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Marines Marli Gniech Karasawa¹, Benedetta Chiancone², Valeria Gianguzzi¹, Ahmed Mohamed Abdelgalel²,
 Roberto Botta³, Chiara Sartor³, Maria Antonietta Germanà¹

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22 Gametic embryogenesis through isolated microspore culture in *Corylus avellana* L.

- ¹Dipartimento di Scienze Agrarie e Forestali. Università degli Studi di Palermo. Viale Delle Scienze, 11. 90128
 Palermo –Italy
- 26 ²Dipartimento di Scienze degli Alimenti. Università degli Studi di Parma, Parma, Italy
- ³ Dipartimento di Colture Arboree, Università degli Studi di Torino. L.go P. Braccini 2, 10095, Grugliasco (TO)
- 28 mariaantonietta.germana@unipa.it
- 29

30 Abstract

31 Haploid technology is a valuable tool to obtain homozygosity particularly in woody plants, in order to support 32 plant breeding. Hazelnut, the fourth worldwide nut tree, is a monoecious, anemophilous species and it is 33 characterized by a sporophytic incompatibility system (Germain and Dimoulas, 1978), not allowing to obtain 34 homozygous plants through conventional methods, involving several self-pollination. In this study, gametic 35 embryogenesis was applied to hazelnut and, particularly, isolated microspore culture was performed on four 36 different cultivars. Two culture media were tested and four temperature stress treatments were applied to the 37 isolated microspores, cultivated at the vacuolated stage of development. For the first time in our knowledge, 38 early embryos were recovered in hazelnut via isolated microspore culture in all cultivars. The assessment of the 39 zygosity condition of embryos carried out using SSR markers showed that analysed embryos were homozygous 40 indicating that they developed from haploid microspores. The response to the culture treatments, was, however, 41 genotype-dependent, as previously reported in experiments regarding male gamete embryogenesis in other fruit 42 crops. 43

- 4344 Keywords: Haploid, hazelnut, gametic embryogenesis, temperature stress.
- 45
- 46

