Vitis 57, 35–40 (2018)

DOI: 10.5073/vitis.2018.57.35-40

Regulation of cluster compactness and resistance to Botrytis cinerea with B-aminobutyric acid treatment in field-grown grapevine

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Summary

Our paper offers unique information regarding the effects of DL-β-amino-n-butyric acid (BABA) on grape cluster compactness and Botrytis bunch rot development. The impact of treatment was investigated on a native Hungarian grapevine cultivar, 'Királyleányka' (Vitis vinifera L.) during three seasons. The highly sensitive cultivar with thin skinned berries provided excellent samples for *Botrytis* bunch rot studies. Our objective was to study if BABA treatment contributes to decrease Botrytis infection by promoting looser clusters. For this purpose, the female sterility effect of BABA in grapevine flowers was also examined, which may result in looser clusters. Cluster compactness was characterized with two different indexes; bunch rot incidence was assessed in percentages. Ovaries of flowers were examined under epifluorescent microscope. Significant differences were found in the compactness index (berry number/bunch length) between the control (2.87 \pm 0.83 no./cm) and treated bunches in the case of 2.0 g·L⁻¹ (2.18 \pm 0.77 no./ cm) and 3.0 g·L⁻¹ (1.90 \pm 0.72 no./cm) concentrations. Bunch rot incidence, however, was highly dependent on disease pressure influenced by the precipitation during ripening. In year 2013 all treatments gave significantly lower infection incidence, while the extremity of rain in 2012 and 2014, resulted in no epidemic or high infection, respectively. The treatment with 2.0 g·L⁻¹ BABA concentration decreased cluster parameters and led to the lowest disease incidence. Microscopic studies proved that successful treatments on cluster structure can be traced back to the female sterility caused by BABA. Our results presented clear evidence for the effectiveness of BABA treatment on Botrytis bunch rot by promoting looser clusters.

Key words: BABA (DL-β-amino-n-butyric acid); Botrytis bunch rot; cluster compactness; epifluorescent microscopy; female sterility.

Introduction

Botrytis bunch rot of grape is a particular problem in the continental temperate regions of Europe with occasionally occurring wet macroclimate during bloom and berry ripening, that is favorable for disease development. However, several other variables play a direct or indirect role in development of the infection, e.g. susceptibility of the berries, cluster architecture, microclimate of the clusters (VAIL and Marois 1991), canopy management (Werner et al. 2008), or plant nutrition (Keller et al. 2001, Cabanne and Donéche 2003, Valdés-Gómez et al. 2008). Keller et al. (2003) confirmed bloom as a critical developmental stage for infection, followed by latency until the berries begin to ripen. However, the correlation between the primary infection of flowers and the secondary infection of berries is not clear yet (Elmer and MICHAILIDES 2004). Anatomical parameters of berries, like structure of epidermis, cuticle and wax content, play a more important role in cultivar resistance to bunch rot, than the antifungal host defence mechanisms (GABLER et al. 2003). The above mentioned surface characters are influenced by the berry-to-berry contact, consequently by the cluster compactness which has high impact on Botrytis epidemics (HED et al. 2009). There are several attempts, mainly in table grape cultivation, to find the proper chemical, that favourably influences cluster structure. A range of chemical components was evaluated in terms of effectiveness on cluster structure (Weaver and Pool 1971, Schildberger et al. 2011). Gibberelic acid treatment was successfully applied to loosen the clusters (Teszlák et al. 2005, Spies and Hill 2008). Foliar application of the non proteinogenic DL-β-amino-n-butyric acid (BABA) in table grape gave promising results in maintaining quality and controlling bunch rot infection under cold storage (EL-METWALLY et al. 2014). This chemical is an effective inducer of resistance against biotic and abiotic stresses (JAKAB et al. 2001, 2005). Several studies showed, that BABA increased defense capability of the plants via fast hypersensitive response, callose deposition, lignin accumulation and PR protein synthesis (HAMIDUZZAMAN et al. 2005, Ton et al. 2005, Cohen et al. 2011). It induced local and systemic resistance against downy mildew in grape leaves (Cohen et al. 1999). Besides the priming of pathogen-specific defense responses by BABA, it may have direct fungicidal effect as well. Treatment on field-grown grapevine controlled the downy mildew infection through inhibition of sporulation (Reuveni et al. 2001), and inhibited mycelial growth and germination of Botrytis cinerea on agar medium (FISCHER et al. 2009). Furthermore, application of BABA during fertilization may reduce berry number through

