In silico analysis of MeJA-induced comparative transcriptomes in Brassica oleraceae L. var. capitata

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Abstract Brassica oleracea var capitata is a member of the Brassicaceae family and is widely used as an horticultural crop. In the present study, transcriptome analysis of B. oleraceae L. var capitata was done for the first time using eight-week old seedlings treated with 50 µM MeJA, versus mock-treated samples. The complete transcripts for both samples were obtained using the GS-FLX sequencer. Overall, we obtained 275,570 and 266,457 reads from seedlings treated with or without 50 µM MeJA, respectively. All the obtained reads were annotated using biological databases and functionally classified using gene ontology (GO), the Kyoto Encyclopedia of Genes and Genomics (KEGG). By using GO analyses, putative transcripts were examined in terms of biotic and abiotic stresses, cellular component organization, biogenesis, and secondary metabolic processes. The KEGG pathways for most of the transcripts were involved in carbohydrate metabolism, energy metabolism, and secondary metabolite synthesis. In order to double the sequenced data, we randomly chose two putative genes involved in terpene biosynthetic pathways and studied their transcript patterns under MeJA treatment. This study will provide us a platform to further characterize the genes in B. oleracea var capitata.

Keywords Brassica oleracea, EST, Gene ontology, KEGG pathway, Methyl jasmonate

Introduction

B. oleraceae L. var. capitata (cabbage) is one of the most important vegetable crops belong to the species B. oleraceae L., and grouped under the family Brassicaceae. It is an herbaceous, biennial and dicotyledonous flowering plant with leaves forming a characteristic compact head. Cabbage has a positive impact on human health and is, in addition to being a source of vitamins and fiber, connected with secondary metabolites called glucosinolates, which are known to possess anti-carcinogenic properties (Sarikami et al. 2009). Brassicaceae vegetables are a good source of antioxidants because of their high phenolics and glucosinolate content. Throughout the growth and developmental stages, plants are ordinarily exposed to various environmental biotic and abiotic factors. Biotic and abiotic stresses are major concern for the sustainable production of these crops. Each type of biotic and abiotic stresses function through different types of molecular mechanisms that affect plants, and eventually cause damage (Dixon and Lamb 1990). In particular, some biotic and abiotic elicitors can activate specific secondary metabolite production. Under these conditions, a number of signaling pathways can be pre-activated by salicylic acid (SA), jasmonic acid (JA), ethylene or abscisic acid pathways which are generally, involved in the defense responses. Efforts to develop plants resistant to biotic and abiotic stresses is an important task in biotechnology and functional genomics study of which is one of the most important tool for identifying potential genes related to stress resistance.

Genome sequence of B. oleracea was not available till now. An alternative and efficient method for analyzing transcriptome can be done using ESTs. ESTs are short, unedited, randomly selected single pass sequence read derived from cDNA libraries, also called as ‘poor’ man’s genome and proven to be a valuable tool in molecular biology ( Nagaraj et al. 2007). Recently, EST sequencing is a widely used application tool to study gene expression pattern in response to a given...
environmental stimulus and a transcriptomics study of a plant at various stages of development under different experimental conditions with various plant tissues (Kim et al. 2003). EST technology was primarily introduced in the human genome project, and used widely to clone new genes, determine tissue-specific gene expression profiles, annotate functional genome sequences and so on (Ohlrogge and Benning 2000; Brandle et al. 2003). Raw sequences obtained from high-throughputput screening are quite impossible to annotate manually. In these cases, computational methods are used to process EST sequences, such as sequence cleaning, vector masking, clustering, assembly and annotation to yield biological information to putative sequences. The use of a bioinformatics tool for homology-based functional annotation and statistical information is simple for ESTs from a range of organisms in the public non redundant databases (Falgueras et al. 2006). Gene ontology (GO), are liable biological annotation schema that is globally accepted by biological communities, was developed with three structured vocabularies (i.e., ontologies) to describe genes and proteins in terms of cellular components, biological processes, and molecular function (Pal 2006).

