



## Novel promoters that induce specific transgene expression during the green to ripening stages of tomato fruit development

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# **Novel promoters that induce specific transgene expression during the green to ripening stages of tomato fruit development**

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1 **Abstract**

2

3 Fruit-specific promoters have been used as genetic engineering tools for studies  
4 on molecular mechanism of fruit development and advance in fruit quality and  
5 additional value by increasing functional component. Especially fruit-ripening  
6 specific promoters have been well utilized and studied in tomato; however, few  
7 studies have reported the development of promoters that act at fruit developing  
8 stages such as immature green and mature green periods. In this study, we  
9 report novel promoters for gene expression during the green to ripening stages of  
10 tomato fruit development. Genes specifically expressed at tomato fruit were  
11 selected using microarray data. Subsequent to confirmation of the expression of  
12 the selected 12 genes, upstream DNA fragments of the genes LA22CD07,  
13 Les.3122.2.A1\_a\_at and LesAffx.6852.1.S1\_at which specifically expressed at  
14 fruit were isolated from tomato genomic DNA as promoter regions. Isolated  
15 promoter regions were fused with the *GUS* gene and the resultant constructs  
16 were introduced into tomato by agrobacterium-mediated transformation for  
17 evaluation of promoter activity in tomato fruit. The two promoters of LA22CD07,  
18 and LesAffx.6852.1.S1\_at showed strong activity in the fruit, weak activity in the  
19 flower and undetectable activity in other tissues. Unlike well-known fruit-  
20 ripening specific promoters, such as the E8 promoter, these promoters exhibited  
21 strong activity in green fruit in addition to red-ripening fruit, indicating that the  
22 promoters are suitable for transgene expression during green to ripening stages  
23 of tomato fruit development.

24

25 **Keywords:** *fruit-specific promoter, tomato, green stage, red stage, fruit*  
26 *development*

27

28 **Key Message**

1 Novel fruit specific promoters have been identified and are suitable for transgene  
2 expression during green to ripening stages of tomato fruit development.

3

4 **Abbreviations:** GUS, beta-D-glucuronidase gene

## 1 **Introduction**

2

3 The tomato (*Solanum lycopersicum*) is one of the major Solanaceae crops and one  
4 of the most widely eaten fruits in the world. Genetic engineering has been used  
5 in an effort to improve the quality of the tomato fruit (Butelli et al. 2009;  
6 Dharmapuri et al. 2002; Le et al. 2006; Lewinsohn 2001; Mollet et al. 2008;  
7 Rosati et al. 2000; Schijlen et al. 2006, 2007; Wang et al. 2008).

8         The tomato also serves as a vehicle for the production of useful proteins.  
9 For example, we reported the overexpression of the miraculin gene and the  
10 production of miraculin protein in the tomato fruit (Hirai et al. 2010; Hiwasa-  
11 Tanase et al. 2012; Sun et al. 2007; Yano et al. 2010). Chen et al. (2009) reported  
12 the production of thymosin alpha1, an immune booster that plays a role in the  
13 maturation, differentiation and function of T-cells, in the tomato fruit. Zhang et  
14 al. (2007) described the expression of human coagulation Factor IX in the tomato  
15 fruit.

16         The cauliflower mosaic virus 35S promoter (35S promoter) is a  
17 constitutive promoter that is widely used for the expression of foreign genes in  
18 higher plants. However, in some cases the 35S promoter is not suitable for gene  
19 expression because of the possibility that 35S promoter-driven constitutive gene  
20 expression could be damaging to plant growth and development.

21         To overcome the problem of the 35S promoter, tissue-specific promoters  
22 have been isolated. Fruit-specific promoters have been isolated as tools for fruit-  
23 specific gene expression. In the tomato, promoters from ethylene response genes,  
24 such as E8 and E4, have been well studied as fruit-specific promoters (Cordes et  
25 al. 1989; Coupe and Deikman 1997; Deikman et al. 1992, 1998; Deikman and  
26 Fischer 1988; Kneissl and Deikman 1996; Lincoln et al. 1987; Montgomery et al.  
27 1993a; Xu et al. 1996). Polygalacturonase (Montgomery et al. 1993b; Nicholass et  
28 al. 1995) and lipoxygenase promoters (Beaudoin and Rothstein 1997) have also

1 been reported as fruit specific in the tomato. These classical promoters have been  
2 reported to act during the late ripening stage of fruit development. On the other  
3 hand, information of promoters that act at fruit expanding stage (immature  
4 green), mature green stage and throughout the developmental stage are much  
5 less common than the fruit-ripening specific types, although recently Estornell et  
6 al. (2009) reported some promoters driving gene expression preferentially in the  
7 fruit with different activity ranges.

8 Many promoter variations expand the capability of intended use depending  
9 on the purpose. Therefore, in this study we attempted to isolate novel fruit-  
10 specific promoters with different activity from classical promoters. We selected 12  
11 genes which showed high expression in fruit tissues using microarray data  
12 obtained from tomato cultivar 'Micro-Tom', which has become a model plant of  
13 the Solanaceae family (Matsukura et al. 2008). Upon confirmation of the  
14 expression of the selected genes, cloning of the promoter regions, and the  
15 promoter analysis using *GUS* gene, we finally identified two promoters with  
16 fruit-specific activity. Unlike some classical fruit-specific promoters, these  
17 promoters were driven *GUS* gene expression throughout the fruit development in  
18 the green to ripening stages.

19

## 20 **Materials and methods**

21

### 22 **Identification of candidate genes from microarray data**

23 Tomato genes which show fruit-specific expression were selected by using  
24 gene expression data from following three sources; (i) a dataset available in  
25 MiBASE (old version, <http://www.kazusa.or.jp/jsol/microtom/>) using 'Micro-Tom'  
26 cDNA array produced by Japan Solanaceae genomics consortium (Yano et al.,  
27 2006), (ii) a dataset GSE19326 available in Gene Expression Omnibus  
28 (<http://www.ncbi.nlm.nih.gov/gds>) (Ozaki et al. 2010), and (iii) datasets 'Wild type

1 tomato fruit development (set 1 and set 2)' available in Tomato Functional  
2 Genomics Database (<http://ted.bti.cornell.edu/cgi-bin/TFGD/miame/home.cgi>)  
3 (Alba et al. 2005). Sequences of LA15CA04, LA22CD07, LC09AH08, LC04DC11,  
4 LA12AA05, LA14AD08 and FB14DB02 were obtained from MiBASE  
5 (<http://www.pgb.kazusa.or.jp/mibase/>). Consensus sequences of unigenes from  
6 which Les.331.1.S1\_at, Les.3122.2.A1\_a\_at and LesAffx.6852.1.S1\_at probes  
7 were designed were obtained from Affymetrix website  
8 (<http://www.affymetrix.com>). Consensus sequences of TC115787 and TC116003  
9 were obtained from Dana-Farber Cancer Institute Tomato Gene Index  
10 (<http://compbio.dfci.harvard.edu/cgi-bin/tgi/gimain.pl?gudb=tomato>).

11

## 12 **RNA isolation and Real-time PCR (RT-PCR) analysis**

13 Total RNA was isolated from the leaves, flowers, stems, roots, and green  
14 and red fruits of 3-month-old 'Micro-Tom' plants using TRIzol® (Invitrogen, USA)  
15 according to the manufacturer's instructions. One microgram of total RNA from  
16 each sample was treated with RQ1 RNase-Free DNase (Promega, USA) and was  
17 used for first-strand cDNA synthesis with a poly-T primer and SuperScript II  
18 Reverse Transcriptase (Invitrogen, USA) according to the manufacturer's  
19 instructions.

