Functional characterization and developmental expression profiling of gibberellin signalling components in *Vitis vinifera*

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Abstract

Gibberellins (GAs) regulate numerous developmental processes in grapevine (*Vitis vinifera*) such as rachis elongation, fruit set, and fruitlet abscission. The ability of GA to promote berry enlargement has led to its indispensable use in the sternospermocarpic (‘seedless’) table grape industry worldwide. However, apart from VvGAI1 (VvDELLA1), which regulates internode elongation and fruitfulness, but not berry size of seeded cultivars, little was known about GA signalling in grapevine. We have identified and characterized two additional DELLAs (VvDELLA2 and VvDELLA3), two GA receptors (VvGID1a and VvGID1b), and two GA-specific F-box proteins (VvSLY1a and VvSLY1b), in cv. Thompson seedless. With the exception of VvDELLA3-VvGID1b, all VvDELLAs interacted with the VvGID1s in a GA-dependent manner in yeast two-hybrid assays. Additionally, expression of these grape genes in corresponding *Arabidopsis* mutants confirmed their functions in planta. Spatiotemporal analysis of VvDELLAs showed that both VvDELLA1 and VvDELLA2 are abundant in most tissues, except in developing fruit where VvDELLA2 is uniquely expressed at high levels, suggesting a key role in fruit development. Our results further suggest that differential organ responses to exogenous GA depend on the levels of VvDELLA proteins and endogenous bioactive GAs. Understanding this interaction will allow better manipulation of GA signalling in grapevine.

Key words: F-box proteins, GA receptors, gibberellins (GAs), gibberellin signalling, grapevine (*Vitis vinifera*), VvDELLA proteins.

Introduction

Bioactive gibberellins (GA) are phytohormones involved in major physiological processes (Fleet and Sun, 2005; Yamaguchi, 2008). The most common bioactive GAs in higher plants are GA₁, GA₃, and GA₇, a small subset of the more than 136 GAs identified (MacMillan, 2001). The richest sources of GAs in most plants are seeds (Dennis Jr. and Nitsch, 1966), inflorescences, and nodes (Kaufman *et al.*, 1976). In seeds, GA is synthesized in the embryo (Kaneko *et al.*, 2003), while the tapetum of anthers, stamens, flower receptacles, and rosette leaves constitute specific synthesis sites in flowering plants (Kaneko *et al.*, 2003; Hu *et al.*, 2008). The metabolic processes regulating GA biosynthesis and/or deactivation in these organs are tightly controlled by its concentration, as well as developmental, hormonal, and
environmental cues (Yamaguchi, 2008). Generally, GA levels are elevated in reduced GA-response mutant backgrounds, and vice versa (Thomas et al., 1999; Ueguchi-Tanaka et al., 2005; Griffiths et al., 2006).

The involvement of GAs in grapevine (Vitis vinifera) berry development and size determination was first described by Coombe (1960). In the berries of seeded cultivars, GAs are produced by the seeds (Lavee, 1960). Application of GA to seeded grape cultivars prior to anthesis increases fruit set and induces parthenocarpy (Chundawat et al., 1971); causes lignification and contortion of the rachis (Agüero et al., 2000); and induces berry enlargement in some cultivars (Nijjar and Bhatia, 1969), while reducing berry size in others (Chundawat et al., 1971).

In stenospermocarpic cultivars, which represent the majority of the 'seedless' commercial cultivars, fertilization is followed by endosperm abortion, which occurs early during fruit development and leads to cessation of seed development. In these cultivars, seeds serve as the primary source of GA only prior to abortion (Conde et al., 2007). Thus, the berries of these cultivars are usually small with relatively low levels of GA (Iwahori et al., 1967). To enhance berry size and viticulturally important traits such as rachis elongation and berry abscission, seedless cultivars have been treated with GA since the late 1950s (Weaver, 1958). However, GA application at other developmental stages can have negative effects on reproductive development in grapes: when applied at anthesis GAs may induce formation of shot-berry, and enhance berry abscission (Mosesian and Nelson, 1968). In addition, early application of GA to shoots may reduce fruitfulness due to the development of uncommitted primordia to tendrils (Srinivasan and Mullins, 1981).

Seedless cultivars show a wide range of sensitivity to GA treatment (Wolf and Loubser, 1992). Parallel to these varietal differences, differential GA sensitivity has also been reported in tissues/organs within the cultivar (Mullins et al., 1992; Agüero et al., 2000), although the basis for these differences remains unknown. To understand the underlying mechanism of GA response in grapevine in general, and in the berry in particular, it is imperative to elucidate the grapevine GA signalling components. Recent studies showed that GA receptors and the early GA signalling pathway are highly conserved in higher plants (Hirano et al., 2008; Sun, 2011). The GA signal is perceived by its nuclear receptor GA INSENSITIVE DWARF1 (GID1). GA activates its signalling pathway by promoting interaction between GID1 and DELLA, a GA signalling repressor. Binding of GA-GID1 to DELLA induces recognition of DELLA for ubiquitination by a specific F-box protein (SLY1 in Arabidopsis and GID2 in rice), a subunit of the ubiquitin E3 ligase SCF complex. Once polyubiquitinated, DELLA is degraded rapidly by the 26S proteasome. Thus far, the only characterized GA signalling component in grapevine has been VvGAII (Boss and Thomas, 2002), which is a DELLA protein that represses internodes and rachis elongation, but has no effect on berry size. This implies that other component(s) of the GA signalling cascade may regulate grape berry size.

In this study, we isolated and functionally characterized all the grapevine homologues of DELLA, GID1, and SLY1. We also carried out spatiotemporal analysis of VvDELLA, VvGID1, and VvSLY1 transcripts, and of VvDELLA proteins, and investigated their regulation in response to GA3 and the GA biosynthesis inhibitor [paclitaxelazol (PAC)] treatments. One of the major findings of this study is that the response of an organ to GA3 application depends on the total amount of VvDELLA proteins and bioactive GAs in the organ.

Materials and methods

Field experiment and sampling

The experiments were conducted in the 2010 and 2011 growing seasons, on 10-year-old Thompson seedless vines (Vitis vinifera L.) growing in a vineyard in Moshav Lachish, Israel (N31°33′33″; E34°51′26″). The planting system was composed of Y-shaped trellises with planting distances of 3 × 1 m on Richter 110 (V. berlandieri × V. rupestris) rootstock. Standard cultural practices were applied in the vineyard. All experiments were replicated three times using eight-vine plots arranged in a randomized complete-block design.

