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ORIGINAL ARTICLE

Polyamines and methyl jasmonate in bulb formation of in vitro propagated tulips

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Abstract Effects of polyamines (PAs), their precursors, ornithine (Orn) and arginine (Arg), as well as methyl jasmonate (MeJA) were studied on bulb formation in vitro with four tulip cultivars differing in bulbing ability. The endogenous PA profiles of the shoots and bulbing shoots were monitored during successive micropropagation stages: S1, sixth week of the standard multiplication subculture; S2, fifth week of the last multiplication subculture; S3, 5 weeks later, just before the cold treatment; S4, at the end of the 13-week-cold treatment; and S5, 6 weeks after the end of cold treatment (shoot base swelling/bulb formation phase). Putrescine (Put), spermidine (Spd) and spermine (Spm) (50 and 100 µM) as well as Arg and Orn (500 and 1000 μ M) were added to the medium at various bulbing stages (S3, S4 and S5). Positive effects of these compounds on bulb formation were found for the cultivar 'New Beet' (NB), with the lowest bulbing ability. Both PAs and their precursors, when used at lower concentrations at the beginning of bulbing process (S3), together with standard MeJA treatment, significantly enhanced bulb numbers in this cultivar. Putrescine was the major endogenous polyamine at the stages of shoot multiplication (S1-S2). Spd levels were the highest in 'Fringed Black' (FB), the cultivar of extremely high bulbing ability, and this PA dominated over Put during the entire micropropagation process in this cultivar. Endogenous Spd contents manifested similar dynamics in all cultivars showing two peaks: at shoot multiplication (S2) and at the end of cooling (S4). The contents of all the PAs were the lowest at the

Keywords *Tulipa* · Polyamines · MeJA · Micropropagation · Bulb formation · In vitro

Introduction

In the majority of geophyte species, the formation of bulbs, corms or tubers is an essential stage in the micropropagation process because only the storage organ shows high rooting ability and good field performance as, e.g. in lily (Langens-Gerrits et al. 2003; Ishimori et al. 2007; Ascough et al. 2008) and tulip (Podwyszyńska and Sochacki 2010). In our tulip micropropagation system, the bulbing process is directly induced in shoots multiplied in vitro during several cycles over the course of 1–3 years (Podwyszyńska and Sochacki 2010). Bulb development takes 8-10 months and comprises three phases. In the first phase, shoots are prepared for bulbing at the temperature of 18 °C. They develop dormancy and become sensitive to low temperature treatment, and their endogenous level of abscisic acid (ABA) increases and level of indole-3-acetic acid (IAA) decreases (Podwyszyńska et al. 2004). In the second phase, bulb development is induced by treatment of the shoots with a low temperature of 5 °C for 12-14 weeks. During



final bulbing stage (S5) except the Spm level in cultivar NB in which it increased at this time. It is postulated that the high bulbing competence may be related to a transient increase in Spd content in response to low temperature treatment (S4), followed by its rapid decrease during bulb growth and maturation (S5, 6 weeks after the end of cooling) when developing bulbs begin to enter dormancy. The low levels of all the PAs at final bulbing stage as well as low Put/Spd ratio may also aid in the bulb formation of tulip in vitro.

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this phase, shoot dormancy is released, endogenous ABA levels fall while IAA levels rise. In the third phase, which takes 2–3 months, shoots are maintained on sucrose-rich medium at 18 °C and bulbs form.

Under this sequence of treatments, shoots of most tulip genotypes produce bulbs successfully. However, some genotypes have low bulbing capacity under these conditions (Podwyszyńska 2006b; Podwyszyńska et al. 2014). Other tulip genotypes, such as the one used for this study, the breeding line P14, produce high numbers of very small bulbs with an average mass below 60 mg. Such small bulbs grown in soil emerge and form new daughter bulbs only in about 30 % of cases. In turn, another tulip genotype "New Beet" (NB) forms relatively large bulbs in vitro but with a low efficiency of 50 %.

Therefore, several attempts have been undertaken in order to improve bulb formation of in vitro cultured tulip. It was found that modification and prolongation to 14 weeks of the last multiplication subculture, prior to cooling combined with the application of NAA and gibberellin biosynthesis inhibitors, enhance shoot's bulbing capacity (Podwyszyńska 2006b). Bulbing efficiency was markedly increased by treatments with MeJA at the last phase of this process, 6 weeks after the end of cooling shoots (Podwyszyńska 2006a). Recently, it has been proved that the replacement of thidiazuron (TDZ) (used as standard for shoot multiplication) with isopentenyladenine (iP) in the last multiplication subculture significantly enhanced bulb formation (Podwyszyńska et al. 2014). In the case of the cultivar NB, with low bulbing ability, such treatments, however, were found to be insufficient for enhancing the efficiency of this storage organ development.

For a better understanding of the mechanism of bulb formation, hormonal status was analysed in successive stages of this process for many bulbous species, both in natural and in vitro conditions. The important role of decreasing endogenous GAs during the bulb induction phase was shown in Allium wakegi (Yamazaki et al. 2002). Their findings clarify why gibberellin inhibitors stimulated bulb formation of micropropagated lily (Kim et al. 1994) and tulip (Podwyszyńska 2006b). The role of ABA level in dormancy development (increasing level) and release (decreasing level) closely associated with an in vitro bulb formation was studied in lily (Gerrits et al. 1992; Kim et al. 1994), tulip (Podwyszynska et al. 2004) and onion (Yamazaki et al. 2002). Also, jasmonates, jasmonic acid (JA) and methyl jasmonate (MeJA) have been considered to play an important role in the process of storage organ formation, including bulb formation in vitro of Allium sativum and A. cepa (Ravnikar et al. 1993; Koda 1997; Kim et al. 2003), Tulipa gesneriana (Podwyszyńska 2006b) and Narcissus triandus (Santos and Salema 2000).

However, little attention has been paid to the action of polyamines (PAs) in bulb formation. First, their role was studied in the development of tubers in potato (Mader 1995; Kumar et al. 1996). Recently, their beneficial effects have been reported for the in vitro formation of gladiolus corms (Kumar et al. 2011; Kumar and Palni 2013) and yam tubers (Ondo Ovono et al. 2010a, b). Polyamines (PAs): putrescine (Put), spermidine (Spd) and spermine (Spm) occur ubiquitously in plants, especially in actively growing tissues, and are considered plant growth regulators influencing several physiological and developmental processes (Kakkar et al. 2000; Bais and Ravishankar 2002; Kaur-Sawhney et al. 2003; Kosson and Prange 2005; Hussain et al. 2011; Anwar et al. 2015). They are involved in cell division, somatic embryogenesis, stem elongation and flowering, root growth, tuber development, senescence and plant responses to abiotic and biotic stress tolerance. However, the mechanism of action remains to be elucidated. PAs are compounds with amino groups. The amino acids (AA) arginine (Arg) and ornithine (Orn) are precursors of putrescine. Spermidine is synthesised from putrescine and spermine from spermidine S-adenosylmethionine in the presence of the spermine synthase. There is little information on polyamine action in the process of in vivo bulb formation in tulip (Le Nard and Fiala 1990; Kollöffel et al. 1992). So far the role of polyamines was not studied in tulip cultured in vitro.

In recent years importance of methyl jasmonates in polyamine metabolism has been investigated, and many studies indicate that application of methyl jasmonates can alter polyamine metabolism, resulting in accumulating of polyamines in tissue. This effect is attributed to influencing gene expression resulting in changes in activity of polyamine biosynthesis genes: arginine decarboxylase (ADC), ornithine decarboxylase (ODC) and S-adenosylmethionine decarboxylase (SAMDC) (Walters et al. 2002).

The aim of this study was to determine the effects of polyamines, their precursors and methyl jasmonate on the efficiency of in vitro bulb formation in tulip. Analysis of the endogenous levels of polyamines during this process was also included.

