An exogenous cytokinin supply in the ornamental fern *Asplenium nidus* L. induces an unusual post-transplant biomass accumulation

Alberto Pagani¹, Danilo Carnelos¹, Jorge Molinari¹, Ernesto Giardina¹, Adalberto Di Benedetto¹,²*

¹ Universidad de Buenos Aires, Buenos Aires-BA, Argentina.
² Universidad Nacional de Mar del Plata, Provincia de Buenos Aires, Argentina

Abstract

Ferns are ornamental plants with a low relative growth rate and long production cycles, which are grown at small pot volumes to optimize the commercial space for sale. However, the root restriction effects under this plant management can limit biomass accumulation and frond area. Since an exogenous spray with cytokinin (6-benzyl aminopurine (BAP)) has been suggested as a tool to override the root restriction in plants grown in pots, this study aimed to evaluate the effect of different BAP doses (5, 50, 100 or 200 mg L⁻¹) once (7 days after transplant), twice (7 and 30 days after transplant) or three times (7, 30, and 60 days after transplant), on plant growth and frond area development in spore-propagated *Asplenium nidus* fern plants grown in pots. Both increasing the BAP doses and number of applications led to an unusual response: an excessive decrease in the frond plastochron and a significant increase in the frond number initiated at the apical shoot meristem. This large frond number was not sustained due to the low net photosynthetic rate of the younger fronds and significantly limited outward appearance.

Keywords: growth regulator, leaf growth, ornamental foliage plants, pot root restriction.

Introduction

Ornamental plants with a low relative growth rate and long production cycles (24 months or more), such as ferns (Page, 2002; Liao et al., 2017), are grown in small pot volumes and frequently transplanted before sale. However, *Asplenium nidus* fern plants grown in small pots suffer from root restriction (Pagani et al., 2020), in agreement with previous reports in other ornamental potted plants (Di Benedetto et al., 2020a). In this regards, Pagani et al. (2020) showed that the use of large pots (1500-cm³) in *A. nidus* increases fresh weight (FW), dry weight (DW) and frond area (the main commercial aesthetic trait). They also found higher frond appearance rate, frond area expansion, frond thickness, relative growth rate and net assimilation rate.

One of the main alternatives to override the root restriction syndrome in ornamental plants (Di Benedetto et al., 2020a) and vegetables (Di Benedetto et al., 2020b) is to use a synthetic cytokinin such as 6-benzyl aminopurine (BAP), which is able to regulate plant metabolism under abiotic stresses. The main effects of exogenous cytokinin on fern growth in culture have been described in detail (Kosakivska...
et al., 2016). In particular, Menendez et al. (2010) found that BAP induces bud formation and sporophyte production in A. nidus plants. Park et al. (2020) found that kinetin and BAP induce new shoots from the apical shoot meristem (SAM), while inhibiting rhizophore formation. In agreement, Souheil and Rola (2014) found that A. nidus plants grown in a solid medium supplemented with BAP and kinetin showed increased numbers of shoots and leaves and increased stem height. However, there are no reports on the effects of BAP during the commercial growth of A. nidus or other related spore-propagated ferns.

The fern A. nidus (Bird’s nest fern), a common epiphytic ornamental, has a short, erect rhizome and a rosette of simple fronds. Leaf stalks are stout and almost black and can be as long as 5 cm, whereas leaf length can be up to 150 cm or more and up to 20 cm wide. The plant narrows gradually, tapering both towards the pointed tip and towards the base. Although the life cycle of a fern includes two alternating generations: a diploid sporophyte and a haploid gametophyte, only the former has commercial significance. The fern A. nidus is native to tropical southeastern Asia and eastern Africa and it is in great demand as an ornamental plant.