Treatment with GA3 and PAC

Groups of 25 uniform 15 cm shoots and 25 inflorescences [E-L 15, on the Modified Eichhorn and Lorenz system (Coombe, 1995)] were selected on vines of similar vigour for internode and rachis experiments, respectively. Inflorescences at E-L 17 and clusters at E-L 27 were selected for carpel and berry experiments, respectively. Organs received a single Triton X-100 (0.025%)-formulated GA3 (Pro-Gibb 4%/Abbott Laboratories, Chicago, USA) or 0.8 mM PAC (CULTAR 25 SC, Syngenta AG, Basel, Switzerland) application. Rachises and internodes were treated with 121 µM GA3, while carpels and berries were treated with 90 µM GA3. To allow effective inhibition of GA biosynthesis, PAC was applied 4 days before GA3 and control treatments. PAC-GA treatment was included as well, where samples received GA3 application 4 days after PAC treatment. This combined treatment was carried out to verify that growth inhibition by PAC is mainly due to decreased GA biosynthesis, and thus can be reversed if exogenous GA is applied. Triton X-100-treated organs served as controls. All organs were treated either by dipping or spraying to the point of run-off.

Morphological response of organs to GA3 and PAC

Pre-treatment lengths of rachises, and weights of berries, were recorded. Increments in the lengths of rachises and the newest internodes arising after treatment were monitored at 5-day intervals, while berry weights were assessed at 10-day intervals for 30 days.

Sampling for GA response/signalling analyses

Organs and tissues were sampled 6 h after GA treatments. All samplings were done before 14:00 to minimize circadian effects on gene expression. Sampled organs and tissues were immediately frozen in liquid nitrogen in the vineyard, and stored at ~80°C prior to analyses.

Sampling for temporal and spatial analyses, and GA quantitation

Sampling of young internodes was carried out from the most distal internodes from the base of young shoots at E-L 15, while young rachises were sampled from inflorescences at E-L 15. Tissues and organs at similar developmental stages were marked and sampled at véraison and defined as mature internodes and rachis. Young leaves and tendrils were defined as those borne on the first and second nodes (from the shoot tip), while mature leaves and tendrils were sampled from the 12th node. Carpels were sampled at E-L 17, while berries were sampled at E-L 27, and subsequently at 10 and 30 days after the first sampling, herein referred to as 0, 10 and 30 days after fruit set (DAF), respectively. Roots were obtained from single node cuttings immersed in water for about 21 days. Samples were collected before 09:00 to minimize circadian effects on gene expression.
**Phylogenetic analysis of genes associated with GA signalling**

Multiple alignment of protein sequences of the *Arabidopsis*, rice and *V. vinifera* families were generated using the CLUSTAL W alignment algorithm (Thompson et al., 1994) using AlignX of the Vector NTI suite (Lu and Moriyama, 2004). Phylogenetic unrooted trees were constructed using the neighbour-joining (NJ) method in the web-based Phylogeny.fr software (http://phylogeny.lirmm.fr/phyllo.cgi/index.cgi; last accessed: 22 December 2014) (Dereeper et al., 2008).

**Gene cloning and plasmid construction**

Total RNA was extracted using the CTAB protocol as previously described (Acheampong et al., 2010). First-strand cDNA was synthesized using Moloney Murine Leukemia Virus Reverse Transcriptase (M-MLV RT) (Promega Corporation, Madison, WI, USA) according to the manufacturer's instructions. Full-length ORFs of all genes were PCR-amplified from cDNA from different organ primer sets listed in Supplementary Table S1. PCR fragments were amplified with primers having the recommended Gateway overhangs, cloned into pENTR/D-TOPO or pENTR/SD/D-TOPO vectors (Invitrogen, Carlsbad, CA, USA), and subsequently cloned into the GATEWAY-based, 35S-driven pK7WG2.0 expression vector (Karimi et al., 2002) or pET-DEST42 protein expression vector (Invitrogen). The overexpression constructs were introduced into Agrobacterium tumefaciens strain GV3101 by electroporation. To ensure the specificity of the anti-VvDELLA polyclonal antibodies, sequences encoding different domains were used (VvDELLA1, T101-V205; VvDELLA2, P113-Q226; VvDELLA3, V55-D151). These recombinant His-tagged constructs were expressed in BL21- CodonPlus (DE3) RILP strains (Strategene, Santa Clara, CA, USA). The full lengths of the VvDELLA genes were also PCR amplified, cloned into the Entry and Destination vectors, and transformed into BL21-CodonPlus (DE3) RILP cells.

For the yeast two-hybrid (Y2H) assay, pLexA-NLS and pACTII were used as bait and prey expression vectors, respectively, as described previously (Dill et al., 2004). VvGID1 and VvSLY1 proteins were expressed in pLexA-NLS as fusions with LexA DNA binding domain (DNA-BD), while VvDELLA proteins were expressed in pACTII as fusions with the GAL4 activation domain (AD). Primers for Y2H cloning, with appropriate restriction sites are listed in Supplementary Table S2.

**Arabidopsis transformation of grape genes**

Using the A. tumefaciens-mediated transformation method, 35S promoter-driven grape cDNA constructs were transformed into corresponding *Arabidopsis* mutants for complementation tests. 35S:VvGID1 was transformed into gid1a-2 gid1c-2 mutants (Griffiths et al., 2006). 35S:VvSLY1 was transformed into sly1-10 +/- plants due to severe infertility of the sly1-10 homozygote line (Steber et al., 1998). Later the sly1-10 homozygote allele was identified in T2 generations of VvSLY1 transformants. 35S:VvDELLA was transformed into the gal1-3 rga24 mutant (Silverstone et al., 1998). Isolation of homozygous transgenic lines containing single insertion sites was done as described previously (Hu et al., 2008).

**Y2H assays**

DNA-BD and AD fusion construct co-transformation into Saccharomyces cerevisiae strain L40, growth testing with 3-AT, and β-galactosidase liquid assaying were carried out as described previously (Dill et al., 2004).

**Quantitative real-time PCR analyses**

The transcript levels of VvGID1s, VvDELLAs, and VvSLY1s were measured by quantitative real-time PCR (qRT-PCR) using Takara SYBR-green Premix Ex Taq (Thermo Fischer Scientific, Waltham, MA, USA) on the Rotor Gene 6000 Real-time PCR machine (QIAGEN, Hilden, Germany). The previously characterized VvGAPDH (Reid et al., 2006) was used as a normalizer; its expression in grapevine organs is not GA regulated (Giacomelli et al., 2013), and its orthologue in *Arabidopsis* was found to be unaffected by GA (Dill et al., 2004). The SYBR-Green reaction mixture consisted of 12 µl of 0.5 µM forward and reverse primers (Supplementary Table S1), 6 µl of SYBR-Green, and 3 µl of first-strand cDNA. PCR reactions were performed using the following parameters: 15 min at 95°C, and 40 cycles of 15 s at 95°C, 20 s at 60°C, and 20 s at 72°C. Each sample was analysed six times, comprising three biological repeats with two technical repeats each. To ensure accurate quantitation of transcripts, we selected and used primers of comparable efficiencies. We also synthesized a plasmid containing the amplicons of all qRT-PCR products of the genes cloned into the PUC19 vector, linearized it by digesting with BamHI restriction enzyme (NEB, Ipswich, MA, USA), and used it as a template for gene-specific calibration curves, as briefly described below. From an initial concentration of 23 pmol/µl (428 816 420 copies), 10-fold serial dilutions were made and used as templates. Amplicons were confirmed by agarose gel electrophoresis and sequencing.

