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1	SEXUAL STERILITY is essential for both male and female gametogenesis in tomato
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3	A gene essential for gametogenesis in tomato
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10	Abbreviations:
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1	Abstract: Gametogenesis is a key step in the production of ovules or pollen in higher plants.
2	The molecular aspects of gametogenesis are well characterized in the model plant Arabidopsis;
3	however, little information is known in tomato, which is a model plant for fleshy fruit
4	development. In this study, we characterized a tomato (Solanum lycopersicum L.) x-ray mutant,
5	sexual sterility (Slses), that exhibited both male and female sterility. Morphological analysis
6	revealed that the Slses mutant forms incomplete ovules and wilted anthers devoid of pollen
7	grains at the anthesis stage. Genetic and next-generation sequencing analyses revealed that the
8	Slses mutant carried a 13-bp deletion within the first exon of a homologue of
9	SPOROCYTELESS/NOZZLE (SPL/NZZ), which plays an important role in gametogenesis in
10	Arabidopsis. Complementation analysis in which the complete SISES genomic region was
11	introduced into the Slses mutant fully restored normal phenotypes, demonstrating that
12	Solyc07g063670 is responsible for the Slses mutation. SISES probably act as transcriptional
13	repressor because of EAR motif at the C-terminal region. Gene expression levels of WUSCHEL
14	(SIWUS) and INNER NO OUTER (SIINO), both of which are required for ovule development,
15	were dramatically reduced in the early stages of pistil development in the Slses mutant,
16	suggesting a positive regulatory role of SISES in the transcription of gametogenesis genes and
17	differences in the regulation of INO (SIINO) and integument development by SPL/NZZ (SLSES)
18	between Arabidopsis and tomato. Taken together, our results indicate that SISES is a novel
19	tomato gametogenesis gene essential for both male and female gametogenesis.

2 Keywords: Anther, Gametogenesis, Mutant, Ovule, Sexual sterility

3 Introduction:

4 Tomato is one of the most important crops produced and consumed worldwide and has 5 also been used as a model plant for studying plants in the Solanaceae family and plants bearing 6 fleshy fruits. To date, diverse tomato studies have been conducted, especially tomato genomic 7 studies, given the availability of the complete genome sequence (The Tomato Genome 8 Consortium, 2012), various public databases, and mutant resources that were generated using 9 breeding lines and model tomato lines, including a dwarf cultivar, 'Micro-Tom' (Scott and 10 Harbaugh, 1989; Minoia et al. 2010; Saito et al. 2011).

Angiosperms form reproductive organs, such as anthers and pistils, that bear pollen grains and ovules, respectively. In this reproductive system, gametogenesis plays an important role in producing offspring. To date, several genes involved in gametogenesis have been identified in the model plant Arabidopsis, and their stage-specific roles in the regulation of reproductive development have been elucidated (Schneitz et al. 1997; Wilson and Zhang 2009; Bencivenga et al. 2011; Plackett et al. 2011).

According to Sanders et al. (1999), the development of anthers and pollen in Arabidopsis is divided into 15 stages. During stages 1 to 4, when the anther primordium emerges and establishes a bilateral structure, *EXCESS MICROSPOROCYTES1/EXTRA*

1	SPOROGENOUS CELLS (EMS1/EXS), a putative leucine-rich repeat receptor-like kinase, and
2	TAPETAL DETERMINANT1 (TPD1), a ligand for EMS1/EXS, both function to determine
3	archesporial cell number. Both the Arabidopsis ems1/exs mutant and the tpd1 mutant are male
4	sterile and form extra meiocytes caused by additional L2 layer cells, and the tapetal and middle
5	cell layers are absent in these mutants (Canales et al. 2002; Zhao et al. 2002; Yang et al. 2003).
6	Differentiation of the endothecium, middle layer, tapetum and microspore mother cells initiate
7	at stage 5. Then, microspore mother cells undergo meiosis between stages 5 and 7 (Sanders et
8	al. 1999), producing microspores or pollen grains within the tapetal cell layer.
9	DYSFUNCTIONAL TAPETUM1 (DYT1) encodes a putative basic helix-loop-helix (bHLH)
10	transcription factor that is predicted to act downstream EMS1/EXS, and this action is required
11	for the normal expression of ABORTED MICROSPORES (AMS) and MALE STERILITY1
12	(MS1), which are essential for normal tapetal development and viable pollen production during
13	these stages (Wilson et al. 2001; Sorensen et al. 2002; Zhang et al. 2006).
14	On the other hand, many genes related to ovule development have also been identified
15	in Arabidopsis (Schneitz et al. 1997; Skinner et al. 2004). Based on cytological aspects, ovule
16	primordia are divided into 3 elements: the funiculus, chalaza and nucellus. Genes specifically
17	expressed within each part have been identified (Schneitz et al. 1997; Sieber et al. 2004a; Kelly
18	and Gasser 2009). The site-specific expression of the homeobox gene WUSCHEL (WUS) within
19	the nucellus is critical for integument development and the formation of normal mature ovules