Based on the hypothesis that an exogenous supply of BAP to the A. nidus sporophyte shoot would increase biomass accumulation by overriding root restriction, the aim of our work was to study the effect of different doses of BAP to the plant. Park et al. (2020) found that kinetin and BAP induced new shoots from the apical shoot meristem (SAM), while inhibiting rhizophore formation. In agreement, Souheil and Rola (2014) found that A. nidus plants grown in a solid medium supplemented with BAP and kinetin showed increased numbers of shoots and leaves and increased stem height. However, there are no reports on the effects of BAP during the commercial growth of A. nidus or other related spore-propagated ferns.

Materials and Methods

Two experiments were carried out in a greenhouse located at the School of Agronomy, University of Buenos Aires, Argentina (34°35’59”S, 58°22’23”W and altitude 25 m), in successive years, from October 10th 2017 to February 9th 2018, and from October 8th 2018 to February 11th 2019 (Experiments 1 and 2, respectively).

Plantlets from spores grown in 128-plug-cell trays of A. nidus L. were obtained from a commercial propagator and then transplanted into rigid 1500-cm³ plastic pots (one plant per pot). The pots were filled with a 40:40:20 (v/v/v) mix of Sphagnum maguellanicum peat:river waste: perlite.

Plants were irrigated as needed with high-quality tap water (pH: 6.64 and electrical conductivity of 0.486 dS m⁻¹) using intermittent overhead mist, and one weekly fertigation (1N:1P:1K:1Ca v/v/v/v) (50 mg L⁻¹ N) was included.

Half hourly averages of the air temperature were measured using a HOBO H08-001-02 data logger (Onset Computer Corporation, MA, USA) protected from direct radiation by aluminum foil shades. The mean air temperatures ranged between 22.7 °C and 26.08. The greenhouse was covered with a black shade cloth (for 50% full sunlight) and mean photosynthetic active radiation during the experiment ranged between 7.10 and 10.60 mol photons m⁻² day⁻¹. The plants were arranged at a density of 6 plants m⁻² to avoid mutual shading.

In Experiment 1, plants were sprayed with 0 (distilled water control), 5, 50, 100 or 200 mg L⁻¹ 6, benzyl amino purine (BAP) solutions (Sigma-Aldrich Co., St. Louis, MO, USA) 7 days after transplanting. In Experiment 2, plants were subjected to different numbers of BAP applications, by spraying them with 0 (distilled water control), 5, 50, 100 or 200 mg L⁻¹ BAP solutions at different times. Applications were performed once (7 days after transplant), twice (7 and 30 days after transplant) or three times (7, 30, and 60 days after transplant). The BAP concentrations were chosen from the results of a preliminary experiment not included in this work. In both experiments, all leaves were sprayed to run-off at sunset (approximately 50 mL pot⁻¹). BAP was first diluted in 80% ethanol and no surfactants were added.

Plants for destructive measurements were harvested at transplant and at 30-day intervals in Experiment 1, and at transplant and at 30-day intervals after the last BAP spray in Experiment 2. Roots were washed, and root, stem and frond FWs were recorded. DWs were recorded after drying roots, stems and fronds to constant weight at 80°C for 96 hours. The number of fronds was recorded and each frond area development was determined using the ImageJ® (Image Processing and Analysis in Java) software.

The rate of leaf appearance (RLA), the relative growth rate (RGR), the rate of leaf area expansion (RLAE), the mean partitioning rate (LAR), the specific leaf area (SLA), the leaf area ratio (LAR), the specific leaf area (SLA), the leaf area partitioning (LAP), the allometric coefficients (β) between roots: shoots and fronds: stems were calculated as previously (Di Benedetto and Tognetti, 2016).

The experimental design was a randomized one with four BAP applications for Experiment 1 and four BAP concentrations and three times of BAP application for Experiment 2. Data were subjected to a one-way and two-way analysis of variance (ANOVA) using STATISTICA 8 software (StatSoft) after checking ANOVA assumptions for Experiments 1 and 2 respectively. Means were separated by Tukey’s tests (p ≤ 0.05). Slopes from straight-line regressions of RLA, RGR, NAR, LAR, LAP and allometric β coefficients were tested using the SMATR package.