**Antibody production**

Cloning of constructs for protein induction was as described above. Proteins were expressed from constructs encoding differential regions of each VvDELLA protein using 0.5 mM IPTG and 200 µg ml⁻¹ rifampicin as previously described (Kudernova et al., 1999). The His-tagged recombinant proteins were subsequently purified using a QIAGEN protein purification kit as detailed in the protocol (QIAGEN). The expressed proteins were sequenced by mass spectrometer to ensure sequence integrity, and used as antigens to produce affinity-purified rabbit anti-VvDELLA polyclonal antibodies. Production and purification of polyclonal antibodies were contracted to GenScript USA Inc (Piscataway, NJ, USA). Recombinant full-length VvDELLA proteins, used as sizing standards to locate endogenous VvDELLA proteins, were also expressed and purified as described above, and quantified using BSA standards.

**Protein extraction and immunoblot analyses of VvDELLA proteins**

Total plant protein was extracted from organs as previously described (Wang et al., 2006) with slight modifications. Samples were first homogenized in liquid nitrogen in the presence of polyvinyl polypyrrolidone (PVPP). The protein pellets obtained were dissolved in SDS-PAGE sample buffer containing 0.15 M Tris (pH 6.8), 1.2% SDS, 30% glycerol, 2.14 M β-mercaptoethanol (Sigma Aldrich, St Louis, MO, USA). Extracted proteins were quantified by band intensities confirmed by fractionating on 10% SDS-PAGE gel, and staining with Coomassie protein staining buffer (0.1% Coomassie Brilliant Blue R-250, 50% methanol and 10% glacial acetic acid). Equal amounts of proteins were separated by 10% SDS-PAGE and transferred to PROTEAN nitrocellulose transfer membrane (Whatman GmbH, Dassel, Germany) using the Mini-Protein Transfer system (Bio-Rad Laboratories, Hercules, CA, USA). Detection of protein was as described by Yamamoto et al. (2010) with slight modifications. Blotted membranes were blocked in 3% protein solution (3% milk powder, 50 mM Tris-HCL pH 7.5, 150 mM NaCl, 0.05% Tween 20) for 1 h, and the primary antibody titres were 1:2000, 1:1000, and 1:5000 for VvDELLA1, VvDELLA2, and VvDELLA3, respectively. Purified full-length recombinant proteins of VvDELLA1 (3.75 ng), VvDELLA2 (3.75 ng), and VvDELLA3 (37.5 ng) were used to identify and quantify endogenous VvDELLA proteins for each blot. Chemi-luminescence images of blots were captured by MF-ChemiBIS 2.0 (DNR Bio-Imaging Systems Ltd, Jerusalem, Israel).

**Quantification of endogenous gibberellins**

Triplicate samples (0.5 g) of freeze-dried organs/tissues were weighed, homogenized in liquid nitrogen, and extracted with 80% methanol containing 1% acetic acid and 3H-labelled GAs as internal standards (Supplementary Table S3), for 1 h at 4°C.
Samples were centrifuged at 3000g for 10 min, and filtered through LRC-2 Frits Bond Elut Reservoir (Varian Inc, Palo Alto, CA, USA) to remove residual plant materials. The solvent (80% methanol, 1% acetic acid) extraction was repeated for 10 min, and samples centrifuged and filtered as before. The two extracts were combined and evaporated to dryness at 35°C using Savant SpeedVac Concentrators (Thermo Fischer Scientific). Dried samples were dissolved in 1 ml of 80% acetonitrile, 1% acetic acid. The acetonitrile was removed by evaporation in vacuo.

Endogenous bioactive GAs, their precursors, and deactivation products were measured as previously described (Plackett et al., 2012). The prominent ions were analysed by a liquid chromatography tandem mass spectrometry system consisting of a triple quadrupole mass spectrometer (Agilent 6410; Agilent Technologies, Santa Clara, CA, USA) and an Ultra High Performance Liquid Chromatography (UHPLC) system equipped with an octylphenyl column (ZORBAX XDB-Phenyl, 2 × 50 mm, 1.8 μm; Agilent Technologies). The endogenous GA contents were calculated from the peak area ratios of the endogenous GA to internal standards spiked during the extraction process.

Statistical analyses

Unless otherwise stated, all experiments were designed with a completely randomized distribution. Data are presented as the mean ± standard error (SE). Statistical significance for data from field experiments was determined by one-way analysis of variance (ANOVA) followed by Tukey multiple comparison tests (JMP 7.01 software, SAS Institute, Cary, NC, USA), and significance values set at α = 0.05 as indicated in the text and figure legends.

Accession numbers

Sequence data can be found on the NCBI GenBank data libraries under accession numbers: VvGID1a (AM468374), VvGID1b (AM479851), VvDELLA1 (AM459432.1), VvDELLA2 (AM470304.2), VvDELLA3 (AM484828.1), VvSLY1a (AM4455694.2), VvSLY1b (AM450967), OsGID1 (Os05g0407500), AtGID1a (At3g05120), AtGID1b (At3g63610), AtGID1c (At5g27320), AtGAI1 (At1g14920), AtRGAI (At2g01570), AtRGL1 (At1g66350), AtRGL2 (At3g03450), AtRGL3 (At5g17490), SLR1 (Os03g0707600), AtSLY1 (At4g24210), and OsGID2 (Os02g0580300). Descriptions of the grapevine genes are listed in Supplementary Table S4.

Results

Grapevine GA signalling genes

Bioinformatic analyses were performed on the V. vinifera cDNA database (http://compbio.dfci.harvard.edu/tgi/; last accessed: 20 July 2012) and two independent genome sequence projects (Jaillon et al., 2007; Velasco et al., 2007), using sequence information of the characterized orthologues in rice and Arabidopsis from NCBI (http://www.ncbi.nlm.nih.gov/; last accessed: 23 December 2014). These databases revealed two putative homologues encoding GA receptors (VvGID1a and VvGID1b), two F-box proteins (VvSLY1a and VvSLY1b), and two putative DELLA genes (VvDELLA2 and VvDELLA3), in addition to the previously characterized VvGAI1 (Boss and Thomas, 2002), herein referred to as VvDELLA1 (Supplementary Table S4).

