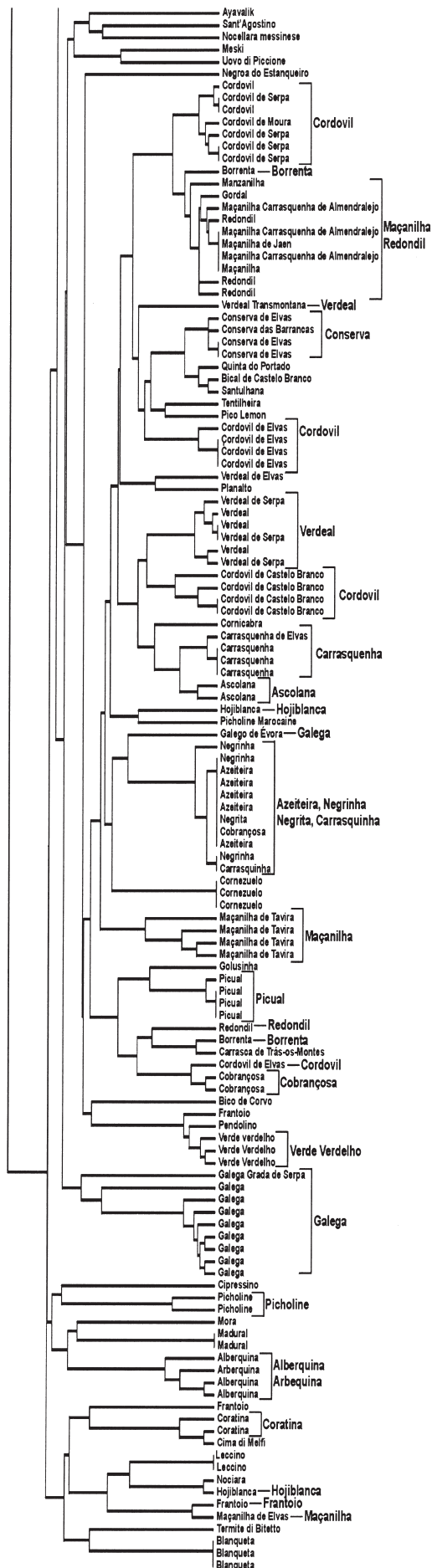
'Alberquia', also know as 'Arbequina' according to the FAO's olive germplasm database, is one of the most widely used Spanish cultivars for oil production. Genotypes of three samples with the denomination 'Alberquia' were analyzed and compared with one sample of 'Arbequina'. Although the genotypes of the four accessions differed from one another at one to four alleles, the 'Alberquia' and 'Arbequina' samples grouped together to the exclusion of all other cultivars in the phenogram, forming a cultivar family.

Similarly, intra-cultivar variability with differences in up to four alleles was observed in 'Picual', 'Conserva de Elvas', 'Verde Verdelho', 'Ascolana', 'Coratina', 'Cobrançosa' and 'Picholine'.

An interesting case is 'Galega', the most widely used cultivar for oil production in Portugal, accounting for about 80% of the orchards. Eight individuals under this denomination were analyzed and showed unique SSR profiles. Seven of the eight genotyped accessions differed from each other in only 1 to 2 alleles, while the other sample differed from them in up to 10 alleles. Nevertheless these 8 accessions and another 'Galega' accession, 'Galega Grada de Serpa', which differed from the other 'Galegas' in up to 15 alleles out of 28, clustered in the same group. 'Galego de Évora', also differs from the 'Galega' group in up to 15 alleles, but did not cluster within the 'Galegas', although

Table 2. Genetic parameters of the studied loci in 130 samples. The table shows the number and size range of alleles detected, the probability of finding identical genotypes for each locus and for all loci combined, expected and observed heterozygosity, and the probability of null alleles at each locus.