1	in Arabidopsis and tomato (Gross-Hardt et al. 2002; Sicard et al. 2008a). Accordingly, WUS
2	expression is generally confined to the nucellus during ovule development based on the
3	regulation by specific genes or hormones in Arabidopsis. For example, the bell1 (bel1) mutant
4	and the corona (cna)/phabulosa (phb)/phavoluta (phv) triple mutant, which showed abnormal
5	ovule development, exhibit an aberrant WUS expression pattern that prevents normal
6	integument formation (Brambilla et al. 2007; Bencivenga et al. 2012; Yamada et al. 2016).
7	Furthermore, WUS expression expands into the chalaza with exogenous treatment with the plant
8	hormone cytokinin N^6 -benzylaminopurine (BAP), and the treatment results in the formation of
9	new primordia (Bencivenga et al. 2012). In addition, precise mRNA localization of
10	AINTEGUMENTA (ANT), which encodes an AP2 family transcription factor, and INNER NO
11	OUTER (INO), which encodes a plant-specific YABBY family transcription factor, at the
12	chalaza is required for normal ovule integument development (Elliott et al. 1996; Baker et al.
13	1997; Villanueva et al, 1999).

There are several reports that describe anther and ovule development in tomato. For example, tomato *male sterile 10³⁵ (MS10³⁵)* has been isolated as a homologous gene of *AtDYT1*, and the *ms10³⁵* mutant exhibits dysfunctional meiosis and aberrant tapetum formation (Jeong et al. 2014). The *parthenocarpic fruit (pat)* mutant forms aberrant ovules caused by a failure of integument development (Mazzucato et al. 1998). In addition, the mini zinc finger gene *INHIBITOR OF MERISTEM ACTIVITY (IMA)* regulates ovule development by activating D-

class gene of ACBD model of flower development expression in tomato and Arabidopsis
 (Sicard et al. 2008ab). However, gametogenesis genes and their associated mechanisms remain
 largely unknown in tomato.

SPOROCYTELESS/NOZZLE (SPL/NZZ) plays a key role in gametogenesis, both in 4 anther and ovule development, in Arabidopsis (Schiefthaler et al. 1999; Yang et al. 1999), and $\mathbf{5}$ loss of function of SPL/NZZ results in abnormal germ cell production, causing strong sterility 6 in both male and female reproductive organs. SPL/NZZ transcripts are observed in the stamen 78 primordia and developing stamens as well as in ovule primordia, which is induced by AGAMOUS (AG) or brassinosteroids (Ito et al. 2004; Ye et al. 2010). AG is a member of 9 MADS-box gene and it is necessary for carpel and stamen development (Pinyopich et al. 2003). 10 Additionally, SPL/NZZ physically binds with CINCINNATA (CIN)-like TEOSINTE 11 BRANCHED1/CYCLOIDEA/PCF (TCP) transcription factors or YABBY family transcription 12factors through the N-terminal domain and the TOPLESS/TOPLESS-RELATED (TPL/TPR) 13co-repressor at the EAR motif (Sieber et al. 2004b; Chen et al. 2014; Wei et al. 2015). Then, 14SPL/NZZ functions as an adaptor-like transcriptional repressor through interaction with 15TPL/TPR (Wei et al. 2015). However, very little is known about the function of SPL/NZZ in 16 tomato. In this study, we report the identification and characterization of SlSES, a homologue 1718of SPL/NZZ, as an essential factor of gametogenesis development. This study demonstrates that pollen and ovule development were arrested in the very early developmental stages of each
 reproductive organ in the *Slses* mutant.