VvGID1 genes and proteins

A BLAST search showed VvGID1a and VvGID1b protein sequences with high sequence homology to Arabidopsis AtGID1 (Griffiths et al., 2006; Nakajima et al., 2006) and rice OsGID1 (Ueguchi-Tanaka et al., 2005). While each VvGID1 parologue encodes a 344 amino acid protein, and has a single intron 38 nucleotides downstream of the start codon, there are differences in their genomic architecture. VvGID1a is located on chromosome 14, and its intron size is 900bp meaning it is encoded by a 1935bp sequence, while VvGID1b is located on chromosome 7, and is encoded by a 1817bp sequence containing a 782bp intron. The amino acid sequences of the two VvGID1 paralogues are 79.1% identical, and share sequence homology with AtGID1a-1c and OsGID1 in conserved GA-binding residues (Supplementary Figure S1A). Like most angiosperms, VvGID1a is clustered with the ‘GID1ac’ group, whereas VvGID1b is clustered with the ‘GID1b’ group (Voegel et al., 2011) on the phylogenetic tree (Fig. 1A). Similar to OsGID1 (Ueguchi-Tanaka et al., 2005) and AtGID1s (Nakajima et al., 2006), both VvGID1s have the conserved HGG and GxSxG motifs characteristic of hormone-sensitive lipases (HSLs), and also possess two (Ser-190 and Glu-288) of the three amino acids (Ser, Glu, His) that constitute the so-called catalytic triad of the HSL family. Most other functionally important residues previously described for GID1 (Ueguchi-Tanaka et al., 2005; Nakajima et al., 2006; Murase et al., 2008) are present in both paralogous. A difference should be noted in position 128, where Thr in VvGID1a is replaced by Tyr in VvGID1b (Supplementary Figure S1A).

VvDELLA genes and proteins

Similar to Arabidopsis RGA and rice SLR1, there are no introns in the genes encoding the 64.8kDa, 66.1kDa, and 58.6kDa proteins of VvDELLA1, VvDELLA2, and VvDELLA3, respectively. VvDELLA2 and VvDELLA3 share sequence similarity with the already characterized VvDELLA1 (Boss and Thomas, 2002), and the orthologues in Arabidopsis (RGA, GAI, RGL1, RGL2, and RGL3) and rice (SLR1) (Fig. 1B, Supplementary Figure S1B). Whereas the deduced amino acid sequences of VvDELLA1 and VvDELLA2 were 66% identical, both paralogues were ~36% identical to VvDELLA3. All three VvDELLAs possess the distinct triad of motifs required for GID1-DELLA interactions (Murase et al., 2008); DeLLaΦLxYxV, MαxVαxLExLExΦ, and TVhnPnxLxWxM, albeit with sequence variations in specific residues (Supplementary Figure S1B). The third Leu in the DeLLaΦLxYxV motif of VvDELLA3 is substituted by another aliphatic hydrophobic amino acid, Ala, while Val-40 (representing Φ) in VvDELLA1 and 2 is replaced by Gly-32 in VvDELLA3. Glu-36 (VvDELLA1) and Glu-47 (VvDELLA2) in this motif is also replaced by hydrophobic amino acid Gly-28 in VvDELLA3. The first Met in the MαxVαxLExLExΦ motif of VvDELLA1 and VvDELLA2 is substituted by Leu in VvDELLA3, while Met-75 and Met-56 in VvDELLA2 and VvDELLA3, respectively, are replaced by Ile-64 in VvDELLA1. Thr-79 and Thr-90 in the TVhnPnxLxWxM motif of VvDELLA1 and VvDELLA2, respectively, are substituted by non-polar Val-70 in VvDELLA3, whereas Val-80 and Val-91 in the former are replaced by Leu-71 in the latter. Tyr-82 (VvDELLA1) and Tyr-93 (VvDELLA2) are substituted by Cys-73 in VvDELLA3.
VvSLY1 genes and proteins

VvSLY1a and VvSLY1b are both 184 amino acid proteins, encoded by intron-less genes, located on chromosomes 7 and 18, respectively. VvSLY1a is a 20.2 kDa protein, while VvSLY1b is 20.9 kDa. Amino acid similarity between the VvSLY1 proteins is 60%, and identity to Arabidopsis and rice orthologues is 44 and 25% for VvSLY1a, and 51 and 28% for VvSLY1b, respectively (Fig. 1C). VvSLY1s alignment with AtSLY1 (McGinnis et al., 2003) and OsGID2 (Sasaki et al., 2003) shows two variable domains (one at the N-terminus and the other close to the C-terminus), conserved N-terminus F-box domains (52–80% sequence identity with AtSLY1 and OsGID2), and GGF (54–86% identity with AtSLY1 and OsGID2) and LSL (35–73% identity with AtSLY1 and OsGID2) domains located at the C-terminus (Supplementary Figure S1C).

VvGID1s, VvDELLAs, and VvSLY1s are functional in transgenic Arabidopsis and in Y2H assays

We tested the functions of these genes in Arabidopsis mutants. 35S:VvGID1a and 35S:VvGID1b were introduced separately into the Arabidopsis gid1a gid1c double mutant, which is a semi-dwarf with reduced fertility (decreased seed number per siliqua, and siliqua length) (Griffiths et al., 2006). Expression of either VvGID1a or VvGID1b completely rescued all phenotypes of gid1a gid1c (Fig. 2A, B; Supplementary Figure S2A, B). 35S:VvDELLA1, 35S:VvDELLA2, and 35S:VvDELLA3 were transformed into the Arabidopsis gal-3 rga-24 mutant, which is a semi-dwarf (Dill and Sun, 2001). Expression of each VvDELLA reduced the plant height of gal-3 rga-24 (Fig. 2C, D), indicating that all three VvDELLAs were functional to rescue the rga-24 defect. Similarly, 35S:VvSLY1a and 35S:VvSLY1b were introduced into the Arabidopsis sly1-10 mutant, which is a semi-dwarf with reduced fertility (McGinnis et al., 2003). Both VvSLY1a and VvSLY1b rescued the sly1-10 mutant defects (Fig. 2E, F; Supplementary Figure S2C, D).

