Doubled haploid production in alfalfa (Medicago sativa L.) through isolated microspore culture

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Rapid production of doubled haploids (DHs) through isolated microspore culture is an important and promising method for genetic study of alfalfa. To induce embryogenesis in alfalfa, isolated microspores were submitted to abiotic stresses during their initial culture in order to stimulate them to form embryos and plantlets. ‘Baoding’ and ‘Zhongmu No 1’ alfalfa cultivars supported reproducible and reliable proliferation response irrespective of any stress treatment of microspores. The microspore developmental stage for isolated microspore culture was studied and we found that uninucleate microspores were best to initiate culture. Exposure of microspores to appropriate low temperature or heat shock stresses were able to increase the efficiency of embryogenesis. The most effective low-temperature treatment was 4°C for 24 h and the frequency of plantlets induction was 20.0%. The most effective heat shock treatment was 32°C for 2 d and the frequency of plantlets induction was 14.17%. The analysis of ploidy level performed by flow cytometer revealed that the majority of 278 regenerated plantlets were haploid (65.83%) or doubled haploid (33.81%). This is the first report of haploid production in alfalfa through isolated microspore culture.

Development of haploid plants through isolated microspore culture is an efficient method that are useful in genetic studies. The haploid plants have the gametic chromosome number (n) and when the chromosome number of a haploid have undergone spontaneous or induced chromosome duplication by chromosome doubling agents, it is termed as a doubled haploid (DH) with chromosome number 2n. For basic research the haploids have many important advantages. Haploids provide beneficial tools for genome research which aim to elucidate the structure function and evolution of past and present genomes. Doubled haploids (DHs) are genetically normal and it provides a unique opportunity to screen gametophytic variation caused by recombination and segregation during meiosis. Even the recessive variations can express itself in the absence of their dominant counterparts in such DHs plants. Isolated microspore culture and anther culture are the two major methods used for haploid production. The regenerated plants produced by anther culture are composed of mixoploids and it is difficult and laborious to detect ploidy of the plants. The more advantageous method is regenerating plants by isolated microspore culture because haploids or DHs are produced. Isolated microspore culture has recently been used more frequently in many species, e.g., tobacco, wheat, barley, canola, rice, cabbage, pepper, and maize, for which efficient induction protocols are available.

Alfalfa (Medicago sativa L.) is one of the most important legume forage which plays a vital role in livestock production. According to the statistics of the National Alfalfa & Forage Alliance, an estimated 32 million ha of alfalfa were grown annually in the world. Success in the production of haploid plants from tetraploid alfalfa was achieved by crossing tetraploids (4X) with a diploid (2X) pollen parent to yield haploid progeny. Further work by Zagorska et al. demonstrated conclusively that haploid plants could also be generated from alfalfa through anther cultures. In their studies, anthers were pretreated for four days at 4°C and treated with 1 Gy gamma radiation prior to culture. Of 100 regenerants, a total of only four haploid plants were produced. The frequency of haploid production in alfalfa by the 4X-2X cross and anther culture methods is very low. Efficient microspore embryogenesis in culture is successfully induced by the application of abiotic stress and other pretreatments in a number of species, including legumes. Therefore, rapid production of haploid plants through microspore

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The genotype, growth conditions of donor plants, the developmental stage of microspores, culture medium, and different stress treatments have been reported to influence the efficiency of embryogenesis in isolated microspore culture. The microspore developmental stage has been extensively studied with late uninucleate and early binucleate microspores being most responsive. The positive effects of modifying the osmotic pressure of the medium on microspore embryogenesis have been observed previously in many species, including legumes. Short heat shock treatments (32°C for 24–72 h) during the early stage of isolated microspore culture have been required to induce microspore embryogenesis in *Brassica*. Exposure of microspores to a low temperature (4°C for 24–48 h) preceding the heat shock treatment reportedly promotes microspores embryogenesis. Furthermore, the additions of a high level of sugar, activated charcoal, and colchicine in the induction medium were found to increase microspore embryogenesis. In conclusion, apart from selection of genotype with embryogenic capacity and a suitable microspore development stage, it is important for microspores to be subject to the action of a single stress or several kinds of stress to induce embryos and plant formation.