3

4 **Results:**

5 The identification of the sterile Slses mutant

6 To uncover the regulatory mechanism underlying reproductive organ development, we screened for mutants that were defective in normal reproductive development. From the tomato $\overline{7}$ 8 mutant populations that were generated by gamma ray-irradiated lines (Saito et al. 2011), one mutant producing seedless fruits was isolated. To characterize the cytological aspects of the 9 mutant, we first counted the number of each type of flower organ. In both the WT and mutant, 10 the inflorescence was typically composed of 5 sepals, petals and anthers and a single pistil 11 (Table 1). Although the WT formed an inflorescence with swollen anthers containing mature 12viable pollen grains at the anthesis stage, the mutant formed clearly defective anthers; they 13appeared wilted and small, and no pollen grains were visible at this time (Fig. 1A-D). To 14confirm the fertility of the ovules and pollen of the mutant, we crossed WT pollen to the mutant 15stigma or mutant pollen to the WT stigma and examined whether viable seeds were produced. 16No pollen was visually observed in the mutant. As the result, no viable seeds were produced 17from either type of cross-pollination; however, the mutant occasionally produced seedless fruit 18(data not shown). These results indicated that the mutant showed both male and female sterility. 19

Based on this phenotype, the mutant was designated as sexual sterility (Slses).

2	Then, to clarify genetic nature of the Slses mutation, we conducted crossing analysis
3	with wild type Micro-Tom. Given that the Slses homozygous mutant exhibited both complete
4	male and female sterility, heterozygous Slses plants were crossed with WT, and the resulting F_1
5	seeds and the following F2 offspring were obtained (Supplementary Fig. S1). The
6	morphological phenotype and fertility of 17 F ₁ plants were almost equivalent to those of WT,
7	indicating that the Slses mutation was recessive (Table 2). The phenotypes of the F ₂ offspring
8	segregated to 118 (phenotype; WT) : 40 (phenotype; Slses), which approximately fit the
9	theoretical ratio of the WT and <i>Slses</i> phenotypes 3 : 1, suggesting that the <i>Slses</i> mutation was
10	monogenic (Table 2).

Next, time course analysis of cross-sections during anther development was conducted. 11 In this analysis, flower buds at different developmental stages, which were defined by the size 12of flower buds, were obtained, and cross-sections of the anther structure were examined under 13light microscopy. There was no obvious difference in timing of flowering and flower 14development between WT and the Slses mutant. At the stage when the flower buds were 1 to 2 15mm and 2 to 3 mm, normal formation of the vascular bundle (VB), epidermis (Ep), endothecium 16 (En), middle layer (ML), tapetum (T), and pollen mother cells (PMC, 1-2 mm bud) or meiotic 17cells (MC, 2-3 mm bud) were observed in WT (Fig. 2E, F). In contrast, the formation of the En, 18ML, T, PMC and MC were not clearly evident, and we only observed the vascular bundle and 19

1	epidermis at the stage of 1 to 2 mm and 2 to 3 mm buds in the Slses mutant (Fig. 2I, J).
2	Microspores (Msp) were observed at the bud stage of 4 to 5 mm in WT but not in the Slses
3	mutant (Fig. 2G, K). At anthesis stage, when the bud size is 6 to 7 mm, many mature pollen
4	grains were clearly observed in the anther locules in the WT, whereas the Slses mutant never
5	underwent pollen grain formation inside the anther (Fig. 2A-D, H, L). The anther structure
6	appeared wilted and less colored in the mutant compared with the WT (Fig. 1A, B, Fig. 2A-D).
7	Next, we observed the pistil in the WT and the Slses mutant (Fig. 1E-I). The size of
8	the ovary at anthesis in the Slses mutant was clearly smaller than that in the WT (Fig. 1I). At
9	the 1 to 2 mm bud size stage, we observed ovule primordia around the placenta in both the WT
10	and the Slses mutant (Fig. 3E, I). At the 2 to 3 mm and 4 to 5 mm bud stages, ovule integuments
11	and megaspore mother cells (MMCs) were observed, and the ovule primordia enlarged as the
12	stages proceeded in WT. We could not observe the ovule integuments and MMC, which are
13	essential for complete ovule development, and the enlargement level of ovule primordia was
14	quite low in the Slses mutant (Fig. 3 F, G, J, K, M-O). From this stage, ovule development was
15	arrested in the Slses mutant. We observed cross-sections of mature ovules at the anthesis stage
16	in WT. The ovules were smaller, and their structure appeared disorganized in the Slses mutant
17	(Fig. 3A-D, H, L).