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the induction of female sterility (Jakab *et al.* 2001). In the case of *Arabidopsis* flowers, BABA treatment as soil drench induced callose deposition in the micropylar region of the ovules, inhibiting the fertilization process (Kocsis and Jakab 2008). BABA application to flowering tomato plants resulted in a strong reduction in fruit set (Vaknin 2016), although as a root drench it did not repress seed development of tomato (Luna *et al.* 2016).

In our study, BABA treatment was tested to regulate cluster compactness and consequently to control bunch rot on field-grown grapevine. The study was carried out on one of the native cultivar in Hungary, 'Királyleányka' (synonyms: 'Königstochter', 'Feteasca regale'), whose clusters are highly susceptible to Botrytis infection. The cultivars indigenous in the Carpathian Basin are more susceptible to *Botrytis* bunch rot infection than the worldwide ones. Among them the studied cultivar is widely used for wine production in this region. Based on previous studies, we hypothesized that BABA may be capable of decreasing grapevine susceptibility to Botrytis infection through different ways, namely as potent inducer of resistance, as direct fungicide (direct ways) and as inducer of female sterility (indirect way). According to our knowledge, the effects of BABA applied on grapevine flowers against bunch rot infection were analysed for the first time in this study.

Material and Methods

Experimental vineyard and mainten a n c e: Field experiments were conducted in three consecutive seasons, from 2012 to 2014 on field-grown grapevines (Vitis vinifera L. 'Királyleányka') grafted onto Berl. x Rip. T. F. SO4. The experimental vineyard was located at the Institute for Viticulture and Oenology, Pécs, Hungary. The vineyard was planted in 1987, with vine-by-row spacing of 3.0 x 1.2 m in north-south orientation. All vines in this study were not sprayed especially for Botrytis bunch rot, although the following standard spray program was applied for control of other fungal diseases uniformly on whole experimental area: three applications of mancozeb (Dithane DG at 2.0 kg·ha⁻¹); two applications of folpet (Folpan 80 WDG at 1.25 kg·ha⁻¹); two applications of copper hydroxide (Copac Flow at 2.0 L·ha⁻¹); four applications of sulfur (Kumulus S at 2-4 kg·ha⁻¹); one application of proquinazid (Talendo at 0.25 L ha⁻¹). Furthermore, in 2012 one application of tebuconazole + triadimenol + spiroxamin (Falcon at 0.3 L·ha⁻¹); one application of metrafenon (Vivando at 0.25 L ha⁻¹); in 2013, one application of meptil-dinokap (Karathane Star at 0.6 L·ha⁻¹); one application of miklobutanil (Talentum at 0.1 L·ha⁻¹) and in 2014, one application of metiram + ametoktradin (Enervin at 2.5 kg·ha⁻¹); two applications of meptil-dinokap (Karathane Star at 0.6 L·ha⁻¹); two applications of metrafenon (Vivando at 0.25 L·ha-1); one application of tetraconazole + proquinazid (Talendo Extra at $0.25~L\cdot ha^{-1}$); one application of difenoconazole + cyflufenamid (Dynali at 0.5 L ha⁻¹); one application of boszkalid + krezoxim-metil (Collis SC at 0.4 l/ha).

