Combined inoculation with dark septate endophytes and arbuscular mycorrhizal fungi: synergistic or competitive growth effects on maize?

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Abstract

Background: Effects on maize were assessed of dual inoculation with arbuscular mycorrhizal fungi (AMF) and dark septate endophytes (DSE) isolated from other plant species.

Methods: Suspensions of DSE isolated from Stipa krylovii were prepared at different densities (2, 4, and $8 \times 10^5$ CFU mL$^{-1}$) and inoculated separately (AMF or DSE) or together (AMF + DSE), to explore their effects on maize growth.

Results: Inoculation with AMF or medium and high densities of DSE and combined inoculation (AMF + DSE) increased plant above-ground growth and altered root morphology. Differences in plant growth were attributable to differences in DSE density, with negative DSE inoculation responsiveness at low density. AMF promoted plant above-ground growth more than DSE and the high density of DSE promoted root development more than AMF. Combined inoculation might lead to synergistic growth effects on maize at low density of DSE and competitive effects at medium and high DSE densities.

Conclusions: AMF and DSE co-colonized maize roots and they had positive effects on the host plants depending on DSE density. These findings indicate the optimum maize growth-promoting combination of AMF and DSE density and provide a foundation for further exploration of potentially synergistic mechanisms between AMF and DSE in physiological and ecological effects on host plants.

Keywords: Arbuscular mycorrhizal fungi, Dark septate endophytes, Maize, Inoculation

Background

Dependence on chemical fertilizers and pesticides results in pesticide residues in crops and soils and increases production costs. Microbial technology has been widely used to solve this problem by inoculating roots with endophytic fungi to exploit their potentially symbiotic associations and stimulate plant growth to provide pollution-free production systems.

Arbuscular mycorrhizal fungi (AMF) are important soil fungi that inhabit the roots of most terrestrial plant species with which they form potentially symbiotic associations. Under appropriate conditions the AMF can increase plant nutrient uptake and carbon fixation by photosynthesis, and enhance plant tolerance to biological or abiotic (e.g., drought, salinity, and low temperatures) stresses [1–5]. For example, the extraradical mycelium of AMF form an important bridge to transport nutrients...
outside the roots to the intraradical mycelium [6] and can promote an effective relationship between plants, bacteria and beneficial fungi [7]. The concentrations of glomalin in soil have often been correlated with AMF biomass and the formation, deposition, and/or decomposition of glomalin in soils seem to be largely dependent on a multitude of interactions among plants, AMF, and other soil microorganisms, including prokaryotes [8]. Simulation experiments indicate that inoculation of plants with AMF increases plant biomass and alleviates mechanical damage to the root system [9]. AMF inoculation also increases the abundance of certain beneficial bacterial species and decreases that of certain pathogenic fungi in the rhizosphere [10].

Dark septate endophytes (DSE) are a miscellaneous group of asexually propagating fungal endophytes that colonize living root tissues intracellularly and intercellularly in a range of extreme ecosystems [11, 12]. DSE have a wide host range and ecological distribution including mycorrhizal [13, 14] and non-mycorrhizal [15, 16] plants. In contrast to arbuscular mycorrhizal fungi, DSE not only grow inter- and intra-cellularly within the root cortex but can extend into the vascular tissue [17, 18]. Typical micro structures of DSE such as dark septate hyphae and microsclerotia are formed with different degrees of melanism [14, 19, 20]. DSE melanin can assist host plants in resisting adverse environmental conditions by protecting fungal mycelium from abiotic stresses such as high temperatures, drought, and potentially toxic elements [21, 22]. Numerous studies suggest that DSE can promote the uptake and transformation of mineral and organic nutrients and increase the adaptability of the host or the nutritional status of plants to increase resistance to diseases and other stresses [11, 23–26].

DSE may form a root-fungal association with the host [14], modify the mycorrhizal status of the plant, and thus modulate a different symbiotic association in the rhizosphere [27]. A previous study using DSE with other microorganisms such as Trichoderma viride assessed the effects of combined inoculation on plant growth and substantiated their positive influence [28].