To further characterize the biochemical properties of VvGID1s, VvDELLAs, and VvSLY1s, we performed Y2H assays. Previous studies have shown that interactions between GID1 and DELLA from rice and Arabidopsis are enhanced in yeast cells in the presence of bioactive GAs (Ueguchi-Tanaka et al., 2005; Griffiths et al., 2006). We found that VvGID1a displayed GA-dependent binding to each of the three VvDELLAs (Fig. 3A). VvGID1b also interacted with VvDELLA1 and VvDELLA2 in a GA-dependent manner. Interestingly, VvDELLA–VvGID1a interactions were stronger, and VvGID1b did not interact with VvDELLA3, even in the presence of GA3. We also tested SLY1-DELLA
interaction using Y2H assays. Similarly to *Arabidopsis* SLY1 (Dill et al., 2004), both VvSLY1s interacted with all three VvDELLAs individually (Fig. 3B), although VvSLY1b showed stronger binding to VvDELLAs than VvSLY1a as indicated by the enhanced expression of the reporter genes (growth in higher 3-AT concentrations and higher β-gal activities). The results of the *in planta* functional assay and the Y2H assay indicate that these VvGID1s, VvDELLAs, and VvSLY1s function similarly to the *Arabidopsis* and rice orthologues in GA perception and signalling.

### GA regulation of VvGID1 and VvSLY1 transcript levels

Results from several studies (Ueguchi-Tanaka et al., 2005; Griffiths et al., 2006; Voegele et al., 2011) point to GA-induced feedback inhibition of *GID1* transcripts. To test whether similar regulation occurs in *Vitis*, we analysed the changes in expression of the *VvGID1* genes in different organs subjected to GA3 and PAC treatment regimes (Fig. 4A, B). Based on our results, it appears that: (i) with the exception of *VvGID1a* in rachis (Fig. 4A), GA3 treatment downregulated both *VvGID1* transcript levels, while PAC treatment enhanced their levels; (ii) the degree of change was lower in response to GA3 than in response to PAC, with the exception of *VvGID1b* levels in carpels and berries; (iii) generally, GA3 treatment resulted in greater downregulation in transcript levels of *VvGID1b* than *VvGID1a*. The most dramatic difference was in the rachis where GA3 treatment reduced *VvGID1b* mRNA levels by ~7-fold, but did not alter *VvGID1a* expression. (iv) Similarly, PAC treatment produced a greater increase of transcripts of *VvGID1b* than *VvGID1a*, resulting in a 3-, 10-, 1.5-, and 2-fold difference between both paralogues in internodes, rachis, carpels, and berries, respectively. Upregulation of *VvGID1a* and *VvGID1b* expression was maximal in the PAC-treated internodes (~3- and 6.5-fold, respectively).

*VvSLY1* expression displayed a similar pattern to that of *VvGID1* (Fig. 4C, D). GA3 treatment downregulated transcript levels of both *VvSLY1* homologues, while PAC treatment increased their levels, compared to the respective control samples. Unlike *VvGID1*, the degree of regulation of expression of *VvSLY1a* and *VvSLY1b* was similar, and the effect of GA was more pronounced than that of PAC. There were only

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**Fig. 2.** Grapevine GA signalling genes rescue the phenotype of corresponding *Arabidopsis* mutants. (A, C, E) Gross morphology of gid1a-2 gid1c-2, ga1-3 rga-24, and sly1-10 *Arabidopsis* mutants and representative transgenic plants transformed with *VvGID1* s, *VvDELLA* s, and *VvSLY1* s, respectively. Bar: 5 cm in (A), (E); 1 cm in (C). Whole-plant pictures of *VvGID1*, *VvDELLA*, and *VvSLY1* transformants were taken at 51, 70, and 60 days, respectively. (B, D, F) Average final plant heights of wild type (WT), mutant, and transgenic plants of (A), (C), and (E). The height of each individual transformant is significantly different from the corresponding mutant (*n ≥ 8; P < 0.01). In contrast, the height of 3SS:GUS gid1a gid1c lines was not significantly different from gid1a gid1c (*n ≥ 15*). Parameters for *VvGID1*, *VvDELLA*, and *VvSLY1* transformants were measured at 51, 81, and 87 days, respectively. Asterisk: zero plant height due to the lack of inflorescence stem elongation.
marginal differences when different organs were compared, and VvSLY1b seems to be unaffected by PAC treatment in carpels.

**GA regulation of VvDELLA proteins**

We did not find any distinct expression pattern of transcripts of VvDELLAs in response to GA$_3$ or PAC treatments (Supplementary Figure S3). To test the effect of GA on the stability of VvDELLA proteins, we analysed levels of VvDELLA1, VvDELLA2, and VvDELLA3 in GA$_3$- and PAC-treated rachises, internodes, and berries using antibodies specific for VvDELLA1, VvDELLA2, and VvDELLA3, separately. The immunoblot analyses showed that the level of all three VvDELLA proteins decreased with GA$_3$ application, while PAC treatment appeared to only slightly increase the VvDELLA protein levels in the rachis (Fig. 5A; Supplementary Figure 4S).

**Spatial and temporal expression of GA signalling components**

Having identified and functionally analysed the different paralogues of putative VvDELLA, VvGID1, and VvSLY1 genes, we reasoned that they may exhibit different spatial and/or temporal expression profiles, which may be related to their function and/or to the quantities of endogenous bioactive GA species in these organs. Indeed, qRT-PCR analyses of the GA signalling components in selected organs revealed differential expression patterns (Fig. 6). VvGID1a and VvGID1b were redundantly expressed in all organs analysed, and their transcript levels were highest in the roots (Fig. 6A). However, VvGID1a expression was in general lower than VvGID1b, ranging from 1.5-fold lower in young berries (0 d) to 37-fold lower in roots. Interestingly, mRNA levels of VvGID1s increased in mature tissues compared to young tissues, except in berries where both VvGID1 transcripts levels decreased 2-fold from fruit set to 10 DAF, followed by a 2- and 5-fold increase for VvGID1a and VvGID1b, respectively, from 10 to 30 DAF.

As shown in Fig. 6B, all three VvDELLAs were expressed in all tissues analysed, albeit at different quantities. Generally, VvDELLA2 had the highest expression while VvDELLA3 had the lowest expression in all organs, except berries in which VvDELLA1 was lowest. Similarly to VvGID1s, roots contained relatively high levels of all three VvDELLAs, with VvDELLA3 having the highest levels compared to all other organs. In rachis and tendrils, VvDELLA1 and VvDELLA2 mRNA levels increased in mature tissue. VvDELLA3 expression also increased in mature rachis but decreased in mature tendrils. Intermediate levels of all VvDELLAs were detected in internodes, without a clear age-dependent pattern. Interestingly, similar to both VvGID1s, VvDELLA3 expression decreased slightly from fruit set to 10 DAF followed by a 2-fold increase at 30 DAF. An inverse expression profile was observed for VvDELLA2 over the same period of berry development.