47 Introduction

- 48 Hazelnut (*Corylus avellana* L.), belonging to the Betulaceae family of the order Fagales and native to European
- 49 regions, Turkey and the Caucasus mountains (Kafkas and Dogan, 2009), is the sixth worldwide tree nut, with a
- 50 production of more about 858,000 tons (FAOSTAT, 2013), behind cashew, Persian walnut, chestnut, almond
- and pistachio. All cultivated forms of hazelnut are diploid with a monoploid number of chromosomes n=x=11. It
- is a monoecious, dichogamous, self-incompatible and wind pollinated plant (Rovira et al. 1993). The
 incompatibility system is sporophytically determined and controlled by a single locus with multiple alleles
- 54 (Martins 2010). Incompatibility is a main factor limiting genetic improvement, and it prevents to reach
- (containe 2010). Intering whether is a state of the intering generation in protonic to
- bomozygosity by conventional methods, involving several self-pollination cycles.
- 56 Gametic embryogenesis is recognized as an important tool for plant breeding, making possible to develop, in
- 57 only one step, completely homozygous lines, from heterozygous parents (Bueno et al. 2006; Seguì-Simaro and
- 58 Nuez 2008; Islam and Tuteja 2012; Datta 2005; Solis et al. 2008; Germanà 2011a). This opportunity is
- 59 particularly useful for breeding in woody species, in which it is not possible to obtain homozygosity through
- 60 conventional methods (Germanà 2011b), because of the long juvenile phase, the high degree of heterozygosity,
- 61 the large size and, often, of the self-incompatibility (Germana` 2006; 2009).
- 62 Haploid (Hs) and doubled haploid (DHs) plants, are very useful to fix parental lines to produce F1 hybrids, for
- 63 rapid introgression of new characteristics through backcrossing, to develop population molecular maps, to fix
- 64 characteristics through transformation and mutagenesis (Datta 2005; Szareijo and Foster 2006; Soriano et al.
- 65 2013) and to do reverse breeding (Dirks et al. 2009; Ferrie and Möllers 2011). In addition, haploids and double-
- haploids are important for genomic studies and genome sequencing (Aleza et al. 2009; Dirks et al. 2009; Ferrie
- 67 and Möllers 2011; Foster et al. 2007; Germanà et al. 2013; Talón and Gmitter 2008), for physical mapping
- 68 (Leeuwen et al. 2003; Kunzel et al. 2000; Wang et al. 2001), genetic mapping (Houssain et al. 2007; Chu et al.
- 69 2008; Zhang et al. 2008), and for the integration between physical and genetic mapping (Kunzel et al. 2000;
- 70 Wang et al. 2001).
- Gametic embryogenesis represents a unique system where, thanks to the plant totipotency, a single immature
 gamete switches from the gametophytic pathway toward the sporophyte formation (Prem et al 2012). Gametic
- raise embryogenesis, that can be achieved from female (gynogenesis) or male (microspore embryogenesis) gametes,
- 74 rarely occurs in nature (Silva 2012), and it is usually induced subjecting immature gametes to some kinds of
- 75 stress. Particularly, temperature stresses, hot and cold, are the most effective and commonly used ways to
- 76 promote microspore embryogenesis in several species (Ferrie and Caswell 2010; Germanà 2011 a; Shariatpanahi
- et al. 2006; Silva 2012). Specifically, the involvement of heat shock proteins (HSPs), when temperature
- 78 treatments are applied in the microspore embryogenesis, has been described, but the molecular mechanism of
- 79 the induction and the relationship between different stresses remain unclear. However, it is supposed that the
- 80 increased levels of HSPs may be associated with the embryogenic potential acquisition (Karami and Saidi 2010;
- 81 Silva 2012).
- 82 In many crops, anther culture is often the method of choice for H and DH production because of its simplicity
- 83 (Sopory and Munshi 1996). Nevertheless, the technique of isolated microspore culture, performed by removing
- 84 somatic anther tissue and requiring better equipment and more skills compared to the first technique, provides a
- 85 better system for investigating cellular, physiological, biochemical and molecular processes involved in pollen
- 86 embryogenesis (Nitsch 1977; Reinert and Bajaj 1977; Germanà, 2011b).

- 87 In this study, the technique of isolated microspore culture was applied to five hazelnut genotypes, testing two
- 88 basal media and four temperature stresses, with the aim to obtain micropore-derived embryo regeneration.
- 89
- 90 Materials and Methods
- 91

92 Plant material and pollen developmental stage

93 Immature catkins (Fig. 1a) were harvested in February from trees of the following cultivars of Corvlus aveilana 94 L.: 'Tonda Gentile Romana' (2 clones: TR and GR), 'Imperatrice Eugenia' (IE), 'Minnulara' (M) and 95 'Carrello' (C).. Catkins with anthers (Fig. 1a) mostly containing microspores at the vacuolated/uninucleated 96 stage (Fig. 1b) were selected, assuming that, like in many other fruit crops (Germana 2011 a; 2011b), this stage 97 of development is the most responsive for gametic embryogenesis also for hazelnut. For this reason, a 98 preliminary investigation was carried out to select the appropriate catkin size staining anthers with few drops of 99 4',6-diamidino-2-phenylindole (DAPI) solution (1 mg/mL) and observing them under fluorescence microscope 100 (Zeiss, Axiophot, Germany). Catkins with anthers containing microspores at vacuolated/uninucleate stage were 101 selected and, as common pre-treatment, were placed in darkness at 4 °C for two weeks.

102

103 Catkins sterilization and microspore isolation and culture

104 After two weeks of 4 °C pretreatment, 5 catkins for each genotype were surface sterilized by immersion for 3 105 min in 70% (v/v) ethyl alcohol, followed by immersion for 20 min in 25% (v/v) sodium hypochlorite solution 106 (about 1.5% active chlorine in water) with few drops of Tween[®] 20, and then rinsed three times with sterile 107 distilled water. After that, the material was maintained in immersion in gentamicin antibiotic solution (0.2%) for 108 30 minutes. The antibiotic solution was removed and the microspore isolation was performed following the 109 procedure reported by Kumlehn et al., (2006) with limited modifications. Isolated microspores were cultured at 110 the concentration of 100,000 microspores per mL and a final volume of 1.0 mL per each Petri dish (3001-type 111 Petri dishes, BD Biosciences) was placed.