Low colonization rates by AM fungi at the pre-symbiotic stage may be compensated by DSE by some weak competitive or antagonistic interactions between the two fungal groups [29, 30]. A succession of dominant fungal colonizers from AMF to DSE has been observed in the roots of the grass Deschampsia flexuosa and may be related to the different nutrient acquisition strategies of the two fungal groups [31]. Numerous separate studies of AMF and DSE have drawn widespread attention to their favorable ecological functions [32] but combined inoculation of plants with AMF and DSE have been little studied and remain poorly understood [33].

The objective of the present study was to obtain insights into the interactions between AMF and DSE and to extend their potential use in future field application. Suspensions of DSE isolated from Stipa krylovii were prepared at different densities and AMF and DSE were used to inoculate maize separately (AMF or DSE) and together (AMF+DSE) to investigate their effects on the host plant. The work addressed the questions of whether DSE and AMF can co-colonize maize roots in vitro and whether inoculation with DSE and AMF has synergetic or competitive effects on the growth of the host plant?

Results

Plant growth

Inoculation with AMF and DSE had significant effects on plant growth (Table 1). Plant height, ground diameter, and leaf area in combined treatments AM+MD (64.4 cm) and AM+HD (65.6 cm) were higher than

| Treatments | Height (cm)  | Ground diameter (mm) | Leaf area (cm²) | Shoot fresh biomass (g) | Root fresh biomass (g) | Shoot dried biomass (g) | Root dried biomass (g) | Inoculation responsiveness (%) |
|------------|-------------|----------------------|-----------------|------------------------|------------------------|------------------------|------------------------|-----------------------------|
| CK         | 35.47 ± 4.83b | 0.54 ± 0.02d         | 192.72 ± 33.32d | 8.89 ± 3.54c           | 1.7 ± 0.82e            | 1.32 ± 0.39c           | 0.74 ± 0.29d           | None                        |
| AM         | 63.8 ± 4.62a  | 1.07 ± 0.08ab        | 767.67 ± 107.50abc | 30.68 ± 3.88ab         | 8.05 ± 0.51bc          | 4.11 ± 0.70ab          | 2.92 ± 0.39abc         | 70.72                       |
| LD         | 39.97 ± 4.20b | 0.56 ± 0.02d         | 1764 ± 27.44d   | 6.38 ± 1.90c           | 1.26 ± 0.43e           | 1.06 ± 0.31c           | 0.72 ± 0.12d           | −16.16                      |
| MD         | 45 ± 4.50b    | 0.75 ± 0.05c         | 283 ± 27.99d    | 10.31 ± 1.94c          | 3.94 ± 0.82d           | 1.99 ± 0.30bc          | 1.83 ± 0.56c           | 46.07                       |
| HD         | 59.1 ± 4.76a  | 0.99 ± 0.01b         | 644.19 ± 63.73bc| 23.71 ± 4.05b          | 6.22 ± 0.13c           | 3.86 ± 0.33b           | 2.63 ± 0.30b           | 68.28                       |
| AM+LD      | 61.33 ± 5.69a | 1.06 ± 0.07ab        | 635.04 ± 52.19c | 25.89 ± 2.35b          | 8.92 ± 0.85ab          | 4.11 ± 0.98ab          | 3.38 ± 0.63ab          | 72.50                       |
| AM+MD      | 64.37 ± 2.76a | 1.08 ± 0.03ab        | 789.88 ± 8.98ab | 28.06 ± 1.19ab         | 10.67 ± 0.31a          | 3.81 ± 0.42b           | 3.65 ± 0.50ab          | 72.37                       |
| AM+HD      | 65.63 ± 5.28a | 1.16 ± 0.11a         | 873.51 ± 45.56a | 35.38 ± 4.97a          | 10.78 ± 0.98a          | 6.12 ± 1.59a           | 4.15 ± 0.79a           | 79.94                       |