Experimental performance: In the year 2012, treatments included 2.0 g·L⁻¹ BABA (DL- β -amino-n-

butyric acid, Sigma) concentration, while the next two years three concentrations (1.0 g·L⁻¹, 2.0 g·L⁻¹, 3.0 g·L⁻¹) were dissolved in water and sprayed on the flowers of 'Királyleányka' at BBCH 65 phenological stage (Lancashire *et al.* 1991). The applied concentrations of BABA treatment were based on our preliminary results on grey mould (CSIKÁSZ-KRIZSICS *et al.* 2013) and Reuveni *et al.* (2001) field experiment in grapevine against *Plasmopara viticola*. Treatments were applied in the early morning (at 8-9 a.m.) before opening of the flowers. The bunches were collected and evaluated at harvest time (BBCH 89).

Data collection for the experiments: Treated and control clusters were removed and taken to the laboratory just before harvest. Compactness was assessed by the total number of berries per centimeter of the bunch (Pommer et al. 1996) in the year 2012. Bunch length included the length of the main rachis and that of the lateral branches. Rachis length was measured from the first lateral branch to the bottom of the cluster. In addition, the measured parameters were completed with the weight of the clusters and that of the berries in the year 2013. Thus, another compactness index derived by dividing bunch weight with the squared bunch length was calculated (Tello and Ibánez 2014). Bunch rot incidence was determined in percentage by visual assessment of the health status of the clusters (% of bunches infected) at harvest time.

Microscopic analysis of grapevine flowers: Inflorescences in the phenological stage BBCH 68 were harvested and stored in ethanol, glycerol and water (1:1:1 v/v) until use. The staining procedure was made according to Lu et al. (2011) with minor modifications. The flowers were fixed in 200 μL acetic acid and ethanol (3:1 v/v) for 1.5 h. To soften the tissues, they were submerged in 1 N NaOH for 15 min in 60 °C thermostat, then washed 3-times with distilled water and stained with 200 μL aniline blue (0.01 %) for 5-10 min. Callose depositions and callose-rich structures, e.g. pollen grains and pollen tubes fluoresced yellow under the epifluorescent microscope (Nicon Eclipse 80i microscope with UV light – adapted system, with illumination from an Osram HBO 100 W/2 mercury lamp). Micrographs were taken by the Spot Basic 4.0 software.

Statistical analyses: Data from three repetitions each with 10 clusters per treatments (n = 30) were used for evaluation of cluster compactness. Bunch rot incidence was calculated in percentages, based on 50 randomly selected clusters per treatments, visually observed as healthy or infected. For microscopic analysis 5-5 pollinated flowers per 4 clusters per treatments (80 flowers in all) from 2013 were used. Average values and standard deviation (SD) data were calculated using Microsoft Excel 2010 software. The significance of differences was assessed using Student's t-test.

Results

Effects of BABA on cluster compactness and on ovary fertilization: The applied BABA concentration in 2012 was 2.0 g·L⁻¹. Control bunches were 21.4 ± 9.1 cm bearing 96.1 ± 36.6 berries, which resulted in 4.67 ± 1.39 cluster compactness index calculated as

berries per cm. The BABA treatment completed the expected effect in so far as the compactness index of treated bunches decreased by 15 % (3.95 \pm 1.12). Next year the concentration dependent efficiency of BABA treatments was investigated, when 1.0 g·L⁻¹, 2.0 g·L⁻¹ and 3.0 g·L⁻¹ concentrations of BABA were applied (Table). Although the treatment with 1.0 g·L⁻¹ did not result in significant differences in the measured parameters to the control, its compactness index calculated with greater impact on bunch length gave significantly lower value. The 2.0 g·L⁻¹ concentration decreased significantly these parameters except the bunch length. The weight of the berries and consequently that of the clusters were decreased also by the 3.0 g·L⁻¹ concentration, compared to the control. Only this concentration affected bunch length, while berry number was not changed significantly. Cluster compactness of control bunches was significantly lower in the year 2013 than in the previous year because of the unfavorable fruit set. The treatments with 2.0 g·L⁻¹ and 3.0 g·L⁻¹ concentrations further loosened the clusters based on both indexes (Table, Fig. 1).