As observed for VvGID1s and VvDELLAs, the expression of VvSLY1a and VvSLY1b (Fig. 6C) was high in the roots. Apart from roots, VvSLY1a had the highest expression in mature rachis, and VvSLY1b in berries at 30 DAF. Both paralogues showed distinctly opposing expression patterns during tissue development: whereas one paralogue was upregulated during tissue development, the other was either downregulated or unchanged. VvSLY1a levels increased with development of internodes and rachis (3- and 1.5-fold, respectively, compared to young tissues), remained unchanged during leaf development, and decreased as tendrils and berries developed. VvSLY1b, on the other hand, was upregulated during development of leaves, tendrils, and berries, representing 3- and 6-fold increases in the first two tissues when they matured, and a 3-fold increase in 30-d-old berries, compared to 0 d. VvSLY1b expression remained unchanged during rachis aging, and was downregulated as internodes matured.

**Temporal and spatial profiles of VvDELLA proteins**

We carried out immunoblot analyses to examine whether different VvDELLA proteins show unique temporal or spatial
expression patterns. In agreement with their mRNA expression profiles, all three VvDELLA proteins were present in the tissues analysed (Fig. 5B). Correlation between temporal transcript and protein profiles was evident for all VvDELLAs in tendrils and at the transition from carpels to berries, and also for VvDELLA1 and VvDELLA3 in leaves (Figs 5B and 6B). The protein profiles in all organs except berries also indicate that: (i) levels of VvDELLA1 and VvDELLA2 were lowest in mature leaves, and highest in young rachis; (ii) levels of VvDELLA1 and VvDELLA2 increased with tendril development, and decreased with leaf and rachis age; (iii) VvDELLA2 levels decreased whereas VvDELLA1 was unaffected by internode age; (iv) levels of VvDELLA3 considerably decreased as all organs aged.

Significant observations in VvDELLA protein profiles during berry development are: (i) levels of VvDELLA3 decreased steadily; (ii) compared to other VvDELLAs, levels of VvDELLA2 were induced dramatically at the transition from carpels to berries; (iii) VvDELLA1 and VvDELLA2 were lowest at 10 days of fruitlet development.

Endogenous GA quantities in developing grapevine organs

The fact that GA application reduced the expression of VvGID1s (Figs 4A and B), and caused degradation of VvDELLA proteins (Fig. 5A) prompted us to explore the hypothesis that the developmental expression and protein
profiles of these genes may be determined, at least in part, by the endogenous GA quantities in the organs. To this end, the content of the bioactive GAs, GA1, and GA4, and their precursors and deactivation products were quantified (Table 1).

The 13-hydroxylation pathway, which leads to the biosynthesis of GA1 (Supplementary Figure S5), was characterized by the high levels of GA10 in all organs (Table 1). GA1 was present only in tendrils and carpels. Interestingly, GA1 levels reduced by 4-fold at fruit set, and further decreased as berries developed. GA3, the deactivation product of GA1, was detected in most organs, and showed a similar temporal profile as GA1. Notably, GA3 levels were elevated during carpel–berry transition, but markedly declined during berry development. The non-13-hydroxylation pathway that produces GA4 was characterized by relatively lower quantities of GA intermediates. GA3 was undetectable in all samples except young internodes. GA4 quantities in carpels decreased towards fruit set. It is important to note that from fruit set to 10 DAF, quantities of both GA4 and GA14 (the deactivation product of GA3) in berries increased. As berries developed, GA14 levels remained constant while GA4 levels decreased. In all but tendrils, carpels and 0 d berries, GA4 quantities were higher than GA1, whereas the opposite was true (except in tendrils and older berries) of GA14 and GA8. Generally, the levels of the different GA species either decreased or remained constant as tissues/organs developed.

Effect of GA3 and PAC application on organ development

Different grapevine organs exhibit markedly different responses to GA application (Weaver, 1958; Mullins et al., 1992; Agüero et al., 2000). To investigate whether such differential responses to GA3 application correlate with quantities of signalling components and/or endogenous GAs, we evaluated growth rates of internodes, rachis, and berries after GA3 and PAC treatments (Fig. 7). Our results show that all organs analysed displayed a GA-related growth response. However, different organs showed varying degrees of GA responses: rachis and berries responded more dramatically to GA3, while internodes had the greater response to PAC treatment. Compared to the control, PAC treatment resulted in marked (>40-fold) attenuation of internode elongation (Fig. 7A). About 40% of the PAC-treated shoot tips dropped, making it impossible to measure internode elongation on those shoots. GA treatment, on the other hand, resulted in an insignificant increase in internode growth (Fig. 7A). Both GA3 and PAC treatments resulted in ~2-fold change in rachis length (Fig. 7B), and in similar changes in berry weight (Fig. 7C). The GA3-treated berries obtained an elongated shape while PAC-treated and control berries presented a rounded shape. Furthermore, GA3 treatments partially or completely rescued the suppressive effect of PAC in all organs (PAC-GA treatments), confirming that the effect of PAC was GA associated.

Discussion

The GA signalling cascade has been extensively described in a number of plant species. However, apart from VvDELLA1 (VvGA11), characterized by Boss and Thomas (2002), little was known about the GA signalling components in grapevine, even though worldwide commercial cultivation of seedless table grapes involves intensive GA treatment for horticulture practices. This study is the first in-depth functional characterization of the major GA signalling components of grapevines.

Structural and functional analyses of GA signalling genes

Functional studies in the Arabidopsis gid1a gid1c mutant (Fig. 2A, B; Supplementary Figure S2A, B) and Y2H analyses (Fig. 3A) indicate that both VvGID1s are active
Table 1. Quantities of selected GA species in 13-hydroxylated and non-13-hydroxylated pathways in V. vinifera cv. Thomson seedless