CK, treatments with sterilized DSE and AMF; AM, sterilized DSE with AMF; LD, low concentration of DSE with sterilized AMF; MD, medium concentration of DSE with sterilized AMF; HD, high concentration of DSE with sterilized AMF. AM + LD → AM + HD, different concentration of DSE and AMF. Data followed by different letters in the same column are significantly different at *P* < 0.05.
with separate inoculants and the uninoculated control. Shoot fresh/dried biomass in treatment AM + HD were significantly higher than in other treatments ($P < 0.05$). Root fresh and dried biomass in all combined inoculation treatments (AM + LD, AM + MD, AM + HD) were significantly higher than in separate inoculation treatments and the control ($P < 0.05$). The maximum values overall occurred in treatment AM + HD. Moreover, the mean inoculation responsiveness in separate and combined inoculation treatments was 19.3 and 32.9%, respectively. The average positive inoculation responsiveness was 57.2% (range 46.1–79.9%) with the maximum observed in AM + HD. Interestingly, the minimum plant biomass and leaf area occurred in treatment LD and was not significantly different from the control. The inoculation responsiveness in LD was −16.2%.

Shoot total P and K concentrations were also significantly increased by AMF inoculation (Fig. 1). Plant TP and TK increased significantly with increasing DSE density with a minimum value in LD, and combined inoculation with AMF showed similar results with a maximum value in AM + HD.

**Root system morphology and fungal colonization**

Root system morphology was significantly affected by the inoculation treatments (Table 2). Separate and combined inoculation significantly increased total root length, root surface area and total root volume except in separate treatment LD ($P < 0.05$), with maximum
inoculation responsiveness observed in AM + HD (63.3, 54.3 and 70.2%, respectively). The root mean diameter in all inoculation treatments was significantly larger than in the control \( (P < 0.05) \). Interestingly, the root morphology inoculation responsiveness of LD was negative.

Roots in the inoculation treatments were highly colonized with AMF and DSE but no colonization was observed in control roots (Fig. 2). The total colonization and colonization intensity of AMF and DSE were significantly different in the different inoculation treatments. With DSE inoculation alone the maximum values occurred in treatment HD. Total colonization and colonization intensity of AMF increased in AMF + DSE combined inoculation compared with AMF inoculation alone and DSE total colonization also increased compared with DSE inoculation alone. DSE colonization intensity was higher in treatment AM + LD than in LD treatment and lower in AM + MD and AM + HD than in the corresponding separate inoculation treatments.

### Photosynthesis and leaf chlorophyll content
Photosynthesis was significantly greater with AMF and DSE inoculation. Net photosynthetic rate and transpiration rate were significantly higher in treatment AM + HD than in the other treatments (Fig. 3, \( P < 0.05 \)). The maximum and minimum intercellular \( \text{CO}_2 \) concentrations were observed in LD (502 μmol \text{CO}_2 \text{ mol}^{-1}) and AM + HD (223 μmol \text{CO}_2 \text{ mol}^{-1}). Stomatal conductance was not significantly different between the separate treatments and the combined treatments and control, with the maximum observed in treatments AM, AM + MD and AM + HD. Plant leaf chlorophyll content ranged from 21.4 to 26.1 and followed the sequence: AM + MD > AM + HD > AM + LD > AM + M D > HD > LD > CK.

### Plant endogenous hormones
Inoculation with AMF and DSE significantly affected the endogenous hormones (IAA, CTK, GA, and ABA) of the shoots and roots (Fig. 4). AMF inoculation significantly increased root CTK accumulation. DSE treatment MD significantly increased CTK and IAA accumulation in the shoots and roots and reduced ABA accumulation. Treatment HD and combined inoculation (AM + LD, AM + MD, AM + HD) significantly increased the accumulation of the four endogenous hormones. The maximum IAA, CTK and GA were observed in AM + HD and the maximum and minimum ABA were observed in HD and LD in the shoots and roots.