In this study, in silico approach was used to classify B. oleracea L.var. capitata ESTs based on GO vocabularies, quantification of transcript abundance, classification of biochemical pathways based on KEGG pathway, and alternatively spliced transcripts. Putative functional protein domains were also analyzed using biological databases. Using similarity search, unique sequences was assigned as putative sequences and further analysis of those sequences aid to understand a complex gene network expression.

Materials and methods

Plant materials and MeJA treatments

B. oleracea seeds were germinated in soil under in vitro conditions at 25°C with continuous light and dark conditions (16h/8h). Each pot had five cabbage seedlings. After eight weeks, seedlings were treated with 50 µM MeJA and without (mock). For treatment, sterilized filter paper (Whatman No. 6, ø185 mm) was dipped in 5 mL of 40 % ethanol (control) or 50 µM MeJA placed in pots on the top of cabbage seedlings without touching its leaf surfaces. After treatment, pots were covered by polyethylene bags and wrapped well as shown in Figure 1. All the plants were grown at 25°C, 16h light/8h dark condition for 3 days. Cabbage leaves harvested and frozen immediately with liquid nitrogen and stored at -70°C which was used for sequencing. For expression analysis, eight week old seedlings were treated with and without 50 µM MeJA for up to 9 days. Seedlings were collected every alternative day (3rd, 5th, 7th, and 9th day) and immediately frozen with liquid nitrogen and stored at -70°C until required.

RNA isolation and cDNA library construction

The total RNA was isolated via the aqueous phenol extraction procedures using TRIzol® Reagent (Thermo Fisher Scientific, Catalog No. 15596-026). Messenger RNA was isolated from 50 µg of the total RNA sample using the FastTrack™ MAG Micro mRNA Isolation Kit (Life Technologies, cat# K1580-01). The kit was used following the protocol as provided by the manufacturer. The isolated mRNA was quantified using Agilent RNA 6000 NANO CHIP (Agilent, cat.# 5067-1511) and 200 ng of this material was used in the cDNA Rapid Library Preparation protocol. The first step of cDNA Rapid library preparation is fragmentation of mRNA using ZnCl2 and heat treatment. The cleaved RNA fragments primed with random hexamers were reverse transcribed into first strand cDNA, and double-stranded (ds) cDNA was generated after removing the RNA template and synthesizing a replacement strands. The double-stranded cDNA ends were polished (blunted), and short adapters were ligated onto both ends. The adapters provide priming sequences for both amplification and sequencing of the sample library fragments, as well as the “sequencing key”, a short sequence of 4 nucleotides used by the system software for base calling and, following purification and size selection using AMPure beads. Finally the quality of the library of cDNA fragments was assessed using 2100 BioAnalyzer (Agilent), and the library was quantitated to determine the optimal amount of the library to use as input for emulsion-based clonal amplification.

Emulsion PCR and sequencing run

Single “effective” copies of template species from the DNA library to be sequenced were hybridized to DNA Capture Beads. The immobilized library was then resuspended in the amplification solution, and the mixture is emulsified, followed by PCR amplification. After amplification, the DNA-carrying beads were recovered from the emulsion and enriched. The second strands of the amplification products were melted away as part of the enrichment process, leaving the amplified single-stranded DNA library bound to the beads. The sequencing primer is then annealed to the immobilized amplified DNA templates. After amplification, the DNA-carrying beads were set into the wells of a PicoTiterPlate device (PTP) such that wells contain single DNA beads. The loaded PTP was
then inserted into the Genome Sequencer FLX Instrument, and sequencing reagents were sequentially flowed over the plate. Information from all the wells of the PTP is captured simultaneously by the camera, and can be processed in real time by the onboard computer. Samples were sequenced by Macrogen Inc, Korea (http://www.macrogen.com)

RNA Isolation and semi quantitative PCR analysis

RNA was extracted from B. oleraceae that were subjected to the experimental treatment using the RNeasy kit (Qiagen, USA) according to the manufacturer’s instructions. The quality and concentration of RNA was measured using a spectrophotometer (GE nanovalue, USA). To obtain the first strand cDNA, 1.5 µg of total RNA was used, and cDNA was synthesized using a Power cDNA kit (Invitrogen, USA) following the manufacturer’s instructions. We performed RT-PCR using optimum PCR conditions: 92°C for 2 min, 92°C for 40 sec, 54.5 ~ 60°C for 30 sec, 72°C for 1 min, 72°C for 10 min for 28 cycles, gene specific primers along with optimum annealing temperatures were listed in Table S1. The housekeeping gene encoding actin was used as a standard for all samples. For the analysis of all transcripts, untreated samples were used as negative control.

