Petal abscission in fragrant roses is associated with large scale differential regulation of the abscission zone transcriptome

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Flowers of fragrant roses such as Rosa bourboniana are ethylene-sensitive and undergo rapid petal abscission while hybrid roses show reduced ethylene sensitivity and delayed abscission. To understand the molecular mechanism underlying these differences, a comparative transcriptome of petal abscission zones (AZ) of 0 h and 8 h ethylene-treated flowers from R. bourboniana was performed. Differential regulation of 3700 genes (1518 up, 2182 down) representing 8.5% of the AZ transcriptome was observed between 0 and 8 h ethylene-treated R. bourboniana petal AZ. Abscission was associated with large scale up-regulation of the ethylene pathway but prominent suppression of the JA, auxin and light-regulated pathways. Regulatory genes encoding kinases/phosphatases/F-box proteins and transcription factors formed the major group undergoing differential regulation besides genes for transporters, wall modification, defense and phenylpropanoid pathways. Further comparisons with ethylene-treated petals of R. bourboniana and 8 h ethylene-treated AZ (R. hybrida) identified a core set of 255 genes uniquely regulated by ethylene in R. bourboniana AZ. Almost 23% of these encoded regulatory proteins largely conserved with Arabidopsis AZ components. Most of these were up-regulated while an entire set of photosystem genes was prominently down-regulated. The studies provide important information on regulation of petal abscission in roses.

Organ abscission is an important developmental process that regulates the detachment of leaves, flowers, flower parts, fruits etc. from main body during the course of development. Controlled abscission of old, diseased or surplus organs is necessary to conserve resources for developing organs to maintain healthy growth while abscission of fruits and seeds ensures reproductive success and survival through dispersal. Although abscission leads to separation of an entire organ from a plant, the processes leading to separation are controlled by cells of a small zone, the abscission zone (AZ). The AZ, only a few cell layers thick, shows differential sensitivity to hormones and differential regulation of genes that trigger abscission. Hormones that either promote or inhibit abscission have to be maintained in a fine balance. Ethylene is one of the most important of these in initiating abscission in most plants including trees. Mutants of ethylene perception and signaling show delayed organ abscission in Arabidopsis and tomato. Not surprisingly, components of the ethylene biosynthesis and signal pathway are often expressed in an AZ-specific manner.

Unlike ethylene, auxin inhibits or delays abscission in leaves, flower and fruits. Components of auxin transport and signalling including auxin response factors (ARFs) play an important role in abscission. Other

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hormones like abscisic acid (ABA) and jasmonic acid (JA) also influence abscission although their involvement at the molecular level is much less studied [31–34].

Although abscission is known in several plants, the key determinants of abscission have primarily been identified in the model plant Arabidopsis where genes like HAESA and HAESA LIKE2 encoding LRR type receptor kinase [35–37], IDA [38–39], AGL15 [39], AGL18 [39], MAP kinases MPK3 and MPK6 [40], NEVERSHEDE [41], EVERSHED [42], FOREVER YOUNG FLOWER [43], have been characterized. In rice, studies on the process of seed shattering have led to identification of SH4 [44], qSH1 [45], SHAT1 [46] and SH5 [46], while abscission of tomato has been shown to be controlled by a MADS box gene, JOINTLESS [47,48]. In most other plants, regulatory genes controlling abscission remain to be identified.

The availability of microarray at the turn of the century and NGS techniques over the last 10 years has enabled large scale gene expression analysis of even fine tissues such as organ AZ in model plants like Arabidopsis and tomato [49–52] as well as a few other plants like citrus [53], apple [54], melon [55], olive [56], litchi [57], sugarcane [58]. Nevertheless, the difficulties associated with isolation of the abscission zones limit studies on organ abscission in most plants even by NGS techniques.

Rose is an important flower in the floriculture and fragrance industries. The fragrant variety of rose (such as *Rosa bourboniana*) is popular but sensitive to ethylene. It has a short vase life of 1 to 2 days post-pollination that decreases its commercial value. In contrast, the non-fragrant *R. hybrida* is less sensitive to ethylene and has greater vase life of several days. These differences are partially associated with differences in sensitivities of the ethylene pathway [59–61] which in turn affect expression of cell wall modifying genes and programmed cell death-like processes [31–34]. Nevertheless, the molecular determinants that are responsible for regulation of petal abscission in fragrant and non-fragrant roses remain unclear and need to be identified to increase flower life in roses.

In the present study we show, through transcriptional profiling of petal AZ of ethylene-treated flowers of *R. bourboniana* and *R. hybrida*, a prominent alteration in expression of a large number of regulatory genes that include transcription factors, kinases/phosphatases and F-box proteins besides many other pathways. We also show the activation of the ethylene pathway but a suppression of the JA, auxin and light-governed pathways in the course of abscission. The study provides important information on abscission regulation in roses that could be used for improvement of ornamentals.

Results

Illumina-based sequencing and assembly reveals large scale transcriptomic changes during ethylene-induced petal abscission in roses. To understand the processes governing ethylene-induced petal abscission in rose, a comparison of the transcriptome data was carried out between 0 h (ethylene-untreated) and 8 h ethylene-treated (representing the mid-point of abscission) petal AZs from *R. bourboniana* on an Illumina Hi-Seq2000 in three independent replicates. Further comparisons were also carried out between 8 h ethylene-treated *R. bourboniana* petal AZ with ethylene-treated whole petals and with 8 h ethylene-treated petal AZs of *R. hybrida*. A total of 32,961 genes were found expressing in the rose petal AZ transcriptome out of 43,301 genes predicted in the genome. Very little variation in expression was observed in FPKM plot comparisons of biological replicates in contrast to comparisons of control and experimental samples indicating high reproducibility of the experimental data set (Fig. S1). Following assembly, a comparative analysis was performed to identify differentially expressed genes (DEGs) in abscission that were defined as significantly up- or down-regulated based on a log2-fold change (|FC| ≥ 1 and ≤ −1 respectively) and P value ≤ 0.05.