### Soil physicochemical properties
EEG, TG and ALP were significantly lower in the control than in the inoculated treatments (Table 3). DSE inoculation significantly increased EEG, TG and ALP by 1.26, 0.39 and 16.1%, respectively, under separate inoculation and by 2.86, 1.35 and 8.6%, respectively, in the combined inoculation treatments. The maximum EEG and TG were observed in AM + HD and the maximum AP, AK and ALP in LD. Inoculation with AMF and DSE significantly reduced the activity of soil urease.

### Correlation analysis
Pearson’s correlation analysis shows significant relationships between DSE colonization intensity, plant growth and soil variables (Fig. S1). SEM was used to quantify the relative effects of DSE colonization intensity, plant CTK, photosynthetic rate, plant biomass, root volume, plant total P, soil AP, ALP and glomalin content using the correlation coefficients \( (R \text{ values}) \). DSE infection intensity increased the accumulation of plant total P by directly increasing plant CTK content and soil glomalin, which indirectly increased plant biomass in separate

### Table 2
Effects of different treatments on root morphological traits and inoculation responsiveness of maize

| Treatments | Total Length (cm) | Inoculation responsiveness | Surf Area (cm²) | Inoculation responsiveness | Avg Diam (mm) | Inoculation responsiveness | Root Volume (cm³) | Inoculation responsiveness |
|------------|------------------|---------------------------|------------------|---------------------------|---------------|---------------------------|-------------------|---------------------------|
| CK         | 1357.08 ± 90.4 cd | 0                         | 966.68 ± 41.15b | 0                         | 1.27 ± 0.13b | 0                         | 30.27 ± 1.81c     | 0                         |
| AM         | 1932.01 ± 117.87bc | 30                        | 1680.12 ± 109.32a| 42.46                     | 1.54 ± 0.06b | 17.53                     | 35.76 ± 1.87c     | 15.35                     |
| LD         | 1072.06 ± 48.85d | −26.59                    | 923.38 ± 15.34b | −5                        | 1.51 ± 0.01b | 15.89                     | 29.22 ± 2.06c     | −3.59                     |
| MD         | 1991.39 ± 321.94b | 31.85                     | 1619.53 ± 121.46a| 40.31                     | 1.59 ± 0.07ab| 20.13                     | 35.23 ± 1.91c     | 14.08                     |
| HD         | 2610.07 ± 116.98b | 48.01                     | 1687.04 ± 113.26a| 42.7                      | 1.61 ± 0.07ab| 21.12                     | 41.21 ± 0.68c     | 26.55                     |
| AM + LD    | 2189.76 ± 182.43bc| 38.03                     | 1828.51 ± 78.34a | 47.13                     | 1.53 ± 0.07b | 16.99                     | 65.62 ± 3.8b      | 53.87                     |
| AM + MD    | 2536.56 ± 134.44b | 46.5                      | 1873.19 ± 185.75a | 48.39                     | 1.71 ± 0.15ab| 25.73                     | 78.14 ± 4.06b     | 61.26                     |
| AM + HD    | 3701.72 ± 363.28a | 63.34                     | 2114.4 ± 154.32a | 54.28                     | 2.1 ± 0.18a  | 39.52                     | 101.58 ± 10.07a   | 70.2                      |

CK, treatments with sterilized DSE and AMF; AM, sterilized DSE with AMF; LD, low concentration of DSE with sterilized AMF; MD, medium concentration of DSE with sterilized AMF. AM + LD → AM + HD, different concentration of DSE and AMF. Data followed by different letters in the same column are significantly different at \( P < 0.05 \).
inoculation treatments \( (\chi^2 = 18.329, \text{df} = 15, P = 0.246, \text{RMSEA} = 0.098, \text{GFI} = 0.87, \text{AIC} = 78.329; \text{Fig. 5A}) \). DSE infection intensity increased plant biomass indirectly by increasing plant photosynthetic rate and root volume in combined inoculation treatments \( (\chi^2 = 11.083, \text{df} = 9, P = 0.270, \text{RMSEA} = 0.099, \text{GFI} = 0.90, \text{AIC} = 65.083; \text{Fig. 5B}) \). Moreover, soil AP was negatively correlated with plant biomass and root volume in across treatments.