Materials and methods

Plant material and micropropagation conditions

Four tulip genotypes differing considerably in capacity of in vitro bulb formation were used in the study: 'Fringed Black' (FB) (late flowering Fringed Group) with extremely high bulbing ability, in which shoots form some bulbs even without low temperature treatment; P20 (mead-season



flowering, Fringed Group) forming bulbs with relatively high efficiency; 'New Beet' (NB) (Single Late Group) with the lowest ability to form bulbs; and P14 (late flowering, Lily-flowered Group) forming numerous but very small bulbs. In vitro shoot cultures of these cultivars were established, multiplied by adventitious regeneration, and prepared for bulb formation, essentially following our previously published protocol (Podwyszyńska and Sochacki 2010), except when polyamines, amino acids and MeJA were added in the particular experiments. Briefly, at S1 the shoots were continuously multiplied in vitro with 6 week subculture periods in 330 ml jars containing 40 ml of multiplication medium (Podwyszyńska and Sochacki 2010) supplemented with 1 µM TDZ, 20 µM iP and 1 µM NAA solidified with 6 g l⁻¹ agar (Difco). In the last multiplication subculture, iP was added at 25 µM but no TDZ. After 5 weeks of this last multiplication subculture (S2), a liquid medium (35 ml per jar) was added containing Murashige and Skoog (1962) (MS) mineral and organic compounds and 5 µM paclobutrazol, and the shoots were grown on the resulting two-phase medium for a further 5 weeks (S3) (thus the last subculture period before cooling was prolonged to 10 weeks). Then, the shoots were transferred in undivided clusters to modified MS medium (60 ml per jar) containing ammonium nitrate at the concentration reduced to the half, 7 % sucrose and 0.1 % activated charcoal but no cytokinins, auxins nor paclobutrazol and the jars were kept at 5 °C for 13 weeks to induce the bulbing process. Finally (after the end of cooling, S4), to initiate bulb growth, the shoots were cultured (on the same medium) for 13 weeks at 16 °C with 16/8 h photoperiods (illuminated at a photosynthetic photon flux density of 55 μmol m⁻² s⁻¹ by warm-white fluorescent lamps during the light phases). Shoot base swelling, i.e. bulb development start approximately 6 weeks after the end of cooling (S5). In order to accelerate bulb formation, except when mentioned, MeJA (Duchefa Biochemie BV, The Netherlands) solution was added twice: seven and ten weeks after the end of cooling. MeJA solution was added under the sterile conditions (5 mL of 0.1 % MeJA solution per 100 mL of a culture medium) (Podwyszyńska and Sochacki 2010). A solution of 0.1 % MeJA was dissolved (v/v) in 10 % ethanol.

In order to evaluate the effects of PAs and their precursors on bulb formation, four experiments were performed in which these compounds were added to the medium (before autoclaving) at different stages. The interactions of PAs and AAs with MeJA were also estimated in the cultivars NB and P14, when stated.

Experiment 1

The aim of this experiment was to determine the effect of MeJA and its interaction with PAs (Put and Spm at

 $100~\mu M$) and AAs (Arg and Orn $1000~\mu M$) on bulb formation of the cultivar NB (of the lowest bulbing ability). The compounds were added to the bulbing medium (without PGRs and with 7 % sucrose) just before cooling (S3). MeJA solution was added two times: 7 and 10 weeks after the end of cooling (as described above) to the half of the containers with bulbing shoots.

Experiment 2

All four tulip cultivars were used and PAs (Put and Spm) and AAs (Arg and Orn) were added to the bulbing medium just before cooling (S3). The concentrations of PAs and AAs were selected based on the preliminary experiment. Arg and Orn were used at 500 and 1000 μ M, and Put and Spm at 50 and 100 μ M. MeJA solution was added two times, as in the standard procedure described above.

Experiment 3

The genotypes NB and P14 were used. PAs, Put, Spm and Spd, as well as Arg and Orn, were added in water solutions to the shoot cultures just after the end of cooling (S4). The compounds were added in sterile water solutions in the way that the final concentration of each polyamine was 50 μ M and concentration of each amino acid was 500 μ M. MeJA effects, depending on PA/AA treatments, were studied only for the cultivar NB. MeJA was added (as in standard procedure) to the half of the NB bulbing shoot cultures of each AA/PA treatment.

Experiment 4

The genotypes NB and P14 were used. PAs (Put, Spm and Spd) as well as AAs (Arg and Orn) were added in water solutions to bulbing shoot cultures 6 weeks after the end of cooling (S5). The compounds were added in the same way as in the third experiment: PAs at the final concentration in the medium of $50~\mu M$ and AAs at $500~\mu M$. MeJA was added (as in standard procedure) to the half of the containers with bulbing shoots treated with AA/PA treatments.

Control shoot cultures were not treated with either PAs or AAs.

Thirteen weeks after the end of cooling (in each experiment), bulbs were taken out of containers, rinsed with tap water, dried and stored at 20 °C for 5 weeks. At this time, the total number of bulbs and the number of regular bulbs (properly formed and covered with tunic) obtained from each jar as well as mean bulb mass were recorded. Five or six jars, each containing three clumps of 3–4 shoots (10 shoots per jar), were used in the in vitro bulb formation efficiency experiments. The bulb



formation data were subjected to analysis of variance and treatment means were compared by Duncan's test at P = 0.05.

Determination of polyamines

The endogenous polyamine profiles were monitored by the following sampling, extraction and high-performance liquid chromatography HPLC procedures. Samples of basal parts of shoots (ca. 1 cm long and 1 g fresh weight) were taken at the following stages of micropropagation/bulb development: S1, after 6 weeks of the standard multiplication subculture; S2, at the fifth week of the last multiplication subculture; S3, 5 weeks after the addition of liquid medium containing paclobutrazol (just before the cold treatment); S4, after 13 weeks (the end) of cold treatment; and S5, 6 weeks after the end of cold treatment (during the shoot base swelling/bulb formation phase. The samples were frozen at -20 °C, lyophilised and stored at -20 °C until analyses.

Tissue contents of PAs were also compared during the last two bulbing stages (S4 and S5) between two cultivars that were markedly differing in bulbing ability (easily forming bulbs cultivar FB and recalcitrant cultivar NB) depending on exogenous AA (Arg and Orn each at 500 $\mu M)$ and PA (Put and Spm, each at 50 $\mu M)$ treatments given before cooling (S3).

Four samples containing plant material were collected from four jars (one mixed sample from one jar) representing each treatment. Free polyamines were analysed according to basic procedure of Flores and Galston (1982) with some later modifications (Kosson and Elkner 2010; Horbowicz et al. 2011).

Briefly, plant tissues were homogenised in 5 % (v/v)perchloric acid and amines were derivatised with benzoyl chloride. Bonzoyl derivatives of amines were extracted with ethyl acetate and pooled acetate fractions were evaporated to dryness at 40 °C in stream of air. The residue was dissolved in mobile phase used for HPLC analyses. HPLC analysis was performed with a liquid chromatograph (Agilent Technologies, model 1200 Series). The mobile phase was a mixture of acetonitrile-water (45:55, v/v) at a flow rate of 1.0 mL min⁻¹. Benzoylated amines were eluted isocratically at 30 °C using an RP column Eclipse XDB— C_{18} analytical (4.6 × 150 mm, 5 µm particle size) and C₁₈ guard column. The benzoyl polyamines were detected at 245 nm, DAD detector, and amines contents were calculated from standard curves of commercially available polyamines.

Data were subjected to analysis of variance and treatment means were compared by Duncan's test at P = 0.05.

Results

Bulb formation

The studied PAs and their precursors significantly influenced bulb formation of micropropagated tulips but their effects were strongly differentiated depending on genotype, term of treatment and MeJA application.