| GA species | Internode | Rachis | Leaves | Tendrils | Carpels | Berries |
|------------|-----------|--------|--------|----------|---------|---------|
|            | Young     | Mature | Young  | Mature   | Young   | Mature  | 0 d    | 10 d   | 30 d   |
| GA<sub>63</sub> | 2.6±0.3   | n.d.  | n.d.   | n.d.     | n.d.    | n.d.    | 1.7±0.1 | 0.8±0.2 | n.d.   |
| GA<sub>44</sub> | n.d.       | 0.4±0.1| n.d.   | n.d.     | n.d.    | n.d.    | 1.1±0.0 | 0.8±0.1 | 0.1±0.0 | n.d.   |
| GA<sub>19</sub> | 26.5±3.2  | 0.7±0.1| 5.0±1.2| 24.7±0.5| 15.2±5.5| 23.9±5.0| 28.8±1.7| 11.4±0.1| 1.3±0.3| 0.1±0.0|
| GA<sub>20</sub> | n.d.       | n.d.   | n.d.   | 4.6±0.2  | 0.7±0.1 | n.d.    | 3.0±0.1 | 0.8±0.1 | n.d.   | n.d.   |
| GA<sub>29</sub> | n.d.       | 1.0±0.3| n.d.   | 4.3±0.4  | n.q.    | n.q.    | 2.8±0.2 | 1.6±0.0 | 0.2±0.0 | 0.1±0.0|
| GA<sub>1</sub>  | n.d.       | n.d.   | n.d.   | n.d.     | 2.9±1.2 | 0.9±0.2 | 2.8±0.2 | 0.7±0.2 | 0.4±0.1| n.d.   |
| GA<sub>6</sub>  | 2.2±0.1   | 1.4±0.5| n.d.   | 5.6±0.8  | 1.1±0.5 | n.q.    | 9.8±1.0 | 14.8±0.8| 1.0±0.2| 0.2±0.0|
| GA<sub>12</sub> | 1.8±0.1   | n.d.   | n.d.   | n.d.     | n.d.    | n.d.    | n.d.   | n.d.   | n.d.   | n.d.   |
| GA<sub>24</sub> | 5.4±0.3   | 0.4±0.1| n.d.   | 0.9±0.1  | 0.1±0.0 | n.d.    | n.d.   | n.d.   | n.d.   | n.d.   |
| GA<sub>4</sub>  | 0.6±0.1   | 0.2±0.0| 0.1±0.0| 2.0±0.5  | 2.1±0.5 | n.d.    | 1.0±0.2 | 0.4±0.1 | 0.8±0.1| 0.2±0.1|
| GA<sub>34</sub> | 0.8±0.1   | n.d.   | n.q.   | 1.6±0.1  | 0.5±0.0 | 1.0±0.3 | 0.2±0.0 | 0.4±0.0 | 3.0±0.5| 2.7±0.2|

13-hydroxylated pathways are shown in the top half of the table and non-13-hydroxylated pathways in the bottom half. Values represent mean amounts of GA species (ng g<sup>−1</sup> FW) determined in three biological replicates ± SE; n.d., undetected or could not be reliably quantified due to low abundance; n.q., detected, but could not be quantified due to co-migration of impurities or undetected internal standard (IS).

**GA receptors.** Because Thr-128 in GID1a is essential for the direct interaction with Φ of the DeLLaΦLxYxV motif (Murase et al., 2008), it is likely that substituting this residue with the aromatic residue Tyr-128 in VvGID1b could account for the lower binding affinity of VvGID1b compared to VvGID1a with VvDELLA1 and VvDELLA2, as well as the lack of interaction between VvGID1b and VvDELLA3 in our Y2H tests. Expression of *AtGID1s* is downregulated by GA (Griffiths et al., 2006). We observed a similar regulatory mechanism for both *VvGID1s* (Fig. 4A, B) in grapevine.

Since VvDELLA1 does not regulate berry enlargement (Boss and Thomas, 2002), we speculated that other DELLA family member(s) may modulate GA-related berry growth and responses. The two additional grapevine DELLA genes identified in this study, VvDELLA2 and VvDELLA3, were biologically functional in planta (Fig. 2C, D), and in Y2H analyses (Fig. 3). Nonetheless, sequence variations in functionally important residues such as Ala-33, Gly-32, Val-70, Leu-71, and Cys-83 of VvDELLA3, predicted to be required for the direct interactions of VvDELLA with VvGID1 and/or stabilization of the VvGID1-VvDELLA complex, may account for the absence of interaction between VvDELLA3 and VvGID1b. These residues are critical in the GA-dependent interactions between *Arabidopsis* DELLA proteins and GID1s (Murase et al., 2008). Similar to other model plants (Fu et al., 2002; Itoh et al., 2002; Griffiths et al., 2006), degradation of all three VvDELLA proteins in response to GA treatment (Fig. 5A) further confirms their function as GA signalling repressors.

VvDELLA2 mRNA and protein levels are highest in most tissues examined, suggesting that VvDELLA2 plays a major role in regulating GA responses at most developmental stages. VvDELLA1 transcript and protein levels were relatively abundant in most tissues (e.g. roots, tendrils, and internodes), except in berries, consistent with a previous study (Boss and Thomas, 2002). However, contrary to the above-mentioned report, which did not identify *VvDELLA1* transcripts in berries, we detected low levels of *VvDELLA1* transcripts in berries of Thompson seedless (Fig. 6B). These differences could be due to differences in sensitivity between the detection techniques (qRT-PCR in the current study compared to RNA blotting in the former), or the varietal differences (sternospermocarpic cultivars in the present study compared to seeded cultivars in the former). The expression profile of VvDELLA1 suggests that VvDELLA1 plays a major role in most tissues, but only has a minor role during berry development. VvDELLA3 is expressed at much lower levels than the other two DELLA3, suggesting that DELLA3 plays a minor role in most tissues. This idea should be further tested by functional analysis in planta using a transgenic approach.

To the best of our knowledge, all analysed plants possess only one GA-specific F-box protein in the SLY1/GID2 family (McGinnis et al., 2003; Sasaki et al., 2003). Thus, the presence of two biologically functional paralogues of GA-specific F-box proteins (Fig. 2E, F; and Fig. 3B) in grapevine is unique. These paralogues may have arisen from a duplication event, and their maintenance in the genome may be due to differential specificity of interaction with the various VvDELLA-VvGID1 complexes. Such a possibility is supported by the differences in DELLA-SLY interactions demonstrated in our Y2H analyses. Downregulation of *VvSLY1* expression by GA application (Fig. 4C, D) is supported by the negative feedback regulation of *Arabidopsis SLY1* by GA<sub>3</sub> (Dill et al., 2004). Our results suggest that temporal expression profiles of the *VvSLY1* genes in *planta* may be regulated by the endogenous bioactive GA quantities as well. Moreover, a correlation between the temporal expression pattern of each *VvSLY1* homologue and endogenous GA levels may be indicative of the role each paralogue plays in GA-related organ development. We therefore suggest that *VvSLY1a* be playing a central role in GA-regulated growth of internodes and rachis, while *VvSLY1b* may be important for leaf and tendril development, as well as carpel development, because its expression increased after fruit set as the bioactive GA<sub>3</sub> levels in the organ decreased.
Similar to *Arabidopsis* and other species (Dill et al., 2001; Itoh et al., 2002; Griffiths et al., 2006), we have shown that in grapevine application of GA₃ downregulates *VvGID1* transcripts and results in degradation of VvDELLA proteins (Fig. 4A, B; Fig. 5A). In light of the above, it was important to quantify endogenous GAs in grapevine organs, and determine how they correlate with levels of the signalling components in these organs. The results of our GA quantitative analysis, which show a general decrease in GA species during organ development (Table 1), is consistent with that of wine grapes (Boll et al., 2009), avocado (Raviv et al., 1987), and other species (Itoh et al., 1999; Kaneko et al., 2003). Interestingly, although GA₁₀ levels are higher than its corresponding GA₃₃ counterpart in the non-13-hydroxylated pathway, GA₁₉ was higher than GA₁₃ in most organs. This could be due to higher turnover of GA₁, supported also by higher levels of GA₃ than GA₃₃ in these organs.