- 112 Petri dishes, sealed with Parafilm[®], were placed at 26±1 °C, in the dark for the first 30 days and later under cool
- 113 white fluorescent lamp (Philips TLM 30W/84, France) with a photosynthetic photon flux density of 35 μ mol m⁻¹
- 114 s^{-1} and a photoperiod of 16 light hours.
- 115

116 Experimental design

- 117 In order to evaluate the influence of medium composition on the embryogenic induction of the five genotypes,
- isolated microspores were cultured on two different media: P (Germanà et al. 1996) and N6 (Germanà and
- 119 Chiancone 2003). Moreover, because it is well known that usually stress enhances microspore response, isolated
- 120 microspores cultured in both media, were subjected to the following thermal stress, just after isolation:
- 121 1) 35 °C for 30 minutes (H),
- 122 2) 40 °C for 60 minutes (H+),
- 123 3) -20 °C for 30 minutes (F),
- 124 4) -20 °C for 60 minutes (F+).
- 125 Height Petri dishes (repetitions) were prepared per each combination (genotype, medium, thermal stress).
- 126

127 Data collection and statistical analysis

128 Isolated microspores in culture were weekly observed by a stereo-microscope (Leica MZ 125) and by an 129 inverted microscope (Zeiss West Germany). In addition, examinations were performed by a fluorescence 130 microscope (Zeiss, Axiophot, Germany), after staining with 4',6-diamidino-2-phenylindole (DAPI). Because 131 changes in the microspore development distribution were observed for a long time, it was decided to report the 132 final monitoring of the different features, registered after 20 months of culture. At this time, per each genotype and per each combination medium/thermal stress, 450 microspores/structures (three repetitions with 150 133 elements each) were counted, in order to individuate the different structural features: uninucleated microspores, 134 135 rarely binucleated with two different nucleus sizes beginning their normal gametophytic pathway, binucleated 136 with two equal-size vegetative-type nuclei just starting their sporophytic pathway, trinucleated, and also 137 multinucleated microspores. Moreover, the number of embryos and calli produced per each Petri dish were 138 registered. These values were used to calculate means and to perform the statistical analysis. Differences among 139 genotypes were tested by one-way analysis of variance (ANOVA) with Conover-Inman's test ($p \le 0.05$). To individuate the influence of the factors "Medium" and "Thermal treatment", means were analysed by two-way 140 141 ANOVA, and mean separation was performed using Tukey's multiple comparison tests at ($p \le 0.05$).

142

143 Detection of homozygosity

Simple Sequence Repeat (SSR) markers were used to assess the genetic condition of embryos obtained frompollen microspore culture and to determine their origin (gametic or somatic).

146 DNA extraction

Embryos were collected from the culture medium by an insulin syringe, frozen in liquid nitrogen and ground
using steel beads in a Tissuelyser (QIAGEN®, Germany). DNA extraction for genotyping was performed as
described in Doyle and Doyle (1987) with minor modifications.