A total of 5638 significant DEGs could be identified in the comparison between 0 h versus 8 h ethylene-treated AZ samples of *R. bourboniana* while 12,191 DEGs were seen in the comparison between 8 h ethylene-treated petal AZ versus 8 h ethylene-treated whole petal samples of *R. bourboniana* and 11,467 DEGs in the comparison between 8 h ethylene-treated AZ samples of *R. bourboniana* versus 8 h ethylene-treated AZ samples of *R. hybrida* (Table S2). Of this, a total of 2349, 5982 and 4266 genes were significantly differentially up-regulated and 3289, 6209 and 7201 genes were significantly down-regulated in *R. bourboniana* petal AZ in the 0 h versus 8 h ethylene-treated AZ (R. bourboniana), 8 h ethylene-treated petal AZ versus whole petal (R. bourboniana) and 8 h ethylene-treated *R. bourboniana* AZ versus *R. hybrida* AZ transcriptome comparisons, respectively. Of the significant DEGs in all comparisons, a further analysis of genes at log2FC > 1 and ≤ −1 was carried out. A total of 1518, 2912 and 1772 genes were significantly differentially up-regulated and 2182, 3376 and 3593 genes were significantly down-regulated in *R. bourboniana* petal AZ in the 0 h versus 8 h ethylene-treated AZ (R. bourboniana), 8 h ethylene-treated AZ versus petal (R. bourboniana) and 8 h ethylene-treated *R. bourboniana* AZ versus *R. hybrida* AZ transcriptome comparisons, respectively (Fig. 1). A larger number of genes were down-regulated with the onset of abscission (Fig. 1). These genes (significant at log2FC > 1 and log2FC ≤ −1) were considered for further analysis.

The expression profiles of eight housekeeping genes namely, ELONGATION FACTOR EF1A (JN39925), UBQ10 (JN39997), GLYCERALDEHYDE-3-PHOSPHATE DEHYDROGENASE (JN399920), SERINE/THREONINE PROTEIN PHOSPHATASE 2A (JN399924), SAND (JN39928), TUBULIN (JN39923) and TIP (JN39921) and ACTIN (KP983187) [46,47] revealed no significant variation in the four transcriptome samples used in the analysis (Fig. S2).

Abscission is associated with a substantial change in the expression of transcriptional and post-translational regulators. An analysis of all DEGs in the 0 h versus 8 h ethylene-treated petal AZ RNA was next performed to identify important biological pathways affected during ethylene-induced petal abscission. The most prominently affected genes in the transcriptome belonged to the transcription factor families, kinases/phosphatases and F-box protein degradation families, hormone signalling components, transporters, biotic/abiotic stress groups, cell wall modification and carbon metabolism (Figs. 2, 3, 4, Tables S3, S4).
Of the 5638 DEGs, 284 genes (122 up, 162 down) encoding putative transcription factors (TFs) were differentially regulated between the 0 and 8 h ethylene-treated petal AZ samples (Fig. 2a; Table S3). Prominent among the differentially regulated TF groups were zinc finger proteins (52 up, 70 down), MYBs (13 up, 18 down), MYCs (9 up, 11 down), homeobox domain genes (6 up, 10 down), ERFs (8 up, 7 down), NACs (7 up, 8 down), WRKYs (4 up, 9 down) and BTB/POZ (2 up, 9 down). Other groups included GRAS (4 up, 4 down), Leucine zipper genes (3 up, 4 down), ARFs (2 up, 3 down), SBP (4 down), heat shock factors (HSFs; 2 up, 1 down), MADSs (2 down) and LOBs (2 up).

Post-translations modifications are an important form of regulation of a large number of proteins and enzymes including TFs during development. One of the most prominent modifications includes phosphorylation/dephosphorylation brought about by kinases and phosphatases. Not surprisingly, genes encoding these accounted for 289 DEGs in the rose abscission transcriptome (Fig. 2b; Table S3). Of these, kinases accounted for 203 DEGs, of which 81 were up-regulated while 122 were down-regulated. The majority encoded serine threonine protein kinases (25 up-, 52 down-regulated) and LRR kinases (12 up- and 16 down-regulated). The histidine kinase group (12 up, 3 down) showed an unusually staggered up-regulation and included three ethylene receptors ETR3, ETR7 and ETR8, one cytokinin receptor and three phytochrome genes, PhyA, PhyB and PhyE. Wall-associated receptor kinases (4 up- and 5 down-regulated), mitogen-activated protein kinases (1 up- and 5 down-regulated) and bulb-type lectin kinases (8 up- and 10 down-regulated) were also abundant. Compared with kinases, phosphatases accounted for a much smaller yet substantial proportion of the DEGs with 86 genes (25 up-regulated, 61 down-regulated). The PP2C group of phosphatases accounted for the majority of genes and were down-regulated (Fig. 2b; Table S3).

In addition, proteins that control turnover of key regulatory proteins in signal transduction pathways and other important cellular processes also play an important role in development. A group of 75 DEGs encoding F-box proteins, involved in proteasomal degradation of important proteins, were observed. These belonged to different groups containing additional domains such as WD repeat (29), LRR (23), Kelch (10) etc. (Fig. 2c). Of the 75, 41 genes were up-regulated while 34 genes were down-regulated in 8 h ethylene-treated petal abscission zones. In addition, 100 genes encoding peptidases and members of the ubiquitin E3 ligase complex (that include cullin and ubiquitin-conjugating enzymes responsible for protein ubiquitylation and degradation) were differentially regulated. Of these, 37 genes were up-regulated while 63 genes were down-regulated (Fig. 2c; Table S3). Amongst both groups, the proteasome components were largely up-regulated. Processes associated with nucleic acid degradation and programmed cell death (PCD) (23 genes), senescence (5 genes) and autophagy (3 genes) were also differentially regulated with most members of nucleic acid degradation being up-regulated during abscission (Fig. 2c; Table S3).

