Optimization of Cucumber Doubled Haploid Line Production Using In Vitro Rescue of In Vivo Induced Parthenogenic Embryos

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ABSTRACT. Homozygous doubled haploid lines (DHLs) from new cucumber (Cucumis sativus L.) accessions could be useful to accelerate breeding for resistant varieties. DHLs have been generated by in vitro rescue of in vivo induced parthenogenic embryos. The protocol developed involves the following: 1) induction of parthenogenic embryos by pollinating with pollen irradiated with a Co\(^{60}\)\(\gamma\)-ray source at 500 Gy; 2) in vitro rescue of putative parthenogenic embryos identified by their morphology and localized using a dissecting scope or X-ray radiography; 3) discrimination of undesirable zygotic individuals from the homozygous plants using cucumber and melon SSR markers; 4) determination of ploidy level from homozygous plants by flow cytometry; 5) in vitro chromosome doubling of haploids; and 6) acclimation and selving of selected lines. Codominant markers and flow cytometry confirmed the gametophytic origin of plants regenerated by parthenogenesis, since all homozygous lines were haploids. No spontaneous doubled haploid plants were rescued. Chromosome doubling of haploid plants was accomplished by an in vitro treatment with 500 \(\mu\)M colchicine. Rescue of diploid or chimeric plants was shown by flow cytometry, prior to their acclimation and planting in the greenhouse. Seling of colchicine-treated haploid plants allowed for the perpetuation by seed of homozygous lines. The high rate of seed set, 90% of the lines produced seed, facilitated the recovery of inbred lines. Despite some limiting factors, parthenogenesis is routinely used in a cucumber-breeding program to achieve complete homozygos-ity in one generation. Breeding for new commercial hybrid cultivars will be accelerated. DHLs are ideal resources for genonomic analyses.

Production of haploids through androgenesis or parthenogenesis is frequently used in modern breeding programs of several vegetable (Dolcet-Sanjuan et al., 1997; Morales et al., 2002; Oliver et al., 2000) and ornamental species (Dolcet-Sanjuan et al., 2001). Chromosome doubling of haploids allows for the production of homozygous doubled haploid lines (DHLs), which, once perpetuated by seed, can be used in hybrid production. Breeding for resistance to diseases in new cucumber hybrid varieties can benefit from a reliable method to produce a high number of DHLs.

The effect of pollination with \(\gamma\)-irradiated pollen on cucumber ovule development was first reported by Truong-Andre (1988). After being developed for melon (Cucumis melo L.) by Sauton and Dumas de Vaulx (1987), the excision of 3-week-old parthenogenic embryos followed by in vitro rescue was applied to cucumber by Sauton (1989) and Niemirowicz-Szczytt and Dumas de Vaulx (1989). Detailed analysis of cucumber haploid plants, as well as evaluation of the genotype, pollen irradiation dose, and season effects on frequency of haploid embryo development, were first reported by Przyborowski and Niemirowicz-Szczytt (1994), Çaglar and Abak (1996), and reviewed in Przyborowski (1996). Later, efforts were dedicated to improve the haploid production efficiency, measured as haploid plants regenerated per fruit, and the production of doubled haploid plants via chromosome doubling, in order to develop cultivars (Çaglar and Abak, 1997, 1999). These reports indicated that the number of haploid lines generated by in situ induced parthenogenesis ranged from 0.2 to 0.9 per 100 seeds. However, no rates were reported for the number of fertile DHLs perpetuated by seeds.

The application of gynogenesis (Dirks, 1996; Gémesné-Juhász et al., 1997) and androgenesis (Kumar et al., 2003), through in vitro ovary and anther culture, were described as alternative but not more efficient approaches to generate haploids in cucumber than parthenogenesis. These three reports measured the production of haploid plants but gave no details about the recovery of fertile DHLs, essential for the use of these techniques in breeding programs.

Herein, a protocol for the production of cucumber DHLs is described, which has proved to be valuable for the generation of short- and long-fruit-type commercial hybrids. The protocol is based on in situ induced parthenogenesis and in vitro embryo rescue. Optimization of the different steps, from parthenogenic embryo induction to perpetuation of DHLs by seed, is discussed. Flow cytometry and microsatellites are used to assess the ploidy level and the homozygosity of the generated lines.