In the cultivars easily forming bulbs (FB, P20 and P14), the commencement of bulb enlargement started approximately 6 weeks after the end of cooling and was associated with leaf yellowing. In NB, such symptoms occurred 4 weeks later than in other cultivars and leaf yellowing was visible, even later, about 12–13 weeks after cooling (Fig. 1a).

Results of the *first experiment* with the cultivar NB and PAs/AAs application before cooling (S3) showed that MeJA did not influence bulb numbers and decreased the bulb mass of control shoots (Table 1). Instead, Put and Spm (100 μ M) combined with MeJA treatment significantly enhanced total and regular bulb numbers (Fig. 1b). Regular bulb numbers increased from 7.2 observed in control to ca. 10.5 for Put and Spm treatments.

Results of the second experiment conducted with all four cultivars, in which AAs and PAs were added to the medium before cooling (S3) and standard MeJA treatment was applied, are presented in Table 2. In cultivar NB, all of the studied PAs and AAs markedly enhanced the total bulb number compared to control (9.4). The highest total bulb numbers (21.0-23.8) were noted with ornithine, arginine and putrescine used at lower concentrations. In this cultivar, the numbers of regular bulbs were also ca. twofold higher for Orn and Arg (12.5 and 11.0, respectively) applied at low concentrations compared to control. In the cultivar NB, the mean bulb mass was enhanced significantly (by ca. 36 %) by the application of spermine at 100 μM. In cultivar FB, only ornithine at 1000 μM increased significantly mean bulb mass (584.9 mg) by ca 57 % compared to control. In the cultivar P14 producing numerous small bulbs, AAs and PAs applied at lower concentrations did not influence bulb formation, except for ornithine, which considerably enhanced the total bulb number. Instead, the addition of AAs or PAs at higher concentrations significantly decreased both bulb numbers and mean bulb mass. In cultivar P20, these compounds generally did not have any positive effect on bulb formation. Arg at 500 μM and Put at 50 μM significantly reduced the number of regular bulbs to the half compared to the control.

The results of *the third experiment*, in which the cultivars NB and P14 were used and PA/AA were applied just after the end of cooling (S4), confirmed that MeJA



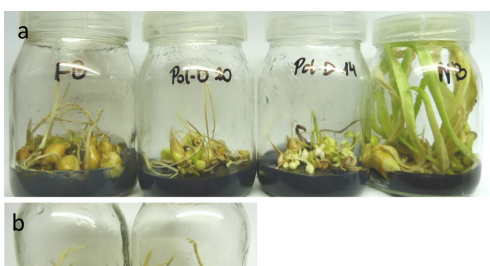




Fig. 1 Differences among micropropagated tulip cultivars in bulbing ability. **a** Bulbs/bulbing shoots 9 weeks after the end of low temperature treatment cultured on a control bulbing medium with standard MeJA treatment; from the left 'Fringed Black' of extremely high bulbing ability (bulb formation process finished), P20—high ability (bulb formation finished), P14—producing numerous small

bulbs (bulb are formed but not yet covered with tunics) and 'New Beet'—delayed and occasionally bulbing (leaves without senescence symptoms). **b** Bulbing shoots/bulbs of 'New Beet' 13 weeks after the end of low temperature treatment with standard MeJA treatment, cultured on a control bulbing medium (left) and with Spm (50 μ M) (right) added to the medium before cooling, at S3

generally decreased bulb numbers, except when applied for the Orn treated shoots (Table 3). In this treatment, the highest total number (21.8) and regular bulb number (18.0) were obtained that differed significantly from the control MeJA+ (13.8 and 12.5, respectively), but were similar to the control MeJA-. The application of Orn, Arg, Put and Spm at S4 without MeJA treatment caused a significant increase of NB bulb mass to ca. 298-332 mg from ca. 200 mg, which was observed in control MeJA-. MeJA treatment, in general, slightly enhanced the bulb mass compared to PA/AA treatments without MeJA application. At the same time, bulb mass (306.4 and 328.5) obtained at MeJA treatment combined with Orn and Spm was significantly higher compared to control MeJA—. In the cultivar P14 (shoots untreated with MeJA), the application of Orn, Put and Spd at S4 caused a marked increase in bulb numbers (Fig. 2).

In the fourth experiment, the cultivars NB and P14 were used and AAs/PAs were applied to the shoot cultures 6 weeks after the end of cooling (S5) and the effects of these compounds on bulb formation were studied depending on MeJA treatment (Table 4). In the cultivar NB, MeJA generally decreased bulb numbers. In P14 shoot cultures treated with Arg or Put, MeJA application also

significantly decreased the total number of bulbs. In this cultivar, however, MeJA combined with Arg resulted with the highest number (9.5) of regular bulbs larger than 100 mg and the highest mean bulb mass (80.4 mg). Besides, both of these values differed significantly from the control MeJA+ (6.8 bulbs of 53.7 mg mean bulb mass, respectively). Orn at 500 μM combined with MeJA also enhanced bulb mass (75.1 mg) in P14 shoots compared to the control MeJA+.

Endogenous polyamine contents

Polyamine patterns and dynamics during the micropropagation process depending on genotype

In standard culture conditions (control medium), significant differences in endogenous PA levels and patterns were noted between genotypes and developmental phases.

Putrescine was generally the most abundant polyamine in three cultivars, NB, P14 and P20 (Fig. 3a). In the cultivar NB, with the lowest bulbing capacity, Put levels were extremely high (1177.9 and 1700.9 nmol g⁻¹ FM) at S1 (shoot multiplication stage) and S2 (5th week of the last

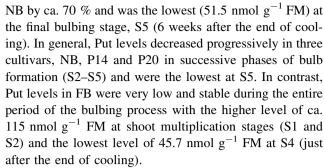


Table 1 Bulb formation in vitro of the tulip cultivar NB following treatments with polyamines and amino acids given before low temperature treatment (S3) depending on MeJA treatment; data recorded 13 weeks after the end of cooling; probability value (P) of F test from two-factor analysis of variance of the bulb formation parameters

Treatment (µM)	MeJA		
	(-)		(+)
Total bulb number			
Control	14.6 ± 3	3.97 bc*	$14.8 \pm 3.42 \text{ bc}$
Ornithine 1000	16.2 ± 3	2.59 abc	$18.8 \pm 5.93 \; abc$
Arginine 1000	14.2 ± 3	3.35 bc	$19.2 \pm 3.56 \text{ ab}$
Putrescine 100	13.6 ± 3	3.58 с	$20.2 \pm 2.59 \text{ a}$
Spermine 100	18.3 ± 4	4.03 abc	$20.8\pm4.03~a$
Effect		P	
AA/polyamines		0.09	
MeJA (-/+)		0.00**	
AA/polyamines × MeJA	(-/+)	0.36	
Regular bulb number			
Control		$8.6\pm1.82~abc$	$7.2 \pm 2.28 \text{ bc}$
Ornithine 1000		9.4 ± 2.01 ab	$7.0 \pm 1.22 \text{ bc}$
Arginine 1000		$7.8 \pm 1.30 \text{ bc}$	$9.2 \pm 2.49 \; abc$
Putrescine 100		6.6 ± 1.95 c	10.4 ± 1.67 a
Spermine 100		$8.8\pm0.96~abc$	10.5 ± 1.73 a
Effect	P		
AA/polyamines	0.27		
MeJA (-/+)	0.22		
AA/polyamines \times MeJA $(-/+)$	0.00		
Mean bulb mass (mg)			
Control	645.	$8 \pm 184.2 \text{ a}$	$430.2 \pm 103.0 \text{ b}$
Ornithine 1000	533.	$7 \pm 102.8 \text{ ab}$	$454.0 \pm 105.2 \text{ b}$
Arginine 1000	509.	$3 \pm 101.3 \text{ ab}$	$483.3 \pm 89.5 \text{ b}$
Putrescine 100	524.	$4 \pm 74.3 \text{ ab}$	$443.5 \pm 102.8 \text{ b}$
Spermine 100	430.	$7 \pm 127.7 \text{ b}$	$474.8 \pm 79.8 \ b$
Effect		P	
AA/polyamines		0.50	
MeJA (-/+)		0.00	
AA/polyamines × MeJA (-/+)		0.12	