**Phytohormone pathways undergo prominent changes with the onset and progression of abscission.** Phytohormones regulate almost every aspect of plant growth and adaptation. They also regulate and are known, in turn, to be regulated in response to abscission. To examine phytohormone involvement during abscission in rose, genes encoding biosynthesis/signaling components of key abscission hormones like ethylene, auxin, jasmonic acid (JA), abscisic acid (ABA), salicylic acid (SA), GA, cytokinin and brassinosteroid were analyzed. Among a total of 111 hormone pathway-related DEGs, the auxin and ethylene signalling pathways were the largest groups with 36 and 24 DEGs, respectively (Table S4). The majority of auxin pathway genes encoding...
Figure 2. Functional cataloguing showing the proportion of DEGs encoding regulatory proteins in the ethylene-treated *R. bourboniana* petal AZ transcriptome (a) transcription factors, (b) kinases and phosphatases, (c) F-box and proteolysis components. Analysis was performed at $\log_2|FC| \leq -1$ and $\geq 1$, Q-value < 0.05. Percentage (%) reflects the percentage of DEGs at $\log_2|FC| \leq -1$ and $\geq 1$ at Q-value < 0.05 out of total mapped genes at any fold change and Q-value.
tryptophan synthase, AUX/IAA proteins, auxin response factors and small auxin up RNA were down-regulated while a few genes encoding an auxin efflux carrier (RC4G0446200), two AUX/IAA proteins (RC6G0552600, RC6G0462900), two auxin response factors (RC5G0517400, RC6G0512800) and five small auxin up RNAs (from RC2G0105500 to RC3G0216500) were up-regulated during abscission (Fig. 3c). In contrast, the majority of ethylene pathway genes were up-regulated during abscission. These included 7 genes encoding 1-aminocyclopropane-1-carboxylic acid oxidase (ACO; RC6G0383500), ethylene receptors (ETRs; from RC4G0031800-RC5G0395000,) and signalling components like CONSTITUTIVE TRIPLE RESPONSE (CTR; RC3G0234100) and ETHYLENE INSENSITIVE 3 (EIN3; RC1G0141700). A gene encoding the ethylene biosynthesis enzyme 1-aminocyclopropane-1-carboxylic acid synthase (ACS; RC7G0182700) was down-regulated while 7 of the 14 differentially regulated ERFs were up-regulated and 7 down-regulated (Fig. 3a). Interestingly, genes encoding components of the JA pathway were prominently down-regulated. These included genes encoding homologues of phospholipase A and D (PLA, RC2G0309000; PLD, RC1G001300), lipoxygenases (LOXs; RC4G0229200, RC5G0078700, RC5G0078800), allene oxide cyclases (AOCs, RC7G0443300, RC2G0577100), acyl-coenzyme A oxidase (RC2G0274500) and an OPDA reductase (OPR, RC2G0094500) involved in JA biosynthesis (Fig. 3b). As an exception, an acyl-coenzyme A oxidase gene (RC2G0274500) was up-regulated. The expression of four JASMONATE-ZIM-DOMAIN PROTEIN genes (JAZs, RC2G0102100, RC2G0111000, RC4G0339000, RC5G0132900), encoding key repressors of the JA-pathway, was down-regulated. One jasmonic acid carboxyl methyltransferase (JMT, RC4G0447400) and one MYC transcription factor homologue (RC7G0296400) that are known to govern jasmonate-regulated plant responses in Arabidopsis were also down-regulated. Among DEGs of the ABA pathway, the expression of the gene encoding the rate-limiting ABA biosynthesis enzyme, 9-cis-epoxycarotenoid dioxygenase (NCED; RC5G0132400) was lower in 8 h ethylene-treated petal AZ. Simultaneously, the expression of genes encoding abscisic acid 8′-hydroxylases (CYP707A; RC6G0562100 and RC6G0114600) that are key ABA catabolism enzymes, was up-regulated. On the other hand, two genes encoding homologues of abscisic aldehyde oxidase (AAO; RC5G0431800) and abscisic acid deficient 4 protein (ABA4; RC7G0271400) possibly involved in ABA biosynthesis and de novo ABA synthesis respectively, were up-regulated. Many protein phosphatase 2Cs (PP2Cs; RC1G0573600, RC5G0571100 and RC6G0083700) which may function as global negative regulators of ABA signalling were down-regulated suggesting that the balance between ABA biosynthesis and catabolism is tightly controlled during rose petal abscission (Fig. S3a). The cytokinin pathway also showed a similar profile with DEGs encoding proteins of cytokinin biosynthesis such as cytokinin synthase