**Discussion**

**AMF and DSE colonization**

AMF are known to be influenced by the activities of other soil microorganisms and share ecological niches with DSE [27]. Previous extensive studies on the physiological and ecological functions of AMF or DSE and their respective inoculation effects have been widely reported [34–37]. However, understanding of the effects of both fungal groups colonizing plants together is lacking. Here, the effects of combining AMF and DSE at various concentrations on maize were investigated.

Typical AMF and DSE root structures were observed, indicating that AMF and DSE can grow together and colonize the roots simultaneously to form a combined potentially symbiotic structure. According to our experimental data, AMF total colonization and colonization intensity and LD colonization intensity were higher in AMF + DSE combined inoculated plants compared with separate AMF or LD inoculation. However, DSE colonization intensities in AM + MD and AM + HD were lower than in the corresponding separate DSE inoculations. We therefore speculate that AMF and low-density DSE colonization were promoted by combined inoculation but niche competition might exist between AM and MD or HD.

**Effects of AMF or DSE alone on plant growth**

Separate inoculation with AMF or DSE increased maize growth (plant height, ground diameter, leaf area, plant biomass) and root system morphology (total root length, root surface area, root volume) compared to the control. DSE inoculation did not significantly affect root diameter and this is consistent with the observations of Li et al. [22]. The plant growth promoting efficacy of AMF was greater than that of DSE inoculation at all three densities but the root system morphology effects of AMF were lower than those of HD. The inoculation responsiveness of LD was negative. We suggest that AMF may significantly promote above-ground plant growth more than DSE but DSE at high density had a greater effect than AMF on root development.

Similarly, DSE inoculum at high density significantly increased shoot and root endogenous hormone levels. This supports the conclusions of Liu et al. [38] who report that DSE inoculation promoted root growth by
Fig. 3 Changes in plant photosynthesis and leaf chlorophyll content. CK, treatments with sterilized DSE and AMF; AM, sterilized DSE with AMF; LD, low concentration of DSE with sterilized AMF; MD, medium concentration of DSE with sterilized AMF; HD, high concentration of DSE with sterilized AMF. AM + LD → AM + HD, different concentration of DSE and AMF. Different letters above the columns are significantly different at P < 0.05.
regulating the content and percentage of endogenous hormones to resist drought stress of the host plant. Soil available P and ALP were maximum in LD inoculation but total P in roots, stems and leaves was minimum, indicating that DSE may have released more phosphatase promoting the accumulation of soil available P and supplied the nutrient to itself rather than to maize at LD [39]. Thus, the inoculum density of DSE may be a key factor determining whether the relationship between DSE and the host is mutualistic.

Table 3 Edaphic variables under different treatments

| Treatments | AP (mg/kg) | AK (mg/kg) | TN (mg/kg) | EEG (μg/g) | TG (μg/g) | ALP (μg/g/h) | U (μg/g/h) |
|------------|------------|------------|------------|------------|------------|--------------|------------|
| CK         | 2.84 ± 0.17ab | 124.43 ± 2.87abc | 11.51 ± 0.25bc | 99.47 ± 0.29b | 626.21 ± 2.3c | 53.22 ± 0.91b | 13.22a |
| AM         | 2.94 ± 0.14ab | 114.03 ± 1.42cd | 11.27 ± 0.49c | 100.45 ± 0.95ab | 625.76 ± 1.29c | 60.07 ± 1.62ab | 12.81 ± 0.65a |
| LD         | 3.65 ± 0.2a | 132.18 ± 3.92a | 12.82 ± 0.21ab | 100.40 ± 0.21ab | 627.12 ± 0.92bc | 63.97 ± 2.32a | 10.11 ± 0.69b |
| MD         | 2.91 ± 0.16ab | 122.57 ± 1.52abc | 11.76 ± 0.27bc | 101.07 ± 0.28ab | 627.5 ± 1.94bc | 63.43 ± 2.04a | 9.13 ± 0.29bc |
| HD         | 3.14 ± 0.56ab | 112.60 ± 3.1cd | 12.43 ± 0.09abc | 100.75 ± 0.15ab | 631.35 ± 0.76abc | 62.82 ± 3.89ab | 7.98 ± 0.32bc |
| AM + LD    | 2.34 ± 0.13b | 129.32 ± 2.01ab | 12.75 ± 0.24ab | 100.99 ± 0.58ab | 630.11 ± 2.68abc | 59.37 ± 0.99ab | 7.65 ± 0.55c |
| AM + MD    | 2.53 ± 0.01ab | 117.33 ± 2.43bcd | 13.59 ± 0.41a | 103.08 ± 0.96a | 636.32 ± 2.63ab | 57.27 ± 1.85ab | 7.54 ± 0.11c |
| AM + HD    | 2.59 ± 0.42ab | 108.2 ± 0.98d | 12.25 ± 0.01abc | 103.14 ± 0.24a | 637.81 ± 1.79a | 57.98 ± 1.89ab | 7.21 ± 0.22c |

