Role of abscisic acid in strigolactone-induced salt stress tolerance in arbuscular mycorrhizal *Sesbania cannabina* seedlings

Cheng-Gang Ren, Cun-Cui Kong and Zhi-Hong Xie*

**Abstract**

**Background:** Strigolactones (SLs) are considered to be a novel class of phytohormone involved in plant defense responses. Currently, their relationships with other plant hormones, such as abscisic acid (ABA), during responses to salinity stress are largely unknown.

**Results:** In this study, the relationship between SL and ABA during the induction of H$_2$O$_2$–mediated tolerance to salt stress were studied in arbuscular mycorrhizal (AM) *Sesbania cannabina* seedlings. The SL levels increased after ABA treatments and decreased when ABA biosynthesis was inhibited in AM plants. Additionally, the expression levels of SL-biosynthesis genes in AM plants increased following treatments with exogenous ABA and H$_2$O$_2$. Furthermore, ABA-induced SL production was blocked by a pre-treatment with dimethylthiourea, which scavenges H$_2$O$_2$. In contrast, ABA production was unaffected by dimethylthiourea. Abscisic acid induced only partial and transient increases in the salt tolerance of TIS108 (a SL synthesis inhibitor) treated AM plants, whereas SL induced considerable and prolonged increases in salt tolerance after a pre-treatment with tungstate.

**Conclusions:** These results strongly suggest that ABA is regulating the induction of salt tolerance by SL in AM *S. cannabina* seedlings.

**Keywords:** Strigolactones, Abscisic acid · arbuscular mycorrhizal, *Sesbania cannabina*, Photosynthesis, Salt stress

**Background**

Saline-alkali stress is a serious ecological problem that negatively impacts plant survival, development, and productivity [1]. To survive such stress, plants have established beneficial associations with several microorganisms present in the rhizosphere that can alleviate the stress symptoms [2]. One of the most intensively studied and widespread mutualistic plant–microorganism associations involves arbuscular mycorrhizal (AM) fungi. About 80% of terrestrial plants, including most leguminous species, are able to establish this type of symbiotic relationship with fungi of the division Glomeromycota [3]. Indeed, AM symbiosis increases resistance to soil salinity in diverse host plants, such as maize, tomato, and lettuce [4], although the underlying mechanisms are not well characterized [5]. *Sesbania cannabina*, which is a soil-improving legume, is highly adaptable to different adverse climatic conditions, including salinity, drought, and waterlogging stresses. Therefore, developing *S. cannabina*–AM fungi symbiotic relationships might represent a good strategy for improving the fertility of saline soils to enable the growth of agriculturally important crops [6].

Establishing a functioning symbiotic relationship with an AM fungus requires a precise coordination between the partners based on a finely regulated molecular dialogue [7]. The molecular dialogue occurring during the so-called pre-symbiotic stage starts when the host plant produces and exudes strigolactones (SLs) into the rhizosphere. Strigolactones are recognized by AM fungi with an uncharacterized receptor that stimulates hyphal growth and branching, thereby increasing the probability of encountering the host roots [8]. In accordance with their role as signaling molecules in the rhizosphere, SLs are...
mainly produced in the roots and have been detected in the root extracts of monocotyledonous and dicotyledonous plants [9]. Strigolactones are biosynthetically derived from carotenoids [10] via a sequential oxidative cleavage by two carotenoid cleavage dioxygenases (CCD7 and CCD8), and are apocarotenoids [11], as is abscisic acid (ABA). Once synthesized, SLs are transported acropetally to the shoot or are exuded into the rhizosphere by the ABC transporter PDR1, which was first identified in Petunia hybrida [12]. The F-box protein MAX2 and the α/β-fold hydrolase D14/DAD2 are the main candidate components of the SL-perception complex in higher plants [13, 14]. Additionally, MAX2 has diverse roles, including in plant responses to abiotic stress [15]. The importance of SLs during the initial stages of mycorrhizal colonizations is widely recognized. Moreover, emerging evidence indicates that SLs may also affect the subsequent steps of the symbiotic relationship developing in response to environmental stimuli, such as salt and drought stresses [4, 16]. We recently observed that SLs can induce salt stress tolerance in AM S. cannabina seedlings that is regulated by the generation of H2O2 as a signaling molecule [17].