Figure 3. Expression profile of differential AZ genes belonging to ethylene (a), JA (b) and auxin (c) pathways. Bars represent the relative fold expression change after ethylene treatment as calculated from transcriptome data using the expression in all three biological replicates (log2 |FC| ≤ −1 and ≥1, Q-value < 0.05). Expression of respective genes in controls was taken as one and shown as a black line across genes for comparison.
**Figure 4.** Functional cataloguing showing the proportion of DEGs associated with defense, wall modification and transport (a) Defence-related components (b) wall modification proteins, (c) transporters. Analysis was performed at \([\log_2|FC| \leq -1 \text{ and } \geq 1]\), Q-value < 0.05. Percentage (%) reflects the percentage of DEGs at \([\log_2|FC| \leq -1 \text{ and } \geq 1]\) at Q-value < 0.05 out of total mapped genes at any fold change and Q-value.
(RC4G0245000), cytokinin riboside 5′-monophosphate phosphoribohydrolase (LOG, RC6G0513700), catabolism genes such as cytokinin dehydrogenase (RC6G0597300 and RC1G0050300) and a histidine kinase gene (RC4G0454300) associated with cytokinin signalling all being up-regulated simultaneously. This suggested that the cytokinin pathway is under complex control during abscission (Fig. S5b). For the GA pathway, one of the primary genes involved in GA biosynthesis encoding a GA20 oxidase (RC1G0310000) was up-regulated while another encoding a GA2 oxidase (RC5G0037300) that inactivates GAs, was down-regulated. GIBBERELLIN INSENSITIVE DWARF1 (RC6G0417300), which initiates GA signalling by promoting degradation of the GA-inhibitory DELLA proteins was also up-regulated suggesting a possible increase in the GA response although other members of the pathway were not affected much (Fig. S5c). Some DEGs related to the SA pathway also underwent change during abscission (Fig. S5d).

**Abscission affects the expression of defence and stress pathways and regulates carbon metabolism and transport.** A large number of stress and defense-associated pathways were also regulated during abscission. Within stress, those encoding pathogenesis-related proteins (39 genes, encoding chitinases, thaumatin-like proteins, defensins, Toll/interleukin-1 receptor, glutamate receptor) and oxidative stress proteins (55 genes, encoding peroxiredoxins, glutaredoxins, thioredoxins and glutathione S-transferases) were abundant. (Fig. 4a; Table S3). Heat shock proteins, which are also reported to express upon wounding and defence, showed a pattern unlike other groups in that 22/28 HSPs were up-regulated and only 6 were down-regulated during abscission (Fig. 4a; Table S3). Likewise, 17/20 DEGs in abiotic stress were up-regulated while 23/29 DEGs associated with biotic stress were down-regulated.

The phenylpropanoid pathway, involved in secondary metabolism and defence and regulated by light, exhibited a disproportionate change with 41/50 DEGs being down-regulated within 8 h of ethylene treatment. These included key pathway genes like phenylalanine ammonia lyase, chalcone synthases, chalcone isomerases, caffeoyl CoA O-methyl transferases etc. (Table S3). The cytochrome P450 oxidase family which regulates the phenylpropanoid, alkaloid and terpenoid pathways was also differentially affected with 44 DEGs (20 up and 24 down-regulated) (Fig. 4a; Table S3).

Wall hydrolysis is an important component that enables separation of the abscising organ while cell wall reinforcement is needed to protect the tissue exposed after organ separation. In keeping with the complexity of the cell wall, 55 DEGs involved in biosynthesis and modification of cell wall components were differentially regulated. These included genes encoding xyloglucan endotransglucosylase/hydrolases (XTHs), pectinesterases, expansins and pectate lyase etc. Almost 65% of these were prominently down-regulated upon abscission (Fig. 4b; Table S3).

Quite strikingly, a large group of genes (245/1475), encoding transporters for ions (33), amino acids (25), water/aquaporins (11), sugar/sugar phosphates (28), ATP-binding cassette (ABC) transporters (22), heavy metals (13), drug transporters (20) and others (61) were differentially expressed during abscission (Fig. 4c). Most of the sugar, ion, metal and ABC transporter genes were up-regulated in 8 h ethylene-treated AZ, while other transporters related to water, phosphate, amino acid, mitochondrial electron transport, nucleotide and vesicle-mediated membrane transport were down-regulated (Fig. 4c; Table S3).

Another rather surprising observation was how abscission affected the light-regulated, photosynthesis-related genes involved in light reactions in plastids. A large majority of these (49/56) encoding components of photosystems I and II, chlorophyll ab-binding proteins, oxygen evolution, electron transport and oxido-reductases were strongly down-regulated while a chlorophyllase was up-regulated suggesting suppression of several plastid activities. This affected carbohydrate metabolism with 42/70 DEGs involved in glycolysis, gluconeogenesis, TCA cycle, PPP pathway and starch degradation being down-regulated (Table S3).

**Validation of differentially regulated genes in ethylene-treated petal abscission zones.** We next performed a qRT-PCR validation of expression using 12 randomly selected DEGs from the above categories that were differentially regulated (7 up and 5 down) in the 8 h ethylene-treated petal AZ (Table S5). A more detailed time-course study was performed by including samples at 0, 4, 8 and 12 h after ethylene treatment for *R. bourboniana* and additionally at 24, 36 and 48 h for *R. hybrida* and 0 and 8 h for ethylene-treated petals. In agreement with the transcriptome data, the expression of genes encoding a putative endochitinase 2-like protein (RC6G0304900), beta-galactosidase (RC2G0081600), a palmitoyltransferase (RC2G0652900), a putative EIN3-binding F-box protein (RC2G0369000), a lysine-histidine transporter-like 8 protein (RC6G0339000), a wall-associated receptor kinase (RC3G0340200) and a putative amine oxidase (RC4G0245000) was induced by ethylene in *R. bourboniana* AZ as seen in the transcriptome and increased from 0 to 12 h post-ethylene treatment during the abscission time-course (Fig. 5). The fold change in real time PCR validation was usually higher than that seen in the transcriptome except for the amine oxidase gene (Table S5). For most, expression also increased in *R. hybrida* AZ but at a later stage and to a lesser extent compared to *R. bourboniana*. In contrast, the transcript abundance of genes encoding a putative AUX/IAA protein (RC4G0454200), a dCTP diphosphatase (RC7G0053900), a putative NRT1/PTR FAMILY 6.4 protein (RC2G0538400) and a putative aquaporin (RC1G0470300) decreased strongly in 8 h ethylene-treated petal AZs of *R. bourboniana* suggesting suppression of these genes by ethylene. While this decrease was also seen in *R. hybrida* for the first two genes, expression of the aquaporin-like gene increased considerably in *R. hybrida* petal AZ and *R. bourboniana* petals while that of the NRT1/PTR FAMILY 6.4 gene increased slightly. The expression of another down-regulated zinc finger protein gene was unusual in that it showed a decline in expression in 8 h ethylene-treated petal AZ, as seen in the transcriptome, but this was preceded by an ethylene-induced increase of fivefold within 4 h.