AP Available phosphorus, AK Available potassium, TN Total nitrogen, EEG Easily extractable glomalin-related soil protein, TG Total glomalin-related soil protein, ALP Activity of alkaline phosphatase, U Activity of soil urease. CK, treatments with sterilized DSE and AMF; AM, sterilized DSE with AMF; LD, low concentration of DSE with sterilized AMF; MD, medium concentration of DSE with sterilized AMF; HD, high concentration of DSE with sterilized AMF. AM + LD → AM + HD, different concentrations of DSE and AMF. Data followed by different letters in the same column are significantly different at P < 0.05.
Interactions between AMF and DSE in plant growth
Combined inoculation with AMF and DSE had significant direct effects on plant growth, root development, plant endogenous hormone levels, and soil properties relative to the control and separate inoculation. The inoculation responsiveness of AM + LD in terms of plant biomass, total root length, root surface area and root volume was greater than that of AM and LD summed but MD and HD showed the opposite trend. This suggests that combined inoculation might lead to synergistic plant growth effects at the low DSE density and to competitive effects at medium and high densities. This supports the hypothesis in AMF and DSE colonization. Nevertheless, AM + MD and AM + HD showed higher growth-promoting efficacies of combined inoculation.

Inoculation responsiveness values of AM + HD to plant growth, root development, plant shoot and root endogenous hormone (ABA, GA, CTK) levels, and EEG and TG in soil were 79.9, 56.8, 31.3, 3.5 and 1.8%, respectively. Correlation analysis shows that EEG and TG were positively correlated with plant growth and physiological indices. It is well known that the soil organic carbon (SOC) pool is an important regulator of carbon fluxes between the atmosphere and the biosphere. Two BRSP fractions (EEG and TG) have been widely reported to be correlated with SOC in previous studies as found in a range of environments [40–42]. The contribution of glomalin to the stocks of soil carbon has also been confirmed [8]. Here, in combined inoculation treatments DSE may have stimulated AMF to release EEG and TG and increase soil carbon for fungal growth. We suggest that the beneficial effects of AMF + DSE on plant growth might be explained by soil carbon accumulation and nutrient exchange with plants and the increases in phytohormone production that they promote [38].

Fig. 5 Structural equation model (SEM) showing the causal relationships among DSE colonization intensity, plant growth and soil variables. The final model fits the data well: maximum likelihood.
Conclusions
Here, we explore associations between AMF and DSE (derived from the roots of *Stipa krylovii*) colonizing maize. Inoculum of AMF and of three densities DSE led to the combined colonization of roots and formed a compound potentially symbiotic association. DSE inoculum at medium and high densities increased plant above-ground growth and root morphology and at a low density gave negative effects. The degree of DSE colonization (or the density of DSE inoculum) might be a key factor determining whether the relationship between this fungal group and its plant host is mutualistic. The combination of AMF and DSE significantly and positively influenced plant above-ground growth and root morphology. These findings support the hypothesis that AMF + DSE combined inoculation has a synergistic effect in promoting the growth of the host plant at the low DSE density and a competitive effect at medium and high DSE densities. Treatments AM + MD and AM + HD exerted the greatest effects on host plant growth and root morphology. Future studies investigating the allocation of nutrient resources between both fungal groups and plants would increase our understanding of dual interactions.