19 The Slses mutant produces seedless fruit

1	Despite male and female sterility, the Slses mutant occasionally bore red mature fruits
2	and the size and weight of these fruits were significantly reduced compared with WT fruits (Fig.
3	4A-D). Most of mature fruits produced in the WT contained 3 locules (average 3.2 locules per
4	fruit), whereas those in the Slses mutant contained 2 locules (average 2.2 locules per fruit),
5	indicating that the number of locules was significantly reduced in the Slses mutant (Fig. 4E).
6	The mature red fruits produced in the Slses mutant were completely seedless, indicating that
7	the Slses mutant induces parthenocarpy. However, molecular mechanism of how the Slses
8	mutation induces parthenocarpy is still unclear.
9	
10	SISES encodes SPL/NZZ
11	To map the Slses mutation, positional cloning was performed using SolCap infinium
12	carrying a large number of SNPs that could discriminate tomato genotypes (Sim et al. 2012),
13	with 2098 markers available for distinguishing between 'Micro-Tom' and 'Ailsa Craig'. The
14	SolCap array coupled with 28 F_2 mapping populations that were derived from crosses between
15	Slses heterozygous plants and Ailsa Craig allowed for narrowing down the candidate area to a
16	2.3 Mbp region corresponding to the physical position between 62,465,682 base pairs (bps) to
17	64,759,097 bps on chromosome 7 based on the protein annotation extracted from version
18	SL2.40 from the SGN database (Fig. 5A). The 2.3 Mbp region spans 334 open reading frames
19	(ORFs), and none of these ORFs are involved in tomato gametogenesis

1	(http://www.kazusa.or.jp/tomato_sbm/). Then, we obtained the whole genome sequence of the
2	Slses mutant using Illumina HiSeq 2000 next-generation sequencing, and its genome
3	information was compared with the tomato reference genome sequence according to Ariizumi
4	et al. (2014). Given that the Slses mutant was a monogenic recessive mutation, we exclusively
5	focused on homozygous mutations specifically found in the Slses mutant. The genome sequence
6	analysis identified 94 homozygous mutations consisting of 9 point mutations and 85 insertion
7	and deletion (indel) mutations within the 2.3 Mbp region. The 9 point mutations included 1
8	nonsynonymous, 1 synonymous, and 7 intergenic mutations. The 85 indels included 2
9	mutations within exon causing modification of amino acid sequence of putative ORFs and 19
10	mutations within intron or UTR regions. In addition, 64 mutations were present within the
11	intergenic region (Supplementary Table S1). Therefore, 1 missense and 1 deletion mutation,
12	which were found in Solyc07g063670, and 1 insertion mutation, which was found in
13	Solyc07g065700, were considered candidates for the SISES gene. Among them, one deletion
14	mutation, which caused a 13-bp deletion from the first exon of Solyc07g063670, was primarily
15	selected because the Slses mutant was produced by gamma-ray irradiation and caused massive
16	flame shift rather than the insertion mutation. Gamma-ray irradiation often causes genome
17	insertion or deletion rather than SNPs, and we often obtain flame shift caused mutant. The
18	deletion caused a frame shift at the 123 rd nucleotide position from the start codon, which was
19	considered the 1 st position. As the results of frame shift the putative amino acid was shortened

from 353 to 57 in the Slses mutant (Fig. 5B, C).