Materials and Methods

PLANT MATERIAL AND POLLINATION WITH IRRADIATED POLLEN. Ten gynoeccious cucumber F\(_1\) hybrids with long fruit (CS9701, CS9901 to CS9903, CS0001 to CS0004, CS0103 and CS0104) and two of short fruit (CS0101 and CS0102) were used as mother plants. A pure line, LP1, with thorns on fruits, stems, and leaves, as a dominant morphological character, was used as pollen donor. LP1 had a SSR pattern distinctive from the mother F\(_1\) hybrids.

Materials and Methods

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LP1, were irradiated at 250 or 500 Gy, with a Co\textsuperscript{60} γ-ray source. Pistillate flowers of the F\textsubscript{1} hybrids were bagged 1 d before the corolla opened, to avoid undesirable crosses. Next morning, while stigmas were receptive, pollination with irradiated LP1 pollen was performed and flowers were bagged again. Three to 5 weeks after pollination fruits were harvested and embryos cultured in vitro.

**Detection of putatively parthenogenic embryos by X-ray radiography.** Eighty-four fruits of four cucumber hybrids (CS0001 to 4) were opened under aseptic conditions, 3 to 5 weeks after pollination with γ-irradiated pollen, and 12,000 seeds were selected and plated on a gel blotting paper. Seeds were dried for 1 h, at 25 °C, in a flow hood chamber, and inserted in a sterile Omnitray (Nalge Europe Ltd., Hereford, U.K.) with lid. This dish was placed on a black and white Polapan 52 film (Polaroid Corp., Luton, U.K.) and introduced in a Faxitron model MX-20 cabinet (Faxitron X-ray Corp., Wheeling, Ill.). In preliminary trials, best results were obtained setting the voltage of the X-ray tube to 16 kV, and the exposure time to 70 s. Each radiography allowed for the imaging of ≈100 to 150 seeds. Cucumber seeds from fruits harvested 3 weeks after self-pollination were used under the same conditions to determine the reliability of the X-ray radiography in detecting zygotic embryos.

**In vitro rescue of putative parthenogenic embryos.** Three to 5 weeks after pollination fruits were harvested, seeds extracted under sterile conditions, and in situ induced parthenogenic embryos were rescued in vitro by culture in E20H8 medium. Fruits were disinfected by spraying their surface with 96% (v/v) ethanol. One-third of the fruit, the proximal part to the pedicel, was discarded since preliminary results indicated that it contained no embryos. Parthenogenic embryos were identified by their morphology, such as absence or nonsymmetric displacement of cotyledons or other abnormalities. The E20H8 medium was based on the E20 (Sauton and Dumas de Vaulx, 1987), supplemented with 7.9 mM CaCl\textsubscript{2}, 2H\textsubscript{2}O, 0.17 mM CoCl\textsubscript{2}, 6H\textsubscript{2}O, 0.10 mM FeEDTA, 20 g·L\textsuperscript{−1} sucrose, and 8 g·L\textsuperscript{−1} Bacto agar (Difco Laboratories, Detroit). The pH was adjusted to 5.9 before autoclaving. During the first two subcultures this medium was supplemented with 0.06 μM IAA and 15 μM silver thiosulfate (STS), to promote root growth and overcome vitrification, respectively.

Embryos were extracted from seeds and plated in petri dishes (6 cm diameter) with 10 mL of E20H8 medium. Cultures were incubated at 28 °C with a 16-h photoperiod (50 μmol·m\textsuperscript{−2}·s\textsuperscript{−1}). A maximum of four subcultures to fresh medium were done every 2 weeks. Developed plantlets were transferred to tubes, and later on to flasks, with E20H8 medium, following micropropagation by axillary branching. During micropropagation, cultures were incubated at 28 °C with a 16-h photoperiod (70 μmol·m\textsuperscript{−2}·s\textsuperscript{−1}). Plantlets with a good root system and four to six normal leaves were acclimated in Jiffy-7 peat plugs (Clause Tézier SA, Portes-les-Valence, France).