Methods
Isolation and identification of DSE
Roots of *Stipa krylovii* were surface-sterilized in 75% ethanol and 5% sodium hypochlorite for 5 min, rinsed three times in deionized water, and then transferred to potato dextrose agar (PDA) culture medium with antibiotic supplements (ampicillin and streptomycin sulfate) and incubated at 27°C [43]. The isolated DSE was identified as *Alternaria* sp. by molecular identification and deposited in the general microbiology center of China National Committee (CGMCC, address: 3, Courtyard 1, Beichen West Road, Chaoyang District, Beijing), with the preservation number CGMCC No.17463. *Diversispora epi-gaea* (formerly *Glomus versiforme*) was used as the AMF experimental material.

Preparation of AMF and DSE inocula
The DSE inoculum was prepared by aseptic growth in flasks with Modified Melin-Norkra (MMN) medium. The inoculated flasks were oscillated at 170rpm and incubated at 27°C in the dark for 2 weeks. The AMF *Diversispora epi-gaea* was provided by the Institute of Plant Nutrition and Resources, Beijing Academy of Agriculture and Forestry Sciences, and the inoculum contained spores, external mycelium and mycorrhizal root fragments with a spore density of 26g⁻¹, a colonization rate of 87%, and a hyphal length of 3.12mg⁻¹. Controls received DSE and AMF inocula sterilized by autoclaving at 121°C for 30 min.

Greenhouse experiment
Maize seeds were acquired from the Zhongnongzuo Technology Development Co. Ltd. of Beijing and surface-sterilized in 70% (v/v) ethanol for 5 min and in 10% (v/v) NaClO for 10 min, rinsed several times with deionized water and placed in a culture dish with moist sterile filter paper in the dark at 25°C for 3 days. Three maize seeds were sown per pot and one plant was retained at the three-leaf stage. Plants were inoculated with *Alternaria* sp. or *D. epigaea* or inoculated combined (AMF inoculation × 3 DSE densities) with four replicates of each treatment. The AM treatment consisted of 50g *D. epigaea* (1300 spores) and sterilized DSE per pot. The three DSE densities were low (LD, 2 × 10^5 CFU mL⁻¹), medium (MD, 4 × 10^5 CFU mL⁻¹) and high (HD, 8 × 10^5 CFU mL⁻¹) with sterilized AMF, respectively. Combined inoculations of AMF + DSE consisted of AM (50g) + LD (50mL), AM (50g) + MD (50mL), or AM (50g) + HD (50mL). A control consisted of equal amounts of sterilized AMF and DSE medium. A total of 32 pots (8 treatments × 4 replicates) were grown for 60 days before harvest.

Plant measurement and fungal infection
Plant height and ground diameter were recorded every 10 days. Gas exchange was measured using a portable photosynthesis system (Li-6400; Li-cor Inc., Lincoln, NE), comprising photosynthesis (Pₚ), stomatal conductance (Gₛ), transpiration rate (Tₛ) and intercellular CO₂ concentration. The relative chlorophyll content of leaves was measured with a SPAD-502 leaf chlorophyll meter (Minolta, Osaka, Japan). Plant roots and shoots were harvested and weighed fresh and dry (60°C, 48h), and oven-dried samples were used to determine plant nutrient (phosphorus and potassium) contents and the phytohormone gibberellic acid (GA), the hormones abscisic acid (ABA), indole-3-acetic acid (IAA), and cytokinin (CTK) were determined after 60 days. Fine roots selected from the soil were collected to determine colonization by DSE and AMF. The endogenous hormones (GA, CTK, IAA, and ABA) and fungal colonization were measured according to the methods of Bi et al. [9]. Leaf area was determined by plotting the shape of leaves on cardboard and calculating the leaf area through a square grid before drying. The contribution of fungi to biomass was evaluated in terms of inoculation responsiveness, calculated as (total biomass of inoculated maize — total biomass of non-inoculated maize) / total biomass of inoculated maize × 100%.