^{*} Mean separation within each parameter of growth by Duncan's multiple range test. The mean \pm SD (n = 5) followed by the same letter do not differ at P=0.05

multiplication subculture), respectively. A relatively high Put level at S1 (ca. 580 nmol g⁻¹ FM) was also noted for the cultivar P14. At S1, the Put content in NB was 14.6-fold higher than in FB (ca. 116 nmol g⁻¹ FM). Then, at S3 (just before cooling), Put level significantly decreased in



Spermidine highest levels (ranging from ca. 169 to 397 nmol g⁻¹ FM) were observed in P-14 (the cultivar producing numerous but small bulbs), whereas the lowest levels (from 63.4 to 115.1 nmol g⁻¹ FM) were found in the cultivar P20, with high bulbing ability. In other genotypes, FB and NB, the levels of this compound were slightly lower than in P14 (Fig. 3b). The trends in Spd dynamics were generally similar in all of the genotypes. In P14 at S1 (the shoot multiplication stage), Spd content was 330 nmol g⁻¹ FM. Then, it rose to 389 nmol g⁻¹ FM at S2 (fifth week of the last multiplication subculture) and declined again to 288 nmol g⁻¹ FM at S3 (just before cooling). At S4 (just after the end of cooling), Spd content reached the highest value (397 nmol g⁻¹ FM) to reduce to the lowest content six weeks later (S5) (ca. 170 nmol g⁻¹).

Spermine levels and dynamics differ significantly between genotypes. In general, this polyamine occurred at the lowest concentrations compared to the endogenous contents of Put and Spd. The highest levels (96 and 86 nmol g⁻¹) were noted in P14 at S2 and FB at S3, respectively (Fig. 3c). Similar trends in Spm dynamics were observed for the cultivars P14, P-20 and NB during the stages S1–S4. The contents increased slightly from S1 to S2, and then lowered at S3 (before cooling). During the cooling period, the levels in these three cultivars did not change significantly. Subsequently, between S4 and S5, the contents declined drastically in P14 and FB from ca 72 nmol g⁻¹ to ca. 20 nmol g⁻¹ FM but in NB increased. In genotype P20, Spm contents (13.8–28.7 nmol g⁻¹ FM) were very low and generally stable through the micropropagation stages.

Putrescine to spermidine ratio

The high predominance of Put, especially at shoot multiplication stages, was observed both in the cultivar with low bulbing ability, NB, and easily forming bulbs, P20 (Table 5); Put/Spd ratios ranged from 4.67 to 5.0. In contrast, Spd dominated in the cultivar FB of extremely high bulbing ability in which Put/Spd ratio was during all the phases below 1. Put/Spd ratios of all the cultivars were the high during multiplication stages, S1 and S2. Subsequently, when shoots enter dormancy at S3 (before cooling), Put/Spd ratios decreased markedly and continued to decrease in subsequent



^{**} P value in italic means that effect is significant

Table 2 Bulb formation in vitro in four tulip cultivars depending on exogenous polyamines and their precursors added to a medium before low temperature treatment (S3); all bulbing shoots were treated with MeJA; data recorded 13 weeks after the end of cooling; probability value (*P*) of *F* test from one way-analysis of variance of the bulb formation parameters

Treatment (µM)	Genotype				
	NB	FB	Pol-14	Pol-20	
Total bulb number	•				
Control	$9.4 \pm 2.97 \text{ e*}$	$13.3 \pm 1.26 \text{ ab}$	$25.8 \pm 5.32 \text{ b}$	$25.0 \pm 2.45 \text{ ab}$	
Ornithyne 500	$21.0 \pm 5.10 \text{ abc}$	$10.7 \pm 3.08 \text{ bc}$	$34.0 \pm 4.71 \text{ a}$	20.5 ± 4.12 abcd	
Ornithyne 1000	$16.0 \pm 4.47 \text{ cd}$	$10.5 \pm 4.28 \text{ bc}$	8.3 ± 1.26 e	$18.5 \pm 3.87 \text{ cd}$	
Arginine 500	22.8 ± 6.46 ab	$11.8\pm3.13~ab$	$23.4 \pm 4.45 \text{ bc}$	$19.3 \pm 3.77 \text{ bcd}$	
Arginine 1000	17.8 ± 2.77 abcd	$12.0 \pm 0.82 \text{ ab}$	$17.0 \pm 4.76 d$	20.0 ± 3.65 bcd	
Putrescine 50	$23.8 \pm 5.85 \text{ a}$	$11.2 \pm 2.86 \text{ bc}$	$21.8 \pm 3.95 \text{ cd}$	$15.0 \pm 2.00 \text{ d}$	
Putrescine 100	$16.7 \pm 5.50 \text{ bcd}$	$15.8 \pm 3.20 \text{ a}$	$15.5 \pm 4.20 \text{ d}$	$25.8 \pm 4.21 \text{ a}$	
Spermine 50	$16.2 \pm 5.15 \text{ cd}$	$10.3 \pm 1.97 \text{ bc}$	$23.3 \pm 4.92 \text{ bc}$	$24.0 \pm 4.69 \text{ abc}$	
Spermine 100	$13.7 \pm 2.25 \text{ de}$	$14.3 \pm 3.30 \text{ ab}$	$20.3 \pm 6.55 \text{ bcd}$	$26.0 \pm 4.97 \text{ a}$	
Effect	P = 0.00**	P = 0.11	P = 0.00	P = 0.00	
Regular bulb numb	ber				
Control	$6.8 \pm 1.79 \text{ c}$	$10.0 \pm 3.16 \text{ ab}$	$18.5 \pm 2.89 \text{ a}$	$18.0 \pm 2.16 \text{ ab}$	
Ornithyne 500	12.5 ± 1.91 a	$8.8 \pm 2.48 \text{ b}$	$19.0 \pm 4.69 \text{ a}$	$13.0 \pm 6.38 \text{ bcd}$	
Ornithyne 1000	$8.0 \pm 2.83 \text{ c}$	$9.5 \pm 3.78 \text{ ab}$	4.0 ± 2.16 e	$14.0 \pm 2.94 \text{ bcd}$	
Arginine 500	$11.0 \pm 4.24 \text{ ab}$	$7.8 \pm 2.56 \text{ b}$	$13.6 \pm 3.65 \text{ a-d}$	$11.3 \pm 3.30 \text{ cd}$	
Arginine 1000	$6.4 \pm 3.05 \text{ c}$	$10.0 \pm 2.94 \text{ ab}$	$8.5 \pm 2.08 \text{ de}$	14.5 ± 2.38 abcd	
Putrescine 50	$7.7 \pm 2.50 \text{ c}$	$9.3 \pm 3.39 \text{ ab}$	$14.5 \pm 4.93 \ abc$	$9.5 \pm 1.00 \text{ d}$	
Putrescine 100	$9.0 \pm 2.45 \text{ bc}$	12.3 ± 2.99 a	$9.3 \pm 3.40 \text{ cd}$	$18.4 \pm 3.29 \text{ a}$	
Spermine 50	$8.2 \pm 2.86 \text{ c}$	$8.0 \pm 1.79 \text{ b}$	15.8 ± 4.27 ab	16.0 ± 2.16 abc	
Spermine 100	$7.5 \pm 2.43 \text{ c}$	$10.8\pm1.50~ab$	$12.0 \pm 3.46 \text{ bcd}$	$16.8 \pm 3.40 \text{ ab}$	
Effect	P = 0.00	P = 0.05	P = 0.00	P = 0.01	
Mean bulb mass (n	ng)				
Control	$310.4 \pm 42.90 \text{ b}$	$372.6 \pm 45.0 \text{ bcd}$	$113.3 \pm 43.57 \text{ ab}$	257.3 ± 60.01 a	
Ornithyne 500	$279.3 \pm 82.87 \text{ b}$	$474.8 \pm 52.2 \ ab$	$72.0 \pm 28.62 \text{ a-c}$	$256.5 \pm 72.08 \; a$	
Ornithyne 1000	$243.2 \pm 67.00 \text{ b}$	$584.9 \pm 46.3 \text{ a}$	$65.1 \pm 30.45 \text{ bc}$	257.6 ± 35.36 a	
Arginine 500	$234.2 \pm 71.13 \text{ b}$	$456.2 \pm 95.9 \text{ ab}$	120.2 ± 39.72 a	300.6 ± 43.88 a	
Arginine 1000	$326.1 \pm 151.04 \text{ b}$	$422.0 \pm 70.5 \text{ b}$	$90.3 \pm 37.93 \text{ a-c}$	248.6 ± 45.33 a	
Putrescine 50	$277.0 \pm 50.15 \text{ b}$	$408.4 \pm 96.7 \ \mathrm{bc}$	$86.9 \pm 44.92 \text{ a-c}$	282.2 ± 61.56 a	
Putrescine 100	$252.3 \pm 67.97 \text{ b}$	$445.9 \pm 113.2 \text{ b}$	$71.2 \pm 29.00 \text{ a-c}$	231.57 ± 85.52 a	
Spermine 50	$246.5 \pm 48.64 \text{ b}$	$283.8 \pm 77.5 \text{ cd}$	$66.4 \pm 18.59 \text{ bc}$	$230.8 \pm 64.59 \text{ a}$	
Spermine 100	423.1 ± 36.75 a	$261.7 \pm 36.5 \text{ cd}$	$48.4 \pm 22.23 \text{ c}$	260.5 ± 156.29 a	
Effect	P = 0.00	P = 0.00	P = 0.05	P = 0.95	