Results and Discussion

In Experiment 1 of the present study, a single BAP spray on A. nidus spore-propagated sporophytes led to significant changes in aesthetic traits such as a decrease in frond size and an increase in SAM (Figure 1).
When BAP dose increased, SAM differentiated a significantly higher number of lateral growth areas (290.91% compared to controls) (Figure 2), while RLA increased (120.89%), but RLAE (100.20%) and SLA (70.60%) decreased (Table 1).

In homogenate cultures from sporophytes of *A. nidus*, both gametophyte and sporophyte regeneration take place in a hormone-free medium (Fernández et al., 1993), which indicates that explants have enough endogenous hormones to start to grow. However, it has been shown that, during the micro-propagation of many ferns, sporophyte formation increases by adding BAP (Ravi et al., 2015). In addition, previous reports on sporophytes of *A. nidus* cultured in a medium supplemented with BAP have shown that rhizomes become swollen due to bud proliferation induced by the synthetic cytokinin (Fernández et al., 1993). However, few studies have evaluated the effects of exogenous cytokinin on spore-propagated sporophyte biomass accumulation.

At the end of Experiment 2, during which four BAP concentrations at three times of application were tested, total frond area decreased as both the BAP concentration and number of applications increased (Figure 3).

**Figure 1.** Control *Asplenium nidus* plants (A) and *Asplenium nidus* plants sprayed with a single 200 mg L\(^{-1}\) 6-benzylaminopurine (BAP) (B) at the end of Experiment 1. Horizontal lines indicate the 1 cm scale.

**Figure 2.** Effects of five 6-benzyl aminopurine (BAP) concentrations (0, 5, 50, 100 or 200 mg L\(^{-1}\)) on the number of lateral vegetative meristems (SAM) at the end of Experiment 1 in *Asplenium nidus* plants.
Table 1. Effects of five 6-benzylaminopurine (BAP) concentrations (0, 5, 50, 100 or 200 mg L⁻¹) on total and individual frond area and specific leaf area (SLA) on a fresh weight basis at the end of the experiment, rate of leaf appearance (RLA) and relative leaf expansion rate (RLAE) in *Asplenium nidus* plants. Different lower case letters indicate significant differences (*p* < 0.05) between treatments according to the Tukey’s Test (number of frond and frond area) or SMATR analysis (RLA, RLAE and SLA).

| BAP (mg L⁻¹) | Number of fronds plant⁻¹ | Frond area (cm² frond⁻¹) | RLA (fronds week⁻¹) | RLAE (cm² cm⁻² day⁻¹) | SLA (cm² g⁻¹) |
|--------------|---------------------------|--------------------------|---------------------|------------------------|--------------|
| 0            | 44.25d                    | 321.69a                  | 1.958e              | 0.0099a                | 65.99a       |
| 5            | 59.60b                    | 201.10b                  | 2.811b              | 0.0075b                | 36.70b       |
| 50           | 78.90a                    | 153.48c                  | 3.883a              | 0.0075b                | 28.41c       |
| 100          | 57.00b                    | 134.60d                  | 2.667c              | 0.0068b                | 21.59d       |
| 200          | 51.60c                    | 62.80e                   | 2.367d              | -0.0002c               | 19.04d       |

Figure 3. Effects of five 6-benzylaminopurine (BAP) concentrations (0, 5, 50, 100 or 200 mg L⁻¹) applied once (7 days after transplant) (I), twice (7 and 30 days after transplant) II or three times (7, 30, and 60 days after transplant) (III) on total frond area at the end of Experiment 2 in *Asplenium nidus* plants. Bars indicate standard errors.