20 The first-strand cDNA was subsequently used as a template for the  
21 expression analysis of the selected genes. RT-PCR reactions were performed with  
22 25 to 30 cycles for the gene expression analysis using designed gene-specific  
23 primers (Table 1). After the PCR reaction, an equal volume of each amplified PCR  
24 product was subjected to electrophoresis on a 1% TAE agarose gel and was  
25 visualized using ethidium bromide.

26

## 27 **Quantitative real-time PCR (qRT-PCR)**

28 For the analysis of LA22CD07 and LesAffx.6852.1.S1\_at expression

1 during fruit development and ripening, total RNA was isolated from the ovary,  
 2 young (12, 15, and 18 days after flowering) and mature green fruits, orange fruits,  
 3 and red fruits using the RNeasy plant mini kit (Qiagen, Japan) according to the  
 4 manufacturer's instructions. The first-strand cDNA was synthesized from 0.75 µg  
 5 of total RNA using the Superscript VILO cDNA synthesis kit (Invitrogen, USA). A  
 6 ten-fold dilution of the first strand cDNA was used as a template for the qRT-  
 7 PCR using SYBR Premix Ex Taq II (Takara-Bio Inc., Otsu, Japan) in a Thermal  
 8 Cycler Dice Real-Time System TP800 (Takara-Bio Inc., Otsu, Japan) according to  
 9 the manufacturer's instructions. The thermal cycling parameters were set at  
 10 95°C for 10 min to denature, followed by 40 cycles at 95°C for 5 sec and 68°C for  
 11 30 sec. The relative quantification of the target gene expression was calculated  
 12 using the tomato *ubiquitin3* gene (X58253) as an internal control. The following  
 13 primer sequences were used: LA22CD07 forward, 5' -  
 14 GATCAAACCTATTGCTGCCAG-3', and reverse, 5'-  
 15 CTCTTCCTTGCTTCCACTCCAA-3'; LesAffx.6852.1.S1\_at forward, 5'-  
 16 CTGAAATGTCCCGTGATGATGC-3' and reverse, 5'-  
 17 CGCTTGCAGGTTCTCTGTTC-3'; *ES* forward, 5'-  
 18 TGGAAAGCCCTAGAGTTGAGGA-3' and reverse, 5'-  
 19 GAATCAACAAGTCCTTTAACAC-3'; and *ubiquitin3* forward, 5'-  
 20 CACCAAGCCAAAGAAGATCA-3' and reverse, 5'-TCAGCATTAGGG CACTCCTT-  
 21 3'.

22

### 23 Isolation of promoter regions

24 Genomic DNA was extracted from the tomato cultivar 'Moneymaker' using  
 25 the CTAB method (Murray and Thompson 1980). Each 5' flanking region of  
 26 LA22CD07 and LesAffx.6852.1.S1\_at was isolated from genomic DNA using the  
 27 GenomeWalker™ Universal Kit (Clontech, USA) as the putative promoter  
 28 regions. The promoter regions were obtained from a second PCR reaction using

1 the GenomeWalker™ Universal Kit, purified using the Wizard(R) SV Gel and  
2 PCR Clean-Up System (Promega, USA), and directly sequenced. The ATG start  
3 codons were predicted using ORF Finder  
4 (<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>), and the sequences were compared  
5 with homologs of other plant species, such as *Arabidopsis*. Approximately 2 kb  
6 of 5' upstream regions from the predicted ATG start site were re-amplified from  
7 the 'Moneymaker' genome using KOD Plus (TOYOBO, Japan). The amplified  
8 products were cloned into the pCR®-Blunt II-TOPO® Vector (Invitrogen, USA)  
9 and sequenced.

10

### 11 **Transient promoter assay**

12 The promoter region in the pCR®-Blunt II-TOPO® Vector was digested with  
13 restriction enzymes and ligated in front of the *GUS* gene in the pBI121 vector to  
14 replace the 35S promoter. The constructs containing the promoter region or  
15 pBI121 as a control was transformed into *Agrobacterium tumefaciens* strain  
16 GV3101 through electroporation and was used in a transient promoter assay. The  
17 assay was performed using green fruit of 'Micro-Tom' as previously described  
18 (Orzaez et al. 2006). The agrobacterium containing the construct was injected  
19 into green fruit and incubated 4 days at 25°C under long-day conditions (16 h  
20 light and 8 h dark). The total protein from the infected fruit was subjected to a  
21 quantitative GUS activity assay using 4-methylumbelliferyl-beta-D-glucuronide  
22 (4-MUG) as a substrate.

23

### 24 **Production of transgenic tomato**

25 The transformed *A. tumefaciens* was also used for the production of a  
26 transgenic tomato with 'Micro-Tom' cultivar. Transformants were produced  
27 according to Sun et al. (2006). The presence of the promoter-GUS fusions in the  
28 regenerated plants was confirmed by PCR using genomic DNA isolated from the

1 regenerated plants as templates.

2

### 3 **GUS assay**

4 For the quantitative analysis, GUS activity was assayed using the  
5 substrate 4-MUG according to Jefferson et al. (1987) with slight modifications  
6 (Moon and Callahan 2004). Tomato tissue was crushed using liquid nitrogen, and  
7 the protein was extracted in extraction buffer (Moon and Callahan 2004). The  
8 protein concentration was measured using the Bradford method (Bradford 1976).  
9 Approximately 100 µg of protein was used for the GUS assay. The reaction  
10 product 4-methylumbelliferone (4-MU) was measured with Safire (Tecan,  
11 Switzerland).

12 The histochemical GUS analysis was performed using 5-bromo-4-chloro-3-  
13 indolyl-β-D-glucuronide (X-Gluc) according to Jefferson et al. (1987) with slight  
14 modifications to the assay buffer. To reduce the background from GUS staining,  
15 100 mM phosphate (pH 8.0) was used instead of 50 mM phosphate (pH 7.0) in the  
16 assay buffer. For the analysis of the red fruit in Fig. 3B, 20% methanol (final  
17 volume) (Kosugi et al. 1990) was added to the assay buffer to further reduce the  
18 background staining. The tomato tissues were incubated in assay buffer at 37°C  
19 for 16 or 6 h. After staining, the sample was washed with 70% ethanol to  
20 terminate the reaction.

21

## 22 **Results and Discussion**

23

### 24 **Identification of promoter candidate genes from microarray data for expression in** 25 **green fruit**

26 To obtain candidates for novel fruit-specific promoters with unique  
27 activities compared to classical promoters, such as the E8 promoter, which  
28 mainly acts in the fruit late-ripening stage, we employed two strategies. The first

1 strategy was to identify highly expressed genes in green fruit, and the second  
2 was to uncover novel fruit-specific genes.