Against this background, the objectives of this study were to test the effects of genotype, microspore development stage, low temperature, and heat shock on microspore embryogenesis in alfalfa and to establish an efficient protocol for haploid production in alfalfa using isolated microspore culture.

**Results**

**Microspore embryogenesis.** Flower buds (Fig. 1A) were harvested from young inflorescences. Then isolated microspores were cultured in NLN-13 liquid medium. Initial development of the microspores in culture was visible as an enlargement of the cells within 5–8 days of culture. Some of these swollen microspores (Fig. 1B) continued apparent gametophytic development whereas other moderately swollen ones showed sporophytic cell division, forming multicellular embryogenic clusters after approximately 15 days in culture. Well-developed embryos visible to the naked eye were formed after approximately 21 days of culture (Fig. 1C). Embryos with two or three leaves were placed on solid B5 medium. Plantlets with one to three leaves (Fig. 1B) were transferred to soil and acclimatized in a greenhouse.

**Effect of genotype on alfalfa microspore embryogenesis.** A range of genotypes was tested during our experiments. Significant variation in microspore embryogenesis among genotypes was found (Table 1). Initial microspore division was observed for alfalfa genotypes at different frequencies, and resulted in the formation of microcalluses. For some genotypes of alfalfa, growth was arrested very early. Ultimately, out of the 10 genotypes studied, only ‘Baoding’ and ‘Zhongmu No. 1’ alfalfa (Table 1) proliferated first to calluses (Fig. 1D) and then to plantlets (Fig. 1E). The greatest frequency of plantlet induction was found in ‘Baoding’ alfalfa and the frequency of plantlets induction was 7.5% (Table 1). The others remained recalcitrant, whatever the medium, treatment or culture conditions, indicating that genotype is the main factor governing androgenetic capacity in these species. A high frequency of plantlet formation was achieved for ‘Baoding’ alfalfa in microspore culture, therefore this genotype was used as plant material in the following experiments.

**Effect of microspore developmental stage on alfalfa microspore embryogenesis.** The developmental stage of microspores (tetrad, early uninucleate, late uninucleate, and binucleate stages) was assessed for improving embryogenic yield in isolated microspore culture. The best embryogenic development was from uninucleate microspores (Fig. 1F; Table 2) when flower buds were 6.02–6.20 mm long and 1.50–1.72 mm wide (Table 3). Earlier stages contained mainly tetrads that did not survive in culture, while at later stages microspores were already unable to shift from gametophytic to sporophytic development required for proliferation conductive to the ultimate formation of embryos.

**Effect of low-temperature treatment on microspore embryogenesis.** The effect of low-temperature treatment applied to isolated microspores on embryogenesis of ‘Baoding’ alfalfa is displayed in Table 4. The best proliferation responses were obtained with isolated microspores that were cultured at 4°C for 24 h in the dark. After treatment with this temperature, the frequency of plantlets induction was 20.0%, whereas the frequency of plantlets induction was 10.0% in the control treatment combination. Isolated microspores that had been subjected to cold stress at 4°C for 24 h showed increased viability during the culture. However, it was observed that cold treatment for more than 24 h had no positive effect on the percentage of microspore division.

**Effect of heat shock stress on microspore embryogenesis.** Performance of ‘Baoding’ alfalfa in isolated microspore culture after heat shock stress treatment at 32°C is shown in Table 5. We found that an appropriate heat shock stress treatment had a positive effect on isolated microspore cultures. The most effective stimulant was the use of 32°C for 2 d. From isolated microspores treated with a heat shock stress for 2 d, the frequency of plantlets induction was 14.17%, whereas by establishing isolated microspore culture without the heat shock stress treatment, the frequency of plantlets induction was 7.5%.