Figure 4 shows an example for 5 mg L⁻¹ BAP repeatedly sprayed plants. The results from Experiment 2 showed that frond number and RLA significantly increased according to the BAP concentration and number of applications (Table 2), while, at the same time, RLAE and SLA decreased (Table 2).
The fern *A. nidus* has an apical cell-based meristem with a distinct single apical initial cell, which is always formed during, and is responsible for, the early development of the gametophyte (Bartz and Gola, 2018). However, the unusual increase in the shoot apical meristem shown in Figures 1 and 4 was the result of both biomass stem accumulation and lateral meristem differentiation (Figure 2). In this regards, Romanenko et al. (2019) showed that during *in vitro* propagation of the fern *Dryopteris filix-mas*, cytokinin inhibits gametophyte development, reduces the size, or causes multiple overgrowth and deformation of the heart-shaped thallus, and affects the formation of gametes. Although a common BAP-spray response is a change in photo assimilate partitioning which favor stems (Di Benedetto et al., 2020a), there are no previous reports of that extreme response in fern sporophytes or other ornamental plants grown at commercial pot growing.

The main changes observed in plant attributes during Experiments 1 and 2 were related to an increase in the number of fronds per plant and a decrease in total frond area (Table 1 and Figure 3). These results may be explained by RLA (as an estimator of frond initiation) and RLAЕ (as an estimator of frond expansion). On the other hand, SLA (as an estimator of frond thickness) decreased as well, involving an increase in frond thickness (Tables 1 and 2). As a result, the higher number of new fronds initiated were small and thick.

**Table 2.** Effects of five 6-benzyl aminopurine (BAP) concentrations (0, 5, 50, 100 or 200 mg L⁻¹) applied once (7 days after transplant) (I), twice (7 and 30 days after transplant) (II) or three times (7, 30, and 60 days after transplant) (III) on frond number, total frond area and specific leaf area (SLA) on a fresh weight base at the end of Experiment 2, rate of leaf appearance (RLA) and relative leaf expansion rate (RLAE) in *Asplenium nidus* plants. Different lower case letters indicate significant differences (*p* < 0.05) between treatments according the Tukey’s Test (number of frond and frond area) or SMATR analysis (RLA, RLAЕ and SLA).

| BAP    | Number of fronds | RLA (fronds week⁻¹) | Frond area (cm² frond⁻¹) | RLAЕ (cm² cm⁻² day⁻¹) | SLA (cm² g⁻¹) |
|--------|------------------|---------------------|--------------------------|-----------------------|---------------|
| (mg L⁻¹) | plant⁻¹          |                     |                          |                       |               |
| 0      | 16.8d            | 0.556f              | 106.28a                  | 0.0133a               | 23.74a        |
| I-5    | 22.3d            | 0.622f              | 84.98b                   | 0.0118b               | 22.71b        |
| I-50   | 30.1c            | 0.861e              | 66.38c                   | 0.0099c               | 22.13b        |
| I-100  | 33.7b            | 0.861e              | 67.06c                   | 0.0092c               | 21.14b        |
| I-200  | 35.1b            | 1.083d              | 71.45c                   | 0.0098c               | 19.84b        |
| II-5   | 23.5d            | 0.761e              | 72.64c                   | 0.0126b               | 22.83b        |
| II-50  | 30.4c            | 0.967d              | 71.90c                   | 0.0093c               | 22.03b        |
| II-100 | 34.7b            | 1.067d              | 48.91d                   | 0.0094c               | 21.66b        |
| II-200 | 36.6b            | 1.222b              | 54.10d                   | 0.0091c               | 21.15b        |
| III-5  | 24.1d            | 0.811e              | 66.49c                   | 0.0102c               | 22.16b        |
| III-50 | 31.6c            | 1.044d              | 50.72d                   | 0.0098c               | 21.83b        |
| III-100| 35.2b            | 1.122c              | 44.97d                   | 0.0101c               | 21.18b        |
| III-200| 38.9a            | 1.306a              | 41.47c                   | 0.0094c               | 18.94c        |

