Exogenous gibberellin can effectively and rapidly break intermediate physiological dormancy of *Amsonia elliptica* seeds

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Accelerated global warming is leading to the loss of plant species diversity, and *ex situ* preservation of seeds is becoming an increasingly important aspect of species conservation. However, information on dormancy and germination is lacking in many endangered species. *Amsonia elliptica* (Apocynaceae) is the only *Amsonia* species native to Korea, and the South Korean Ministry of Environment has designated it Class II endangered wildlife. Nevertheless, the dormancy class and the dormancy breaking method for seeds of this species for germination are not precisely known. We identified the structure of *A. elliptica* seeds and the causes of dormancy, which inhibits germination. In addition, we tried to develop an effective germination promotion method by testing the wet stratified condition, which breaks dormancy, and the form of gibberellin that can replace it. Fresh seeds of *A. elliptica* imbibe water, but the covering layers (endosperm and seed coat) inhibit germination by mechanically restricting the embryo. Initial germination tests confirmed low embryo growth potential and physiological dormancy (PD). Restriction due to the covering layer was eliminated by seed scarification, and abnormal germination was observed. After 12 weeks of cold moist stratification at 4°C, only 12% of seeds germinated. However, 68.8% of seeds subjected to 8 weeks of warm moist stratification followed by 12 weeks of cold stratification germinated, indicating that warm stratification pretreatment before cold stratification is effective in breaking dormancy. *A. elliptica* seeds exhibited intermediate PD. Furthermore, 61.3% of seeds soaked in 500 mg/L GA₄+₇ for 14 days and incubated at 25/15°C germinated. Therefore, GA₄+₇ rapidly broke the
dormancy of *A. elliptica* seeds compared with warm plus cold stratification treatment, thus providing an efficient method for seedling production.

**KEYWORDS**

*Amsonia elliptica*, physiological dormancy, seed scarification, dormancy break, moist stratification

**Introduction**

Ecosystems are rapidly changing owing to various environmental problems and global warming, and one of the most important issues is the loss of plant species diversity (Walther et al., 2002). Various *in situ* and *ex situ* efforts have been made worldwide to conserve plant diversity (Hawkins et al., 2008; Li and Pritchard, 2009). A common *ex situ* conservation strategy for plant species involves storing seeds in seed banks, which allows several genetic resources to be protected in a relatively small space (Cochrane et al., 2007; Kramer and Havens, 2009; Hay and Probert, 2013). However, collecting and storing seeds does not necessarily solve the conservation issue because the seeds must eventually germinate if they are to be used for restoration. For many species, however, we do not have information about their dormancy-breaking and germination requirements, which are essential for plant propagation from seeds. Therefore, understanding the dormancy and germination characteristics of seeds is as important for restoration as collecting and storing a diverse and large number of species.

Seed dormancy means germination does not occur even if environmental conditions are suitable for germinating non-dormant seeds (Hilhorst, 1995; Baskin and Baskin, 2014). Dormancy may be due to characteristics (blocks) of the embryo (e.g., underdevelopment and/or a physiological inhibiting mechanism) and/or the seed/fruit coat, and these blocks must be removed or overcome before germination can occur (Baskin and Baskin, 2014). To break dormancy, seeds must be exposed to certain environmental conditions, which vary depending on the dormancy type (class). There are five classes of dormancy, which may be subdivided into levels, e.g., physiological dormancy (PD) is divided into three levels (Nikolaeva, 2004; Baskin and Baskin, 2021). Therefore, knowledge of the dormancy classes and how they can be overcome is useful for plant propagation and utilization and is especially essential for species conservation.

*Amsonia* is a genus in the Apocynaceae family that is widely distributed from the southern United States to northwestern Mexico and eastern Asia. *Amsonia elliptica* is the only *Amsonia* species in Korea, and it is a plant with great ornamental and medicinal potential (Sauerwein et al., 1991; Kim et al., 2006; Yang et al., 2012). However, *A. elliptica* has been designated Class II endangered wildlife by the South Korean Ministry of Environment because of its rarity in nature and the loss and destruction of its habitat. Therefore, information on the seed dormancy break and germination requirements of this species is important for its conservation and restoration.