150

151 PCR amplification and SSR allele sizing

- 152 Five fully characterized SSR loci were used for assessing the origin of the embryos: CaT-B107, CaT-B503,
- 153 CaT-B504, CaT-B505, CaT-B507 (Boccacci et al., 2005). PCR was performed, as described by Boccacci et al.
- 154 (2008), in two steps in a total volume of 10 μl containing 20 ng of DNA, 0.25 U of KAPA Taq DNA
- 155 polymerase (KAPABIOSYSTEMS, Wilmington Massachusetts, USA) 1 μl of 10X PCR buffer, 200 μM
- 156 nucleotide mix and 0.5 mM of each primer. PCR conditions were as follows: an initial denaturation step at 95°
- 157 C for 3 min followed by 32 cycles of denaturation (30 s at 95°C), annealing (45 s at 52° C), and extension (90 s
- 158 at 72° C). The final elongation step was at 72°C for 30 min. Two µl of the amplification product were used as
- template for a second PCR, carried out for 28 cycles with the same conditions of the first one.
- 160 PCR products were analyzed on a 3130 Genetic Analyzer (Applied Biosystems, Foster City, California, USA).
- 161 Data were processed using GeneMapper Software (ver. 4.0; Applied Biosystems) and alleles were defined by
- their size in base pairs, by comparison with the size standard (GeneScan-500 LIZ, Applied Biosystems).
- 163 Data obtained were compared with the hazelnut database of DISAFA.
- 164

165 Results and Discussion

- 167 The microspore culture monitoring carried out by fluorescence microscope after 20 months of culture and after
- 168 DAPI staining, showed uninucleated (Fig. 1b), binucleated presenting nuclei with asymmetrical division (Fig.
- 169 2a), binucleated revealing nuclei with symmetrical division (Fig. 2b), trinucleated (Fig. 2c), and multinucleated
- 170 stage (Fig. 2d); the performances of the different genotypes are reported in table 1-6. Actually, the presence of
- binucleated microspores with asymmetrical division of the nucleus, indicating of the gametophytic pathway,
- 172 was very sporadic: never more that 3% and in three genotypes, (C, IE and M), less than 1% of the total (Table
- 173 1).
- 174 The statistical analysis carried out to study the *in vitro* isolated microspore development in the five hazelnut 175 genotypes tested, showed different responses among the genotype (Table 1). For this reason, the influence of the two parameters "Medium" and "Thermal treatment" on hazelnut microspore development was carried out 176 177 separately per each genotype. Particularly, regarding the "Medium" influence on the multinucleated microspores 178 production, the P medium was the most valuable in the genotypes C, M and TR (Tables 2, 5 and 6). In the other 179 genotypes, differences between the two media were not statistically significant. Two way ANOVA evidenced 180 also that in the cultivars C and M, the formation of multinucleated microspores was mainly influenced by the 181 factor "Medium", while different results were recorded for the genotypes GR, IE and TR, where a significant interaction was observed between the two factors. IE, GR and C, were the genotypes with the highest 182
- 183 percentages of multinucleated microspores.
- For GR, the factor "Treatment" was the main source of variability in the multinucleated microspore induction (Table 3). Particularly, statistically significant differences were observed between the treatment F+ and the treatment H, while the other treatments induced intermediate responses. The analysis of interaction evidenced a statistically significant difference in the thermal treatment H, where N6 induced better response than P (32.0% *vs* 19.8%) (data not shown).
- Diverse response was observed in the cultivar IE concerning the same parameter, in which none of the factors
 had a predominant influence (Table 4), but differences were observed in the response to thermal treatments
 between the two media: P better than N6 in H+ (+34.7% vs 24.7%) and in F (40.0 % vs 23.3%), whereas N6
- 192 better than P in F+ (30.0% vs 22.7%) (data not shown).
- 193 On the contrary, for TR, both factors ("Medium" and "Treatment") had a significant influence on the 194 multinucleated microspore induction. Analyzing separately the factors, P medium induced a statistically
- significant higher percentage of multinucleated microspores than N6; while, regarding the thermal treatments,
- two groups were individuated: Co and H induced statistically higher response than H+, F and F+ (Table 6).
- 197 Furthermore, regarding the interaction, main differences were observed in the Control where the P medium
- induced a statistically higher response than N6 (37.3% vs 22.0%)
- 199 After three-four months of culture, stereo- microscope observations revealed new structures: light brown calli
- 200 (Fig. 3) that increased in quantity and volume and, also, the formation of globular embryos was detected (Fig. 4
- a). Moreover, different shapes of embryos were observed: elongated (Fig. 4 b) and also often embryos with a
- suspensor-like structure (Fig. 4c). This kind of structure was observed also in Citrus in the embryos obtained
- through isolated microspore culture, but not by anther culture.
- 204 Previously, in fruit tree crops, Hofer et al (1999) reported embryo regeneration, through isolated microspore
- 205 culture, in apple and, recently (Chiancone et al., 2015), the production of gamete-derived embryos obtained by
- in the same way in Citrus was reported.