Locus	Alleles (no.)	Size range	Probability of identity (PI)	Expected heterozygosity ( $H_e$ )	Observed heterozygosity ( $H_o$ )	Probability of null alleles
<i>ssrOeUA-DCA1</i>	4	204-228	0.227	0.581	0.816	-0.148
<i>ssrOeUA-DCA3</i>	11	226-246	0.073	0.804	0.981	-0.098
<i>ssrOeUA-DCA4</i>	14	128-188	0.148	0.688	0.52	0.099
<i>ssrOeUA-DCA5</i>	6	195-207	0.413	0.417	0.5	-0.06
<i>ssrOeUA-DCA7</i>	8	127-169	0.194	0.615	0.275	0.211
<i>ssrOeUA-DCA8</i>	8	122-150	0.132	0.705	0.971	-0.156
<i>ssrOeUA-DCA9</i>	13	161-206	0.054	0.824	0.971	-0.08
<i>ssrOeUA-DCA11</i>	11	125-167	0.149	0.697	0.49	0.122
<i>ssrOeUA-DCA13</i>	8	117-153	0.132	0.726	0.677	0.028
<i>ssrOeUA-DCA14</i>	5	169-187	0.301	0.502	0.634	-0.087
<i>ssrOeUA-DCA15</i>	4	242-264	0.181	0.66	0.767	-0.064
<i>ssrOeUA-DCA16</i>	16	121-191	0.043	0.849	0.98	-0.071
<i>ssrOeUA-DCA17</i>	15	101-183	0.146	0.67	0.738	-0.041
<i>ssrOeUA-DCA18</i>	12	163-203	0.1	0.77	0.95	-0.102
PI for all loci:	135		$9.4 \times 10^{-13}$	0.679	0.734	-0.014



its name would suggest an affiliation within the group.

A different pattern was found in the ‘Maçanilha’ cultivar. ‘Maçanilha de Jaen’, originally from Spain, is a cultivar grown in Alentejo (Portugal). It is used with ‘Maçanilha Carrasquenha de Almedralejo’ for the production of canned fruits. The allelic profiles of ‘Maçanilha de Jaen’ and of two individuals of ‘Maçanilha Carrasquenha de Almedralejo’ as well as that of ‘Maçanilha’, a cultivar widely distributed in Spain, were identical. Another sample of ‘Maçanilha Carrasquenha de Almedralejo’ revealed an SSR profile very similar to the other samples under the same name, differing in only one allele at locus *ssrOeUA-DCA17*. ‘Manzanilha’, the Spanish counterpart of ‘Maçanilha’, had a different allelic profile from its Portuguese synonym at locus *ssrOeUA-DCA13*. ‘Maçanilha de Tavira’, grown in Algarve (Southern Portugal) and ‘Maçanilha de Elvas’, grown in Alentejo (Central/Portugal) differ substantially from each other and the other ‘Maçanilhas’.

For a survey of the genetic diversity within the ‘Verdeal’ group, three ‘Verdeal Alentejana’, three ‘Verdeal de Serpa’, one ‘Verdeal de Elvas’ and one ‘Verdeal Transmontana’ were genotyped. ‘Verdeal Transmontana’ is grown in the North of Portugal, while the other ‘Verdeal’ samples are grown in Alentejo in Central/Southern Portugal. In this case the SSR profiles obtained were different for each sample within and between cultivars, except for identical genotypes for one ‘Verdeal Alentejana’ and one ‘Verdeal de Serpa’ accession and thus the ‘Verdeal’ accessions were placed in three clades in the phenogram.

Another interesting case is ‘Cordovil’, which has different denominations according to its place of origin. In this study one ‘Cordovil de Moura’, three ‘Cordovil’, four ‘Cordovil de Castelo Branco’, four ‘Cordovil de Elvas’ and four ‘Cordovil de Serpa’ were genotyped. In FAO’s olive germplasm database ‘Cordovil de Serpa’ is described as a synonym of ‘Bico de Corvo’, from which one accession was analyzed. Within ‘Cordovil’ denominations, several different allelic profiles were obtained. ‘Cordovil’ samples displayed three genotypes, with one accession identical to one of the ‘Cordovil de Serpa’ genotypes, another accession closely related to the latter two (two alleles difference) and the third accession differing in up to 17 alleles from the ‘Cordovil’ samples. ‘Cordovil de

Fig. 1. UPGMA phenogram of genotyped olive samples. Cultivar families, synonymous and homonymous cultivars are highlighted.

Elvas’ showed two genotypes differing in two alleles, ‘Cordovil de Serpa’, had three genotypes differing at only one allele and ‘Cordovil de Castelo Branco’ also displayed three genotypes, with differences of up to six alleles. In consequence of the genetic variability of the ‘Cordovil’ cultivar family, the accessions were placed in five different clades in the phenogram. The synonymy of ‘Bico de Corvo’ with ‘Cordovil de Serpa’ could not be confirmed.

Several other accessions with identical names were revealed to be homonymous in that they showed little genetic similarity and thus were placed in different positions in the phenogram. For example, individuals of ‘Borrenta’, ‘Frantoio’ and ‘Hojiblanca’ differed in 14, 26, and 23 alleles respectively. Homonymy was also found in accessions of ‘Redondil’, as three samples, differing in only one allele each, clustered together, while a fourth sample differing in 14 alleles was grouped separately.