The general inoculation effect was evaluated by fungal total colonization and colonization intensity in the root system [44]. Each treatment was examined using the glass
slide in which 30 randomly-selected 0.5-cm-long root segments were cleared with 10% (w/v) potassium hydroxide and stained with 0.5% (w/v) acid fuchsin [45]. Fungal total infection (%) is expressed as the percentage of infected fine root segments in each root sample: infection intensity (%) = (infected length of root segments / total length of infected root segments) × 100. Root system morphology, comprising total root length, number of root tips, root surface area, and average root diameter and volume, were evaluated using RootSnap software (CID Bio-Science, Camas WA).

**Soil properties**

Soil available phosphorus (AP), available potassium (AK), total phosphorus (TP), and total potassium (TK) were determined by inductively coupled–plasma emission spectroscopy (ICP-OES, Optima 5300DV, Perkin Elmer, Waltham, MA). The total nitrogen (TN) contents were determined by the Kjeldahl method [46]. Soil phosphatase was determined by the method of Tarafdar and Marschner [47] and soil urease activity by that of Hoffmann and Teicher [48]. Soil glomalin was quantified as glomalin-related soil protein (GRSP). Easily extractable BRSPs (EE-BRSPs, EEG) and total BRSPs (T-BRSPs, TG) were determined by the methods of Wright and Upadhyaya and Janos et al. [49, 50] using bovine serum albumin as the standard to determine the GRSP concentration in the extracts by Bradford assay.

**Statistical analysis**

The effects of the treatments on the measured variables were evaluated by one-way analysis of variance (P < 0.05) and differences between mean values by Tukey’s multiple-range test (P < 0.05) using the SPSS v. 21.0 for Windows software package (IBM Corp., Armonk, NY). Pearson’s correlation analysis and the structural equation model (SEM) were used to test the effects of DSE on plant growth using the SPSS software package (version 21.0, Amos Development Corp., Meadville, PA).

**Abbreviations**

AMF: Arbuscular mycorrhizal fungi; DSE: Dark septate endophytes; PDA: Potato dextrose agar; MMN: Modified Melin-Norkra; Pn: Photosynthesis; Gs: Stomatal conductance; Tr: Transpiration rate; IAA: Indole-3-acetic acid; CTK: Cytokinin; ABA: Hormone abscisic acid; GA: Gibberellic acid; AP: Available phosphorus; AK: Available potassium; TN: Total nitrogen; TP: Total phosphorus; TK: Total potassium; EEG: Easily extractable glomalin-related soil protein; EE-BRSPs: Easily extractable BRSPs; TG: Total glomalin-related soil protein; ALP: Activity of alkaline phosphatase; U: Activity of soil urease; CK: Treatments with sterilized DSE and AMF; AM: Sterilized DSE with AMF; LD: Low concentration of DSE with sterilized AMF; MD: Medium concentration of DSE with sterilized AMF; HD: High concentration of DSE with sterilized AMF; AM+LD: Low concentration of DSE with AMF; AM+MD: Medium concentration of DSE with AMF; AM+HD: High concentration of DSE with AMF.

**Supplementary Information**

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**Authors’ contributions**

Data curation, LX and YB; Formal analysis, LX; Funding acquisition, YB; Investigation, LX, SM and QH; Methodology, LX and YB; Supervision, YB; Visualization, LX; Writing and language editing, LX, JS and PC. All authors have read and approved the manuscript.

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**Availability of data and materials**

All data generated or analyzed during this study are included in this manuscript and its supplementary information files, and the datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Supplementary Information**

Additional file 1.

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