**Homozygosity and ploidy determinations.** The origin of lines, obtained by in vitro embryo rescue and micropropagation, was determined by the microsatellite (SSRs) patterns complemented with the ploidy level, determined by flow cytometry. Lines that were heretozygous for any of the SSRs used (CMGA165, CSAT425, CMAG59, and CMSSR1), therefore originated from zygotic embryos, were eliminated. Zygotic embryos could originate from viable pollen of LP1 unaffected by γ-ray irradiation or from uncontrolled pollinations. Lines homozygous for all SSRs, presumably originated from parthenogenic embryos, were maintained in vitro and their ploidy level was determined. Flow cytometry was used to determine the DNA content of each line, during in vitro micropropagation and after in vitro colchicine treatment. Young leaves from actively growing plants were finely chopped and macerated for 5 min into 1 mL LB01 lysis buffer (Dollet-Sanjuan et al., 1997), with 2% (v/v) Triton X-100 to release intact nuclei. The supernatant was filtered through a 56-μm nylon mesh. The nuclei were labeled with propidium iodide, and red fluorescence was analyzed using a 675-nm band-pass filter. Results were presented in a histogram expressing the distribution of nuclei according to the red fluorescence intensity, which is proportional to the DNA content.

**SSR analysis.** Four SSR markers were used for the homozygosity test of the haploid lines: CMGA165 and CSAT425 (Damin-Poleg et al., 2001), CMAG59 (Katirz et al., 1996), and CMSSR1 (Garcia-Mas, unpublished). CMGA165, CMAG59, and CMSSR1 are melon SSRs, and CSAT425 is a cucumber SSR. PCR reactions were performed in a final volume of 15 μL with 1 μL Taq buffer [20 mM (NH\textsubscript{4})SO\textsubscript{4}, 75 mM Tris-HCl pH 8.8, 0.01% (v/v) Tween 20], 2 mM MgCl\textsubscript{2}, 166 μM dNTPs, 2 pmol of each primer (one labelled with the fluorochrome IRD-800), 1 unit of Taq DNA polymerase (Applied Biosystems, Foster City, Calif.), and 60 ng of DNA. The cycling conditions were as follows: an initial cycle at 94 °C for 1 min followed by 35 cycles at 94 °C, 30 s; 40–60 °C, 30 s; 72 °C, 1 min; and a final cycle at 72 °C for 5 min. SSR fragments were separated in a LI-COR IR\textsuperscript{2} (LI-COR, Lincoln, Nebr.) automatic sequencer. Five microliters of loading buffer [95% (v/v) formamide, 20 mM EDTA, 0.05% (w/v) bromophenol blue, 0.05% (w/v) xylene cyanol] were added to the PCR mix, samples were denatured at 100 °C for 5 min and 0.8 μL were loaded on to a LI-COR IR\textsuperscript{2} sequencer using 25-cm plates with 6% (w/v) acrylamide in 1× TBE and 7.5 M urea. Electrophoresis was performed at 1500 V, 35 mA, and 31 W at 50 °C until the PCR products were visible. The molecular weight of each microsatellite band was estimated by comparing its migration with the IRD-labelled STR molecular size marker (LI-COR).

**Chromosome doubling of haploid plants.** As shown by flow cytometry, most of the rescued lines, homozygous for all SSRs, were haploid. Micropropagation of haploid lines was required before their chromosome number was doubled in vitro with a colchicine treatment. This was performed on 100 to 200 cuttings, with one or two axillary buds each, per line. Explants were either cultured in petri dishes (10 cm diameter) or polystyrene RA40 Microboxes (Combines, Gent, Belgium) with E20H8 medium supplemented with 500 μM colchicine, dissolved in dimethyl sulfoxide (DMSO). Forty-eight hours later, explants were transferred to colchicine-free E20H8 medium. Four to 8 weeks later, new shoots grew up from some explants, out of most of the parthenogenic lines. Twenty shoots per line were rescued and individually micropropagated in E20H8 medium. Doubled haploid lines, confirmed by flow cytometry, were maintained in vitro until they were acclimated and transferred to the greenhouse for self-pollination.