- 207 Because of the data distribution, it was not possible to apply the two-way ANOVA to the average number of
- 208 calli and embryos registered per Petri dish, but for the novelty and relevance of these results, it is interesting to
- 209 describe the performance of the different cultivars. The statistical analysis of the average calli number per Petri
- dish confirmed the morphogenic potential of TR, that, together with GR, produced the highest number of calli
- (Fig. 5a). Actually, the highest average callus number per Petri dish was recorded in the genotype 'Tonda
 Gentile Romana' TR, in the treatment H+ in combination with the N6 medium (13.7) (Fig. 5f). As reported for
- Gentile Romana' TR, in the treatment H+ in combination with the N6 medium (13.7) (Fig. 5f). As reported forthe microspore development, also for calli production, the response to the different media and to the thermal
- treatments was genotype-dependent. The cold thermal treatments positively influenced callus production:
- specifically, F+ treatment induced the highest average number of calli per Petri dish in C. IE (in combination
- with P medium), GR (in combination with both P and N6 media); while, the hot thermal treatments, and
 particularly, the H+, seemed to produce a higher callus number in M (in combination with N6) and in TR (in
 combination with P and N6) (data not shown).
- 219 The embryo achieving does not appear to be influenced by the treatments in the same way than calli. Really,
- 220 with the exception of M, in which the combination of H+ thermal treatment with N6 medium, was proven to be
- suitable both for callus and embryo achieving. For the other cultivars, the results were different: the highest
- average embryo number were recorded in the control, in combination with P medium for C and TR, with N6
- 223 medium for GR, and in the H+ treatment in combination with N6 medium for IE (Fig. 5b-f).
- Generally, regarding the effect of the medium composition, P medium induced a little higher percentage of multinucleated microspores and also a higher number of embryos. Not always at the genotype with the very high number of multinucleated microspores corresponded the higher microspore-derived embryo number (for example, IE).
- Regarding the stress, many research highlighted its importance in inducing microspore embryogenesis.
 However, in this study the higher number of embryos was obtained in both P and N6 control (without additional stresses other that 4 °C before isolation), except for IE.
- 231

232 Detection of homozygosity

Genetic analysis showed that the hazelnut embryos had a single allele at each of the five SSR loci analysed (Figs. 6 and 7) The allele found at each locus matched either one of the 2 alleles of the parent genotype (in the case of the figures 'Minnulara' was the cultivar of origin of the microspores). Being the embryos very small, ploidy level could not be determined; for this reason we can postulate that the embryos are very likely haploids but there is no evidence they may not be double haploids. For sure the embryos analysed originated from a haploid microspore.

- 240 The production of homozygous plants in fruit trees of high commercial value like hazelnut is highly desired to
- help increase the breeding efficiency and reduce the time to obtain new varieties. But, this is not possible in
- woody plants, like hazelnut, characterized by long juvenile cycle, high degree of heterozygosity, large size and
- 243 with incompatible systems. The only way to obtain isogenic lines is through haploid technology. Actually,
- 244 microspore embryogenesis is a promising tool in plant breeding since it permits to obtain full homozygous
- plants in just one step using heterozygous parents (Germanà 2011 a, b). This technology can be of great interest

to increase the research and plant breeding of *Corylus avellana*, a species not very often subject of improvement

247 (Dunstan and Thorpe 1986).