*Figure 4. Asplenium nidus* plants sprayed with 5 mg L⁻¹ 6-benzyl aminopurine (BAP) one (7 days after transplant) (A), two (7 and 30 days after transplant) (B) and three times (7, 30, and 60 days after transplant) (C) at the end of Experiment 2. Lines indicate the 1 cm scale.
for a transient indeterminacy during frond development, usually producing lateral pinnae during a longer period (Cruz et al., 2020). In this regards, class I KNOX genes are known to be expressed in the SAM and frond primordia, as well as in the pinnae of compound fern leaves, suggesting that the same network for compound leaf development might be in place in ferns and seed plants (Vasco and Ambrose, 2020). Since cytokinin has been involved in Class I KNOX gene regulation (Rashotte, 2020), we can hypothesize that an exogenous BAP spray to A. nidus plants over-increases cytokinin levels at the SAM and leads to a negative growth signal (Figures 1 and 4).

At the end of Experiment 1, total DW decreased (43.00%) as BAP concentration increased. Figure 5A shows that, although all plant organs decreased biomass accumulation on a DW basis, fronds were the ones mainly affected. On the other hand, data from Experiment 2 showed that total DW decreased in response to an increase in BAP concentration, which was amplified with consecutive BAP applications (Figure 5B).

**Figure 5.** Effects of five 6-benzyl aminopurine (BAP) concentrations on dry weight partitioned between different plant organs at the end of the experiment 1 (A) and total dry weight at the end of experiment 2 when five BAP concentrations were applied once (7 days after transplant) (I), twice (7 and 30 days after transplant) II or three times (7, 30, and 60 days after transplant) (III) in Asplenium nidus plants. Bars indicate standard errors.

Table 3 shows a decrease in RGR (70.19%) and NAR (23.03%), which explains the decrease in total DW in response to higher single BAP concentrations. A decrease in LAP and the β coefficient from root: shoot allometries explained the higher biomass accumulation in shoots, while the changes in the β coefficient from fronds: stems allometries explained the sudden increase in the SAM observed in Figures 1, 2 and 4.

**Table 3.** Effects of five 6-benzyl aminopurine (BAP) concentrations (0, 5, 50, 100 or 200 mg L⁻¹) on the relative growth rate (RGR), net assimilation rate (NAR), leaf area ratio (LAR), leaf area partitioning (LAP) and β coefficients from roots: shoots and fronds: stems allometries during Experiment 1 in Asplenium nidus plants. Different lower case letters indicate significant differences ($p < 0.05$) between treatments according to Tukey’s Test (number of frond and frond area) or SMATR analysis (RGR and NAR).
During Experiment 2, RGR and NAR decreased with the BAP concentration and the number of applications (Table 4). On the other hand, in agreement with Experiment 1, LAP and β coefficients from the roots: shoots and fronds: stems allometries showed higher photo-assimilate partitioning to shoots and stems respectively with the use of BAP in single or repeated applications (Table 4).

**Table 4.** Effects of five 6-benzyl aminopurine (BAP) concentrations (0, 5, 50, 100 or 200 mg L⁻¹) applied once (7 days after transplant) (I), twice (7 and 30 days after transplant) (II) or three times (7, 30, and 60 days after transplant) (III) on the relative growth rate (RGR), net assimilation rate (NAR), leaf area ratio (LAR), leaf area partitioning (LAP) and β coefficient from the roots: shoots and fronds: stems allometries in *Asplenium nidus* plants. Different lower case letters indicate significant differences (p < 0.05) between treatments according the Tukey`s Test (number of frond and frond area) or SMATR analysis (RGR and NAR).