3 Firstly we analyzed microarray data using mRNA from ‘Micro-Tom’ green  
4 fruit to identify genes that were highly expressed in green fruit and selected  
5 seven genes (LA15CA04, LA22CD07, LC09AH08, LC04DC11, LA12AA05,  
6 LA14AD08 and FB14DB02). Moreover microarray database of several ‘Micro-  
7 Tom’ tissues were available from the Kazusa DNA Research Institute and  
8 Cornell University due to obtain promoter candidate genes for fruit-specific  
9 expression. Consequently, five genes (Les.331.1.S1\_at, Les.3122.2.A1\_a\_at,  
10 LesAffx.6852.1.S1\_at, TC115787 and TC116003) were selected. In total, 12  
11 promoter-candidate genes were identified (Table. 1).

12

### 13 **Expression analysis of the promoter-candidate genes by RT-PCR**

14 To examine whether the promoter-candidate genes uncovered from the  
15 microarray data are expressed in tomato fruit and the specificity, we performed  
16 RT-PCR analysis using the primer sets listed in Table 1.

17 We first examined the seven promoter-candidate genes predicted to have  
18 high expression levels in green fruit. As shown in Fig. 1, the expression was  
19 detected after 25 PCR reaction cycles and was clearly detectable at 27 and 30  
20 cycles using cDNA template derived from green fruits. The expression levels were  
21 different among the promoter-candidate genes. Based on the expression levels at  
22 27 and 30 cycles, we selected LA22CD07, LA12AA05 and LA14AD08, which were  
23 highly expressed in green fruit, for further studies.

24 Next, the organ-specific expression patterns were investigated for the five  
25 promoter candidate genes predicted fruit-specificity to understand which  
26 candidates displayed fruit-specific expression (Fig. 2). In this analysis, the  
27 expression of *ES* gene was also investigated to compare the expression of  
28 promoter-candidate genes with a well-known fruit-specific gene. As a result,

1 Les.3122.2.A1\_a\_at and LesAffx.6852.1.S1\_at exhibited fruit-specific expression.  
2 However, they also exhibited different expression patterns. Les.3122.2.A1\_a\_at  
3 showed specific and high expression in the both green and red fruit stages,  
4 whereas LesAffx.6852.1.S1\_at was highly expressed in the green fruit but was  
5 only slightly expressed in the red fruit. Les.331.1.S1\_at was also highly  
6 expressed in the green and red fruits; however, a low level of expression was  
7 detected in the flower. TC115787 was expressed in the flower, stem and root in  
8 addition to the green and red fruit. TC116003 was expressed throughout the  
9 examined organs except the red fruit. The *ES* gene was highly expressed in the  
10 red fruit but was almost undetectable in the green fruit. This result supports  
11 previous studies, which reported that the *ES* gene was expressed in a ripening-  
12 specific manner (Deikman and Fischer 1998; Kneissl and Deikman 1996; Lincoln  
13 et al. 1987).

14 We uncovered two promoter-candidate genes of Les.3122.2.A1\_a\_at and  
15 LesAffx.6852.1.S1\_at with fruit-specific expression and one gene of  
16 Les.331.1.S1\_at with high expression in the fruit and low expression in the flower.  
17 Notably, these three candidates were highly expressed in the green fruit, in  
18 which *E8* gene expression was almost undetectable. Moreover, the three  
19 candidates were also expressed in the red fruit. These results suggest that the  
20 promoters of the three candidate genes were active in fruit and have different  
21 activities than the *E8* promoter.

22 From these results, six genes, LA22CD07, LA12AA05, LA14AD08,  
23 Les.331.1.S1\_at, Les.3122.2.A1\_a\_at and LesAffx.6852.1.S1\_at, were selected for  
24 subsequent analysis.

25

## 26 **BLASTN analysis of the candidates**

27 To obtain functional information for the promoter-candidate genes, a  
28 BLASTN analysis was performed. The results were summarized in Table 2,

1 which listed the top hits of functionally annotated genes resulting from BLASTN  
2 analysis. The BLASTN analysis showed that LA14AD08 returned a hit for a clp-  
3 like energy-dependent protease from the tomato and stink bell (*Fritillaria*  
4 *agrestis*), indicating that LA14AD08 represents a family of Clp proteases.  
5 Although LA22CD07 and LA12AA05 hit to the tomato full-length cDNA  
6 sequences (Aoki et al. 2010), they did not hit to functionally annotated tomato  
7 gene. However, LA22CD07 and LA12AA05 returned hits for the erythroblast  
8 macrophage protein emp from *Ricinus communis* (XM\_002525023) with an e-  
9 value of 5E-39 and the sufD protein from the *Ricinus communis* (XM\_002534741)  
10 with an e-value of 2E-69, respectively. The result suggest that the two candidates  
11 are homologs of the erythroblast macrophage proteins emp or sufD.

12 Les.331.1.S1\_at returned hits for the tomato LOX gene U13681 (Kausch  
13 and Handa 1995) and tomloxB (U09025) with e-values of 0 (Ferrie et al. 1994).  
14 Ferrie et al. (1994) reported the fruit-specific expression of the LOX gene.  
15 Beaudoin and Rothstein (1997) reported that the LOX gene promoter activity was  
16 active in tobacco and tomato fruits.

17 Les.3122.2.A1\_a\_at returned a hit for tomato gene S66607 (Pear et al.  
18 1993), which has been described as a pectin methylesterase-like sequence,  
19 indicating that Les.3122.2.A1\_a\_at is a member of the pectin methylesterases.  
20 The expression pattern and promoter analysis of S66607 have not been analyzed;  
21 however, it has been reported that some members of the pectin methylesterases  
22 exhibited fruit-specific expression (Gaffe et al. 1997; Hall et a l. 1994).

23 LesAffx.6852.1.S1\_at returned hits for tomato cDNAs with e-values of 0  
24 whose functions have not been reported. LesAffx.6852.1.S1\_at also returned a hit  
25 for a cysteine protease of *Gossypium hirsutum* (AY171099) with 69% identity,  
26 suggesting that the LesAffx.6852.1.S1\_at is a member of the cysteine proteases.

27

28 **Isolation and characterization of selected gene promoters**

1           Because the Les.331.1.S1\_at promoter had been analyzed previously  
2 (Beaudoin and Rothstein 1997), we decided to clone the promoter regions that  
3 have not been analyzed: LA22CD07, LA12AA05, LA14AD08, Les.3122.2.A1\_a\_at  
4 and LesAffx.6852.1.S1\_at.

5           To clone the promoter regions, we performed genome walking based on  
6 the sequence information of the candidates. In consideration of prospective  
7 practical use, the isolation of promoter regions were used genomic DNA from  
8 ‘Moneymaker’ which is cultivated variety. The PCR fragments obtained from  
9 genome walking were directly sequenced. The ATG start codons were predicted  
10 using ORF Finder (<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>), and the  
11 sequences were compared with homologs of other plant species. Subsequently, the  
12 putative promoter regions, which were approximately 2 kb upstream from the  
13 predicted ATG start codon, were re-amplified and sequenced.

14           In order to analyze the activities of the isolated promoters, each promoter  
15 was cloned to replace the 35S promoter in vector pBI121. We first performed  
16 transient assays using ‘Micro-Tom’ green fruit. Significant GUS activity was  
17 obtained from the LA22CD07, Les.3122.2.A1\_a\_at and LesAffx.6852.1.S1\_at  
18 promoters (data not shown). The GUS activities of the LA12AA05 and LA14AD08  
19 promoters were almost the same as that of uninfected green fruit, suggesting  
20 that the two promoter fragments do not function in green fruit.