**Comparative analysis between ethylene-treated *R. bourboniana* petal AZ versus petal and between petal AZs of *R. bourboniana* versus *R. hybrida* for identification of putative AZ-spe-
Figure 5. Comparative real time PCR validation of selected DEGs obtained from the transcriptome comparisons. Expression was studied in AZs at 0 h (untreated), 4, 8 and 12 h after 0.5 µL L⁻¹ ethylene treatment for R. bourboniana and additionally at 24, 36 and 48 h for R. hybrida. For petal, expression was studied at 0 h and 8 h after ethylene treatment. The real time expression data was normalized using ACTIN as reference. Error bars represent ± SE of three biological replicates. Expression values were analyzed by one-way ANOVA and compared using Duncan’s Multiple Range Test (DMRT) for AZ expression analysis. Values on the bar carrying different letters are significantly different. Paired t test was applied to analyze expression values in petal tissue. Error bars represent SE, * indicates P < 0.05, ** indicates P < 0.01, *** indicates P < 0.001.
components (Fig. 6, Table S9). Of these, 13 were down-regulated in the AZ while one chlorophyllase was up-regulated.

Proteins were the major TFs representing abscission-regulated DEGs. Of these 255 DEGs, 60 showed strong differential regulation as the most important form of regulation during abscission. The MYB/MYC/ERF and Zn finger proteins were the major TFs representing abscission-regulated DEGs. Of these 255 DEGs, 60 showed strong differential regulation even at log2 [|FC|≤ − 2 and ≥ 2]. The majority of these (37) were down-regulated while about a third (23) were up-regulated in R. bourboniana. These included six protein kinases, three TFs, one F-box protein, four transporters, three hormone pathway components, ten defence/phenylpropanoid pathway/wall separation components, nine encoding different enzymes and nine with unknown functions. Within the above groups, four of the kinase genes, six defence-related genes and all the four transporter genes were up-regulated. Strikingly, a substantial fraction with 15 members (25% of the total) encoded light reaction and photosystem components (Fig. 6, Table S9). Of these, 13 were down-regulated in the AZ while one chlorophyllase was up-regulated (Table S9). These DEGs represented genes that were specifically regulated in R. bourboniana petal AZ in response to ethylene and abscission cues.

Discussion

Plant organ abscission is an important developmental process that regulates reproductive success and productivity. Understanding its molecular basis has the potential to increase the economic value of several ornamentals by preventing premature flower/petal abscission. In rose, the fragrant variety is highly sensitive to ethylene and loses petals within 1–2 days post-pollination, decreasing its commercial value. In this study, high-throughput sequencing permitted us to understand the changes occurring within the small specialized AZ tissue that determines organ abscission but has only been studied occasionally due to the relative difficulty in its study.

The R. bourboniana transcriptome was assembled using the sequenced R. chinensis genome as a reference. A total of 32,961 genes out of 43,301 were expressed in the AZ. Of these, 3700 genes (representing ~ 8.5% of the total) were differentially expressed at log2 [|FC|≥ 1, ≤ − 1 (P value<0.05) of which almost 60% (2182 genes) showed a reduction in transcription. This is a much higher number than a previous study where AZ samples were collected at different stages of bud development of R. chinensis in absence of ethylene treatment. Our study confirms that ethylene accelerates abscission in fragrant roses by causing a major change in gene expression in the AZ.

The expression of reference genes, identified previously for rose, did not change in AZs even upon ethylene treatment, indicating their utility as reference for abscission studies (Fig. S2).

The major functional categories of the DEGs included regulatory genes (~ 11% DEGs) encoding transcription factors (284/2220), kinases (203/1849), phosphatases (86/503) and F-box proteins (218/2335) while other categories included hormone signalling (111), transporters (245) and, to a lesser extent, defense and stress response pathways. Surprisingly, light harvesting components (56/338) and the light-regulated phenylpropanoid pathway were strongly and disproportionately suppressed (Figs. 2, 3, 4, Tables S3, S4). All these groups have been identified previously as differentially regulated during abscission in other plants including rose.

One of the major findings of this study has been the rapid transcriptional activation of the ethylene pathway and suppression of auxin and JA pathways within 8 h of ethylene treatment. Ethylene accelerates the abscission process. Hence, a rapid induction of genes encoding components of ethylene perception and signalling in the present transcriptome (Fig. 3) was not surprising. Our own detailed analysis of the ethylene pathway confirmed the up-regulation of ethylene biosynthesis and signalling in R. bourboniana during abscission but its suppression in R. hybrida. Unlike ethylene, endogenous auxin prevents abscission by reducing AZ sensitivity to ethylene. A decrease in auxin response in the AZ allows endogenous ethylene to hasten abscission in various organ systems. Thus, down-regulation of most genes of the auxin pathway suggests that it may suppress petal abscission in rose through similar mechanisms (Table S4; Fig. 3).