Abscisic acid is an important plant hormone with a critical role in the regulation of salt stress responses [18]. Exposure to stresses, such as salinity, induces the accumulation of ABA, resulting in increased tolerance [19]. Several lines of evidence have revealed that ABA induces the accumulation of H2O2, which plays an important role in ABA signaling [20]. Moreover, SLs and ABA are critical for the regulation of salt stress responses and the establishment of symbiotic relationships between host plants and AM fungi [4, 21]. Recent studies unveiled the links between ABA and SLs in a number of physiological processes. Exogenous ABA may enhance the accumulation of SLs, especially under stress conditions [22]. Additional studies have demonstrated that SL metabolism and its effects on ABA levels are seemingly opposite in roots and shoots under stress [23]. However, it is unclear whether SL and ABA are involved in a synergistic effect during stress responses. These results suggest the possible relationship between SL and ABA during the induction of plant stress tolerance is complex.

The mechanisms by which SLs enhance plant stress tolerance have so far been largely uncharacterized. Our previous studies confirmed that H2O2 induces SL production and is critical for SL-induced salt stress tolerance in AM S. cannabina seedlings. In the present study, we inoculated S. cannabina seedlings with AM fungi and monitored the H2O2 level, ABA content, and SL accumulation to examine the role of ABA and the relationship between ABA and H2O2 in the AM fungus-induced increase in SL production in response to salt stress conditions.

**Results**

**Effects of ABA levels on strigolactone accumulation**

In S. cannabina seedlings inoculated with an AM fungus, the root colonization rate steadily increased as seedlings grew and differed significantly between sampling times (Table 1). Additionally, the ABA concentration was the highest in salt-treated S. cannabina seedlings and it was higher in salt-treated AM S. cannabina seedlings than in the water-treated controls (Fig. 1), indicating that salt stress may have triggered ABA biosynthesis both in AM and non-AM S. cannabina seedlings. Moreover, a bioassay of SL showed that the P. ramosa germination rate increased significantly in response to the root extract of salt-stressed AM S. cannabina seedlings [17], as did the endogenous SL levels (m/z 383, m/z356, and m/z 317) (Additional file1: Table S1).

To determine whether ABA affects SL generation, we examined the changes in SL levels in AM S. cannabina seedlings after ABA and tungstate treatments. As shown in Fig. 4b, ABA did not accumulate in seedlings treated with tungstate, which prevents the formation of ABA from ABA aldehyde by inhibiting ABA-aldehyde oxidase [24]. First, the ABA treatment increased SL levels in AM seedlings but not in non-AM seedlings. Second, the germination bioassay revealed that SL accumulated in salt-stressed AM S. cannabina seedlings 10 days after the ABA treatment. However, this ABA-induced increase was not observed in seedlings pre-treated with 5 μM TIS108, a SL synthesis inhibitor (Fig. 2). These results suggested that ABA was important for the AM fungus-induced SL synthesis under salt stress conditions. In contrast, when S. cannabina seedlings were treated with an ABA synthesis inhibitor tungstate, the SL content decreased significantly, but the SL level could be recovered by an ABA co-treatment. It also showed that H2O2 could affect SL accumulation in AM S. cannabina seedlings (Fig. 2). Considered together, these results suggested that ABA induced the accumulation of SLs in AM S. cannabina seedlings.

**Abscisic acid affects the expression of strigolactone biosynthesis/signaling genes in S. cannabina seedlings**

To analyze the molecular mechanisms underlying the induction of SL synthesis in AM S. cannabina seedlings

| Time     | 3 WAS | 5 WAS | 7 WAS |
|----------|-------|-------|-------|
| AMF colonization | 8.6 ± 0.54c | 26.3 ± 3.1b | 45.8 ± 3.73a |

Table 1 Colonization of Sesbania cannabina seedling roots by Funneliformis mosseae. Values are presented as the mean ± standard error of triplicate samples. Data were analyzed using Duncan’s multiple range test at p < 0.05. WAS: weeks after sowing.
under salt stress, we examined the effects of ABA and H$_2$O$_2$ on the expression of CCD7, CCD8, and MAX2 homologs in S. cannabina seedlings. The expression levels of two SL-biosynthesis genes, CCD7 and CCD8, were up-regulated by NaCl, exogenous ABA, and H$_2$O$_2$ (Fig. 3). Furthermore, tungstate and DMTU, an H$_2$O$_2$ scavenger, suppressed the expression of CCD7 and CCD8 in salt-stressed AM S. cannabina roots. Additionally, seedlings pre-treated with TIS108, which blocks SL biosynthesis, exhibited up-regulated expression of CCD7 and CCD8. We also observed that salt stress, exogenous ABA, and H$_2$O$_2$ enhanced the expression of the SL-signaling gene, MAX2, in AM S. cannabina shoots. It is worthy to note that CCD7 and CCD8 mainly expressed in root tissue, while MAX2 in shoot tissue.