21           The three promoters from LA22CD07, Les.3122.2.A1\_a\_at and  
22 LesAffx.6852.1.S1\_at that exhibited GUS activity in the transient assay were  
23 further analyzed using stable transgenic tomatoes. We conducted a GUS  
24 histochemical assay of leaves, roots, stems, flowers, green fruits and red fruits in  
25 regenerated T<sub>0</sub> plants. At least three independent T<sub>0</sub> plants per construct were  
26 assayed. The GUS staining pattern was almost identical among the tested plants  
27 containing the same construct, although the staining intensity varied (data not  
28 shown). Fig. 3 shows the results of a typical GUS staining of the various tissues

1 of transgenic plants containing promoter-GUS fusion constructs. Unlike the  
2 transgenic plants containing the 35S promoter, tissue-specific GUS staining  
3 patterns were observed among the transgenic plants containing the LA22CD07  
4 or LesAffx.6852.1.S1\_at foreign promoter regions. Fig. 3a shows the results from  
5 a 16h GUS staining experiment. The transgenic plants containing the LA22CD07  
6 promoter exhibited strong GUS staining in the green and red fruits, weak  
7 staining in the flowers and undetectable staining in the leaves and roots. The  
8 transgenic plants containing the LesAffx.6852.1.S1\_at promoter also displayed  
9 strong staining in the green and red fruits, but the flower staining was stronger  
10 than that of LA22CD07. No staining was detected in the tissues from the  
11 transgenic plants containing the Les.3122.2.A1\_a\_at promoter (data not shown).  
12 In the case *GUS* gene driven by 35S promoter, the GUS staining was detected  
13 everywhere in tomato plant and the staining levels were relatively high. However  
14 in the green fruit the GUS staining levels were almost same between LA22CD07,  
15 LesAffx.6852.1.S1\_at and 35S promoters. In the red fruit the staining levels were  
16 also high in these promoters but non-specific staining was observed in the non-  
17 transgenic plants. Therefore the red fruits were further treated with assay buffer  
18 containing methanol for 6 h. As shown in Fig. 3b, GUS staining was almost no  
19 detected in the wild-type plants and was observed in red fruits of the transgenic  
20 plants containing the LA22CD07 and LesAffx.6852.1.S1\_at promoters. Moreover  
21 the staining levels were relatively high especially in LesAffx.6852.1.S1\_at  
22 promoter compared with 35S promoter. These results indicated that these  
23 promoters were active in both green and red fruits.

24 Quantitative real-time PCR analysis of LA22CD07 and  
25 LesAffx.6852.1.S1\_at were performed to investigate the detail of the promoter  
26 activities during fruit development and to compare the activity of *E8* promoter as  
27 known fruit-ripening specific (Fig. 4). The expression of *E8* gene was slightly  
28 detected in mature green stage and rapidly increased from orange stage. On the

1 other hand, the expression level of LA22CD07 was gradually increased from 12  
2 days after flowering and reached the highest in the red stage. In the  
3 LesAffx.6852.1.S1\_at the expression was already detected in the ovary and then  
4 gradually increased as described at LA22CD07. The result suggested that the  
5 novel two promoters had different activation pattern from E8 promoter and were  
6 active from small green fruit or ovary stages. Although we have not examined the  
7 GUS staining between flowers and green fruits, it might be possible that the two  
8 promoters are active at early stages of fruit development (flower to green fruit)  
9 because the GUS staining was also observed in the both flowers.

10

## 11 **Conclusions**

12 In this study, we isolated novel two fruit-specific promoters from the tomato.  
13 These promoters exhibited activities that were different from classical fruit  
14 ripening-specific promoters, such as the E8 promoter. The activities are detected  
15 throughout during fruit development from ovary to red-ripe fruit. Therefore, the  
16 identified two promoters might outperform some fruit-specific promoters that act  
17 only fruit-ripening stage depending on the intended purpose. The two promoters  
18 will supply us tools to express genes of interest in fruit regardless of the  
19 developmental stage. In this study, we examined only tomato promoters. However,  
20 it might be possible to use these promoters in the fruits of other plants because  
21 BLAST analysis revealed homologs of LA22CD07 and LesAffx.6852.1.S1\_at from  
22 many plant species.

23

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25

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4

5

## 6 **References**

7

8 Alba R, Payton P, Fei Z, McQuinn R, Debbie P, Martin GB, Tanksley SD,  
9 Giovannoni JJ (2005) Transcriptome and selected metabolite analyses reveal  
10 multiple points of ethylene control during tomato fruit development. *Plant Cell*  
11 17: 2954-2965. doi:10.1105/tpc.105.036053

12

13 Aoki K, Yano K, Suzuki A, Kawamura S, Sakurai N, Suda K, Kurabayashi A,  
14 Suzuki T, Tsugane T, Watanabe M, Ooga K, Torii M, Narita T, Shin-I T, Kohara Y,  
15 Yamamoto N, Takahashi H, Watanabe Y, Egusa M, Kodama M, Ichinose Y,  
16 Kikuchi M, Fukushima S, Okabe A, Arie T, Sato Y, Yazawa K, Satoh S, Omura T,  
17 Ezura H, Shibata D (2010) Large-scale analysis of full-length cDNAs from the  
18 tomato (*Solanum lycopersicum*) cultivar Micro-Tom, a reference system for the  
19 Solanaceae genomics. *BMC Genomics* 11:210. doi:10.1186/1471-2164-11-210

20

21 Beaudoin N, Rothstein SJ (1997) Developmental regulation of two tomato  
22 lipoxygenase promoters in transgenic tobacco and tomato. *Plant Mol Biol* 33:835-  
23 46. doi: 10.1023/A:1005773722657

24

25 Bradford MM (1976) A rapid and sensitive for the quantitation of microgram  
26 quantities of protein utilizing the principle of protein-dye binding. *Analytical*  
27 *Biochem* 72:248-254. doi:10.1016/0003-2697(76)90527-3

28

1 Butelli E, Titta L, Giorgio M, Mock HP, Matros A, Peterek S, Schijlen EG, Hall  
2 RD, Bovy AG, Luo J, Martin C (2008) Enrichment of tomato fruit with health-  
3 promoting anthocyanins by expression of select transcription factors. Nat  
4 Biotechnol 26:1301-1308. doi:10.1038/nbt.1506  
5  
6 Chen Y, Wang A, Zhao L, Shen G, Cui L, Tang K (2009) Expression of thymosin  
7 alpha1 concatemer in transgenic tomato (*Solanum lycopersicum*) fruits.  
8 Biotechnol Appl Biochem 52:303-312. doi:10.1042/BA20080054  
9  
10 Cordes S, Deikman J, Margossian LJ, Fischer RL (1989) Interaction of a  
11 developmentally regulated DNA-binding factor with sites flanking two different  
12 fruit-ripening genes from tomato. Plant Cell 1:1025-1034. doi:  
13 10.1105/tpc.1.10.1025  
14  
15 Coupe SA, Deikman J (1997) Characterization of a DNA-binding protein that  
16 interacts with 5' flanking regions of two fruit-ripening genes. Plant J 11:1207-  
17 1218. doi: 10.1046/j.1365-313X.1997.11061207.x  
18  
19 Deikman J, Fischer RL (1988) Interaction of a DNA binding factor with the 5'-  
20 flanking region of an ethylene-responsive fruit ripening gene from tomato. EMBO  
21 J 7:3315-3320.  
22  
23 Deikman J, Kline R, Fischer RL (1992) Organization of ripening and ethylene  
24 regulatory regions in a fruit-specific promoter from tomato (*Lycopersicon*  
25 *esculentum*). Plant Physiol 100:2013-2017. doi:10.1104/pp.100.4.2013  
26  
27 Deikman J, Xu R, Kneissl ML, Ciardi JA, Kim KN, Pelah D (1998) Separation of  
28 cis elements responsive to ethylene, fruit development, and ripening in the 5'-