2	Genetic linkage analysis using a DNA marker (SISES marker-1; Supplementary Table
3	S2) distinguishes this deletion and the F_2 populations derived from a cross between the WT
4	Micro-Tom and the Slses mutant, revealing a perfect linkage between the mutation and the
5	phenotypes (Supplementary Fig. S2). The candidate gene of the Slses mutant, Solyc07g063670,
6	encodes a domain of plant transcription factor NOZZLE (29-215 a.a.; NCBI Conserved Domain
7	Search; Fig. 5C). This protein shares 26.4% identity (39.2% similarity) with the SPL/NZZ
8	protein and 21.1% identity (31.8% similarity) with the TCP interactor containing EAR motif 3
9	(TIE3) protein of Arabidopsis (Fig. 5D; Supplementary Table S3). The candidate protein also
10	had an EAR motif (LxLxL; Kagale and Rozwadowski 2011) at the C-terminal region (345-351
11	a.a.) similar to that found in the SPL/NZZ and TIE proteins of Arabidopsis (Fig. 5E). Further
12	sequence analysis indicated that no paralogues of the candidate gene existed in the tomato
13	genome (Fig. 5D), suggesting that SISES exists as a single gene and plays a non-redundant role
14	in the tomato genome.

15

16 Complementation experiments of the Slses mutant

To confirm whether the candidate gene *Solyc07g063670* was responsible for the *Slses* mutation, the genomic region of the candidate gene, including the 2.5-kb promoter region, 1,310-bp coding sequence, and 700-bp terminator region, were introduced into the offspring of

1	heterozygous Slses plants (Supplementary Fig. S3) such that the transgene randomly integrated
2	into plants that were homozygous, heterozygous or azygous for the 13-bp nucleotide deletion
3	within Solyc07g063670. We developed a PCR genotyping method that allowed for
4	discrimination of 13-bp nucleotide deletion such that we selected the transgenic plants that
5	carried both the transgene and the endogenous homozygous 13-bp nucleotide deletion (SISES
6	marker-2; Supplementary Table S2; Supplementary Fig. S4). Three such transgenic plants were
7	obtained. The plants exhibited swollen anthers, producing fertile pollen grains and fertile
8	ovules; thus, self-pollination produced viable offspring seeds that germinated thereafter
9	(Supplementary Fig. S5), indicating that the phenotypes in the complementation lines were
10	fully restored. These results indicated that the 13-bp deletion found in Solyc07g063670 was
11	responsible for the Slses mutation and that the Arabidopsis SPL/NZZ homologue in tomato
12	(SISES) functions to regulate normal reproductive organ development.
13	
14	SISES is expressed at early stages of reproductive development and regulates SIWUS and SIINO
15	expression
16	qRT-PCR analysis was conducted to examine the expression pattern of the SISES gene
17	(Fig. 6). mRNA was extracted from flower buds 1 mm in size; pistils from developmental stages
18	when the flower bud is 3, 4.5 and 6 mm in size; pistils 1 day before anthesis (DBA); and pistils

19 at the anthesis stage. In this analysis, several genes related to anther or ovule development in

1	tomato or Arabidopsis were also examined regarding the effect of the Slses mutation on the
2	transcriptional regulation of gametogenesis genes. SISES mRNA was highly expressed in 1 mm
3	flower buds and 3 mm pistils. However, mRNA levels decreased along with pistil development
4	and were barely detectable in 6 mm pistils and 1 DBA buds in WT (Fig. 6). mRNA expression
5	of <i>Male sterile</i> 10^{35} (<i>MS</i> 10^{35}), which is involved in both meiosis and programmed cell death of
6	the tapetum during microsporogenesis in tomato (Jeong et al. 2014), was evident in 1 mm WT
7	buds, whereas its mRNA was minimally detectable in the Slses mutant (Fig. 7A). mRNA
8	expression of BARELY ANY MERISTEM1 and 2-like (BAM1/2-like) and EMS1/EXS-like, which
9	are homologues of Arabidopsis BAM1/2 and EMS1/EXS, respectively, and are essential for early
10	anther development (Zhao et al. 2002; Hord et al. 2006), were significantly downregulated in
11	the Slses mutant (Fig. 7B, C). mRNA expression of TOMATO AGAMOUS1 (TAG1), a
12	homologue of Arabidopsis AG1, which is involved in the transcriptional regulation of SPL/NZZ,
13	was decreased in the Slses mutant compared with the WT (Fig. 7D). In addition, mRNA
14	expression of WUSCHEL (WUS) and INNER NO OUTER (INO), which are the important genes
15	for ovule formation, and GOBLET (GOB), a homologue of Arabidopsis CUP-SHAPED
16	COTYLEDONS1 (CUC1) and CUC2, which are essential for the formation of carpel margin
17	meristems (CMMs) (Hendelman et al. 2013; Kamiuchi et al. 2014), was decreased in the Slses
18	mutant compared with WT (Fig. 7E, F, M). The expression of AINTEGUMENTA-like (ANT-
19	like), PIN-FORMED1-like (PIN1-like), BELL1-like (BEL1-like), REVOLUTA (SIREV),