Germination has been reported to improve when seeds of *Amsonia tabernaemontana* are scarified and then sown (Martin et al., 2013). In addition, according to a patent published in Korea (No. 10-2017-007854), the germination of *A. elliptica* is improved by chemical scarification (Choi and Jung, 2017). Thus, based on the available information, we can conclude that the seed coat of *Amsonia* species inhibits germination. Seed coat inhibition of germination could be due to the water impermeability of the seed coat, i.e., physical dormancy (PY), or to the low growth potential of the embryo, i.e., PD.

The hormone gibberellin (GA) is a major regulator of seed germination and is controlled by various factors within the seed (Vishal and Kumar, 2018). Before the germination process begins, abscisic acid levels are downregulated, whilst the GA content is upregulated (Shu et al., 2016; Urbanova and Leubner-Metzger, 2016). This increase in GA occurs with the release of seed dormancy, such as inhibition and stratification. Treatment with exogenous GA can break dormancy in some dormant seeds can use this opportunity to break dormancy, allowing for faster seedling production. Exogenous GA treatment was effective for breaking dormancy in *Cinnamomum mikao* (Chen et al., 2022), *Leymus chinensis* (He et al., 2016), and *Thalictrum Uchiyama* (Lee et al., 2018) seeds, although *Clematis terniflora* (Jang et al., 2023) and *Ferrula asa-foetida* (Hassani et al., 2009) seeds were unaffected. Many previous studies have applied GA to promote dormancy breakage, although responses to this have shown variation with plant species, dormancy class, forms of GA, concentration, and treatment length.

However, to the best of our knowledge, currently no information is available regarding the exact dormancy class of *A. elliptica*. Therefore, our study aimed to classify dormancy in *A. elliptica* seeds and furthermore, to identify the conditions required to break it. Specifically, we determined whether the seeds were water-permeable and evaluated their responses to scarification, cold and warm (moist) stratification, and treatment with GA. Our results provide valuable information for the stable
preservation and restoration of seeds and could potentially be used to prevent the extinction of *A. elliptica*, through stable seedling production and distribution.

**Materials and methods**

**Plant material**

Ripe seeds were collected in November 2019 from *A. elliptica* plants growing in Miwon-myun, Cheongiu-si, Chungcheongbuk-do, Korea. The seeds were cleaned by hand, sieved to remove plant debris, and stored at 4°C until use in the experiments.

**Seed properties and initial germination test**

An imbibition test was performed immediately after collection to determine whether the *A. elliptica* seeds could imbibite water. First, after determining the weight of 4 replicates of 50 seeds each, the seeds were placed in 15-mL tubes with distilled water. After immersion for 24 h, the seeds were removed from the water, dried with a towel, weighed, and immersed in distilled water again. The same procedure was repeated for 7 days. The increase in mass was calculated based on the initial weight. Scarified seeds were also used to double-check for impermeability of the covering layer.

The viability of fresh seeds was tested using 1% tetrazolium (triphenyl tetrazolium chloride, TTC; Sigma-Aldrich, St Louis, MO, USA) and X-ray scanning. The tetrazolium test was performed on 100 seeds soaked in distilled water at 20°C for 2 days. After soaking in water, the seeds were cut in half and placed on filter paper (Whatman no.1, Cytiva, Marlborough, MA, USA) moistened with tetrazolium solution in a 100-mm Petri dish cut-side-down for 1 day. The degree of embryo staining was observed under a stereomicroscope (SZ61; Olympus Corporation, Tokyo, Japan) to confirm seed viability. Both embryos and endosperms were recognized as viable only when stained red.

Seed size (20 seeds) and weight (100 seeds) were measured 4 times using Vernier calipers (Digimatic caliper; Mitutoyo, Kawasaki, Japan) and a microbalance (AS220, Radwag, Radon, Poland), respectively. The seed structures and cross-sections were observed using a stereomicroscope equipped with a CMOS camera (xScope F630; Dixi Sci., Daejeon, Korea). The E:S ratio was calculated from the longitudinal section image of the seeds.

Initial germination tests were performed within 2 months of seed collection. Seeds were surface-sterilized with 1% sodium hypochlorite for 10 min and thoroughly washed with distilled water before sowing in a Petri dish on two layers of filter paper moistened with distilled water.