1 flanking region of the ripening-related E8 gene. *Plant Mol Biol* 37:1001-1011. doi:  
2 10.1023/A:1006091928367  
3  
4 Dharmapuri S, Rosati C, Pallara P, Aquilani R, Bouvier F, Camara B, Giuliano G  
5 (2002) Metabolic engineering of xanthophyll content in tomato fruits. *FEBS Lett*  
6 22:30-34. doi:10.1016/S0014-5793(02)02699-6  
7  
8 Estornell LH, Orzáez D, López-Peña L, Pineda B, Antón MT, Moreno V, Granell A  
9 (2009) A multisite gateway-based toolkit for targeted gene expression and hairpin  
10 RNA silencing in tomato fruits. *Plant Biotechnol J* 7:298-309. doi: 10.1111/j.1467-  
11 7652.2009.00402.x  
12  
13 Ferrie BJ, Beaudoin N, Burkhart W, Bowsher CG, Rothstein SJ (1994) The  
14 cloning of two tomato lipoxygenase genes and their differential expression during  
15 fruit ripening. *Plant Physiol* 106:109-118. doi:10.1104/pp.106.1.109  
16  
17 Gaffe J, Tiznado ME, Handa AK (1997) Characterization and functional  
18 expression of a ubiquitously expressed tomato pectin methylesterase. *Plant*  
19 *Physiol* 114:1547-1556. doi: <http://dx.doi.org/10.1104/pp.114.4.1547>  
20  
21 Hall LN, Bird CR, Picton S, Tucker GA, Seymour GB, Grierson D (1994)  
22 Molecular characterisation of cDNA clones representing pectinesterase isozymes  
23 from tomato. *Plant Mol Biol* 25:313-318. doi: 10.1007/BF00039542  
24  
25 Hirai T, Fukukawa G, Kakuta H, Fukuda N, Ezura H (2010) Production of  
26 recombinant miraculin using transgenic tomato in a closed-cultivation system. *J*  
27 *Agric Food Chem* 58: 6096-6101. doi:10.1021/jf100414v  
28

1 Hiwasa-Tanase K, Hirai T, Kato K, Duhita N, Ezura H (2012) From miracle fruit  
2 to transgenic tomato: mass production of the taste-modifying protein miraculin in  
3 transgenic plants. *Plant Cell Rep* 31:513-525. doi:10.1007/s00299-011-1197-5  
4  
5 Jefferson RA, Kavanagh TA, Bevan MW (1987) GUS fusions: beta-glucuronidase  
6 as a sensitive and versatile gene fusion marker in higher plants. *EMBO J* 6:3901-  
7 3907.  
8  
9 Kausch KD, Handa AK (1995) Molecular cloning and nucleotide sequence of a  
10 lipoxygenase cDNA from ripening tomato fruit. *Plant Physiol* 107:669-670.  
11 doi:10.1104/pp.107.2.669  
12  
13 Kneissl ML, Deikman J (1996) The tomato E8 gene influences ethylene  
14 biosynthesis in fruit but not in flowers. *Plant Physiol* 112:537-547. doi: [http://dx.](http://dx.doi.org/10.1104/pp.112.2.537)  
15 [doi.org/10.1104/pp.112.2.537](http://dx.doi.org/10.1104/pp.112.2.537)  
16  
17 Kosugi S, Ohashi Y, Nakajima K, Arai Y (1990) An improved assay for  $\beta$ -  
18 glucuronidase (GUS) in transformed cells: methanol almost suppresses a putative  
19 endogenous GUS activity. *Plant Sci* 70:133-140. doi:10.1016/0168-9452(90)90042-  
20 M  
21  
22 Le LQ, Lorenz Y, Scheurer S, Fötisch K, Enrique E, Bartra J, Biemelt S, Vieths S,  
23 Sonnewald U (2006) Design of tomato fruits with reduced allergenicity by  
24 dsRNAi-mediated inhibition of ns-LTP (Lyc e 3) expression. *Plant Biotechnol J*  
25 4:231-242. doi: 10.1111/j.1467-7652.2005.00175.x  
26  
27 Lewinsohn E, Schalechet F, Wilkinson J, Matsui K, Tadmor Y, Nam KH, Amar O,  
28 Lastochkin E, Larkov O, Ravid U, Hiatt W, Gepstein S, Pichersky E (2001)

1 Enhanced levels of the aroma and flavor compound S-linalool by metabolic  
2 engineering of the terpenoid pathway in tomato fruits. *Plant Physiol* 127:1256-  
3 1265. doi: <http://dx.doi.org/10.1104/pp.010293>  
4  
5 Lincoln JE, Cordes S, Read E, Fischer RL (1987) Regulation of gene expression  
6 by ethylene during *Lycopersicon esculentum* (tomato) fruit development. *Proc*  
7 *Natl Acad Sci USA* 84:2793-2797.  
8  
9 Matsukura C, Aoki K, Fukuda N, Mizoguchi T, Asamizu E, Saito T, Shibata D,  
10 Ezura H (2008) Comprehensive resources for tomato functional genomics based  
11 on the miniature model tomato Micro-Tom. *Curr Genom* 9:436-443. doi:  
12 [10.2174/138920208786241225](https://doi.org/10.2174/138920208786241225)  
13  
14 Mollet B, Niederberger P, Pétiard V (2008) Novel tomato flavours introduced by  
15 plastidial terpenoid pathway engineering. *Trends Biotechnol* 26:4-6. doi:  
16 [10.1016/j.tibtech.2007.10.004](https://doi.org/10.1016/j.tibtech.2007.10.004)  
17  
18 Montgomery J, Goldman S, Deikman J, Margossian L, Fischer RL (1993a)  
19 Identification of an ethylene-responsive region in the promoter of a fruit ripening  
20 gene. *Proc Natl Acad Sci USA* 90:5939-5943.  
21  
22 Montgomery J, Pollard V, Deikman J, Fischer RL (1993b) Positive and negative  
23 regulatory regions control the spatial distribution of polygalacturonase  
24 transcription in tomato fruit pericarp. *Plant Cell* 5:1049-1062.  
25 doi:[10.1105/tpc.5.9.1049](https://doi.org/10.1105/tpc.5.9.1049)  
26  
27 Moon H, Callahan AM (2004) Developmental regulation of peach ACC oxidase  
28 promoter-GUS fusions in transgenic tomato fruits. *J Exp Bot* 55:1519-1528. doi:

1 10.1093/jxb/erh162  
2  
3 Murray MG, Thompson WF (1980) Rapid isolation of high molecular weight plant  
4 DNA. *Nucleic Acids Res* 10:4321-4325. doi:10.1093/nar/8.19.4321  
5  
6 Nicholass FJ, Smith CJ, Schuch W, Bird CR, Grierson D (1995) High levels of  
7 ripening-specific reporter gene expression directed by tomato fruit  
8 polygalacturonase gene-flanking regions. *Plant Mol Biol* 28:423-435. doi:  
9 10.1007/BF00020391  
10  
11 Ozaki S, Ogata Y, Suda K, Kurabayashi A, Suzuki T, Yamamoto N, Iijima Y,  
12 Tsugane T, Fujii T, Konishi C, Inai S, Bunsupa S, Yamazaki M, Shibata D, Aoki K  
13 (2010) Coexpression analysis of tomato genes and experimental verification of  
14 coordinated expression of genes found in a functionally enriched coexpression  
15 module. *DNA Res.* 17: 105-116. doi:10.1093/dnares/dsq002  
16  
17 Orzaez D, Mirabel S, Wieland WH, Granell A (2006) Agroinjection of tomato  
18 fruits. A tool for rapid functional analysis of transgenes directly in fruit. *Plant*  
19 *Physiol* 140:3-11. doi: <http://dx.doi.org/10.1104/pp.105.068221>  
20  
21 Pear JR, Sanders RA, Summerfelt KR, Martineau B, Hiatt WR (1993)  
22 Simultaneous inhibition of two tomato fruit cell wall hydrolases,  
23 pectinmethylesterase and polygalacturonase, with antisense gene constructs.  
24 *Antisense Res Dev* 3:181-190.  
25  
26 Rosati C, Aquilani R, Dharmapuri S, Pallara P, Marusic C, Tavazza R, Bouvier F,  
27 Camara B, Giuliano G (2000) Metabolic engineering of beta-carotene and  
28 lycopene content in tomato fruit. *Plant J* 24:413-419. doi: <http://dx.doi.org/10.>

1 1104/pp.105.068221  
2  
3 Schijlen EG, de Vos CH, Martens S, Jonker HH, Rosin FM, Molthoff JW, Tikunov  
4 YM, Angenent GC, van Tunen AJ, Bovy AG (2007) RNA interference silencing of  
5 chalcone synthase, the first step in the flavonoid biosynthesis pathway, leads to  
6 parthenocarpic tomato fruits. *Plant Physiol* 144:1520-1530. doi: [http://dx.doi.org/](http://dx.doi.org/10.1104/pp.107.100305)  
7 10.1104/pp.107.100305  
8  
9 Schijlen E, Ric de Vos CH, Jonker H, van den Broeck H, Molthoff J, van Tunen A,  
10 Martens S, Bovy A (2006) Pathway engineering for healthy phytochemicals  
11 leading to the production of novel flavonoids in tomato fruit. *Plant Biotechnol J*  
12 4:433-444. doi: 10.1111/j.1467-7652.2006.00192.x  
13  
14 Sun HJ, Uchii S, Watanabe S, Ezura H (2006) A highly efficient transformation  
15 protocol for Micro-Tom, a model cultivar for tomato functional genomics. *Plant*  
16 *Cell Physiol* 47:426-431. doi: 10.1093/pcp/pci251  
17  
18 Sun HJ, Kataoka H, Yano M, Ezura H (2007) Genetically stable expression of  
19 functional miraculin, a new type of alternative sweetener, in transgenic tomato  
20 plants. *Plant Biotechnol J* 5:768-777. doi: 10.1111/j.1467-7652.2007.00283.x  
21  
22 Wang S, Liu J, Feng Y, Niu X, Giovannoni J, Liu Y (2008) Altered plastid levels  
23 and potential for improved fruit nutrient content by downregulation of the  
24 tomato DDB1-interacting protein CUL4. *Plant J* 55:89-103. doi: 10.1111/j.1365-  
25 313X.2008.03489.x  
26  
27 Xu R, Goldman S, Coupe S, Deikman J (1996) Ethylene control of E4  
28 transcription during tomato fruit ripening involves two cooperative *cis* elements.

1 Plant Mol Biol 31:1117-1127. doi: 10.1007/BF00040829  
2  
3 Yano K, Watanabe M, Yamamoto N, Tsugane T, Aoki K, Sakurai N, Shibata D  
4 (2006) MiBASE: A database of a miniature tomato cultivar Micro-Tom. Plant  
5 Biotechnol. 23: 195-198. doi:10.5511/plantbiotechnology.23.195  
6  
7 Yano M, Hirai T, Kato K, Hiwasa-Tanase K, Fukuda N, Ezura H (2010) Tomato is  
8 a suitable material for producing recombinant miraculin protein in genetically  
9 stable manner. Plant Sci 178: 469-473. doi: 10.1016/j.plantsci.2010.02.016  
10  
11 Zhang H, Zhao L, Chen Y, Cui L, Ren W, Tang K (2007) Expression of human  
12 coagulation Factor IX in transgenic tomato (*Lycopersicon esculentum*).  
13 Biotechnol Appl Biochem 48:101-107. doi: 10.1042/BA20060224  
14

Table 1. Selected genes found from microarray data and summary of their expression

Category	Database	ID	Result of RT-PCR	Forward primer for RT-PCR	Revers primer for RT-PCR
	MiBASE <sup>a</sup>	LA15CA04	Low expression	5'-TCACTCACCAAGCCCTTTCTCTC-3'	5'-TCCTGAGAAGCAGCCTTAGGAAC-3'
	MiBASE <sup>a</sup>	LA22CD07	High expression	5'-CGATCCGCGCTAATCATCGT-3'	5'-AGCCGTGCTCTGCATCTTTG-3'
	MiBASE <sup>a</sup>	LC09AH08	Low expression	5'-TGGTGGTGAGGCTGTTGAGC-3'	5'-CCATGAGTCGGAACCTGTGC-3'
High expression in green fruit	MiBASE <sup>a</sup>	LC04DC11	Low expression	5'-TGGCGTTTTCTTCATCCTCCA-3'	5'-CAGCTGCCCTTATCCTGAACTGA-3'
	MiBASE <sup>a</sup>	LA12AA05	High expression	5'-CGGGGTGTTGATGCTGAAAC-3'	5'-GAGGGGCTTCCATTCAATCAGA-3'
	MiBASE <sup>a</sup>	LA14AD08	High expression	5'-AACCTCGCGGAGCATCAA-3'	5'-TTAATGGATCCCAACTTCTTG-3'
	MiBASE <sup>a</sup>	FB14DB02	Low expression	5'-GCAATAGCTGGTCGGCTAGAACA-3'	5'-ATCGATTGCTGCGGCCTTA-3'
	GEO <sup>b</sup>	Les.331.1.S1_at	Fruit specific expression	5'-ATGTCTTTGGGTGGAATTGGATGCC-3'	5'-CATCTCCTCGCAAAGCTACCAGTTC-3'
	GEO <sup>b</sup>	Les.3122.2.A1_a_at	Fruit specific expression	5'-ATGTATGCTACGACCATTACTGGTAGCC-3'	5'-CAACCCGCTGGATTAATGAGACCAC-3'
Fruit specific expression	GEO <sup>b</sup>	LesAffx.6852.1.S1_at	High expression in fruit, Low expression in flower	5'-GAAAGACCAACTGAGCCTTTTCAGAAG-3'	5'-ATGCCGCCGTTGTTTATCACCCATTC-3'
	TFGD <sup>c</sup>	TC115787	No fruit specific expression	5'-CCACTTGTTGGAATTGGATGGATGTTG-3'	5'-GATCACTGGAGGAGCTGTATAGCC-3'
	TFGD <sup>c</sup>	TC116003	No fruit specific expression	5'-ATGCCGCCGTTGTTATCACCCATTC-3'	5'-GAAAGACCAACTGAGCCTTTTCAGAAG-3'

2 <sup>a</sup> URL: <http://www.pgb.kazusa.or.jp/mibase/>.