The treatments with 2.0 g·L⁻¹ and 3.0 g·L⁻¹ concentrations reduced significantly the weight of berries, while berry numbers were significantly lower only in the case of the 2.0 g·L⁻¹ concentration. A post pollination study of the pistils of flowers treated by the 2.0 g·L⁻¹ concentration offer an explanation for this phenomenon. Inflorescences were

harvested in the phenological stage BBCH 68, in order to let prevail the female sterility induction of BABA (Fig. 2). In the case of control flowers, it was clearly observable that the pollen tube entered the ovules and there was no callose deposition at the micropylar region in 90 % of the pollinated flowers studied (Fig. 2a). Based on our observations, in some flowers of the BABA-treated inflorescences (20 %) pollen tubes did not reach the ovule, but they were interrupted about halfway in the ovary. Additionally, BABA induced callose deposition at the micropylar region and sometimes in the ovules body itself as well (Fig. 2b). Furthermore, there were BABA-treated flowers (about 30 %), which presented both phenomena in the same ovary, namely there were fertilized and non-fertilized ovules, the latter with vigorous callose deposition at least at the micropylar region (Fig. 2c).

Botrytis bunch rot incidence: The cultivar 'Királyleányka' with medium sized, shouldered clusters and with thin skinned berries, provided excellent samples for Botrytis bunch rot studies (Fig. 1a). Infection incidence was 46.3 % in 2013 on control clusters. In this year all BABA treatments gave significantly lower infection incidence compared to the control (Fig. 3). The most effective one proved to be the 2.0 g·L¹ concentration, which limited the 46 % infection to less than 11.6 %. The treatments with 1.0 g·L¹ and 3.0 g·L¹ concentrations resulted 17.1 % and 33.0 % infection incidences, respectively.

Table

Effect of BABA-treatment on different bunch parameters of 'Királyleányka' in 2013

BABA - (g·L·1)	Measured parameters				Compactness index	
	BW (g)	BL (cm)	NbB (no.)	Wb (g)	NbB/BL (no./cm)	BW (g)/ [BL (cm)] ²
1.0	$85.8a \pm 27.7$	$24.5a \pm 4.8$	$61.2a \pm 18.4$	$1.29a \pm 0.21$	$2.62a \pm 1.04$	$0.165b \pm 0.109$
2.0	$60.3b \pm 30.6$	$22.2a \pm 6.5$	$47.7b \pm 20.0$	$1.23b \pm 0.21$	$2.18ab \pm 0.77$	$0.135b \pm 0.072$
3.0	$63.4b \pm 32.3$	$26.2b \pm 7.4$	$50.2a\pm24.0$	$1.25b \pm 0.18$	$1.90b \pm 0.72$	$0.100c \pm 0.055$
Control	$84.2a \pm 26.5$	$21.5a \pm 7.1$	$58.3a \pm 17.7$	$1.46a \pm 0.22$	$2.87a \pm 0.83$	$0.229a \pm 0.129$

Values are means and standard deviations of data from 3 repetitions each with 10 clusters (n = 30). BW: Bunch weight; BL: Bunch length; NbB: Average number of berries per bunch; Wb: Weight of berries; no.: number. Means within a column followed by the same letter are not significantly different based on Student's t-test (p < 0.05).

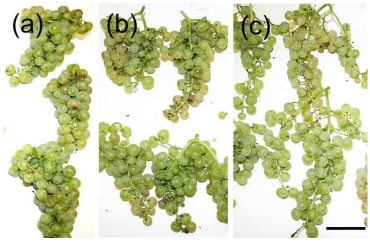


Fig. 1: Control (a), $2.0 \text{ g} \cdot \text{L}^{-1}$ (b) and $3.0 \text{ g} \cdot \text{L}^{-1}$ (c) treated bunches of 'Királyleányka' in the year 2013. (Scale bar: 6 cm).