**Self-pollination and perpetuation of doubled haploid lines.** At least 20 plants from each doubled haploid line were acclimated, hardened, and transferred to the greenhouse for self-pollination. Induction of staminate flowers on gynoecious cucumber lines was achieved by spraying with STS. After a first selloing, lines with a sufficient number of seeds were eliminated from the in vitro stock, and a second self-pollination was performed on the remaining lines. A minimum of 50 seeds was considered necessary to perpetuate a doubled haploid line.
Results and Discussion

Plant material. Inbred lines and hybrids were the source of parthenogenetic embryos in most of the previous reports (Çaglar and Abak, 1996, 1999; Niemirowicz-Szczytt and Dumas de Vaulx, 1989; Przyborowski, 1996; Przyborowski and Niemirowicz-Szczytt, 1994). Herein, and for the first time, in situ induction of haploid parthenogenetic embryos through pollination with γ-irradiated pollen has been reported on parthenocarpic, gynoecious cucumber F1 hybrids of short and long fruits.

During the determination of the optimal irradiation dose (Fig. 1), the genotype had no significant effect (P = 0.074). With a more extensive number of genotypes, the maternal plant was the main factor affecting the rate of parthenogenetic embryos produced (Table 1). Similar results were reported previously, with values ranging from 0.3 to 1.7 embryos per 100 seeds (Przyborowski and Niemirowicz-Szczytt, 1994; Przyborowski, 1996) or as high as 5.3 for one hybrid (Çaglar and Abak, 1996, 1999). These values, apparently lower (Table 1), are not comparable to the latter since they have been obtained from different cucumber types and correspond to the rescued parthenogenic embryos and not to the total number of embryos observed, which is dependent on the morphological criteria used to determine the embryos origin. The reported average number of embryos developed to plants per 100 seeds was 0.3 in Troung-Andre (1988), 0.9 in Çaglar and Abak (1996), and 0.2 to 0.8 in Przyborowski (1996), which are close to the results of rescued parthenogenetic embryos shown in Table 1.

Pollen irradiation dose. The basis to select between the two Co60 γ-irradiation doses (250 or 500 Gy) was to enhance the formation of parthenogenetic embryos and to minimize the formation of zygotic embryos. Without significant interaction (P = 0.635) between genotype and doses of irradiation, a significant increase (P = 0.043) of the number of parthenogenetic embryos produced was observed with 500 Gy (Fig. 1), with no significant difference between the genotypes (P = 0.074). At both irradiation doses the generative nuclei of pollen were inactivated, avoiding the formation of zygotic embryos. Other reports had focused on lowering the dose of 400 or 600 Gy, first reported by Troung-Andre (1988), to improve the rate of parthenogenetic embryo induction (Çaglar and Abak, 1996, 1999; Przyborowski, 1996; Przyborowski and Niemirowicz-Szczytt, 1994). Although they all reported a beneficial effect on the number of embryos, from lowering the dose of irradiation to 200 and 300 Gy, there was no statistical difference observed. In addition, although the number of putative parthenogenetic embryos increased after lowering the irradiation dose, the number of embryos developing into plants was not enhanced (Przyborowski, 1996). Therefore, the 500-Gy dose was chosen to irradiate pollen of the cucumber line LP1 as inducer of parthenogenetic embryos.

Season to induce parthenogenesis. As reported by Przyborowski and Niemirowicz-Szczytt (1994), Przyborowski (1996), and Çaglar and Abak (1996, 1999), summer was the optimal season to induce parthenogenesis through pollination with γ-irradiated pollen, because it is the best season to induce parthenocarpy in the F1 hybrids. In our conditions, parthenogenesis was inducible during the months of May through July, with no differences within this period (data not shown).