3 <sup>b</sup> Gene Expression Omnibus, URL: <http://www.ncbi.nlm.nih.gov/gds>, dataset GSE19326.

4 <sup>c</sup> Tomato Functional Genomics Database, URL: <http://ted.bti.cornell.edu/cgi-bin/TFGD/miame/home.cgi>.

5

6

Table 2. Summary of BLAST analysis.

ID of genes	Category	Organism	Accession	Definition	E-value
LA22CD08	Top hit	<i>Solanum lycopersicum</i>	L38581	<i>Lycopersicon esculentum</i> clp-like energy-dependent protease mRNA, complete cds.	0
	Top hit of functionally annotated genes	<i>Fritillaria agrestis</i>	AF037459	<i>Fritillaria agrestis</i> clp-like energy-dependent protease (clpP) mRNA, complete cds.	1.00E-35
LA12AA07	Top hit	<i>Solanum lycopersicum</i>	AK322312	<i>Solanum lycopersicum</i> cDNA, clone: LEFL1036AH12, HTC in leaf.	0
	Top hit of functionally annotated genes	<i>Ricinus communis</i>	XM_002525023	<i>Ricinus communis</i> erythroblast macrophage protein emp, putative, mRNA.	5.00E-39
LA14AD05	Top hit	<i>Solanum lycopersicum</i>	AK322226	<i>Solanum lycopersicum</i> cDNA, clone: LEFL1035AG05, HTC in leaf.	0
	Top hit of functionally annotated genes	<i>Ricinus communis</i>	XM_002534741	<i>Ricinus communis</i> Protein sufD, putative, mRNA.	2.00E-69
Les.331.1.S1_at	Top hit	<i>Solanum lycopersicum</i>	AK326139	<i>Lycopersicon esculentum</i> lipoxygenase (LOX) mRNA, complete cds.	0
	Top hit of functionally annotated genes	<i>Solanum lycopersicum</i>	U13681	<i>Lycopersicon esculentum</i> lipoxygenase (LOX) mRNA, complete cds.	0
Les.3122.2.A1_a_at	Top hit	<i>Solanum lycopersicum</i>	S66607	<i>Lycopersicon esculentum</i> pectinmethylesterase-like sequence.	0
	Top hit of functionally annotated genes	<i>Solanum lycopersicum</i>	S66607	<i>Lycopersicon esculentum</i> pectinmethylesterase-like sequence.	0
LesAffx.6852.1.S1_at	Top hit	<i>Solanum lycopersicum</i>	AK326008	<i>Solanum lycopersicum</i> cDNA, clone: LEFL2001CF07, HTC in fruit.	0
	Top hit of functionally annotated genes	<i>Gossypium hirsutum</i>	AY171099	<i>Gossypium hirsutum</i> cysteine protease mRNA, complete cds.	2.00E-119

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3

4

1 **Figure captions**

2

3 **Fig. 1**

4 Real-time PCR analysis of the promoter-candidate genes for high expression  
5 levels in green fruits. The expression levels of the genes in green fruits were  
6 analyzed at 25, 27 and 30 cycles of RT-PCR.

7

8 **Fig. 2**

9 Real-time PCR analysis of the promoter-candidate genes for fruit-specific  
10 expression. The tissue-specific expression levels of the candidate, *E8* and *actin*  
11 genes were analyzed using RT-PCR with first-strand cDNAs from the leaves,  
12 flowers, stems, roots, and green and red fruits. L, leaves; F, flowers; S, stems; R,  
13 roots; G, green fruits; R, red fruits.

14

15 **Fig. 3**

16 Histochemical GUS assay of the transgenic plants. The leaves, flowers, roots, and  
17 green and red fruits of T<sub>0</sub> plants were used for the GUS assay. The blue staining  
18 represents GUS activity. (a) Results of the 16h GUS staining of various tissues  
19 (b) Results of the 6h GUS staining of red fruits with buffer containing methanol.  
20 L, leaves; R, roots; F, flowers; G, green fruits; R, red fruits.

21

22 **Fig. 4**

23 Quantitative real-time PCR analysis of LA22CD07 and LesAffx.6852.1.S1\_at. **a**  
24 The developmental stages of the fruits used for these experiments. Bar = 1 mm.  
25 Relative expression levels of LA22CD07 (**b**) and LesAffx.6852.1.S1\_at (**c**) during  
26 fruit development and ripening. The expression level of the E8 gene was analyzed  
27 as a control (**d**). The fruits were harvested at 12, 15, and 18 days after flowering  
28 and at the fruit developmental stages as follows: ovary (OV), mature green stage

- 1 (MG), orange stage (OR), and red ripening stage (RE). The mean values of three
- 2 independent experiments are shown. The error bars represent the standard error.