^{*} Mean separation within columns by Duncan's multiple range test. The mean \pm SD (n = 5) followed by the same letter do not differ at P=0.05

phases; the lowest values were generally noted at the final bulbing phase S5.

Polyamine tissue contents at S4 and S5 depending on exogenous PA/AA treatments and genotype

Exogenously applied AAs and PAs (added to the bulbing medium before cooling) strongly influenced tissue contents of polyamines measured in bulbing shoots of FB and NB at S4 (just after the end of cooling) and six weeks later, at S5.

At S4, *putrescine accumulation* (728.3–817.5 nmol g⁻¹ FM) in NB shoots was markedly enhanced by exogenously applied Orn and Spm. Unexpectedly, the exogenously applied putrescine strongly reduced tissue content of this polyamine (140.0 nmol g⁻¹ FM) compared to the control (302.0 nmol g⁻¹ FM) (Fig. 4a). At the same time, Put tissue contents were much higher in NB compared to FB. Six weeks later, at S5, Put contents drastically decreased in NB; in control, Put level was even twofold lower than in FB (92.9 nmol g⁻¹ FM) (Fig. 4b). However, in PA/



^{**} P value in italic means that effect is significant

Table 3 Bulb formation in vitro of the tulip cultivar NB following the treatments with polyamines and amino acids given after the end of low temperature treatment (S4) depending on MeJA treatment; data recorded 13 weeks after cooling; probability value (P) of F test from two-factor analysis of variance of the bulb formation parameters

Treatment (µM)	MeJA		
	_	+	
Total bulb number			
Control	19.7 abc*	13.8 d	
Ornithine 500	17.6 abcd	21.8 a	
Arginine 500	19.2 abc	16.0 bcd	
Putrescine 50	15.3 cd	14.0 d	
Spermidine 50	19.2 abc	18.3 abc	
Spermine 50	20.7 ab	17.7 abc	
Effect	P		
AA/polyamines	0.01**		
MeJA	0.04		
AA/polyamines × MeJA	0.07		
Total number of regular bulb	S		
Control	16.6 ab	12.5 с	
Ornithine 500	13.4 abc	18.0 a	
Arginine 500	14.6 abc	14.2 ab	
Putrescine 50	12.6 c	10.2 c	
Spermidine 50	16.8 ab	13.3 ab	
Spermine 50	16.5 ab	13.7 ab	
Effect	P		
AA/polyamines	0.09		
MeJA	0.14		
AA/polyamines × MeJA	0.13		
Mean bulb mass (mg)			
Control	198.9 bc	288.8 al	
Ornithine 500	298.9 a	306.4 a	
Arginine 500	248.7 ab	277.8 al	
Putrescine 50	332.4 a	260.3 al	
Spermidine 50	157.2 c	263.5 al	
Spermine 50	298.4 a	328.2 a	
Effect	P		
AA/polyamines	0.00		
MeJA	0.06		
AA/polyamines × MeJA	0.05		

^{*} Mean separation within each bulb growth parameter by Duncan's multiple range test. The mean \pm SD (n = 5) followed by the same letter do not differ at P=0.05

AA treatments, Put tissue contents in NB remained slightly higher than in FB, with the highest value found for Orn treatment (113.1 nmol g^{-1} FM).

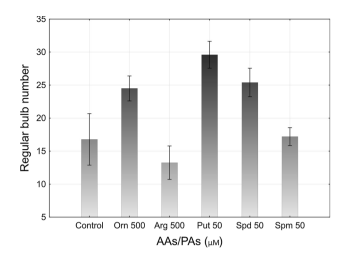


Fig. 2 Regular bulb number of tulip cultivar P14 under treatments with amino acids (Orn and Arg) and polyamines (Put, Spd and Spm) added to the medium just after the end of cooling (S4); Mean \pm SD (n = 5), P = 0.00

At S4, *spermidine* tissue contents were not influenced by AA/PA treatments and were similar in both cultivars, irrespective of exogenously applied AAs/PAs, ranging from ca. 269 to 342 nmol g⁻¹ FM (Fig. 4c). The significant effects of AAs and PAs were visible six weeks later (S5) when Spd levels decreased, especially in FB. In this cultivar, the lowest Spd level (71.8 nmol g⁻¹ FM) was observed with the Put treatment and the highest Spd level (131.8 nmol g⁻¹ FM) was recorded in control (Fig. 4d). In NB treated with exogenous Orn or Spm, the Spd contents decreased slightly at S5 compared to S4 but were still relatively high: 262.4 and 201.1 nmol g⁻¹ FM, respectively.

Spermine tissue contents at S4 were generally higher in FB than in NB. Compared to control, a significant increase in Spm accumulation was noted at S4 and S5 in both cultivars when exogenous Spm was added to the medium (Fig. 4e, f). In NB, the marked increase of Spm tissue content was also recorded for Orn and Arg treatments at S4. Interestingly, Orn application resulted in decrease of Spm at S5. In FB, the significant enhancement of Spm accumulation was visible 6 weeks later (at S5) for all of the AA/PA treatments.