Seeds were incubated in an environment chamber set at 25/15°C, 16/8 h (light/dark), with a white LED (30 µmol·m⁻²·s⁻¹) light source. Seeds were also incubated in continuous darkness by wrapping Petri dishes with aluminum foil. The germination test lasted 30 days, and germination was recorded daily, except under dark conditions, which was not checked until the end of the experiment. All seed properties were investigated within 2 months after collection.

**Effect of seed coat scarification on germination**

The effects of mechanical scarification on germination were determined simultaneously with the initial germination test. *A. elliptica* seeds were cut at the end of the 1-mm long seed coat using a razor blade, and non-scarified seeds were used as controls. After scarification, the seeds were sown on moist filter paper in Petri dishes and tested for germination under light/dark conditions, as described above.

Additionally, scarified seeds were sown in soil to determine whether they could produce normal seedlings. The seeds were sown in sterilized containers (sealed container, 318 × 230 × 175 mm, TAESUNG CNC Co., Ltd., Goyang, Korea) filled with a horticultural substrate (Hanareum; Shinsung Mineral Co., Ltd., Seongnam, Korea) and incubated under a light/dark cycle at 25/15°C. The containers were moistened with distilled water to prevent dehydration. Four replicates of 50 and 25 seeds were used for the Petri dishes and soil containers, respectively. The germination test in the Petri dishes lasted 30 days, and the germinated seeds were counted every day and removed from the dishes. The germination test in the soil was conducted during the same period, and seedling emergence and status were recorded. Germination was recorded when radicle protrusion was > 1 mm, and a normal seedling was defined as the normal emergence of both cotyledons and radicles.

**Dormancy break by stratification**

We determined the effects of cold and warm plus cold stratification on dormancy break in *A. elliptica* seeds. Seeds were warm-stratified in the dark at 15/10°C (16/8 h, autumn) and then cold-stratified in the dark at 4°C (constant, winter). The conditions used in the experiment were set considering the ecological factors present in their natural habitats, such as seed maturation and germination. Seeds were first stratified for 4 or 8 weeks at 15/10°C (warm stratification, WS) and moved to a 4°C (cold stratification; CS) chamber for 4, 8, and 12 weeks. Other seed lots were subjected to CS at 4°C for 4, 8, and 12 weeks without WS. Mesh (filter) bags containing 50 seeds with 4 replications were placed in a sealed plastic box between layers of autoclaved wet pearlite (New Pershine No.2, GFC. Co., Ltd.,
Effect of gibberellins on germination

We evaluated the effect of gibberellins on dormancy break in *A. elliptica* seeds. Various concentrations (100, 500, 1000, and 2,000 mg/L) of GA3 (Cas No. 77-06-5; Sigma-Aldrich) and GA4+7 (2:1, Cas No. 468-44-0/510-75-8; KisanBio, Seoul, Korea) were used to pre-soak *A. elliptica* seeds for 2 days at room temperature (25°C). Thereafter, the seeds were washed in tap water, sown in a Petri dish, and incubated under a light/dark cycle at 25/15°C for 30 days. Four replicates of 25 seeds were used for each concentration, and germination was monitored daily. Based on these results, the next experiment was conducted to determine the type and optimal concentration of GA for breaking dormancy.

In the second experiment, seeds were soaked in 500 mg/L GA4+7 and distilled water at room temperature (25°C) for 2, 4, 7, and 14 days. Five replicates of 20 seeds were used for each treatment, and the seeds were incubated under a light/dark cycle at 25/15°C for 30 days.

Statistical analyses

SAS v9.4 (SAS Institute Inc., Cary, NC, USA) was used to calculate the mean and standard error for each treatment, and the analysis was performed using Duncan’s multiple range test at a significance level of *P* < 0.05.