Unigene dataset generation

The unigene data set contains a set of non-redundant sequences composed of singlets and contigs. Sequence files were produced with the SFF (Standard Flowgram Format) file, and then SFF files were processed using the GS assembler FLX software tool kit (v.2.6) provided by Roche. During the assembly process, the software identified pairwise overlaps between reads, constructed multiple alignments of overlapping reads and divided or introduced breaks into the multiple alignments in regions where consistent differences are found between different sets of reads. The software attempted to resolve branching structures between contigs, and generated consensus base calls of the
contigs by using quality and flow signal information for each nucleotide flow included in the contigs multiple alignments. The contig consensus sequences and corresponding quality scores were generated along with an ACE file of the multiple alignments and assembly metrics files. When paired end data is available, the assembler performed extra steps; organized the contigs into scaffolds using Paired End information to order and orient the contigs and to approximate the distance between contigs, and output scaffolded consensus sequences and corresponding quality scores, along with an AGP file of the scaffolds and specific metrics tables. A de novo assembler project was created for the short cDNA sequence reads with default parameters. Using SeqClean and Lucy tools, we obtained singleton and contig sequences. Low quality and low-complexity sequences were removed using SeqClean (http://sourceforge.net/projects/seqclean/). The Lucy as a sequence cleanup program used for the processes of quality assessment, confidence re-assurance, vector trimming and vector removal. The remaining sequences were used in the functional analysis (Fig. 1).

Gene ontology and KEGG pathway assignment

Gene ontology (GO) term annotation and function-based analysis of unique sequences were performed using Blast2go (B2G), a sequence-based high-through put sequence analysis tool. Using B2G, unigenes were subjected to a BLASTX query against the national center for biotechnology information (NCBI) public non-redundant (NR) database. Based on the BLASTX results, the sequences were putatively named using the BLAST description annotator (BDA) tool embedded in B2G. More collective logic models were embedded in B2G to retrieve GOs, EC numbers, and KEGG maps. GO terms for each of the three main categories (biological process, molecular function and cellular component) were obtained from sequence similarity and BLAST scores (E=10-3) with default parameters. Those annotations were simplified into plant functional categories using the plant GO slim. Interproscan embedded with B2G was used to obtain the protein domain information for the putative sequences (Conesa and Gotz 2008).

Comparative analysis of alternative splicing in MeJA treated samples and control samples

The Arabidopsis genome sequence, annotation and annotated sequence features were downloaded from TAIR (TAIR 10 database release). All transcripts sequences were mapped against the Arabidopsis genome using the C/S version of the BLAT program. We used the BLAT version 35 downloaded from Dr. W.J. Kent’s homepage (http://users.soe.ucsc.edu/~kent/src/). BLAT output contains many suboptimal alignments. Only the best alignment with the highest BLAT score was kept unless we had multiple hits of the same quality. We first grouped transcripts sequences and reference genes into clusters on the genome if they mapped onto the same genomic region, were orientated on the same strand, and had overlapping sequences. Alternative splicing events and differentially expressed gene features were identified using database queries. Only initial and terminal nucleotides were allowed to remain unmapped, >90% of the transcripts sequence had to be involved in perfect matches with the genome, and every exon had to be ≥ 85% identical (or contain at most five errors) to the genome. These transcripts were clustered as described above. Single-exon and unspliced transcripts were discarded as they do not reveal any information about alternative splicing.