**Analysis of ploidy level.** In total, 278 plantlets, obtained through isolated microspore culture, were assessed for their ploidy level. The representative results on flow cytometric DNA analysis of cells in microspore-derived plantlets were shown in Fig. 2. Ploidy level of the plantlets was recorded in Table 6. The majority of plants were haploid and doubled haploid, with 65.83% haploid and 33.81% doubled haploid. In the microspore-derived populations, in addition to haploid and double haploid, there was a low frequency of higher ploidy. One octoploid plant (0.36%) was found in the population.
Discussion

Under certain conditions, microspores can shift development from gametophytic to sporophytic to form embryos and plantlets\(^3\). However, most of legume species are highly recalcitrant to microspore embryogenesis. Difficulties were posed by the generally small anther size and relatively low numbers of microspores per anther in legume crops\(^3\). In the present study, various efforts have been made to induce microspore embryogenesis in alfalfa. To the best of our knowledge, this is the first report of microspore-derived plantlet formation in alfalfa. The haploid and DH plants obtained in our study are promising for applications in genetic studies. In the study, only 277 haploid and DH plants were produced from thousands of flower buds from ‘Baoding’ alfalfa indicating the necessity to improve the protocol for haploid production in alfalfa. Further work is under way to improve microspore responses and extend them to other genotypes.

To establish an efficient protocol for microspore embryogenesis in alfalfa, the effects of genotype, stage of microspore development, low-temperature treatment, and heat shock stress on microspore culture were studied in the present study. Of the 10 genotypes, only the microspores of ‘Baoding’ and ‘Zhongmu No. 1’ alfalfa proliferated to yield calluses and plantlets when microspores were not exposed to any specific treatment, indicating that genotype is crucial for microspore response. Results also demonstrated that ‘Baoding’ alfalfa is a highly responsive cultivar in isolated microspore culture and thus it can be used as experimental material in
future studies. Additionally, the greatest amount microspore-derived plantlet formation in ‘Baoding’ alfalfa were observed when microspores were in the early and late uninucleate developmental stages. Low temperature and heat shock stresses had positive effects on plantlet formation in culture. The most effective stimulant was exposure to 4°C for 24 h or 32°C for 2 d, the frequency of plantlets induction was 20.0% and 14.17%, respectively. Apart from the above-mentioned parameters, addition of active carbon to medium 16, kinds of medium 14,18, culture conditions18, pyramiding electrostimulation17, and osmotic shock17, alone or in combination14, can also affect the

| Genotypes        | Number of plantlets produced | Plantlets induction frequency |
|------------------|------------------------------|------------------------------|
| ‘Zhongmu No. 1’  | 2                            | 1.67% a                      |
| ‘Zhongmu No. 6’  | 0                            | 0 b                          |
| ‘Gongnong No. 1’ | 0                            | 0 b                          |
| ‘Caoyuan No. 2’  | 0                            | 0 b                          |
| ‘Baoding’         | 9                            | 7.5% c                       |
| ‘Longdong’        | 0                            | 0 b                          |
| ‘Aohan’           | 0                            | 0 b                          |
| ‘WL323’           | 0                            | 0 b                          |
| ‘Golden Empress’  | 0                            | 0 b                          |
| ‘Sanditi’         | 0                            | 0 b                          |

Table 1. Performance of different alfalfa genotypes in microspore culture irrespective of the treatment applied. Different letters following survival rate indicate statistically significant differences based on the Duncan’s new multiple range test at \( P < 0.05 \).

| Microspore developmental stage | Number of plantlets produced | Plantlets induction frequency |
|-------------------------------|------------------------------|------------------------------|
| tetrad                        | 0                            | 0 a                          |
| early uninucleate             | 11                           | 9.17% b                      |
| late uninucleate              | 8                            | 6.67% b                      |
| binucleate stages             | 0                            | 0 a                          |

Table 2. Plantlet yield in culture of different developmental stages of microspores from ‘Baoding’ alfalfa plants. Different letters following survival rate indicate statistically significant differences based on the Duncan’s new multiple range test at \( P < 0.05 \).