| BAP (mg L⁻¹) | RGR (g g⁻¹ day⁻¹) | NAR (g cm⁻² day⁻¹) x 10⁻⁴ | LAR (cm² g⁻¹) | LAP (cm² day⁻¹/g day⁻¹) | Shoot: root β | Leaves: stem β |
|--------------|-------------------|--------------------------|---------------|------------------------|---------------|---------------|
| 0            | 0.0182a           | 24.22a                   | 75.14b        | 77.86a                 | 1.12a         | 3.01a         |
| I-5          | 0.0150b           | 17.93b                   | 83.66a        | 68.90b                 | 0.80b         | 1.46b         |
| I-50         | 0.0148b           | 19.69b                   | 75.17b        | 68.05b                 | 0.86b         | 1.14d         |
| I-100        | 0.0129c           | 15.92c                   | 81.03a        | 67.31b                 | 0.69c         | 0.65c         |
| I-200        | 0.0130c           | 15.54c                   | 83.66a        | 66.57b                 | 0.43d         | 0.50f         |
| II-5         | 0.0146b           | 16.18c                   | 90.23a        | 65.80b                 | 0.62b         | 1.29g         |
| II-50        | 0.0118c           | 13.50d                   | 87.41a        | 63.52b                 | 0.57b         | 1.11d         |
| II-100       | 0.0133c           | 16.62c                   | 80.02a        | 63.06b                 | 0.34c         | 0.63c         |
| II-200       | 0.0118c           | 13.67d                   | 86.32a        | 59.65b                 | 0.26c         | 0.34c         |
| III-5        | 0.0130c           | 15.16c                   | 85.75a        | 57.81c                 | 0.61b         | 1.19d         |
| III-50       | 0.0134c           | 16.43c                   | 81.56a        | 56.57c                 | 0.43c         | 0.71e         |
| III-100      | 0.0133c           | 15.90c                   | 83.65a        | 54.92c                 | 0.29d         | 0.62e         |
| III-200      | 0.0120c           | 13.81d                   | 86.89a        | 50.27c                 | 0.22d         | 0.29g         |

According to Page (2002), the main important disadvantages of the biology of ferns are the low-light photosynthetic capacity and the slow plant growth rates. Although differences in control plants were found between experiments (likely related to different spore-propagated *A. nidus* batches), RGR values were higher than those of other related fern species (Liao et al., 2017), with a significant decrease in BAP-sprayed plants (Tables 3 and 4). NAR values changed in the same way as RGR and would explain the results on DW accumulation shown in Figure 5. On the other hand, the distribution of biomass among plant organs is affected by the environment, habit of the plant, life span of the plant, and competitive interactions, and the growth rate is always controlled by the source-sink balance within the plant. This can be illustrated by the LAP growth parameter and both the roots: shoots and fronds: stems allometries, which, in our experiments, favored photo-assimilate partitioning to the SAM (Tables 3 and 4) when a single or repeated BAP spray was applied. As a result, RLA clearly increased and a significantly high number of fronds developed, but could not be sustained by the limited photosynthetic rate at this growth stage. Regarding this issue, Martin et al. (2004) examined numerous physiological parameters in individuals of varying sizes of *A. nidus* and found that the rates of net CO₂ exchange of the fronds measured in *situ* in the field appeared to increase with plant size.

Babenko et al. (2018) indicated that high BAP concentrations inhibited spore germination and gametophyte development of *Polystichum aculeatum*, whereas lower quantities showed slight stimulating effect, which clearly indicates that cytokinin can undoubtedly regulate the growth and development of ferns. On the other hand, Vedenicheva and Kosakivska (2017) indicated that the accumulation of trans-zeatin in fronds of *P. aculeatum* and *Dryopteris flix-mas* was revealed at the intensive growth stage, and that the level of zeatin riboside increased in fronds and rhizomes of both fern species at the stage of sporulation. These authors also found that the root system of both fern species was characterized by a lower level of cytokinin as compared to the aerial part.

**Conclusions**

The unusual results in *Asplenium nidus* when younger sporophytes were sprayed with different BAP doses (Experiment 1) at single or repeated times (Experiment 2) would be explained by a lower endogenous cytokinin level in control plants (such as that found in other ferns) and an excessive cytokinin increase from exogenous
BAP applications, which significantly increased photo-assimilate partitioning to the SAM. This led to an excessive decrease in the frond plastochron and a significantly higher frond number initiated at the SAM. The expansion of these excessive new fronds could not be sustained due to the low net photosynthetic rate of the previously expanded fronds, which limited the outward plant appearance and the sale opportunity.

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