Discussion

Our research on micropropagated tulips showed that the effects of PAs or their precursors were highly dependent on genotype, application term, compound concentration and MeJA treatment. Positive effects of PAs or AAs on bulb formation in vitro were generally found in cultivar NB, with the lowest bulbing ability. All of the studied AAs (Arg and Orn) and PAs (Put and Spm) markedly enhanced bulb



^{**} P value in italic means that effect is significant

Table 4 Bulb formation in vitro of the tulip cultivars NB and P14 following the treatments with polyamines and amino acids given 6 weeks after the end of low temperature treatment (S5) depending on

MeJA treatment; data recorded 13 weeks after cooling; probability value (P) of F test from two-factor analysis of variance of the bulb formation parameters

$Treatment \; (\mu M)$	Genotype					
	NB		Pol-14			
	MeJA					
	_	+	-	+		
Total bulb number						
Control	21.8 ± 3.60 abc*	$16.6 \pm 3.36 \text{ c}$	63.0 ± 13.25 bcd	$71.2 \pm 16.51 \text{ abc}$		
Ornithine 500	$19.3 \pm 2.29 \text{ bc}$	$17.7 \pm 1.03 \text{ c}$	$77.3 \pm 5.50 \text{ ab}$	$66.8 \pm 115.7 \text{ abcc}$		
Arginine 500	$21.8 \pm 4.17 \text{ abc}$	$18.7 \pm 2.16 \text{ bc}$	$82.7 \pm 2.87 \text{ a}$	$64.0 \pm 8.29 \text{ bcd}$		
Putrescine 50	$20.3 \pm 3.30 \text{ abc}$	$18.3 \pm 4.03 \text{ bc}$	$79.0 \pm 12.03 \text{ ab}$	$50.7 \pm 14.35 \text{ d}$		
Spermidine 50	$23.9 \pm 5.61 \text{ ab}$	$20.0 \pm 6.65 \text{ abc}$	$79.8 \pm 13.48 \text{ ab}$	62.4 ± 8.79 bcd		
Spermine 50	25.6 ± 7.18 a	$16.3 \pm 7.31 \text{ c}$	$56.2 \pm 5.63 \text{ cd}$	$59.4 \pm 9.61 \text{ bcd}$		
Effect	P					
AA/polyamines	0.42		0.07			
MeJA	0.00**		0.00			
AA/polyamines \times MeJA	0.35		0.01			
Number of regular bulbs	(>100 mg)					
Control	$11.3 \pm 2.58 \text{ ab}$	$7.6 \pm 2.37 \text{ c}$	6.2 ± 0.84 bcd	6.8 ± 1.48 bcd		
Ornithine 500	$10.6 \pm 1.90 \text{ abc}$	$9.0 \pm 1.79 \text{ bc}$	7.0 ± 2.16 abcd	$7.3 \pm 1.63 \text{ abcc}$		
Arginine 500	$11.0 \pm 1.10 \text{ abc}$	$10.5 \pm 1.64 \text{ abc}$	$5.0 \pm 0.82 \; \mathrm{d}$	$9.5 \pm 3.00 \text{ a}$		
Putrescine 50	$12.0 \pm 4.16 \text{ ab}$	$10.4 \pm 3.21 \; abc$	7.8 ± 2.06 abcd	5.5 ± 1.97 cd		
Spermidine 50	$13.3 \pm 4.11 \text{ a}$	$8.4 \pm 2.56 \text{ bc}$	$9.0 \pm 1.00 \text{ a}$	$7.0 \pm 1.87 \text{ bcd}$		
Spermine 50	11.3 ± 2.56 ab	8.0 ± 2.97 bc	8.0 ± 2.12 abc	5.8 ± 2.28 cd		
	P					
AA/polyamines	0.45		0.54			
MeJA	0.00		0.74			
AA/polyamines × MeJA	0.32		0.01			
Total number of regular i	bulbs					
Control	$20.5 \pm 2.59 \; abc$	$14.1 \pm 4.18 d$	$47.6 \pm 11.48 \text{ cd}$	$52.0 \pm 11.42 \text{ abcd}$		
Ornithine 500	17.0 ± 2.00 abcd	$15.2 \pm 2.40 \text{ cd}$	55.8 ± 5.00 abcd	$54.0 \pm 14.20 \text{ abcc}$		
Arginine 500	$19.0 \pm 3.40 \text{ abcd}$	$18.0 \pm 3.03 \text{ abcd}$	$63.0 \pm 2.45 \text{ ab}$	51.3 ± 13.18 abcc		
Putrescine 50	18.1 ± 3.24 abcd	$15.7 \pm 2.98 \text{ bcd}$	$60.3 \pm 8.62 \text{ abcd}$	$43.0 \pm 12.82 \text{ d}$		
Spermidine 50	$20.9 \pm 5.81 \text{ ab}$	$15.6 \pm 5.78 \text{ bcd}$	67.2 ± 5.17 a	$48.8 \pm 7.90 \text{ bcd}$		
Spermine 50	$22.0 \pm 6.32 \text{ a}$	$13.8 \pm 6.08 \; d$	$43.4 \pm 4.77 \text{ cd}$	$53.6 \pm 17.78 \text{ abcc}$		
	P					
AA/polyamines	0.72		0.08			
MeJA	0.00		0.06			
AA/polyamines × MeJA	0.24		0.12			
Mean bulb mass (mg)						
Control	$194.9 \pm 29.23 \text{ ab}$	190.2 ± 26.66 ab	66.3 ± 15.97 abcd	$52.7 \pm 10.14 \text{ cd}$		
Ornithine 500	$212.7 \pm 24.71 \text{ ab}$	$182.2 \pm 27.52 \text{ b}$	$58.7 \pm 7.77 \text{ bcd}$	$75.1 \pm 20.99 \text{ ab}$		
Arginine 500	212.7 ± 39.80 ab	$185.9 \pm 39.83 \text{ b}$	$49.5 \pm 8.65 \text{ d}$	80.4 ± 16.86 a		
Putrescine 50	197.1 ± 38.86 ab	$218.6 \pm 48.19 \text{ ab}$	$61.9 \pm 1.88 \text{ abcd}$	55.7 ± 10.07 bcc		
Spermidine 50	$245.0 \pm 44.20 \text{ a}$	$186.5 \pm 47.52 \text{ b}$	$62.4 \pm 9.65 \text{ abcd}$	71.1 ± 12.40 abo		
Spermine 50	220.1 ± 20.22 ab	218.3 ± 65.72 ab	$70.6 \pm 10.49 \text{ abc}$	62.5 ± 8.83 abcd		



Table 4 continued

	P	
AA/polyamines	0.60	0.51
MeJA	0.09	0.17
AA/polyamines × MeJA	0.25	0.01

^{*} Mean separation within each bulb growth parameter by Duncan's multiple range test. The mean \pm SD (n = 5) followed by the same letter do not differ at P = 0.05

numbers in this cultivar when used at lower concentrations at the beginning of the bulbing process (before cooling, at S3), together with standard MeJA treatment, and only Spm increased the bulb mass. In other cultivars, FB, P14 and P20, all with high bulbing ability, PAs and AAs substantially did not influence or even markedly reduce bulbing efficiency. As in tulip cultivar NB, bulb formation improvement by the application of different PAs or AAs was obtained for Lilium longiflorum as well (Tanimoto and Matsubara 1995). In this lily species, the most effective was Spm at 100 μM. Also, Varshney et al. (2000) showed that the incorporation of Spm (150 mM) or Put (50 µM) to culture medium stimulated bulblet multiplication of lily Asiatic hybrids. In their study, however, mean bulb mass did not show any significant increase. In our study, the increment of the bulb mass was expected most of all in cultivar P14. That was achieved at Arg treatment given at S5 combined with MeJA application.

Corresponding data for the positive role of Arg in tulip bulb formation were reported by Le Nard and Fiala (1990). They found that high Arg content in mother bulbs (in vivo) before low temperature treatment was correlated with the subsequent more efficient formation of daughter bulbs. In turn, Kollöffel et al. (1992), monitoring levels of free PAs in the various parts the tulip bulbs in vivo during the 12 week-storage at different temperatures (5 or 17 °C), found that irrespective of the temperature all free PA contents increased during storage and Spd dominated in all bulb parts.