Results and Discussion

Sequence and EST assembly

In order to find optimum concentration of MeJA, growth test was done on 10-, 50-, 100 µM of MeJA containing medium using 9-day grown seedling for 3, 5, 7, 9 days respectively (Fig. 2A). Shoot dry weight (Fig. 2B) was measured, and found 50 µM is in biological range and 100 µM is slightly over that range. For the construction of cDNA library, eight-week old B. oleraceae seedlings were treated with 50 µM MeJA and without (control) (Fig. 2C), and previously reported MeJA-inducible glucosinolate biosynthesis-related gene's expression (Mikkelsen et al. 2003; Pozo et al. 2008) was confirmed that the concentration was still working in different aged plant (Fig. 2D). The quality of cDNA library was assessed using 2100 Bio analyser (Agilent). The cDNA library was constructed using GS-FLX sequencer (Roche v2.6) resulted in the total of 275,570 and 266,457 reads respectively. All the obtained reads were subjected to seq clean (http://sourceforge.net/projects/seqclean/) and lucy (http://lucy.sourceforge.net/) with default parameters for removing low quality and vector sequences, and assembled into 458,961 ESTs. Control and MeJA treated ESTs had an average length of 442 bp and 448 bp respectively. The maximum number of ESTs was higher in control compared to the MeJA treated samples. All the assembled sequences were functionally annotated using blast2go (Table 1). EST similarity search against public non-redundant databases

Homology-based functional annotation of putative sequences was obtained through BLAST X queries against public non-
Fig. 2 Physiological status of B. oleracea L. var capitata following MeJA treatment (A) B. oleracea L. var capitata grown for nine days were mock treated or treated with 10, 50, or 100 µM of MeJA. (B) Shoot dry weight (DW) gradually decreased as MeJA concentration and number of treatment days increased. (C) Eight-week old B. oleracea L. var capitata were treated with 50 µM MeJA for 3 days. Relative expression patterns of MeJA-inducible flavin-containing monooxygenase (FMO) and N-hydroxylase for short chain methionine derivative (Cyp79F1) show that the MeJA treatment was done properly

Table 1 EST assembly and annotation reports

| Description                          | No of contigs (Control) | No of contigs (MeJA) |
|--------------------------------------|-------------------------|----------------------|
| Number of contigs                    | 11,248                  | 6,410                |
| Number of Contigs with GO (redundancy) | 96,371                  | 64,952               |
| Number of Contigs with GO (non redundancy) | 9,272                   | 5,414                |
| Number of Contigs without GO         | 1,976                   | 996                  |
| Number of sequence with EC           | 4,879                   | 3,851                |
| Number of Contigs enzyme codes       | 1,098                   | 997                  |

redundant databases with an average E-value of $10^{-3}$ or below, an HSP cut off of 33 and a maximum of 20 blast hits per sequence (Gotz et al. 2008; Mao et al. 2005). The BLAST X search for the control showed 9563 (85.2%) meaningful matches and 1685 (14.9%) didn’t show any significant hits. Similarly, MeJA treated samples showed 5636 (84.3%) matches with known sequences and 1049 (15.6%) didn’t show any similarity matches. The total blast hits for the control samples of B. oleracea var capitata showed 74.89% from the following plants: Arabidopsis lyrata, Arabidopsis thaliana, Brassica rapa, Thellungiella halophila, Zea mays, Brassica napus, Brassica oleracea, and Brassica juncea. The results of both control and MeJA treated samples are displayed separately in Table 2. The abovementioned plants have been relatively well-studied, so there are more experimental data rather than in B. oleracea L. var capitata. Annotations were obtained using BLAST description
Table 2 Classification based on species from BLAST hit sequences for control and MeJA-treated samples

| Species                | Control | MeJA  |
|------------------------|---------|-------|
| *Arabidopsis lyrata*   | 2,656   | 1,528 |
| *Arabidopsis thaliana* | 1,937   | 1,114 |
| *Brassica rapa*        | 1,287   | 734   |
| *Thellungiella halophila* | 495    | 326   |
| *Zea mays*             | 311     | 207   |
| *Brassica napus*       | 268     | 318   |
| *Brassica oleracea*    | 147     | 165   |
| *Brassica juncea*      | 81      | 91    |

annotator (BDA) tool, and all the putative sequences were used for further experimental analysis.