Higher quantities of GA₁ in internode, rachis, leaves, and berries suggest that this molecule is the major bioactive GA promoting growth of these organs. In contrast and consistent with seeded wine grapes (Perez et al., 2000; Boll et al., 2009; Giacomelli et al., 2013), carpels contain relatively high levels of GA₁ and most of its precursors, suggesting a unique role of GA₁ in regulating flower development and fruit set. A unique ‘convex’ profile of GA₁ in berries should be highlighted, where levels of GA₁ are increasing in a certain period of berry development (0–10 DAF) rather than decreasing as is the case for other ageing tissues. This profile may reflect the transition between sources of production of GA₁ at fruit set, and the unique development of the stenospermocarpic berry. We speculate that: (i) prior to anthesis, anthers probably serve as the main source of bioactive GAs (GA₁ and low levels of GA₃), and supply from this source decreases at bloom as stamens fall and fruit set commences; (ii) after fruit set, seeds serve as the main source of GA₄ accumulation, as shown in other plants (MacMillan, 2001). GA₄ increases as seeds develop, and in stenospermocarpic varieties, peaks just before abortion of endosperm (which occurs at about 14 DAF); (iii) After abortion, GA₄ supply from the seed rudiments is decreased.

Endogenous bioactive GAs regulate temporal profiles of *VvGID1* transcripts in a negative feedback fashion, while the GA regulation of temporal profiles of VvDELLA proteins appears to be organ-specific. The expected inverse correlation was observed in tendrils and berries for both VvDELLA1 and VvDELLA2. As bioactive GA quantities decrease during transition from carpel to fruitlet (fruit set), there is a parallel increase in VvDELLA2 accumulation. The subsequent increase in GA₄ levels as fruitlets develop, prior to endosperm abortion, correlates with the decrease in VvDELLA1 and VvDELLA2 quantities during the same period. Reduction in GA₄ quantities after abortion is mirrored by the corresponding significant accumulation of VvDELLA2, and a slight increase of VvDELLA1. This connection between developmental transitions throughout berry development, GA levels, and the VvDELLA protein quantities, may indicate the functional importance of these VvDELLA paralogues in the regulation of GA-mediated berry development.

In all other organs, however, the decrease in endogenous bioactive GA detected in older tissue, compared with a young tissue, was accompanied by decreased levels of
VvDELLAs. These results are consistent with a previous study (Chandler et al., 2002) showing that bioactive GA levels in developing barley leaves correlate with DELLA protein levels. In these organs, it is possible that DELLA proteins may play a more important role in the rapidly growing tissues to regulate growth dynamically. As these organs mature and growth rate declines, DELLA may not be required to regulate growth.

Organ responses to GA3 and PAC depend on both GA signalling components and endogenous GA levels

It can be assumed that a young developing organ with relatively high VvDELLA protein quantities will have a greater response to GA treatment. To test this idea, we compared VvDELLA levels to the response of growing organs to GA3 application. The sum of all three VvDELLA proteins was highest in young rachis, intermediate in young berries, and lowest in young internodes (Fig. 5B). Consistent with our hypothesis, and compared to their respective controls, GA3 application resulted in a 2-fold increase in rachis and berry size, while a similar treatment did not affect internode length (Fig. 7A). PAC treatments, on the other hand, resulted in a 40-fold reduction in internode length, while producing ~1.5-fold reduction in rachis length and berry weight.

High vegetative vigour in grapevines was associated with high endogenous GA levels (Lavee, 1986). Accordingly, it is expected that a minimal amount of GA is required to fulfill the growth potential of an organ under given conditions and allow maximal size enhancement. At such optimal levels of endogenous GA, high response to PAC treatment is expected, and a much subdued or zero response to GA application. Conversely, organs with levels of endogenous bioactive GA that are below this optimum will exhibit a high response to GA treatment, and show little or no response to PAC treatment, depending on the difference between the level of endogenous GA and that required for maximal growth. The data reported in the current study (Table 1; Fig. 7) supports the above scenario and suggest that different organs require different quantities of bioactive GA for maximal size enhancement. The 0.6 ng g⁻¹ FW of GA3 in young internodes was enough to elicit maximal growth in the internodes, as evidenced by the non-responsiveness of the internodes to GA3 application, and the huge response to PAC treatment. In contrast, in young berries, 0.4 ng g⁻¹ FW GA4 (berries at 0 d), and even 0.8 ng g⁻¹ FW (berries at 10 d), were insufficient to ensure development to maximal berry size. Hence, both GA3 and PAC treatments resulted in significant changes in berry size. Similar to our results, GA3-related berry enlargement of Thompson seedless has been widely reported (Harrell and Williams, 1987; Ben-Tal, 1990). In the case of young rachis, 0.1 ng g⁻¹ FW GA3 was only sufficient to produce limited growth, and GA3 application had a significant contribution. Effective GA-related rachis elongation is widely employed by viticulturists to reduce compactness within the cluster and prevent berry rot (Weaver and Pool, 1965; Dokoozlian and Peacock, 2001).

Supplementary material

Supplementary data can be found at JXB online.

Supplementary Table S1. Primers used for gene isolation and gene expression analyses by qRT-PCR.

Supplementary Table S2. Primers used for cloning genes to Y2H vectors.

Supplementary Table S3. Amount of deuterated GA species used as Internal Standard (IS) mix during extraction of endogenous GAs in different organs of grapes.

Supplementary Table S4. GA signalling genes of *V. vinifera* cv. Thompson seedless.

Supplementary Figure S1. Amino acid sequence alignment of major GA signalling genes.

Supplementary Figure S2. Grapevine GA signalling genes rescue silique length and fertility defects of corresponding *Arabidopsis* mutants.

Supplementary Figure S3. GA regulation of expression of *VvDELLA1*, *VvDELLA2*, and *VvDELLA3*.

Supplementary Figure S4. GA3-induced degradation of VvDELLA proteins in cv. Thompson seedless.

Supplementary Figure S5. Schematic representation of major GA metabolism pathways in plants.

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