Results

Seed properties

The *A. elliptica* seed is cylindrical in shape with a length and width of 9.1 ± 0.16 and 2.4 ± 0.05 mm, respectively, and the 100-seed weight was 1.6 ± 0.03 g (Figures 1A, B). Observation of the inner structure of seeds revealed a brown cork-textured seed coat and an embryo at the innermost layer, and an endosperm was identified between the coat and embryo (Figures 1C, D). The initial moisture content (MC) of seeds was 7.2 ± 0.65%, and during imbibition, the MC of both non-scariﬁed and scariﬁed seeds increased rapidly for 2 days, reaching 53.5 and 53.9%, respectively. However, no signiﬁcant increase was observed after 2 days. The viability of seeds tested using tetrazolium solution was 90%, and X-ray analysis revealed that 94% of the seeds were ﬁlled, i.e., they had an embryo (Figures 1E, F).

Effect of pre-soaking duration in GA4+7 on germination

After conﬁrming the effect of GA4+7 on the germination of *A. elliptica* seeds, a moderate concentration of 500 mg/L was used to test the seeds, depending on the soaking period (Figure 7). Seeds soaked in 500 mg/L GA4+7 showed greatly

Dormancy break according to stratification temperature and regime

Cold stratification for 4 weeks signiﬁcantly increased the germination of *A. elliptica* to 13.5% compared with the initial germination of 8.8% (Figure 4). As the length of the cold stratification treatment increased, germination increased to a maximum of only 21.0%; however, there was no signiﬁcant difference between 8 and 12 weeks.

Warm stratification for 8 weeks, followed by 12 weeks of CS, increased the germination of *A. elliptica* seeds to 68.8% (Figure 5). Thus, WS followed by CS was more effective at breaking dormancy than CS alone. In addition, an increased WS period increased germination compared with cold stratification for the same period.
FIGURE 1
Morphological characteristics and viability of *Amsonia elliptica* seeds. (A), arrangement of the seeds in the follicle; (B), longitudinal section of seeds; (C), horizontal section of seeds; (D), the internal structure of seeds; (E), scarified seed after soaking in tetrazolium solution; (F), Fidelity of seeds confirmed by X-ray images. The red arrow represents empty seeds.

FIGURE 2
Initial germination of *Amsonia elliptica* seeds after 30 days. L/D, light/dark condition; D, dark condition; NS, non-scarified; SC, scarified. Bars represent standard error (n = 4). Different lowercase letters indicate a significant difference tested by Duncan’s multiple range test (P < 0.05).
FIGURE 3
Normal and abnormal phenomena observed in germination and seedling emergence after Amsonia elliptica seed scarification. (A), germination of scarified seeds observed in a Petri dish 14 days after sowing; (B), the radicle protrudes first and can grow into normal seedlings; (C), The cotyledon protrudes first, and the radicle fails to protrude, resulting in abnormal growth or death; (D), normal seedlings; (E), abnormal seedlings with arrested cotyledons; (F), abnormal germination.

FIGURE 4
Germination of non-scarified Amsonia elliptica seeds in light/dark at 25/15 °C after cold stratification for 0–12 weeks. CS, cold stratification (4°C). Bars represent standard error (n = 4).
improved germination compared with that of seeds soaked in water for the same period. In particular, seeds soaked for 14 days in GA4+7 exhibited 61.3% germination.

Discussion

*A. elliptica* has physiological dormancy

If seeds with a well-developed embryo and a water-permeable seed/fruit coat sown under suitable germination conditions fail to germinate within approximately 4 weeks, they are either nonviable or have physiological dormancy. *A. elliptica* seeds, whose viability was confirmed, did not germinate within 4 weeks in the initial germination test but produced fully developed embryos (Figure 1B). Thus, seeds of this species have PD, which is a physiological-inhibiting mechanism that prevents germination due to insufficient embryonic growth potential for germination. Therefore, scarification may promote germination by releasing mechanical restrictions on embryos. Patents related to the promotion of *A. elliptica* seed germination indicate that the seeds cannot absorb water, and germination is promoted by

![FIGURE 5](image)

**FIGURE 5**

Germination of non-scarified *Amsonia elliptica* seeds at 25/15°C after combined stratification for indicated periods. WS, warm stratification (15/10°C, 16/8 h); CS, cold (4°C). Bars represent standard error (n = 4).