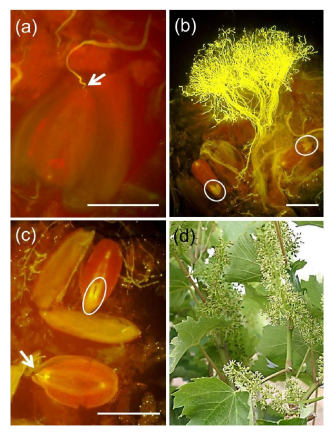


Fig. 2: Microscopic analysis of grapevine flowers in 2013. (a) Fertilization of an ovule in a control flower. Arrows indicate the entrance of pollen tube into the ovule through its micropyle. (b) Interrupted pollen tube growth in a pollinated flower treated by 2.0 g·L⁻¹ BABA solution. Pollen grains and pollen tubes are bright yellow, the unfertilized ovules have callose deposition at the micropylar region (circled). (c) An unfertilized ovule (above) out of the integuments, with callose deposition (circled) in its slim body and a fertilized ovule (below) with a piece of pollen tube, from a flower treated by 2.0 g·L⁻¹ BABA solution. (d) Inflorescences of 'Királyleányka'. (Scale bars: 500 μm)

There was no bunch rot in the year 2012. The year was dry after veraison and during berry ripening, which means that the infection evaluated was mainly the result of primary infection of the flowers. Secondary infection of the clusters was improbable because of the following dry weather (August, September). The treatments of inflorescences in the year 2013, which significantly decreased *Botrytis* incidence, further highlight the significance of primary infection of the flowers. The last study year (2014) gave no significant differences regarding cluster compactness (data not shown), and provided the uneffectiveness of the treatments on *Botrytis* infection (Fig. 3). The unusually high amount of rain in most of the vegetation period in the year 2014 allowed high epidemics in spite of the treatments.

Discussion

The field-experiment showed that the application of BABA on grapevine inflorescences had positive effect on cluster compactness and reduced *Botrytis* infection. Al-

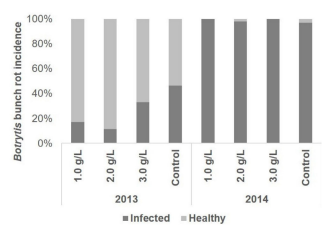


Fig. 3: *Botrytis* infection incidence in response to different BABA concentrations by inflorescence application on a field-grown grape-vine cultivar during the 2013, 2014 seasons. Percentage values are based on 50 clusters per treatments, classified as healthy or infected clusters.

though, disease pressure and weather conditions strongly influenced the results. Correlation between cluster character and bunch rot is well known, because plant growth regulators have long been applied on grapevines for induction of loose clusters and to restrict bunch rot attacks in fungicide-based management (Prior 2006, ZDUNIĆ et al. 2015). Among the many methods of quantifying grape cluster compactness in the literature, we used two different calculations based on three parameters, namely the berry number, bunch weight and bunch length (Tello and IBÁNEZ 2014). The indicator calculated as berry number per centimeter of a bunch was strongly correlated with bunch rot infection (HED et al. 2009), while this metric had low contribution to cluster compactness in characterization of different cluster architecture of several cultivars and clones (VAIL and MAROIS 1991). In our study, we calculated bunch length as the sum of the length of the main rachis and lateral branches, while HED et al. (2009) measured only the rachis length and disregarded berries on the shoulder, because they rarely developed bunch rot in 'Vignoles' grapes. In our case, calculation with the whole bunch length was justified by the bunch architecture of the cultivar, whose lateral branches bear several berries tending to bunch rot. The other compactness indicator we used was calculated with bunch weight and length, proposed by Tello and Ibánez (2014). Regarding compactness indexes, the impact of BABA treatments should be reflected in decreased berry number and lower bunch weight, starting from the female sterility effect of BABA (JAKAB et al. 2001).

Our microscopic studies of grapevine flowers revealed that the mechanisms behind the BABA-induced alteration of cluster architecture were associated with callose formation in the ovules and with inhibited pollen tube guidance in the ovary. These results are in agreement with previous studies that also established callose accumulation in grapevine leaves due to BABA treatment (Hamiduzzaman et al. 2005). Furthermore, inhibited fertilization through induction of callose deposition at the micropylar region and interrupted pollen tubes have also been observed in *Arabidopsis* flowers (Kocsis and Jakab 2008).