According to the FAO olive germplasm database ‘Azeiteira’, an important cultivar for oil production in Portugal, and ‘Blanqueta’, one of the most interesting varieties for the production of oil in Spain, are synonyms. However, genotyping of five ‘Azeiteira’ and three ‘Blanqueta’ individuals refuted synonymy of these cultivars. Samples within each denomination displayed identical microsatellite profiles, but only 12 out of 28 alleles were shared between the denominations. Surprisingly, one accession each of ‘Negrã’, ‘Negrinha’ and ‘Cobrançosa’ showed the same SSR profile as ‘Azeiteira’, suggesting synonymy or extensive mislabeling of our cultivars. The other two ‘Negrinha’ samples are closely related to the Azeiteira group but differ from one another in one allele.

‘Carrasquenha’ and ‘Carrasquinha’, two Portuguese cultivars used for the production of canned fruits are also listed as the same cultivar in the olive germplasm database. Three individuals of ‘Carrasquenha’ showed identical genetic profiles, which differed in 10 alleles from the genotype obtained from ‘Carrasquinha’. ‘Carrasquinha’, for its part, displayed the same genotype as one sample of ‘Negrinha’, an important cultivar for the production of canned fruits in Portugal, and was placed next to the ‘Azeiteira’ group in the phenogram.

‘Carrasquenha’ differs in only one allele from ‘Carrasquenha de Elvas’, indicating a high genetic similarity between the two cultivars. The latter variety originates from Elvas, one of the most important olive growing regions in Portugal.

‘Madural’, ‘Cornezuelo’ and ‘Cornicabra’ are regarded as synonyms by the FAO olive germplasm database. However, this hypothesis was not confirmed in our analysis, as the 3 cultivars displayed SSR profiles differing in up to 17 alleles.

Similarly, ‘Santulhana’ and ‘Gordal’ are considered two denominations for the same cultivar in FAO’s olive germplasm database, while this study revealed a considerable genetic distance between these two cultivars, as 9 out of 28 alleles were different between the two samples.



## Discussion

When considering long term breeding programs in fruit trees, studies dealing with the structure of genetic diversity may give some insights about the selection of cultivated forms, and this may lead to a better management of their diversity (Besnard et al., 2001b). Genetic diversity has been assessed in many species with microsatellites, whose major applications has been the identification and the distinction of cultivars. The level of polymorphism and the associated information content is a crucial criterion for the choice of a particular set of loci. With up to 16 alleles, the number of alleles obtained at the loci used in this study is higher than those reported by other authors (Carriero et al., 2002; Cipriani et al., 2002; Rallo et al., 2000) where three to nine alleles per locus were reported. However, marker polymorphism also varies according to the number and origin of the plants analysed. Co-ancestry of cultivars may reduce the genetic variability of a cultivar group dramatically.

Our study describes intra-varietal polymorphism in a number of olive cultivars, with differences in up to 15% of the analyzed alleles. Cases of cultivars with even higher genetic distances between accessions are ascribed to recent or past mislabeling (homonymy). Intra-cultivar polymorphism in olives has already been reported by Cipriani et al. (2002), where differences in  $\leq 2$  loci out of 30 were considered as somatic mutations occurring in the process of vegetative propagation. With the high level of intra-varietal polymorphism described in the present study, olives pose a stark contrast to another long cultivated, woody crop, the grapevine. While many grapevine cultivars have been propagated vegetatively throughout many centuries, only a few cases of intra-varietal polymorphism (Silvestroni et al., 1997; Vignani et al., 2002) and clonal polymorphism (Boursiquot, 2003) have been reported. While the genotypic stability of grapevine cultivars allowed the reconstruction of ancient crosses (Bowers and Meredith, 1997; Bowers et al., 1999; Lopes et al., 1999; Sefc et al., 1998), the determination of the exact pedigrees of olive cultivars and cultivar families will be complicated by the variability within cultivar lines, as it appears that ancient genotypes cannot necessarily be expected to have been maintained in the olive cultivars without mutation. Differences in the stringency of the selection and propagation procedure between grapevines and olives may account for the observed difference in genotypic stability of those two crops. Alternatively, mutation rates may differ between the two crops and thus produce unequal variability levels under similar agricultural circumstances and in comparable time scales.

Germplasm variability in olives may have been shaped by the century-long practice of local selection and propagation of spontaneous seedlings, and only recently, efforts have been undertaken to describe the existing gene

pool. For the identification of closely related genotypes, synonyms and homonyms, genetic marker analysis will be of indispensable value. The cultivars investigated in this study are only a small part of the world's cultivar ensemble. The genetic characterization of a larger number of cultivars will not only verify and identify more homonymous and synonymous cultivars, but will also reveal more information about the genetic relationship between them.

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