Interactions between ABA and H$_2$O$_2$ related to strigolactone-induced salt tolerance

An earlier study concluded that elevated H$_2$O$_2$ levels are involved in SL-induced salt stress tolerance [17]. Furthermore, several studies have revealed that ABA can induce the increased production of H$_2$O$_2$ (Kwak et al. [20]). Therefore, determining whether ABA helps to increase H$_2$O$_2$ production in AM S. cannabina seedlings is warranted. To test this possibility, we pre-treated S. cannabina seedlings with tungstate, which prevented the accumulation of ABA and considerably decreased H$_2$O$_2$ levels in AM S. cannabina seedlings (Fig. 4). Additionally, ABA-induced SL production was inhibited by the pre-treatment with DMTU (Fig. 2). These results implied that an increase in ABA content is required for H$_2$O$_2$ to induce SL production. In contrast, when we pre-treated seedlings with DMTU, we observed no significant effect on the increase in ABA content under salt stress conditions (Fig. 4b). Thus, ABA appears to function upstream of H$_2$O$_2$ in AM S. cannabina seedlings.

Dependency of ABA biosynthesis for strigolactone-induced salt tolerance

To study the relationship between ABA- and SL-induced salt tolerances, we analyzed the effects of TIS108, a specific inhibitor of SL biosynthesis, and tungstate, an ABA biosynthesis inhibitor, on SL- and ABA-induced salt tolerances in AM S. cannabina seedlings. All plants not treated with salt had similar ΦPSII values (approximately 0.57), while those exposed to 200 mM NaCl had significantly decreased ΦPSII values at 2 and 8 days, especially in seedlings treated with TIS108 and tungstate (Fig. 5a, c). These results suggested that a defect in either ABA or SL accumulation decreased the salt stress tolerance of the seedlings. A pre-treatment with ABA or GR24 (synthetic SL analog) at 2 or 8 days prior to the exposure to salt stress significantly increased ΦPSII values at both control and AM plants. Moreover, GR24 restored the salt tolerance of tungstate-treated plants, while ABA restored the salt tolerance of TIS108-treated plants at 2 days after its application, but was ineffective at 8 days after its application (Fig. 5). Thus, SL was able to rescue the salt stress tolerance of ABA-deficient plants, while...
ABA could only partially and transiently rescue the defective stress tolerance of the tungstate-treated plants. This observation implied that SL might function downstream of ABA in plant stress responses. Furthermore, ABA-induced salt tolerance was effectively blocked by a pre-treatment with DMTU (Fig. 6), while SL-induced salt tolerance was regulated by DMTU [17]. These results suggested that H$_2$O$_2$ was also responsible for the observed ABA- and SL-induced salt tolerances.

Discussion

Plants are sessile in nature. Therefore, to counter biotic and abiotic stresses, such as pathogens, drought, waterlogging, extreme temperatures, salinity, and many others, plants have evolved intricate signaling mechanisms composed of multiple components including plant hormones. Research conducted in the last decade have added SLs to the growing list of plant hormones involved in responses to environmental stresses [25]. In the present study, we revealed that SLs can be regulated by ABA production in $S$. cannabina seedlings (Figs 1, 2).