Besides ethylene and auxin, JA affects abscission of fruits, leaves and flowers, independently from ethylene. JA is known to promote senescence as well as abscission. The JA-defective aos and coil mutants

Scientific Reports | (2020) 10:17196 | https://doi.org/10.1038/s41598-020-74144-3

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of Arabidopsis, show considerably delayed abscission indicating that JA promotes abscission. It also induces anther dehiscence which, like abscission, requires cell separation. In contrast to these studies, a majority of the JA pathway components, from biosynthesis to response, were down-regulated during ethylene-induced abscission in rose. These included JA biosynthesis genes such as PLA, LOX, AOC, OPR, JMT and the JA-responsive MYC homologue that activates JA-responsive genes. Several genes encoding JAZ proteins (negative regulators of JA signalling that are transcriptionally up-regulated during JA signaling) were also down-regulated. The results suggested that JA may suppress rose petal abscission, unlike in Arabidopsis. In this context, the strong down-regulation of the light pathway genes of the photosynthetic machinery (discussed later) is interesting. The initial steps in JA biosynthesis occur in the plastid and the down-regulation of many chloroplastic genes may disrupt plastid functioning and may, in some way, be related to the down-regulation of the JA pathway. Interestingly, the connection between JA and light signalling goes further with many aspects of JA responses requiring functional phytochromes and vice versa. Conclusions about the nature of the involvement of hormones like GA, ABA, cytokinin and SA were difficult, since genes involved in their biosynthesis as well as catabolism were differentially regulated (Fig. S3).

The activation of abscission-specific genes that drive abscission requires the action of TFs and post-translational regulators encoding kinases, phosphatases, receptors and various F-box proteins. Indeed, TFs of different families like ZF, MYB, MYC, ERF, NAC, HB and WRKY accounted for ~5% (284) of the total AZ DEGs highlighting their importance in regulating rose petal abscission (Fig. 2a). Studies of other abscission-related transcriptomes have also revealed their involvement although only a few members have been characterized to date for a detailed role in abscission. Some members of the ERF and MYB/MYC families regulate the ethylene and JA pathways. Interestingly, the ERF family (8 up, 7 down) showed a disproportionately larger number of up-regulated members possibly reflecting the greater involvement of ethylene in abscission while the BTB-POZ family (2 up, 9 down) was more prominently down-regulated as a group suggesting an abscission-inhibitory role. Although very few MADS genes were identified as differentially-regulated, these are important players in determining the AZ development. Two genes RC4G0415400 and RC7G0528400 showing similarity to the Arabidopsis AGL8 and AGL24 were significantly down-regulated at Log2FC < −1 (Table S3). AGL8, and its tomato homologue FUL,2 are known to be involved in seed shattering and abscission, respectively, with FUL2 expression suppressing style abscission in tomato. These genes may perform a similar function in rose petal abscission. AGL24 is the homologue of tomato JOINTLESS, a key abscission regulator required for AZ development. Four other MADS genes were significantly down-regulated but with a lesser change. The

Figure 6. Heat map showing the expression of putative abscission related genes in the comparison of ethylene-treated petal AZ and petals of R. bourboniana and petal AZ of R. hybrida. The 60 DEGs common to the 0 h versus 8 h ethylene-treated petal AZ transcriptomes in R. bourboniana show opposite regulation in the comparison between 8 h ethylene-treated petal AZ versus 8 h ethylene-treated petals of R. bourboniana and 8 h ethylene treated petal AZ of R. hybrida. Analysis was performed at [log2FC| ≤ −2 and ≥ 2)] and Q-value ≤ 0.05.
down-regulation of all these genes in rose probably reflects that these are not needed after formation of the AZ or may inhibit abscission progression.

Protein kinases and phosphatases, with 203 and 86 DEGs respectively (5% of the total DEGs), formed another important group regulating rose petal abscission. Post-translational modifications by these regulate the activity of several enzymes and proteins during developmental processes including abscission90. In Arabidopsis, receptor-like kinases such as HAESA and HAESA-LIKE2 promote abscission while other RLKs like SOMATIC EMBRYOGENESIS RECEPTOR-LIKE KINASE (SERK) positively regulate floral organ abscission in combination with HAE/HSLS91. Two other RLKs—SUPPRESSOR OF BIR1 (SOBIR1/EVERSHEDE) and CASTAWAY control the extent of cell separation during organ shedding91,92. These RLKs, along with MAP kinases, are differentially regulated during the cell separation phase of organ abscission in Arabidopsis96,37,41,49,93,94 and in citrus leaf AZ95.

Almost 40/50 DEGs were strongly down-regulated in the abscission transcriptome thus displaying a considerable indication of the importance of the transport of various compounds to the abscission process. ABC transporters are involved in membrane trafficking of a variety of substrates including lipids, drugs, steroids and other metabolic products9 and hormones100. At least 13 putative ABC lipid transporter genes have been reported to differentially express in the citrus fruit calyx AZ upon ethylene treatment108 whereas six ABC transporter genes were expressed during leaf abscission in sugarcane109. Differential expression of genes involved in transport of sugars, nutrients, metals, nitrate, lipids etc. has also been observed during abscission in olive56,102 and citrus101 while expression of sugar transporter genes was observed in calyx and pedicel AZ transcriptome profiling of citrus101 pyrus66 and tomato52. Collectively these results suggest that transporters may play a major role in membrane trafficking during abscission across plants.