248

249 Conclusions

250

251 In our knowledge, this is the first time that haploid technology was applied to hazelnut and that homozygous 252 early embryos were regenerated through isolated microspore culture. The response to the culture media and 253 treatments, was, however, highly genotype-dependent, as previously reported in experiments regarding tissue 254 culture in hazelnut as well as microspore embryogenesis in other fruit crops. Actually, each genotype presented 255 different behaviour, even if, all of them produced calli and, more important, microspores-derived early embryos. 256 Really, even if it is not possible to individuate the absolute best medium or thermal treatment stress, this result 257 represents a major step forward in the knowledge on gametic embryogenesis in fruit crops. Further studies are, 258 however, necessary to better understand the process of gametic embryogenesis in this species and to optimize 259 the rate of microspore-derived embryos and, in addition, to obtain their germination into plantlets.

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432 Tables

433 Table 1 Influence of genotype on isolated microspore development, after twenty months of culture.

Genotype	Uninucleated microspores	Binucleated microspores (SND)	Binucleated microspores (ASD)	Trinucleated microspores	Multinucleated microspores
	%	%	%	%	%
'Carrello'	27.7 a	28.4 ab	1.9 a	12.9 b	29.1 ab
Tonda G. Romana GR	30.0 a	25.3 bc	2.9 a	11.8 b	30.0 ab
'Imperatrice Eugenia'	23.0 b	23.5 c	0.1 b	22.0 a	31.4 a
'Minnulara'	23.4 b	31.1 a	0.6 b	21.6 a	23.3 b
Tonda G. Romana TR	27.6 a	27.1 ab	0.6 b	19.7 a	25.0 b

434 Per each column, values followed by different letters are statistically different.

435 One-way ANOVA, Conover-Inman's test, $p \le 0.05$.

436 SND: Symmetrical nucleus division

437 AND: Asymmetrical nucleus division

438

439 Table 2 Influence of medium composition and thermal treatment on 'Carrello' isolated microspore development,

440 after twenty months of culture.

Factors	Uninucleated microspores	Binucleated microspores (SND)	Binucleated microspores (ASD)	Trinucleated microspores	Multinucleated microspores
	%	%	%	%	%
P P	27.6 a	25.5 b	2.6 a	12.6 a	31.7 a
Medium (M) N6	27.7 a	31.2 a	1.2 a	13.3 a	26.5 b
M p value	0.962	0.003	0.073	0.593	0.014
Со	23.7 b	29.7 a	1.7 a	13.7 a	31.3 a
Н	34.0 a	26.2 a	2.1 a	11.0 a	26.7 a
Treatment (T)H+	29.4 ab	26.4 a	1.4 a	11.4 a	31.4 a
F	26.7 ab	29.3 a	2.0 a	14.0 a	28.0 a
F+	24.7 ab	30.3 a	2.3 a	14.7 a	28.0 a
T p value	0.025	0.398	0.928	0.400	0.412
M X T p value	0.425	0.330	0.658	0.175	0.145

441 Per each factor and per each column, values followed by different letters are statistically different.

442 Two-way ANOVA, Tukey's test, $p \le 0.05$.

443 SND: Symmetrical nucleus division

444 AND: Asymmetrical nucleus division

445 P medium: Germanà at al 1996; N6 medium: Germanà and Chiancone, 2003.

446 Co: no thermal treatment; H: 35 °C for 30 minutes; H+: 40 °C for 60 minutes; F: -20 °C for 30 minutes; F+: -20

447 °C for 60 minutes.

448

Factors		Uninucleated microspores	Binucleated microspores (SND)	Binucleated microspores (ASD)	Trinucleated microspores	Multinucleated microspores
		%	%	%	%	%
Malling (M)	Р	32.7 a	22.8 b	3.4 a	11.4 a	29.6 a
Medium (M)	N6	27.2 b	27.7 a	2.4 a	12.2 a	30.5 a
M p value		0.010	0.005	0.320	0.598	0.501
	Co	29.0 a	26.3 a	2.0 a	14.3 a	28.3 ab
	Н	34.8 a	23.8 a	4.6 a	10.9 a	25.9 b
Treatment (T)	H+	29.0 a	25.0 a	2.0 a	11.7 a	32.3 ab
	F	28.2 a	25.9 a	3.0 a	11.9 a	30.9 ab
	F+	28.7 a	25.3 a	3.0 a	10.3 a	32.7 a
T p value		0.050	0.852	0.500	0.523	0.023
M X T p value		0.010	0.272	0.157	0.694	0.000

Table 3 Influence of medium composition and thermal treatment on 'Tonda Gentile Romana' GR isolatedmicrospore development, after twenty months of culture.