| Microspore developmental stage | Length of bud size (mm) | Width of bud size (mm) |
|-------------------------------|-------------------------|------------------------|
| tetrad                        | 5.44 a                  | 1.24 a                 |
| early uninucleate             | 6.02 b                  | 1.50 b                 |
| late uninucleate              | 6.20 b                  | 1.72 c                 |
| binucleate stages             | 6.78 c                  | 1.82 d                 |

Table 3. Morphological characteristics of flower buds at different developmental stages of microspores from ‘Baoding’ alfalfa plants. Different letters following survival rate indicate statistically significant differences based on the Duncan’s new multiple range test at \( P < 0.05 \).

| Low-temperature treatments    | Number of plantlets produced | Plantlets induction frequency |
|-------------------------------|------------------------------|------------------------------|
| 4°C for 0 h                   | 12                           | 10.0% a                     |
| 4°C for 3 h                   | 16                           | 13.33% b                    |
| 4°C for 6 h                   | 18                           | 15.0% b                     |
| 4°C for 12 h                  | 17                           | 14.17% b                    |
| 4°C for 24 h                  | 24                           | 20.0% c                     |
| 4°C for 48 h                  | 11                           | 9.17% a                     |

Table 4. Effect of low temperature (4°C) on alfalfa microspore embryogenesis in culture. Different letters following survival rate indicate statistically significant differences based on the Duncan’s new multiple range test at \( P < 0.05 \).
efficiency of embryogenesis in microspore culture of alfalfa. Therefore, the effect of these parameters on induction of microspore embryogenesis in alfalfa will be studied in future studies.

The present study centered on a single factor influencing haploid production efficiency via inducing microspore embryogenesis in alfalfa and low temperature treatment is more efficient than heat shock stress. Pyramiding some stress agents is a promising way forward to trigger the switch of isolated microspores from the gametophytic to sporophytic developmental pathway in extremely recalcitrant genotypes. A combination of osmotic pressure and electric shock significantly improved responses from isolated microspores of some legume species and yielded calluses. reported that the conjunction of cold pretreatment and heat shock stress, as compared to a single stress treatment alone, increased microspore embryogenesis in broccoli. Further efforts will be designed to improve microspore responses by pyramiding different stress agents.

Materials and Methods

Plant materials. Ten different genotypes of alfalfa, ‘Zhongmu No. 1’, ‘Zhongmu No. 6’, ‘Gongnong No. 1’, ‘Caoyuan No. 2’, ‘Baoding’, ‘Aohan’, ‘Longdong’, ‘WL323’, ‘Golden Empress’, and ‘Sanditi’, were used as donor materials. They were obtained from different seed companies. The donor plants were grown in plastic pots in horticultural peat substrate (one plant per pot) in a greenhouse. The alfalfa plants started to flower 9–10 weeks after sowing. Forty flower buds of appropriate size for microspore culture were collected per genotype and three replicates were undertaken.

Table 5. Effect of heat shock (32°C) on alfalfa microspore embryogenesis in culture. Different letters following survival rate indicate statistically significant differences based on the Duncan's new multiple range test at \(P < 0.05\).

| low-temperature treatments | Number of plantlets produced | Plantlets induction frequency |
|----------------------------|------------------------------|-----------------------------|
| 32°C for 0 d               | 9                            | 7.5% a                      |
| 32°C for 1 d               | 12                           | 10.0% b                     |
| 32°C for 2 d               | 17                           | 14.17% c                    |
| 32°C for 3 d               | 13                           | 10.0% b                     |
| 32°C for 4 d               | 9                            | 8.33% a                     |
| 32°C for 5 d               | 1                            | 0.83% b                     |
| 32°C for 6 d               | 0                            | 0% g                        |

Table 6. Determination of ploidy level of microspore-derived plantlets.

| material | Number of tested plantlets | Haploid | Doubled haploid | Octoploid |
|----------|-----------------------------|---------|-----------------|-----------|
| Baoding  | 278                         | 183     | 94              | 1         |