The strong ethylene-induced suppression of the plastidial photosynthetic machinery genes does appear to be a component of changes associated with onset of fruit abscission104,103,108 as well as other AZ transcriptome profiling studies52,42,66. In rose, these genes accounted for 25% of the 60 most strongly regulated abscission-specific genes obtained from further comparisons with ethylene-treated petal AZ (Fig. 4c; Table S3). Transporters selectively govern the access of molecules in or out of the cells, thereby controlling the overall distribution of substances at their site of action or they may enable nutrient transport from the abscising organ to the parent organ prior to organ separation. Although the role of transporters in abscission is not clear, the fact that more than 200 genes encoding amino acid transporters, ABC transporters, ion transporters and drug transporters were differentially regulated during petal abscission in rose is indication of the importance of the transport of various compounds to the abscission process. ABC transporters are involved in membrane trafficking of a variety of substrates including lipids, drugs, steroids and other metabolic products9 and hormones100. At least 13 putative ABC lipid transporter genes have been reported to differentially express in the citrus fruit calyx AZ upon ethylene treatment108 whereas six ABC transporter genes were expressed during leaf abscission in sugarcane109. Differential expression of genes involved in transport of sugars, nutrients, metals, nitrate, lipids etc. has also been observed during abscission in olive56,102 and citrus101 while expression of sugar transporter genes was observed in calyx and pedicel AZ transcriptome profiling of citrus101 pyrus66 and tomato52. Collectively these results suggest that transporters may play a major role in membrane trafficking during abscission across plants.

One of the most prominent sets of DEGs in AZ was associated with the plastid photosystem components. Almost 40/50 DEGs were strongly down-regulated in the abscission transcriptome thus displaying a considerable difference from the 1:1.5 ratio of up/down genes in the AZ. Although the few cell layer thick AZ would not strike one as a major site for regulation of photosystem-related genes, decrease in expression of photosynthetic genes does appear to be a component of changes associated with onset of fruit abscission104,103,108 as well as other AZ transcriptome profiling studies52,42,66. In rose, these genes accounted for 25% of the 60 most strongly regulated abscission-specific genes obtained from further comparisons with ethylene-treated R. bourboniana petals and R. hybrida petal AZ. The strong ethylene-induced suppression of the plastidial photosynthetic machinery genes functioning in photosystem organization, light-harvesting, chlorophyll binding, electron transport and carbon fixation in rose petal AZ (Table S3) appears in some way to be necessary for the abscission process to begin or progress. Chloroplasts are the major source for generation of reactive oxygen species (ROS) and a disruption in chloroplast photosynthetic machinery increases ROS and activates senescence102. The decrease in expression of various photosystem components during petal abscission in rose may similarly disrupt the photosynthetic electron transport machinery leading to increased ROS generation and abscission although this will require more detailed studies. Another major component preferentially expressed in 8 h ethylene-treated petal AZ of R. bourboniana is associated with cell death and disassembly of proteins, nucleic acids and cell wall components. Disassembly of cellular structures have been noted in the context of abscission and observed in previous studies of abscission102,106–108 including our own99,103 where genes encoding cysteine proteases and components of PCD and autophagy were up-regulated. A large number of cell wall remodelling genes encoding XTHs, expansins, cellulase synthases, pectate lyases and arabinogalactan proteins probably contribute to cell separation and fortification of the wall after abscission1,11,12,49,50,55,60–62,109–114 and was also observed in rose. Since organ separation exposes the nascent tissue to pathogens, protection of this tissue is necessary and is associated with expression of a large number of defence-related genes12,13,49,50,115–118. Genes encoding PR-proteins and those associated with...
Further analysis to obtain genes specific to *R. bourboniana* AZ was carried out with comparisons with ethylene-treated *R. bourboniana* petals to exclude genes represented in petals (with which the petal AZ is closely attached) and those represented in ethylene-treated petal AZ of *R. hybrida* (which shows reduced ethylene sensitivity and delayed abscission). The analysis revealed a total of 845 and 790 DEGs respectively (at \(\log_{2}FC \geq 1, \leq -1\), \(P < 0.05\); Tables S7, S8) within which 255 DEGs were identified as being part of a core set of genes that were regulated by ethylene only in *R. bourboniana* petal AZ but not in petals and *R. hybrida* AZ. Strikingly, 58/255 genes (~23%) belonged to the regulatory category consisting of TFs/kinases/phosphatases/F-box proteins. This is much higher than the proportion (11%) observed in comparison between 0 and 8 h ethylene-treated petal AZs and suggests a specific requirement for this group in abscission (Table S9). The TF subset accounting for 10% of these genes included members of the AP2-ERF/MYB/MYC/Zn finger families. About 60/255 DEGs were uniquely regulated in AZs at \(\log_{2}FC \geq 2, \leq -2\) (\(P < 0.05\)). The majority of the kinases, defence-related and the transporter genes in this group were up-regulated (Table S9) while 13 of the 15 photosystem-related genes were strongly down-regulated. The down-regulation of these plastid-associated genes along with the light-regulated phenylpropanoid pathway indicates that light-associated components may have a strong negative effect on progression of abscission and is currently under study. A comparison of this core set of 255 genes with Arabidopsis reveals that at least 58 have homologues in Arabidopsis that show abscission-related expression in Arabidopsis siliques and floral parts as shown in Genevisible (https://genevisible.com/search) and other studies\(^{49,56,89,119–124}\) (Table S9). Interestingly, these include two TFs, WIN1 and MYB94 both involved in wax biosynthesis\(^{120,123}\), MYB117, WRKY33\(^{121}\) (a PAMP and SA responsive TF), seven receptor kinases\(^{89,119}\), eight transporters and several genes encoding different enzymes. The functions of all these genes in the context of abscission are not clear and will require future knock-out and over-expression studies. Nevertheless, the studies suggest that a large fraction of the abscission machinery representing different functional groups is conserved across families as diverse as rose and Arabidopsis, despite tissue differences.

In conclusion, ethylene-induced rose petal abscission is associated with large scale gene expression changes distinct from those in the petal. It appears to be guided by changes in responses of the ethylene, auxin and JA pathways and related ERFs/MYB/MYC that may affect sensitivities to these hormones. The expression of phytochromes and the prominent suppression of light-associated photosystem and phenylpropanoid components suggest a close interaction of light signalling and plastid with abscission. Expression of regulatory genes like TFs, kinases/phosphatases/F-box genes that include homologues of MADS genes *AGL8*, *AGL24* and RLKs like *HSL1* and *RLK7* besides transporters and defense pathways suggest some conservation with Arabidopsis with certain components like the JA pathway being regulated in a unique manner in rose.