Given the common metabolic precursor shared by ABA and SL, a potential crosstalk between these two hormones was recently proposed, although these hormones appear to be negatively correlated. For example, SL levels reportedly decreased, while ABA content increased in tomato and lettuce plants grown under drought conditions in the absence of AM symbiosis [16]. We also observed that ABA levels increased rapidly under salt stress conditions in plants not colonized by an AM fungus, whereas SL production was very low (Fig. 1). Therefore, this negative correlation might represent a general plant strategy to cope with water-related stresses in the absence of mycorrhization. Conversely, a positive correlation between ABA and SLs has been observed in stress-treated mycorrhizal plants [4, 16]. Similarly, increasing ABA and SL levels were detected in $S$. cannabina seedlings under salt stress conditions in the presence of the AM fungus (Fig. 1). Additionally, ABA enhanced $P$. ramosa germination, whereas tungstate had the opposite effect (Fig. 2). The different relationships between SL and ABA in AM and non-AM $S$. cannabina respectively may indicate that the ABA catabolism was

Fig. 3 Relative expression levels of SL-biosynthesis/signaling genes in $Sesbania$ cannabina seedlings after a 10-day treatment. a Relative expression levels of CCD7. b Relative expression levels of CCD8. c Relative expression levels of MAX2. The expression level of each gene was normalized against that of the $S$. cannabina tubulin gene [41] and is provided relative to the control expression level (= 1). Data are presented as the mean ± standard deviation of three biological replicates. Data were analyzed using Duncan’s multiple range test. Different letters above the error bars indicate statistical significance at $p < 0.05$.
intensively altered by AM inoculation. Overall, it seems that the increase in ABA concentration induced by the AM fungus contributed to the increased accumulation of SLs in salt-stressed *S. cannabina* seedlings.

Our previous results suggested that H$_2$O$_2$-induced SL production is due to induced/activated NADPH oxidase and is important for SL-induced salt stress tolerance. In this study, we confirmed that ABA has a key role in the production of H$_2$O$_2$, which regulates the accumulation of SLs. We revealed that H$_2$O$_2$ production was substantially decreased by a pre-treatment with tungstate (Fig. 4a). Moreover, ABA accumulation was unaffected by a pre-treatment with DMTU (Fig. 4b). Previous investigations also concluded that ABA causes H$_2$O$_2$ to accumulate in the apoplast, which is dependent on NADPH oxidase and is important for ABA signaling [20, 26]. These results suggest that ABA may act upstream of H$_2$O$_2$ in the signaling pathway of AM *S. cannabina* seedlings. Furthermore, we provided evidence that H$_2$O$_2$ is important for ABA-induced SL production. We also observed that SL production was considerably decreased by a pre-treatment with tungstate or DMTU (Fig. 2). An ABA treatment partly restored the *P. ramosa* germination rate in DMTU-treated mycorrhizal plantlets. Simultaneously, an H$_2$O$_2$ treatment significantly restored the *P. ramosa* germination rate of tungstate-treated mycorrhizal plantlets (Fig. 2). Taken together, these results implied that ABA induced SL accumulation in AM *S. cannabina* seedlings, which may be partially dependent on increased H$_2$O$_2$ levels.

In this study, we confirmed that ABA is involved in the SL-induced salt tolerance of *S. cannabina* seedlings inoculated with an AM fungus. Exposure to tungstate decreased SL levels and ΦPSII values in salt-stressed AM *S. cannabina* seedlings (Figs 2, 5a, c). Furthermore, an ABA treatment significantly restored the *P. ramosa* germination rate in TIS108-treated mycorrhizal plantlets, indicating that the ABA-induced salt stress tolerance of AM *S. cannabina* seedlings required the accumulation of SL (Fig. 2). The relationship between SL and ABA during plant responses to abiotic stresses has been rarely studied. Many investigations have considered ABA as the ‘abiotic stress hormone’ because its biosynthesis is rapidly induced by exposures to environmental stresses, especially water-related stresses such as drought and salinity [27–29]. The reported increase in the ABA content of AM plants subjected to abiotic stress likely contributes to the enhanced tolerance of these plants to environmental stresses [4, 30]. Additionally, increased SL levels were detected in lettuce plants grown in the presence of the AM fungus *Rhizophagus irregularis* under salt stress conditions [4]. A similar trend has been observed in drought-treated lettuce and tomato plants [16]. These observations combined with our results indicate that SL-induced salt stress tolerance of AM plants may be mediated by a complex set of signal transduction pathways with ABA as a common signaling molecule.