Functional analysis based on gene ontology

Functional gene annotation is a difficult task for newly sequenced non-model plants than human since the plant genome contains numerous genes reflecting adaptations to environmental factors. To simplify this process, gene ontology (GO) has evolved in the field of functional genomics. GO describes the function of gene based on control vocabularies and for elucidating hierarchical relationships between gene groups. Control vocabularies are grouped into three major categories, namely molecular function, biological process, and cellular component (The Gene Ontology 2010). The data obtained through this study were organized into one, a combination of two and belonging to all three categories. The data were then grouped in a Venn diagram (Fig. 3). The total EST sequences were 3,617 (2,322), 943 (498), 31,418 (21,236) for the control and MeJA (in parenthesis) respectively, and grouped into cellular component (CC), molecular function (MF), and biological process (BP). A total of 381 ESTs were annotated into CC and MF groups, 2294 ESTs were annotated into MF and BP, and 5143 ESTs were annotated into BP and CC groups, and a total of 8420 ESTs were mapped into all three GO categories in case of MeJA-treated samples. According to the plant-GOslim, plant-specific GO vocabularies were screened. In our result, a large number of unique sequences were grouped under the first category of biological processes with subcategories such as the responses to cadmium ion, salt stress, oxidation-reduction process, cold, regulation of plant-type hypersensitive response, serine family amino acid metabolic process, protein targeting to membrane, chitin, DNA-dependent regulation of transcription, glycolysis, bacterium, gluconeogenesis, protein phosphorylation and wounding (Table S2). It is noteworthy that the number of wound responsive and Golgi organization-related contigs are more highly ranked by MeJA treatment (Table S2). The second category included cellular component which includes unique sequences associated with plasma membrane, nucleus, cytosol, chloroplast, plasmodesmata, Golgi apparatus, mito- chondria, integral to membrane, chloroplast stroma, chloroplast envelope, vacuolar membrane, cytoplasm, nucleolus and so on (Table S3). The third representative category was molecular function with ATP binding, protein binding, zinc ion binding, sequence-specific DNA binding transcription factor activity, DNA binding, nucleotide binding, protein serine/threonine kinase activity and so forth based on their hit numbers of contigs (Table S4). The GO results rely on previously annotated GO information of other plants.

Generally, various transcription factors and kinases play a crucial role in stress related signaling pathways (Horan et al. 2008). For example MeJA is the important signaling cascade linked to various biotic and abiotic stress mechanisms that modulates various physiological processes in plants, including root senescence, and the defense responses against insect and pathogen attack (Wasternack and Parthier 1997; War et al., 2012). Also MeJA induces or increases the biosynthesis of plant secondary metabolites that play an important role in various environmental conditions (Yukimine et al. 1996; Ramakrishna and Ravishankar, 2011). For instance, MeJA increases the production of soyasaponin in glycyrrhiza glabra cells (Hayashi et al. 2003), saikosaponin in the adventitious roots of *Bupleurum falcatum* (Aoyagi et al., 2001), and ginsenoside production in *P. ginseng* (Shim et al., 2010). For example a putative gene transcript of squalene monooxygenase located in integral to membrane (cellular component), involved in biosynthetic process (biological process), and resulted in nucleotide binding activity (molecular function) (Table S1)