Fig. 1

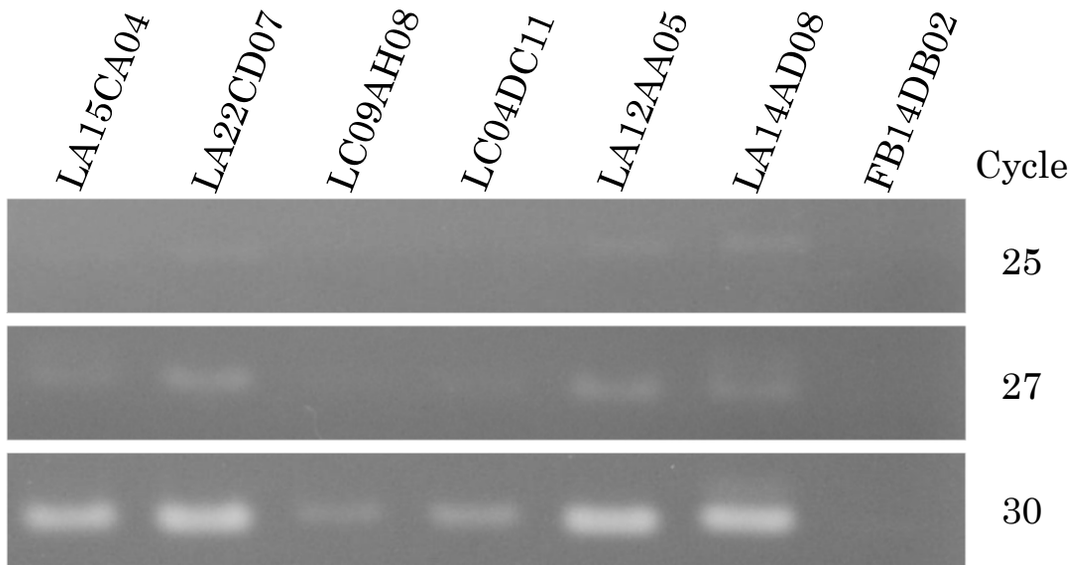


Fig. 2

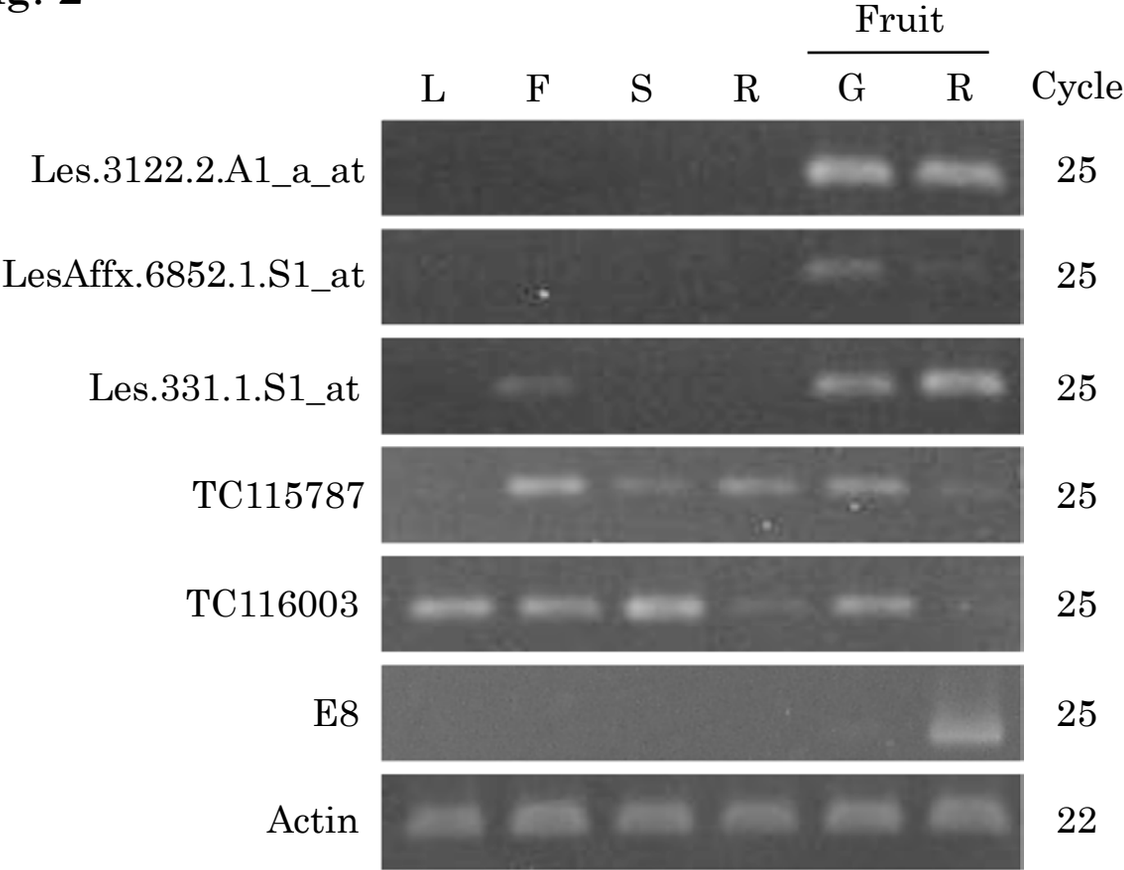


Fig. 3

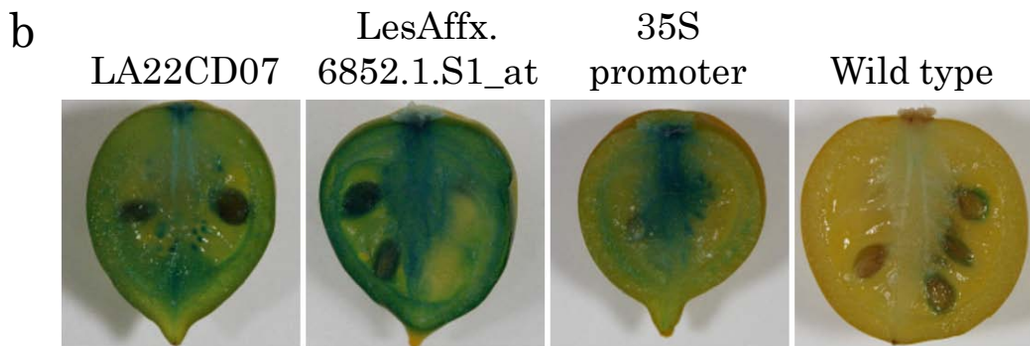
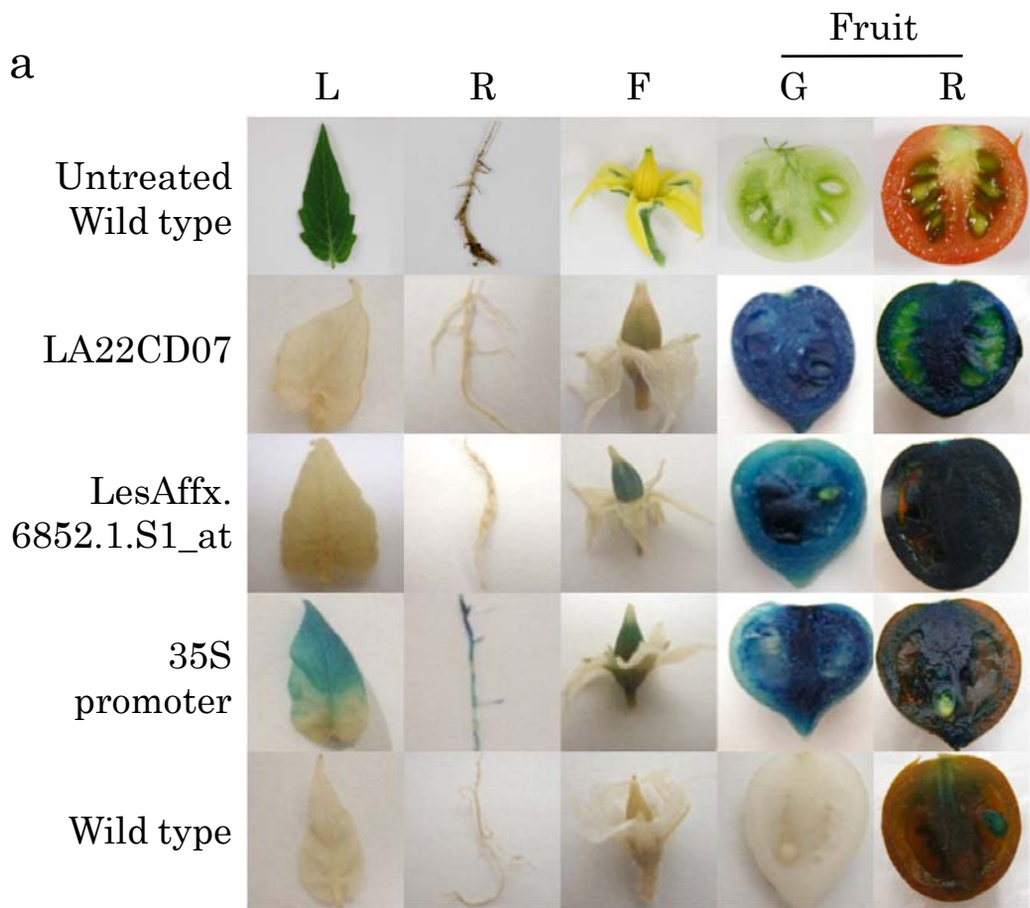


Fig. 4