**Conclusions**

In summary, we uncovered a dynamic interplay between SLs, ABA, and H$_2$O$_2$ produced in response to an AM fungus that contributed to the salt stress tolerance of *S. cannabina* seedlings. Following the perception of the ABA signal, H$_2$O$_2$ was rapidly produced, which then increased the accumulation of SLs, and ultimately enhanced the salt stress tolerance of *S. cannabina* seedlings. This study provides new insights into the complex interplay between ABA and SLs in AM symbiosis under abiotic stress conditions.
stress tolerance of the seedlings. Additional studies are needed to provide the genetic evidence of the involvement of ABA in H₂O₂-induced SL generation and to identify the critical signaling components between SL production and salt stress responses in *S. cannabina* seedlings inoculated with an AM fungus. Such studies will help to elucidate the molecular mechanism underlying the ABA- and SL-induced salt tolerance of AM plants.

**Methods**

**Plant materials and treatments**

*Sesbania cannabina* (Retz.) Pers. seeds were obtained from the Shandong Academy of Agricultural Sciences, Shandong, China. Before sowing, the seeds were surface-sterilized in 5% sodium hypochlorite for 5 min and rinsed several times with distilled water. The seeds were germinated at 28 °C in distilled water and then sown in trays containing autoclaved zonolite. After 2 weeks, individual seedlings were transferred to 1-L pots containing autoclaved zonolite inoculated with 10 g inoculum (approximately 121 spores). The original inoculum [AM fungus *Funneliformis mosseae*] was propagated in pot-cultured *Trifolium repens* plants for 8 weeks, and included infected roots, hyphae, spores, and substrates. The growth conditions were as follows: 12 h photoperiod, 25/17 °C (day/night), and light intensity of 600 μmol m⁻² s⁻¹.

Three-week-old seedlings were used for all treatments with 200 mM NaCl solutions. The reagents used as figures.
specific scavengers or inhibitors [5 mM DMTU (H₂O₂ scavenger) [31], 5 mM sodium tungstate [32], and 2 μM TIS108 (the most potent and specific SL biosynthesis inhibitor) [33]] were purchased from Sigma-Aldrich (St Louis, MO, USA). Additionally, 10 mM H₂O₂, 100 μM ABA, and 1 μM GR24 (synthetic SL analog) were used as exogenous signaling molecules [34]. Plantlet leaves were sprayed with 100 μL exogenous signaling molecule or inhibitor solution. Treatments were completely randomized and replicated three times. An equal volume of distilled water was applied as the control treatment. Unless stated otherwise, inhibitors were applied 24 h before the exogenous signaling molecules were applied.

Photosynthetic parameters
Gas exchange and modulated chlorophyll fluorescence parameters were simultaneously detected using a LI-6400XTR Portable Photosynthesis System (Li-Cor, Lincoln, NE, USA) equipped with a 6400–40 Leaf Chamber Fluorometer (Li-Cor). The leaves were incubated in darkness for 20 min before being analyzed. The minimal fluorescence level of the dark-adapted leaves was measured using a modulated pulse (< 0.05 μmol m⁻² s⁻¹ for 1.8 s), while the maximal fluorescence level was measured after applying a saturating actinic light pulse of 8000 μmol m⁻² s⁻¹ for 0.7 s. The actinic light intensity was increased to 1000 μmol m⁻² s⁻¹ and then maintained for about 30 min. The steady-state fluorescence yield was also recorded. A saturating actinic light pulse of 8000 μmol m⁻² s⁻¹ for 0.7 s was then applied to induce the maximum fluorescence yield by temporarily inhibiting photosystem II (PSII) photochemical activities. The minimum steady-state fluorescence yield was determined during a brief interruption of the actinic light irradiation in the presence of far-red light (λ = 740 nm). Finally, PSII efficiency (ΦPSII) and non-photochemical quenching of chlorophyll fluorescence were calculated according to the method described by Maxwell and Johnson [35].

Measurement of arbuscular mycorrhizal fungal colonization
The percentage of roots colonized by mycorrhizal fungi was calculated using the gridline intersection method [36] after samples were stained with trypan blue [37].

Measurement of H₂O₂
Seedlings were harvested 10 days after treatments to measure the H₂O₂ content. The H₂O₂ concentration was determined by monitoring the absorbance of titanium peroxide at 415 nm following the method of Brennan and Frenkel [38]. One unit of H₂O₂ was defined as the chemiluminescence produced by the internal standard of 1 μM H₂O₂ g⁻¹ fresh weight.