More evidence for the role of PAs in storage organ formation in vitro have been reported for tuberous geophytes. Thus, Put, Spd and Spm at 100 or 500 μM increased the number of gladiolus cormlets, with the maxima observed for Spd treatment (Kumar et al. 2011). That was similar to our results concerning tulip cultivar NB, in which all these compounds enhanced number of bulbs. An advantageous effect of Spd (1 μM) was also demonstrated for potato tuber formation (Mader 1995). In yam, both PAs (Put, Spd and Spm) and AAs (Arg and Orn) added to a medium at various concentrations (1–1000 μM) markedly accelerated tuber development, and significantly enhanced tuber sizes (Ondo Ovono et al. 2010a, b).

Analysis of the endogenous PA contents and patterns have provided additional evidence of polyamine role in bulbing process. We found that Put was the major polyamine in three tulip genotypes at the stages of shoot multiplication (S1-S2), in the presence of TDZ and/or iP. Then, Put levels progressively declined in all of the cultivars and were lowest at the final bulbing stage (S5). Spd manifested similar dynamics in all of the cultivars, showing two peaks: at shoot multiplication (S2) and at the end of cooling (S4). Spm contents were generally higher at multiplication stages (S1-S2), mainly in the cultivars producing bulbs easily (FB and P14). Then, its contents declined to the lowest levels at the final bulbing stage (S5), except for cultivar NB, in which Spm increased at this time. This phase in the cultivars easily forming bulbs (FB, P20 and P14) was associated with the commencement of bulb enlargement and the simultaneous development of dormancy manifesting with leaf yellowing, as previously reported for another easy bulbing tulip cultivar 'Prominence' (Podwyszyńska et al. 2014). In the cultivar NB, such symptoms were not visible at S5 (6 weeks after the end of cooling). Bulbs of this cultivar started to enlarge 4 weeks later than in other cultivars and leaf yellowing was visible even later, about 12-13 weeks after cooling.

As in our study, high endogenous Put level at the multiplication stage was detected in shoots of taro (Colocasia esculenta), while exogenously applied Spd or Spm induced tuberisation (Francisco et al. 2008). Considering these results and our data, we concluded that the transient increase of Spd levels after the low temperature treatment (S4) associated with dormancy release could be involved in the initiation of the bulbing process. This phase of bulb or any other storage organs development (e.g. tubers or corms) characterized with cell differentiation into storage tissue (De Hertogh and Le Nard 1993; Rees 1992; Podwyszyńska 2012). We further observed that despite the fact that Spd content decreased at S5 together with other PAs, Spd was generally the major polyamine at the final bulbing stages (S4 and S5). This Spd prevalence was evidenced by the generally low Put/Spd ratios in all of the cultivars at the final bulbing phase. In three cultivars these ratios were



^{**} P value in italic means that effect is significant

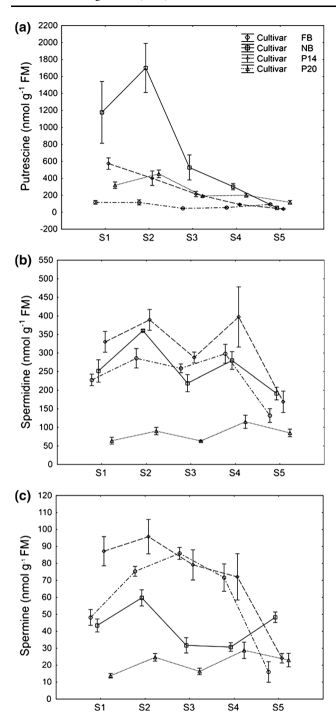


Fig. 3 Contents of endogenous polyamines in the in vitro shoots/bulbing shoots of the tulip cultivars 'Fringed Black' (FB), 'New Beet' (NB), P14 and P20 during successive micropropagation stages: S1, after 6 weeks of the standard multiplication subculture; S2, at the fifth week of the last multiplication subculture; S3, 5 weeks later—just before cold treatment; S4, at the end of the 13-week-cold treatment; S5, 6 weeks after the end of cold treatment (at the shoot base swelling/bulb formation phase); mean \pm SE (n = 4), F and P for ANOVA Phase \times Cultivar are given for each endogenous polyamine: a putrescine: F = 6.8, P = 0.00, b spermidine: F = 2.6, P = 0.01, c spermine: F = 9.22, P = 0.00

below 1 and in one cultivar it was ca. 1.4. Earlier, analogous hypothesis that tuberising conditions in potato characterized with low Put/Spd ratios, below 1, was reported by Mader (1995). All of these observations correspond to the reported findings that Put is the most abundant at cell division, but Spd and Spm are essential for inducing cell differentiation, as reviewed by Bais and Ravishankar (2002) and Anwar et al. (2015).

Our results have provided other interesting evidence on the role of Spd in the bulbing process. In cultivar FB, with extremely high bulbing ability, Spd prevalence was observed at all of the micropropagation stages, even at shoot multiplication. Besides, the shoots of this cultivar, unlike other genotypes, have a tendency to produce some bulbs, even under non-inducing conditions, i.e. during shoot multiplication in the presence of cytokinins, without low temperature treatment.

In turn, Kumar and Palni (2013) noted that the levels of all PAs showed an increase during the middle phase of gladiolus cormlet development in vitro and Spd was found to be the major polyamine. Then PA levels declined during cormlet maturation. We also demonstrated that the contents of all of the PAs decreased in developing bulbs. We suppose that it could be associated with ABA increment and dormancy development of tulip bulbing shoots (Podwyszynska et al. 2004). Furthermore, we postulated that low bulbing ability of the cultivar NB may be related to the up-regulation of Spm biosynthesis at S5, since all PA levels, including Spm, declined in all of the cultivars producing bulbs easily. We suppose that such Spm increase at S5 could interfere in dormancy development in the NB cultivar.

Morphogenic responses of tulip shoots in vitro to polyamines and their precursors can be better understood in light of numerous evidences showing that polyamines and PGRs such as ethylene, auxins, cytokinins and ABA mutually influenced the metabolism of each other, as well as the actions and translocation (Bais and Ravishankar 2002; Anwar et al. 2015). The authors cited numerous reports on interactions between polyamines and PGRs in several morphogenic and physiological processes such as, e.g. cell division and senescence. Since PAs and ethylene derive from and compete for a common precursor S-adenosylmethionine (SAM), and ethylene is known to stimulate senescence, the polyamines are considered to counteract this process, e.g. leaf senescence (Bais and Ravishankar 2002; Anwar et al. 2015). The results of our study indicate that delayed leaf senescence and bulb formation in cultivar NB can be associated with higher levels of endogenous Spm at the final bulbing stage compared to other cultivars. That could reduce the biosynthesis of ethylene, whereas it was found in our earlier study that



Table 5 Rations of Put/Spd in shoots and bulbing shoots of micropropagated tulips of each cultivar, in each phase

Cultivar	Put/Spd ratio Micropropagation phases						
	NB	4.67	4.72	2.42	1.08	0.27	
FB	0.51	0.40	0.18	0.18	0.70		
P14	1.74	1.03	0.74	0.23	0.23		
P20	4.96	5.0	3.0	1.74	1.37		

ethylene promoted tulip bulb growth in vitro (Podwyszyńska 2006a).

Previous investigations revealed that the low bulbing ability and related delayed leaf yellowing of another tulip cultivar 'Blue Parrot' was associated with large CK accumulation during last phase of the bulb formation (Podwyszyńska et al. 2014). Rakova and Romanov (2005) based on their studies with seedling of transgenic Arabidopsis indicated that endogenous PAs can significantly modulate an amplitude of a CK primary response gene expression in vivo. These authors speculated that cytokinin-induced accumulation of PAs (Sergiev et al. 1995) can be a response, alleviating the consequences of plant treatment with excessive CK doses. Such a situation could occur in the NB shoots at multiplication stages (S1-S2) when this cultivar exhibits extremely high Put levels and a subsequent drastic drop at S3, as the response to high cytokinin level at S1.