KEGG biochemical pathway analysis

As an alternative method of grouping unique sequences on the basis of biochemical functions, were assigned to metabolic pathways via Kyoto Encyclopedia of Genes and Genomes...
Table 3 Secondary metabolite-related genes in *Brassica oleraceae* L. var. capitata based on KEGG biochemical analysis

| Pathways                      | Enzymes               | ECNo.       | No. of EST | Pathways                      | Enzymes               | ECNo.       | No. of EST |
|-------------------------------|-----------------------|-------------|------------|-------------------------------|-----------------------|-------------|------------|
| Phenyl propanoid biosynthesis | Peroxidase            | EC:1.11.1.7 | 21         | Phenyl propanoid biosynthesis | Prenylcysteine oxidase| EC:2.3.1.92 | 3          |
|                               | Cinnamoyl-CoA reductase | EC:1.2.1.44 | 13         |                               | Hydroxymethylglutaryl-CoA reductase | EC:1.2.1.68 | 2          |
|                               | Cinnamyl-alcohol dehydrogenase | EC:1.1.1.195 | 6         |                               | Hydroxymethylglutaryl-CoA reductase (NADPH) | EC:2.1.1.104 | 1          |
|                               | Trans-cinnamate 4-monoxygenase | EC:1.14.13.11 | 6         |                               | Flavonol synthase | EC:1.14.11.23 | 9          |
|                               | 4-Coumarate---CoA ligase | EC:6.2.1.12 | 6          |                               | Trans-cinnamate 4-monoxygenase | EC:1.14.13.11 | 6          |
|                               | Beta-glucosidase       | EC:3.2.1.21 | 5          |                               | Dihydrokaempferol 4-reductase | EC:1.1.1.219 | 5          |
| Sinapoylglucose---malate O-sinapoyl transferase | EC:2.3.1.92 | 3          |            |                               | Flavonoid 3-monoxygenase biosynthesis | EC:1.14.13.21 | 3          |
|                               | Coniferyl-aldehyde dehydrogenase | EC:1.2.1.68 | 2          |                               | Caffeoyl-CoA O-methyltransferase | EC:2.1.1.104 | 1          |
|                               | Caffeoyl-CoA O-methyltransferase | EC:2.1.1.104 | 1         |                               | Shikimate O-hydroxycinnamoyltransferase | EC:2.3.1.133 | 1          |
|                               | Caffeate O-methyltransferase | EC:2.1.1.68 | 1          |                               | Leucocyanidin oxygense | EC:1.14.11.19 | 1          |
|                               | Sinapate 1-glucosyltransferase | EC:2.4.1.120 | 1         |                               | Myrccene synthase | EC:4.2.3.15 | 7          |
|                               | Quinate O-hydroxycinnamoyltransferase | EC:2.3.1.99 | 1         |                               | Monoterpene and Diterpene Biosynthesis | EC:1.1.1.207 | 4          |
|                               | Shikimate O-hydroxycinnamoyltransferase | EC:2.3.1.133 | 1         |                               | (+)-neomenthol dehydrogenase | EC:1.1.1.207 | 4          |
| Terpene backbone biosynthesis | 1-deoxot-D-xylulose-5-phosphate reductoisomerase | EC:1.11.1.7 | 2          |                               | (-)-menthol dehydrogenase | EC:1.1.1.207 | 4          |
|                               | Acetyl-CoA C-acetyltransferase | EC:1.2.1.44 | 13         |                               | (R)-limonene synthase | EC:4.2.3.20 | 2          |
|                               | (E)-4-hydroxy-3-methylbut-2-enyl-diphosphate synthase | EC:1.1.1.195 | 6         |                               | (+S)-limonene synthase | EC:4.2.3.16 | 2          |
|                               | 1-deoxy-D-xylulose-5-phosphate synthase | EC:1.14.13.11 | 6         |                               | taxane 13alpha-hydroxylase | EC:1.14.13.77 | 2          |
|                               | 4-hydroxy-3-methylbut-2-enyl diphosphate reductase | EC:6.2.1.12 | 6          |                               | Glucosinolate Biosynthesis | EC:2.6.1.42 | 7          |
|                               | Dimethylallyltransferase | EC:3.2.1.21 | 5          |                               | N-hydroxysphingoimide S-beta-glyosyltransferase | EC:2.4.1.195 | 1          |
|                               |                       |             |            |                               | 9-cis-epoxycarotenoid dioxygenase | EC:1.13.11.51 | 8          |
|                               |                       |             |            |                               | abscisic-aldehyde oxidase | EC:1.2.3.14 | 2          |
|                               |                       |             |            |                               | zeaxanthin epoxidase | EC:1.14.13.90 | 1          |
|                               |                       |             |            |                               | phytoene synthase | EC:2.5.1.32 | 1          |
|                               |                       |             |            |                               | (+)-abscisic acid 8'-hydroxylase | EC:1.14.13.93 | 1          |