Extraction of ABA and HPLC analysis
Seedlings collected for the ABA analysis were immediately frozen with liquid N₂ and stored at −80 °C. Root tissue (0.5 g) was ground in a mortar and homogenized in 5 mL pre-cooled 80% aqueous acetone (4:1, v/v) supplemented with 10 mg/L butylated hydroxytoluene. Extracts were eluted through a Sep Pak C18-cartridge (PerkinElmer, Waltham, MA, USA) to remove polar compounds and then purified using ethyl acetate and NaHCO₃ as described by Sweetser and Vatvars [39]. After centrifuging samples at 14,000 g for 30 min at 4 °C, the supernatant was acidified to pH 3.0 with H₂SO₄ and then treated three times with an equal volume of ethyl acetate. The extract phase was evaporated at 40 °C.
using the RE-100 rotary evaporator (Bibby Sterlin Ltd., Stone Staffordshire, UK). The dried samples were re-dissolved in chromatography-grade acetonitrile with 0.1 M acetic acid and then analyzed by HPLC using a Hypersil-ODS C18 column (4.6 mm × 250 mm, 5 μm). The mobile phase comprised acetonitrile: 0.03 M acetic acid (30:70, v/v) at a flow rate of 1 ml/min. Abscisic acid was detected spectrophotometrically at 254 nm at 25 °C.

The nucleotide sequence reported in this table has been submitted to GenBank with accession numbers MF683952, MF683953 and MF683954.

Quantitative real-time PCR
Root samples from three seedlings were pooled to represent a single biological replicate. Total RNA was extracted from the pooled root samples using TRizol (Invitrogen, Carlsbad, CA, USA) following the manufacturer’s instructions. Three biological replicates were prepared. The extracted RNA was used as the template for reverse transcription reactions completed with the GoScript™ Reverse Transcription System (Promega, Madison, WI, USA). Gene-specific primers were designed using the Primer Premier program (version 5.0) (Premier Biosoft International). The quantitative real-time (qRT)-PCR analysis was conducted with an iCycler iQ Real-time PCR Detection System (Bio-Rad) with three technical replicates. Each reaction was completed in a total volume of 20 μl, which included 0.2 μM primer pairs, 2 μl diluted cDNA, and 10 μl 2× SYBR Green PCR Master Mix (TaKaRa Bio Inc., Dalian, China). The PCR program was as follows: 95 °C for 30 s; 40 cycles of 95 °C for 5 s, 58 °C for 15 s, and 72 °C for 20 s. Tubulin genes were used as the internal controls and relative gene expression levels were calculated using the 2^\(-\Delta\Delta C_t\) method [41]. The sequences of the qRT-PCR primers are listed in Table 2.

Statistical analysis
All data were analyzed using Microsoft Excel (Redmond, WA, USA) and the values for each treatment are presented as the mean ± standard deviation of three replicates. A one-way analysis of variance was completed with the SPSS Statistics 17.0 software (SPSS, Inc., Chicago, IL, USA). Duncan’s multiple range test was used to compare mean values at the p < 0.05 significance level.

Additional file

Additional file 1: Table S1 Effect of different treatments on the relative abundances of three SLs candidates m/z 383, m/z 356 and m/z 317 in S. cannabina seedlings extract. Data were analyzed using Duncan’s multiple range test. Different letters indicate statistical significance at p < 0.05. (XLSX 13 kb)
Abbreviations
ABA: Abscisic acid; AM: Arbuscular mycorrhizal; CCD: Carotenoid cleavage dioxygenases; DMTU: Dimethylthiourea; m/z: Mass to charge ratio;
MAX2: More auxillary growth2; NPK: Non-photochemical quenching of chlorophyll fluorescence; SL: Strigolactone; TIS108: 6-phenoxy-1-phenyl-2-(1H-1,2,4-triazol-1-y) hexan-1-one; QBSII: Photosystem II efficiency

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Availability of data and materials
All data generated or analysed during this study are included in this published article [and its Additional file].

Authors’ contributions
RCG designed and performed the experiment, analysed the data and wrote the manuscript. KCC perform extraction and identification the phytohormone using GC-MS. RCG and KCC participated in revisions of the manuscript. All authors have read the manuscript, given comments and approved the final version of the manuscript.

Ethics approval and consent to participate
Not applicable

Competing interests
The authors declare that they have no competing interests.

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