Ondo Ovono et al. (2010a, b) reported that treatment with Put (10⁻⁵ M) resulted in a dramatic increase in the contents of both endogenous auxin (ten times) and Put in vam tubers, which was associated with better tuber development. Such PA-mediated auxin increment could promote tulip bulb development in the present study, since we found positive effects of exogenous auxins on tulip bulb formation in our previous investigation (Podwyszyńska 2006a, b). Furthermore, Ondo Ovono et al. (2010a, b) demonstrated that exogenous Put at the concentrations of 0.1-10 µM (improving tuber formation) did not influence endogenous Put content at the stage when the tuber was already formed and had already grown. However, the addition of polyamines at higher concentrations (1000 µM) in plant medium enhanced their accumulation in a plant tissue, while it delayed tuber formation. This partly corresponds with our observation referring to the cultivar NB. Thus, shoots of this cultivar characterized with extremely high Put levels at S1 and S2 that could affect subsequent bulb formation retarding this process. This supposition ca be further confirmed by our observations that the exogenous Put decreased its tissue content contrary to the mentioned above results obtained for yam tubers (Ondo Ovono 2010a). Our results indicate a negative feedback regulation of this PA that is in agreement with the known mechanism of maintaining PA homeostasis through one of the mechanisms, e.g. polyamine (Spd and Spm) catabolism catalysed by polyamine oxidases (Ahou et al. 2014) and Put oxidation by copper containing amine oxidase (Tavladoraki et al. 2012). PAs themselves can induce alterations in the expression levels of genes involved in PA metabolism through various regulatory mechanisms, including inhibitory molecules, as reviewed by Ahou et al. (2014) and Anwar et al. (2015).

We observed strong interaction between PAs and MeJA. The cultivar NB of low bulbing ability responded to MeJA application different to other genotypes. Although MeJA was found to significantly improve bulb formation of several micropropagated tulip cultivars (Podwyszyńska 2006a; Podwyszyńska and Sochacki 2010), the NB shoots treated with MeJA alone produced a similar or even smaller number of bulbs compared to untreated shoots. Instead, AA/PA in combination with MeJA treatment enhanced the number of bulbs obtained. Contrary to our results, Ondo Ovono et al. (2010b) reported that both PAs and MeJA had positive effects on in vitro tuberisation of Dioscorea when applied alone and negative when in combination. On the other hand, Biondi et al. (2001) showed that MeJA treatments decreased the endogenous levels of free PAs while increasing the accumulation of their conjugates. These authors also found that depletion of the free PA pool occurred in spite of the activation of all biosynthetic enzymes, such as Arg, Orn and S-adenosylmethionine decarboxylases. They supposed that free PAs were utilised to form conjugates. Moreover, Biondi et al. (2001) revealed that diamine oxidase (DAO) activity was also stimulated by MeJA and DAO was suggested to generate the hydrogen peroxide needed in cell expansion and/or lignification (Angelini et al. 1993, cited after Biondi et al. 2001). In general, cell enlargement is characteristic of the later stages of tuberisation process such as initiation and growth of



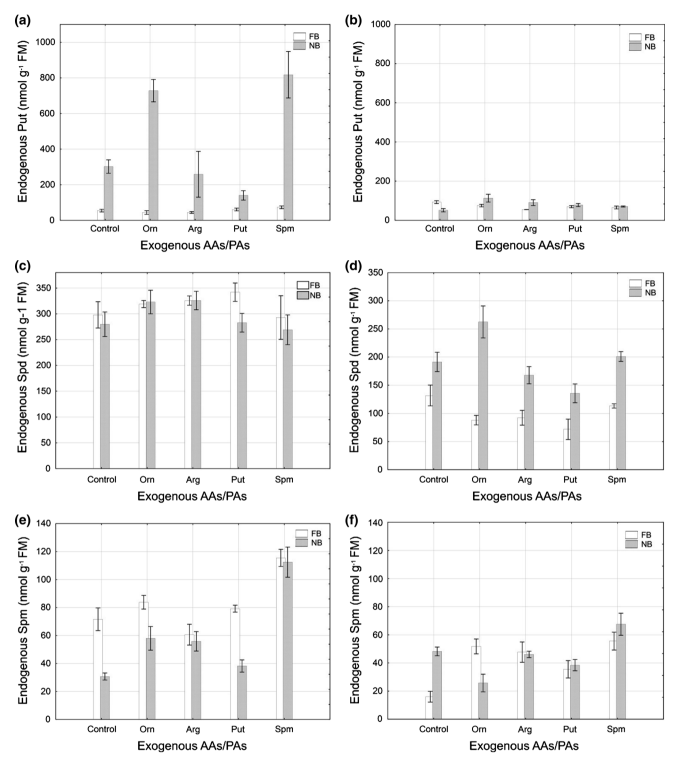


Fig. 4 Tissue contents of polyamines in the in vitro bulbing shoots of the tulip cultivars 'Fringed Black' (FB) and 'New Beet' (NB) at bulbing stages S4 (**a**, **c**, **e**) (at the end of the 13-week-cold treatment) and S5 (**b**, **d**, **f**) (6 weeks after the end of cold treatment) followed treatments with exogenous amino acids (Orn and Arg) and polyamines (Put, Spd and Spm) added to the medium before cold treatment

(S3); mean \pm SE (n = 4), F and P for ANOVA cultivar \times exogenous AA/PA for each tissue content polyamine at S4 or S5 are as follows: **a** putrescine S4: F = 11.2, P = 0.00, **b** putrescine S5: F = 5.4, P = 0.00; **c** spermidine S4: F = 0.6, P = 0.68; **d** spermidine S5: F = 4.3, P = 0.01; **e** spermine S4: F = 3.8, P = 0.01; **f** spermine S5: F = 6.9, P = 0.00



storage organs (De Hertogh and Le Nard 1993; Rees 1992; Podwyszyńska 2012). Besides, storage organ initiation is associated with reorientation of the direction of both cell division and radial expansion in the region of subsequent swelling (Koda 1997; Vreughdenhil et al. 1999), that has been demonstrated to be induced by jasmonates in tuber formation from stolon apex in potato (Koda 1997; Cenzano et al. 2003) and bulbs from leaf sheath in onion or leaf bases in narcissus (Nojiri et al. 1992; Koda 1997; Zaffryar et al. 2007).

Modulation of hormonal status by polyamines during tulip bulb formation is a complex and long-lasting process. PAs positively influenced bulb formation of the low bulbing ability cultivar NB when applied before low temperature treatment. Since PAs were reported to increase IAA and ABA contents (Ondo Ovono et al. 2010a, b; Steiner et al. 2007) and modulate CK contents (Biondi et al. 2003; Rakova and Romanov 2005), they could positively influence bulb formation of the tulip through mutual interactions with these hormones. This assumption is consistent with the general summary of information, referring to the changes in transcriptome linked to the plant hormone/ polyamine interactions presented by Anwar et al. (2015). The authors concluded that Put was positively related to the expression of genes regulating ABA biosynthesis but down-regulated genes of ethylene, jasmonates, and gibberellin biosynthesis, while the Spd action was found to be exactly opposite. Spm enhanced genes for ethylene and jasmonates biosynthesis but down-regulated those for gibberellins and ABA biosynthesis.

Considering all of the acquired data, we postulate that the high bulbing competence may be related to a transient increase in Spd content in response to low temperature treatment (S4), followed by its rapid decrease during bulb growth and maturation (S5, 6 weeks after the end of cooling) when developing bulbs begin to enter dormancy. The low levels of all free PAs at final bulbing stage as well as low Put/Spd ratio may also aid in the bulb formation of tulip in vitro.

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