452 Per each factor and per each column, values followed by different letters are statistically different.

453 Two-way ANOVA, Tukey's test, $p \le 0.05$.

454 SND: Symmetrical nucleus division

455 AND: Asymmetrical nucleus division

456 P medium: Germanà at al 1996; N6 medium: Germanà and Chiancone, 2003.

457 Co: no thermal treatment; H: 35 °C for 30 minutes; H+: 40 °C for 60 minutes; F: -20 °C for 30 minutes; F+: -20 °C for 60 minutes.

459

460 Table 4 Influence of medium composition and thermal treatment on 'Imperatrice Eugenia' isolated microspore461 development, after twenty months of culture.

Factors		Uninucleated microspores	Binucleated microspores (SND)	Binucleated microspores (ASD)	Trinucleated microspores	Multinucleated microspores
		%	%	%	%	%
	Р	23.2 a	22.9 a	0.0 a	20.3 a	34.5 a
Medium (M)	N6	22.4 a	22.5 a	0.0 a	22.1 a	33.9 a
M p value			0.514	0.173	0.486	0.294
	Co	19.3 a	23.7 a	0.0 a	20.7 a	36.3 a
	Н	23.0 a	21.3 a	0.3 a	20.3 a	37.3 a
Treatment (T)	H+	26.7 a	27.3 a	0.0 a	16.3 a	29.7 a
	F	20.0 a	21.7 a	0.0 a	26.7 a	31.7 a
	F+	26.0 a	23.7 a	0.3 a	23.7 a	26.3 a
T p value			0.337	0.570	0.096	0.209
M X T p value			0.823	0.570	0.544	0.024

462 Per each factor and per each column, values followed by different letters are statistically different.

463 Two-way ANOVA, Tukey's test, $p \le 0.05$.

464 SND: Symmetrical nucleus division

465 AND: Asymmetrical nucleus division

466 P medium: Germanà at al 1996; N6 medium: Germanà and Chiancone, 2003.

467 Co: no thermal treatment; H: 35 °C for 30 minutes; H+: 40 °C for 60 minutes; F: -20 °C for 30 minutes; F+: -20 °C for 60 minutes.

469 C 101 00 minutes.

471 Table 5 Influence of medium composition and thermal treatment on 'Minnulara' isolated microspore472 development, after twenty months of culture.

Factors		Uninucleated microspores	Binucleated microspores (SND)	Binucleated microspores (ASD)	Trinucleated microspores	Multinucleated microspores
		%	%	%	%	%
	Р	25.6 a	24.5 b	0.0 a	19.7 a	30.1 a
Medium (M)	N6	29.6 a	28.2 a	1.0 a	19.4 a	21.8 b
M p value		0.067	0.000	0.415	0.245	0.000
	Co	24.3 a	23.0 a	0.3 a	22.7 a	29.7 a
	Н	28.3 a	25.7 a	0.7 a	16.0 b	29.3 a
Treatment (T)	H +	31.0 a	28.0 a	0.3 a	18.3 ab	22.3 a
	F	26.3 a	28.1 a	0.4 a	21.2 a	23.9 a
	F+	28.0 a	27.0 a	0.7 a	19.7 ab	24.7 a
T p value		0.059	0.282	0.289	0.010	0.944
M X T p value		0.008	0.234	0.500	0.008	0.714

473 Per each factor and per each column, values followed by different letters are statistically different.

474 Two-way ANOVA, Tukey's test, $p \le 0.05$.

475 SND: Symmetrical nucleus division

476 AND: Asymmetrical nucleus division

477 P medium: Germanà at al 1996; N6 medium: Germanà and Chiancone, 2003.

478 Co: no thermal treatment; H: 35 °C for 30 minutes; H+: 40 °C for 60 minutes; F: -20 °C for 30 minutes; F+: -20
479 °C for 60 minutes.