Figure 2. Flow cytometric DNA analysis of cells in microspore-derived plantlets. (A) Cells in microspore-derived plantlets. (B) Cells in tetraploid alfalfa (the control).
Isolation and culture of microspores. For all genotypes, flower buds containing microspores at appropriate developmental stages were surface-sterilized in 75% ethanol for 30 s, followed by 10 min in 10% (v/v) bleach, and finally, five sterilized water rinses. Microspores were isolated by crushing the flower buds in B5 liquid medium (pH 5.8)13, with three successive steps of suspension and centrifugation (100 × g. 5 min each) to eliminate debris remaining after serial sieving through 40 μm filters. During the whole isolation procedure all liquids and centrifugation were kept at 4 °C. The microspore concentration was determined with a hemocytometer. After counting, the microspores were finally suspended at a density of approximately 1 × 10^5 microspores ml^{-1} in NLN-13 liquid medium14 and incubated for culture in 60 mm × 15 mm Petri dishes (5 ml per dish). We added 100 ul of activated charcoal (1%) to the medium. The dishes were sealed with Parafilm. Then different stress treatments were applied in the dark, and thereafter cultures were maintained at 25 °C under dark conditions. When calluses reached a minimum diameter of 5 mm, they were transferred onto the induction medium as reported elsewhere for *Medicago truncatula*9. Thus, embryogenesis proceeded through the typical successive stages (i.e. globular, heart, torpedo and cotyledonary) of somatic embryo induction, expression, growth and ultimate conversion to plants. Converted embryos with two or three leaves were placed on solid B5 medium13. Plantlets with four or five leaves were transferred to soil and acclimatized in a greenhouse.

Cytological analysis. During the culture of microspores, morphological characteristics and developmental pathways of microspores were observed by fluorescence microscopy. The microspores were collected by centrifugation, stained with 4',6-diamidino, 2-phenylindole (DAPI) and observed under Olympus SXZ10 fluorescence microscope (Olympus, Japan). The determination of morphological characteristics and developmental pathways was conducted following the procedure described by Barinova et al.9.

Determination of the developmental stage of microspores. Flower buds were harvested from their identification to the opening of petals. Microspores isolated from flower buds at the various developmental stages (tetrad, early uninucleate, late uninucleate, and binucleate stages) were cultured and the optimum developmental stage for proliferation from microspores was identified. Prior to culturing, the stage of microspores was checked under microscope. This was performed by staining with 4% acetocarmine (W/V) in 50% acetic acid (v/v)37. Forty buds from each stage were collected and the lengths of the buds were measured.

Low temperature and heat shock treatments. At the optimum stage of microspore development, they were isolated from flower buds following the procedures outlined previously. Then they were cultured at 4 °C in the dark for 0, 1, 2, 3, 4, 5, 6 d or, alternatively, they were heat-shocked at 32 °C for 0, 3, 6, 12, 24, 48 h. Thereafter, cultures were maintained at 25 °C in the dark. The experiment was conducted with three replicates and each replicate consisted of forty buds. The number of plantlets obtained from isolated microspore cultures were counted after six weeks.

Determination of ploidy. Ploidy level was determined by a flow cytometer and the determination was carried out according to the procedures of Galbraith38. Approximately 20 mg of young leaves were crumbled with a razor blade in 0.1 M citric acid containing 0.5% Tween 20. After adding fluorochrome 4',6-diamidino, 2-phenylindole DAPI at 0.1 mg ml^{-1}, samples were filtered through a 50-μm nylon filter. Measurements were done on a flow cytometer (BD FACSCalibur, USA) using a linear scale. Ploidy level was assessed using tetraploid alfalfa genotype as control.

Data analysis. All experiments were repeated three times, each including at least 10 replicate dishes (6 cm in diameter). The data were analyzed using ANOVA with mean separation by Duncan's or LSD Multiple Range Test at a 95% confidence level.

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Author Contributions
Y. and Z.W. designed the experiments. D.Y., J.S. and Y.S. performed the experiments, analyzed the data, and wrote the manuscript; Z.T. and T.Z. performed the experiments. All authors have read and approved the final manuscript.

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