Detection of putatively parthenogenetic embryos by X-ray radiography. The effectiveness of X-ray radiography to detect and identify seeds containing immature cucumber embryos was 32% (six embryos found by X-rays out of 19 embryos found under a dissecting scope). This efficiency was lower than that indicated by Sauton et al. (1989) for melon. A total of 17 seeds were selected with X-ray radiography as containing embryos (Fig. 2). However, after visualization under the dissecting scope, only six were shown to have embryos, the rest were enlargement of the integuments. In addition, 13 embryos were found by dissection in those seeds not selected by X-ray radiography imaging. Moreover, the viability of the embryos after the required drying and manipulation of seeds to perform the X-ray radiography was reduced to 40%, while that after direct dissection was more than twice this value (see next section). In consequence, since the method was not effective in cucumber, seeds were selected.
and opened under a dissecting scope. Some immature cucumber seed covers are translucent and in other cases the embryos grow out of the seed covers (Fig. 3A), allowing for a first selection of the seeds containing embryos.

**In vitro rescue of putatively parthenogenic embryos.**
In vitro rescue of putatively parthenogenic embryos (Fig. 3) in E20H8 medium, supplemented with IAA and STS, allowed for an average embryo rescue of 83% of the embryos into plants (Table 1). Niemirowicz-Szczytt et al. (2000) found difficulties in conserving in vitro a collection of haploid cucumber plants. The maintenance by micropropagation of haploid lines or doubled haploids, after colchicine treatment, presented no difficulties in the E20H8 medium.

**Homozigosity and ploidy determinations.** Ploidy level determinations by flow cytometry (Fig. 4), of lines derived from putative parthenogenic embryos, showed that 62% of them were haploid plants (Table 2), with <30% of their nuclei being diploid (Fig. 4A). In addition, although 38% of the lines were chimeric (Table 2), with haploid cells and a predominant number of diploid nuclei (Fig. 4B), no complete diploid lines were observed (Table 2). This is in concordance with the spontaneous chromosome doubling observed in some root meristems after micropropagation of haploid lines (Niemirowicz-Szczytt and Dumas de Vaulx, 1989; Przyborowski, 1996; Przyborowski and Niemirowicz-Szczytt, 1994). The use of cucumber and melon SSR markers was useful to discriminate the existence of undesirable zygotic individuals from the homozygous plants (Fig. 5). Panels A, B, and C show that none of the lines inherited the LP1 allele, discarding the possibility of a functional LP1 pollen. SSR patterns in panels D, E, and F show that lines inherited either one of the alleles present in the respective hybrids. A zygotic individual is identified in panel E. This has two alleles: one coming from the pollen donor LP1 and another from the hybrid CS9902. Codominant
markers and flow cytometry confirmed the gametophytic origin of plants regenerated by parthenogenesis, since all homozygous lines were complete haploids or had some haploid cells in the aerial part of the plant. These results validated the effectiveness of the morphological characters used to select the putatively parthenogenic embryos (Fig. 3).

**Chromosome doubling of haploid plants.** Chromosome doubling of haploid plants was accomplished by an in vitro treatment with 500 μM colchicine. In vitro chromosome doubling of haploids allowed for the recovery of viable plants from all the lines. Survival of explants to the colchicine treatment depended on the haploid line, with a normal distribution among the 143 haploid lines treated (Fig. 6). From most of the lines, 20% to 60% of the treated explants produced new shoots. Such recovered shoots had broader and thicker leaves that indicated a possible change in their ploidy level (Fig. 7). After colchicine treatment, 30% of the rescued plants became complete diploids, 55% were

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Table 2. Ploidy level of cucumber (Cucumis sativus L.) parthenogenic lines before and after in vitro colchicine treatment.
flow cytometry and SSRs markers. No spontaneous chromosome doubling was observed among the in vitro rescued lines. In vitro colchicine treatment induced diploid or chimeric plants, with diploid and haploid cells. However, their level of fertility was high enough to perpetuate by seed most of the DHLs. Despite some limiting factors in the rates of DHL production, parthenogenesis is routinely used in cucumber-breeding programs to achieve complete homozygosity in one generation, and accelerate breeding for new commercial varieties. With this protocol, sufficient number of cucumber DHLs can be generated to be selected and incorporated in a breeding program for new resistant varieties, or to be used in hybrid production. So far, DHLs have been selected and used as parental lines for the production of new commercial hybrids.

Literature Cited


Fig. 7. New cucumber shoots growing on explants treated with colchicine (bar = 1 cm).

Fig. 8. Fruits with seed on self-pollinated cucumber doubled haploid lines of long (A) and short (B) fruit type.