480

Table 6 Influence of medium composition and thermal treatment on 'Tonda Gentile Romana' TR isolatedmicrospore development, after twenty months of culture.

Factors		Uninucleated microspores	Binucleated microspores (SND)	Binucleated microspores (ASD)	Trinucleated microspores	Multinucleated microspores
		%	%	%	%	%
	Р	25.6 b	24.5 a	0.0 a	19.7 a	30.1 a
Medium (M)	N6	29.6 a	28.2 a	1.0 a	19.4 a	21.8 b
M p value		0.035	0.303	0.204	0.794	0.002
	Co	24.3 a	23.0 a	0.3 a	22.7 a	29.7 a
	Н	28.3 a	25.7 a	0.7 a	16.0 a	29.3 a
Treatment (T)	H+	31.0 a	28.0 a	0.3 a	18.3 a	22.3 b
	F	26.3 a	28.1 a	0.4 a	21.2 a	23.9 b
	F+	28.0 a	27.0 a	0.7 a	19.7 a	24.7 b
T p value		0.222	0.221	0.709	0.364	0.007
M X T p value		0.010	0.010	0.954	0.330	0.043

483 Per each factor and per each column, values followed by different letters are statistically different.

484 Two-way ANOVA, Tukey's test, $p \le 0.05$.

485 SND: Symmetrical nucleus division

486 AND: Asymmetrical nucleus division

487 P medium: Germanà at al 1996; N6 medium: Germanà and Chiancone, 2003.

488 Co: no thermal treatment; H: 35 °C for 30 minutes; H+: 40 °C for 60 minutes; F: -20 °C for 30 minutes; F+: -20
489 °C for 60 minutes.

490

492 Figure legends

- 493 Fig. 1: a) Catkins and one anther of *Corylus avellana* L., cv. 'Imperatrice Eugenia', at the developmental stage
 494 used for microspore isolation; b) Uninucleated microspore of cv. 'Imperatrice Eugenia' (DAPI staining) (Bar
 495 represents10µm).
- 496 Fig. 2: a) Binucleated pollen, symmetrical division (DAPI staining), cv. 'Imperatrice Eugenia'; b) Trinucleated
- 497 microspore (DAPI staining), cv. 'Imperatrice Eugenia'; c-d) Multinucleated microspore (DAPI staining), cv.
- 498 'Carrello' (c) and 'Tonda Gentile Romana' TR (d) (Bars represent10µm).
- 499 Fig. 3: Microspore-derived calli of cv. 'Tonda Gentile Romana' GR (Bars represent 100μm).
- Fig. 4 Microspore-derived embryos of cv. 'Tonda Gentile Romana' GR: a-c) different stages of development
 (Bars represent 20µm).
- 502 Fig. 5: a) Influence of the genotype on the regeneration of calli and embryos of five hazelnut genotypes (One
- 503 way ANOVA, Conover-Inman's test $p \le 0.05$) Within each parameter (callus or embryo), values followed by
- different letters are statistically different at $p \le 0.05$ according to Conover-Inman's test; b-f): Influence of the
- 505 medium composition (P and N6) and of thermal treatments (35 °C for 30 minutes (H), 40 °C for 60 minutes 506 (H+), -20 °C for 30 minutes (F), -20 °C for 60 minutes (F+)) on the production of calli and embryos of
- 507 'Carrello' (b), 'Tonda Gentile Romana' GR (c); 'Imperatrice Eugenia' (d), 'Minnulara' (e) and 'Tonda Gentile
- 508 Romana' TR (f).
- Fig. 6 Genetic profile at SSR locus CaT-B505 of the cultivar "Minnulara" (above) and of the microsporederived embryo (below). The embryo shows a single allele (128 bp) shared with the parental genotype.
- 511 Fig. 7 Genetic profile at SSR locus CaT-B107 of the cultivar "Minnulara" (above) and of the microspore-
- 512 derived embryo (below). The embryo shows a single allele (114 bp) shared with the parental genotype.
- 513