Bunch structure was modified significantly by different gibberellic acids and gibberellic acid inhibitors, while bunch rot incidence and severity showed no differences in the grapevine cultivar 'Sauvignon blanc' (Mundy *et al.* 2014). There was no significant difference in the induction of loose clusters and *Botrytis* infection using a plant-growth regulator containing prohexadione-calcium (Schildberger *et al.* 2011). Our results confirm these observations, in case all concentrations decreased the disease incidence regardless of the effectiveness of treatments on bunch structure (see treatment with 1.0 g·L¹ concentration). This observation suggests additional mechanisms behind the observed decrease of *Botrytis* bunch rot.

In addition to the supposed effect of BABA treatment on cluster compactness, we trusted its further mechanisms leading to the reduction of *Botrytis* infection. BABA treated plants react faster and more effectively to biotic stress because of the priming of pathogen specific defence responses (Sunwoo et al. 1996, Zimmerli et al. 2001, Ton and Mauch-Mani 2004). The protective effect of BABA can be based not only on induced resistance of the plant, but on direct toxicity on Botrytis cinerea, as it has been reported by Fischer et al. (2009). Inhibition of sporulation, mycelial growth and germination has been observed (REUVENI et al. 2001, Fischer et al. 2009). According to Csikász-Krizsics et al.'s (2013) studies, although BABA can inhibit mycelial growth above 400 µg·L⁻¹ concentration, its fungicidal effect was not observed. Evidence for BABA induced resistance in grapevine against fungal infections (FISCHER et al. 2009, SASEK et al. 2012) or against the Oomycete downy mildew (Cohen et al. 1999, Reuveni et al. 2001, Hamiduzzaman et al. 2005, Slaughter et al. 2008) has been described. The BABA-induced plant defence can explain that the treatment with 1.0 g·L⁻¹ concentration of BABA which did not change the cluster compactness, could significantly reduce *Botrytis* bunch rot incidence. Our study gives explanation for looser clusters and decreased berry weight caused by BABA treatment, because this substance as inducer of female sterility inhibited fully or partly the fertilization of flowers.

However, a successful treatment can not be guaranteed, because timing of treatments and meteorological conditions have decisive impact on the process. GIUDICE et al. (2004) emphasized the importance of timing of treatments with prohexadione-calcium, which decreased fruit set pre bloom and decreased berry weight post bloom. In our case, the treatments were justified at BBCH 65 pheonological stage, when about fifty percent of the flower caps were fallen (LAN-CASHIRE et al. 1991). During this period BABA imposed its female sterility effect on those flowers which were not yet fertilized. Weather conditions around grape bloom also affect fruit set and abscission, which are in strong correlation with bunch rot infection (Molitor et al. 2016). In addition to the flowers, berries after veraison are also very susceptible to bunch rot, while young, immature berries are highly resistant to the disease (Kelloniemi et al. 2015). In accordance with the model of González-Domínguez et al. (2015) our results highlight that primary infection of the inflorescences had significant effect on Botrytis bunch rot. However, secondary infection promoted by high humidity during ripening can further increase harvest lost by bunch rot.

Conclusion

Our paper shows original data on the effect of BABA treatment at full flowering on grape bunch architecture, flower fertility and *Botrytis* bunch rot. We demonstrated that BABA influenced cluster compactness through fully or partly inhibited fertilization in flowers, resulting in decreased number and weight of berries. Furthermore, the observation that all of the treatments decreased disease incidence regardless of bunch structure, emphasized the protective role of BABA through resistance induction or pathogen inhibition. This multiple effect prevailed in the case of 2.0 g·L⁻¹ concentration, resulting decreased cluster parameters and also the lowest disease incidence. In field conditions, however, the outcome of the treatment depends on different environmental factors influencing secondary infection of the clusters.

Acknowledgement

This work was supported by the Hungarian Scientific Grant Agency (OTKA K101430). The present scientific contribution is dedicated to the 650th anniversary of the foundation of the University of Pécs, Hungary.

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Received August 16, 2017 Accepted December 19, 2017