**Material and methods**

**Plant material, treatments and RNA isolation.** Flowers of *Rosa bourboniana* (cv Gruss an Teplitz) and *R. hybrida* (Opening Night) were chosen for study. Excised flowers were treated with 0.5 µl/L exogenous ethylene to trigger abscission which is completed in 16–18 h in *R. bourboniana* but 48–50 h in *R. hybrida*. The 8 h time point represents the mid-point of abscission. RNA was isolated from *R. bourboniana* petals to exclude genes represented in petals (with which the petal AZ is closely attached) and those represented in ethylene-treated petal AZ of *R. hybrida* (which shows reduced ethylene sensitivity and delayed abscission).

**Illumina sequencing.** The cDNA libraries were generated from mRNA of the above samples. The 0 h and 8 h ethylene-treated petal AZ RNA from *R. bourboniana*, the 8 h ethylene-treated petal AZ RNA from *R. hybrida* and ethylene-treated petal RNA of *R. bourboniana* were used for paired-end sequencing (with 100 bp ends) on an Illumina HiSeq2000 sequencing machine (Scigenome, Kochi, India). About 45–67 million reads were generated per sample. Low quality sequences and adaptor sequences were removed and reads trimmed by NGSQC-TOOLKIT (https://www.nipgr.res.in/ngsqctoolkit.html). More than 85% high quality reads with a quality score (Q > 30) value were generated from each library for analysis.

**Assembly and gene expression analysis.** The filtered high quality reads were used in reference-guided assembly using *R. chinensis*\(^{68}\) genome as reference with TopHat v2.1.1\(^{255}\) (https://ccb.jhu.edu/software/tophat/index.shtml) and Bowtie programs\(^{126,127}\) as alignment tools and SAM Tools\(^{128}\) for processing the alignment file. The aligned reads were counted and estimated for gene expression as FPKM using cufflinks v2.2.1\(^{129}\) software\(^{126}\),\(^{127}\) as alignment tools and SAM Tools\(^{128}\) for processing the alignment file. The aligned reads were counted and estimated for gene expression as FPKM using cufflinks v2.2.1\(^{129}\) software\(^{126}\),\(^{127}\) as alignment tools and SAM Tools\(^{128}\) for processing the alignment file. Cuffdiff was used to estimate genes showing significant differential expression using the Cufflinks package\(^{129}\) (http://ial/cole-trapnell-lab.github.io/cufflinks/manual/).

**Functional annotation.** To identify the putative functions, assembled transcripts were annotated using GO\(^{130}\), KEGG\(^{131,132}\) (https://www.genome.jp/kegg/) and InterPro (ftp://ftp.ebiinfo.wsu.edu/species/Rosa_chinesis/Rchinensisgenome.v1.0/functionali/) and integrated with our result using in-house custom scripts in R programming language. A heat map was generated for the differentially regulated contigs using MeV version 4.9 (https://www.tm4.org/mev.html). DEGs with \(\log_{2}FC(\leq -1 \text{ and} \geq 1)\) were grouped under various categories for functional cataloguing (Figs. 2, 3, 4). For analysis of common genes in all three comparisons, DEGs with \(\log_{2}FC(\leq -2 \text{ and} \geq 2)\) were used (Fig. 6).
Validation of mRNA-seq data using qRT-PCR. Real time PCR was performed to quantify the expression of selected DEGs (Table S5) obtained from Illumina sequencing. The cDNAs, used as template, were generated from mRNA using the REVERTEAD MMLV reverse transcriptase (Fermentas). Primers (Table S6) were designed using the primer designing tool at IDT (https://eu.idtdna.com/site) to amplify an amplicon of 80–150 nucleotides with Tm around 60 °C. Reactions were run in triplicates (technical and biological) for each sample using Power-Up SYBR Green on an ABI StepOnePlus real time PCR machine (Applied Biosystems Inc, USA) and the data analyzed was the mean of biological triplicates. The reaction was set up in 20 μl as follows: 1 μl of cDNA, 10 μl SYBR Green Dye master mix (2X), 5 pmol each of forward and reverse primers and water up to 20 μl. The general steps performed during real-time PCR experiment were as follows: step 1, 50 °C, 2 min, step 2, 95 °C, 10 min, step 3 (95 °C 15 s, 60 °C 1 min) × 40 cycles. The specificity of the amplicon was analyzed by a melt curve analysis. The relative mRNA level of the gene in different RNA samples was normalized with respect to ACTIN as the internal control gene\(^6\) and analyzed by 2^−ΔCT method\(^{33}\).

Data access
The sequence data has been submitted to NCBI under the SRA accession: PRJNA594099 (Temporary Submission ID: SUB6655877), Release date: 2021-01-04.

Received: 10 April 2020; Accepted: 8 September 2020
Published online: 14 October 2020

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Acknowledgements

We are grateful to the Council of Scientific and Industrial Research, India for funding the work under the project BSC107 and for Junior and Senior Research Fellowships to PS, SKT, AP Singh, SPP and ASC. We thank Mr Ram Awadh for taking care of the rose plants.

Author contributions

A.P. Sane conceived the experiment. P.S., A.P. Sane, S.K.T., S.P. P performed the bioinformatic analysis. P.S., N.B., A.S.C. and A.K. performed the bioinformatic analysis. P.S. and A.P. Sane analyzed the data and wrote the paper. All authors read and approved the manuscript. CSIR-NBRI manuscript number is ‘CSIR-NBRI_MS/2020/03/01’.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-74144-3.

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