![FIGURE 6](image)

**FIGURE 6**

Percent germination of *Amsonia elliptica* seeds soaked in indicated concentrations of gibberellins (GAs) for 2 days. D.W., seeds soaked for 2 days in distilled water. Bars indicate standard errors (n = 4). Different lowercase letters indicate a significant difference tested by Duncan’s multiple range test (P < 0.05).
treatment with sodium hypochlorite or sulfuric acid to chemically scarify the seed coat (Choi and Jung, 2017). However, in the imbibition test performed in our study, the moisture content of non-scarified A. elliptica seeds increased to 53.5% after immersion in water. Thus, the seeds were water-permeable and did not exhibit physical dormancy.

Germination of the water-permeable seeds of A. tabernaemontana (Martin et al., 2013) and other species, such as Cucumis melo (Welbaum et al., 1990), Fraxinus excelsior (Finch-Savage and Clay, 1997), and Lycopersicon esculentum (Downie et al., 2003) is increased by scarification of the seed coat, which removes the mechanical restriction on the embryo. Nikolaeva (1977) classified dormancy induced by mechanical restriction due to the seed-covering layer as mechanical dormancy. However, according to Baskin and Baskin (2004), suppression of germination by mechanical resistance is a physiologically dormant state in which seeds cannot germinate due to low growth potential ('expansive force') rather than dormancy induced by the covering layer. After the PD is broken, the embryo has sufficient growth potential to overcome the mechanical resistance of the covering layer. However, in the initial germination test, scarified seeds of A. elliptica sown in Petri dishes did not germinate normally, despite the removal of the mechanical resistance of the covering layer (Figure 3A). In normal germination, the radicle usually protrudes first (Figure 3B); however, in scarified seeds, the cotyledons protrude first, and the radicle does not protrude (Figure 3C) or the cotyledons are arrested by the seed coat and grow into abnormal seedlings (Figures 3D–F). For the scarified seeds sown in the soil, although radicles protruded from the seed coat, only a few seeds grew into normal seedlings (Figure S1). This phenomenon was assumed to be caused by the insufficient growth potential of the embryo. Therefore, it was confirmed that artificial scarification could harm both seeds and seedlings. This results in abnormal germination following scarification, indirectly confirming that the cause of germination inhibition is not in the coat but in the seed (embryo).

**Alleviated physiological dormancy by stratification**

The most common dormancy in terrestrial seed plants is PD (Baskin and Baskin, 2014), which is usually alleviated by temperature (cold moist stratification, warm moist stratification, after-ripening), hormones, or chemical treatments (Baskin and Baskin, 2004; Baskin and Baskin, 2014; Cho et al., 2020; Shao et al., 2021; Baskin and Baskin, 2022). Physiological dormancy is divided into two subclasses: regular and epicotyl. In regular PD, the shoot emerges within a few days of radicle emergence, but in epicotyl PD there is a delay of several months between radicle and shoot emergence (Baskin and Baskin, 2022). As confirmed by the initial germination test, A. elliptica seeds have regular PD, subdivided into three levels: non-deep, intermediate, and deep. Dormancy break in seeds with non-deep PD usually occurs during 8–10 weeks of warm or cold stratification, depending on the species. However, cold stratification of A. elliptica seeds resulted in low germination percentages, suggesting that dormancy was either intermediate or deep.

If seeds of temperate-zone species have intermediate PD, warm stratification followed by cold stratification is effective in breaking dormancy, and seeds usually respond to treatment with GA3. Without warm stratification pretreatment, seeds require...
12–16 weeks of cold stratification to germinate. If seeds of temperate-zone species have deep PD, they require 12–24 weeks of cold stratification for dormancy break, and they do not respond to treatment with GA4 (Baskin and Baskin, 2022). For A. elliptica seeds, warm stratification followed by cold stratification dramatically increased the germination percentage (Figure 5). The improved germination of seeds receiving warm + cold stratification compared with that of seeds undergoing only cold stratification suggests that the seeds have an intermediate PD. In the natural habitat, A. elliptica seeds are exposed to warm stratification for at least 4 weeks in autumn before they are exposed to cold stratification in winter.