(KEGG) (Kaneshisa M, 2000). Using enzyme commission (EC) numbers as the basis for assignment all the putative ESTs were subjected to KEGG database query with BLAST score to retrieve KEGG enzyme codes and pathway maps. In this study we identified and categorized KEGG pathways into carbohydrate metabolism, amino acid metabolism, nitrogen metabolism, lipid metabolism, and secondary metabolism (Table S5). Among annotated KEGG pathways, several secondary metabolite-related pathway genes were listed up (Table 3). Phenyl propanoid contributes to all aspects of plant responses including biotic and abiotic stresses (La camera et al. 2004). Phenolic compounds are the biologically active compounds used in traditional medicine as anti-viral, anti-cancer, anti-inflammatory agents (Korkina 2007). Terpenoids pathways are classified into groups based on the number of carbon atoms attached to isoprene units. The major groups of terpenes are monoterpenes (C10), sesquiterpenes (C15), diterpenes (C20), and triterpenes (C30). We also identified ESTs involved in other pathways such as steroid metabolism, glucosinolate pathways, and flavonoid pathways.
Secondary metabolite-related gene expressions and alternatively spliced genes by MeJA

MeJA is a small signaling molecule in the plant kingdom. Environmental stresses such as wounding and pathogen attack can induce MeJA production. Thus, screening MeJA-responsive genes can be a useful pool for the utilization of genetic engineering of crop plants. Several previously reported genes were also screened from MeJA-treated *B. oleraceae* (Fig. 4A). Several known genes including lipoxygenase, myrosinase-associated protein, and lipase, were read abundantly as reported in other plants (Ren et al. 2013). Other uncharacterized genes might be cabbage-specific suitable candidates. In plant, MeJA is well-known to induce monoterpenoid indole alkaloids and isoprenoid biosynthesis. Two selected isoprenoid biosynthesis-related genes, squalene monoxygenase and squalene synthase 1, were up-regulated by MeJA within 3 ~ 5 days respectively using semi-quantitative PCR. The genes selected in the present study are attributed to play an important role in direct or indirect defense mechanisms.

Alternative splicing has recently caught the attention of many plant researchers, as it can be spatially and temporally regulated, and is frequently associated with tissue types and environmental conditions to enhance transcriptome plasticity and proteome diversity. When all the obtained cabbage transcripts were mapped against the Arabidopsis genome using the BLAT program, total 33 transcripts which covered 80% showed different alternative splicing among the control and MeJA-treated samples (Fig. 5A). For example, the exact alternative splicing zone of Acyl-CoA: diacylglycerol acyltransferase (DGAT) and CBL-interacting protein kinase 1 are displayed.

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Fig. 3 Venn diagram of *Brassica oleraceae* data set showing numbers annotated to one, a combination of two, and/or all three GO vocabularies. Numbers in parentheses indicate the results from MeJA-treated samples compared to those of the control (MF, molecular function; BP, biological process; CC, cellular compound).

Fig. 4 MeJA-inducible genes from *Brassica oleraceae* L. var. capitata. (A) Top 10 reads from control and MeJA-treated EST. (B) Expression patterns of selected genes assessed using semi-quantitative PCR analysis. (*SQMO*, squalene monoxygenase; *SS1*, squalene synthase 1)
Fig. 5 Alternatively spliced genes as compared with the Arabidopsis genome. (A) List of a total of 33 alternatively spliced genes. (B) Two of the selected genes show descriptive alternative splicing between the control and MeJA-treated samples (Fig. 5B). Taken together, all the transcripts and differentially transcribed genes in different conditions together with alternatively spliced genes will provide the basis for the functional study of *B. oleraceae L. var capitata*.

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