Gibberellins rapidly break seed dormancy without stratification

Exogenous hormones improve seed germination in many species in past times (Stowe and Yamaki, 1957). In particular, gibberellins are commonly used to improve the germination, growth and flowering of seeds and plants, usually in concert with abscisic acid and auxins, by up- or downregulating genes related to gibberellin biosynthesis (Wolbang et al., 2004; Frigerio et al., 2006; Seo et al., 2006; O’Neill et al., 2010; Yaish et al., 2010). There are many gibberellin structures, and their biological activity differs depending on their form (Mander, 1992; Shi et al., 2016). Among them, GA3 is best known in seed germination and dormancy, and it is also a criterion for classifying PD subdivision according to its response (Baskin and Baskin, 2014). If A. elliptica seeds exhibit an intermediate PD, GA3 is expected to promote germination. However, we found that GA3 did not promote germination, while combining GA4 and GA7 was effective in breaking dormancy and did so in the absence of cold stratification (Figure 6). GA4 and GA7 are synthesized from GA9 reaction with GA3ox, and GA3 and GA7 are the bioactive forms of gibberellin. However, they are not as well known as GA3 in dormancy and germination. GA4 has been identified as the major form of gibberellin in Arabidopsis, Cucurbitaceae, and rice (John et al., 2016). Increased GA4 positively regulates the seed germination of Brassica parachinensis (Chen et al., 2021), Glycine max (Chen et al., 2020), and Syngonanthus verticillatus (Barreto et al., 2020), while reducing GA4 can inhibit germination. The reason A. elliptica seeds do not react to GA3 could be that the 13-hydroxylation pathway negatively affects its biological activity (Magome et al., 2013). Therefore, the percentage germination was even lower than that of the control seeds. Another hypothesis is that GA4 or GA7 contribute to endosperm weakening, which may restrain embryo expansion in A. elliptica seeds (Brown and Bridgall, 1987). Bioactive gibberellins are major contributors to endosperm weakening in Lepidium sativum seeds (Oracz et al., 2012; Voegele et al., 2012). In a study by Oracz et al. (2012), GA4 was also involved in weakening the structure enveloping the embryo. Therefore, inhibition of A. elliptica seed germination is assumed to be eliminated by serial or spontaneous gibberellin actions. However, our study did not investigate the mechanisms, by which gibberellins promote germination by breaking dormancy.

Conclusions

Amsonia elliptica seeds have intermediate PD that is broken by warm conditions followed by cold stratification or by treatment with a mixture of GA4 and GA7. Although the seeds are water permeable, scarification of the seed coat promotes germination by eliminating the mechanical resistance of the embryo. However, a high percentage of the resulting seedlings are abnormal. This phenomenon means that PD in A. elliptica is not simply due to mechanical resistance but is caused by insufficient embryonic growth potential. After dormancy was broken by warm plus cold stratification or by treatment with a combination of GA4 and GA7, the growth of seedlings was normal, indicating that dormancy breaking in A. elliptica seeds increases the growth potential, so that embryos can overcome their mechanical resistance on their own. Dormancy break following warm plus cold stratification in the laboratory simulates the treatment seeds receive in their native habitat due to seasonal changes in temperature between the time of dispersal in autumn and germination in spring. However, this is a long period of time, and we found that normal seedlings can be produced over a relatively short period by soaking the seeds in 500 mg/L GA4+7 for 2 to 14 days and then incubating them under simulated spring temperatures (25/15°C). Thus, treatment with GA4+7 not only breaks intermediate PD but does so within approximately 1 month from the start of seed soaking in GA4+7. Treatment with GA4+7 effectively shortened the period required for dormancy break, enabling rapid germination and seedling production. Our methods can be used to produce A. elliptica for restoration, and the wide use of this species can be expected in various fields, such as medicine and the horticultural industry, through mass propagation.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

Author contributions

SL, KP, and J-SC: conceptualization. B-KJ, and BJ: data curation. SL, HL: methodology. SL, KP: writing–original draft preparation. CB and J-SC: writing–review and editing. All authors contributed to the article and approved the submitted version.
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Conflict of interest

Author SI is employed by The Kiban co. Ltd.,

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.1043897/full#supplementary-material

SUPPLEMENTARY FIGURE 1

Normal seedlings and abnormal germination of scarified seeds of Amsonia eliptica germinated in Petri dishes (A) and soil (B). The black arrow is a normal seedling.
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