1	CORONA-like (CNA-like) and PHABULOSA/PHAVOLUTA-like (PHB/PHV-like), which are
2	homologues of Arabidopsis and are required for ovule formation, were downregulated in the
3	pistil from 4.5 mm and buds in the Slses mutant (Fig. 7G-L).

5 **Discussion:**

6 We identified a 13-bp deletion in the SISES CDS in the Slses mutant, and the deletion caused a frame shift, resulting in the loss of the last 296 amino acids. The plant transcription $\overline{7}$ 8 factor NOZZLE motif and EAR motif are completely lost in the Slses mutant (Fig. 5B, C). In 9 contrast, complementation experiments demonstrated that introduction of the genomic region of Solyc07g063670 with its native promoter and terminator into the Slses mutant fully restored 10 these phenotypes (Supplementary Fig. S5). These results indicate that SISES (Solyc07g063670) 11 12is the gene responsible for the Slses mutant, and the SlSES loss-of-function mutation had a 13drastic effect on the development of reproductive organs.

The SISES protein sequence exhibits significant homology with the SPL/NZZ (Schiefthaler et al. 1999; Yang *et al.* 1999) and TIE proteins (Tao et al. 2013) of Arabidopsis (Fig. 5D). These proteins carry an EAR motif and act as transcriptional repressors due to interaction with TOPLESS through the EAR motif. TOPLESS can repress transcription in vivo (Long et al. 2006; Pan et al. 2010; Causier et al. 2012; Tao et al. 2013; Wei et al. 2015). The putative amino acid sequence of SISES exhibited a highly conserved EAR motif at the C- terminus (Fig. 5E), suggesting that the tomato SISES protein may interact with TOPLESS and
 act as transcriptional repressor.

3	The Arabidopsis homologous gene SPL/NZZ is one of the key genes involved in anther
4	development, especially in the regulation of sporocyte formation in Arabidopsis (Yang et al.
5	1999; Plackett et al. 2011). En, ML, T, PCM and MC development were observed at the 1 to 3
6	mm bud stages in WT but not in the Slses mutant (Fig. 2E, F, I, J). These tissues differentiate
7	from the L2 layer of initial stages of anther development, indicating that differentiation from
8	the L2 layer was aborted in the Slses mutant. As a result, the Slses mutant exhibited wilted
9	anthers and produced no mature pollen grains (Fig. 2C, D). The phenotypes of the Slses mutant
10	were similar to those of the Arabidopsis <i>spl/nzz</i> mutant (Yang et al. 1999), indicating that <i>SlSES</i>
11	is essential for anther primordia formation in tomato. We detected minimal mRNA expression
12	of MS10 ³⁵ and EMS1/EXS-like, which are homologues of Arabidopsis DYT1 and EMS1/EXS,
13	respectively, in the 1 mm bud of the Slses mutant (Fig. 7A, C). Given that both the tomato
14	$ms10^{35}$ mutant and the Arabidopsis $dyt1$ mutant exhibit normal PMC development and
15	abnormal tapetum development during pollen production in the anther (Zhang et al. 2006; Jeong
16	et al. 2014) and that DYT1 acts downstream of SPL/NZZ in Arabidopsis (Zhang et al. 2006), it
17	is suggested that SISES also acts upstream of MS10 ³⁵ . In addition, MS10 ³⁵ mRNA expression
18	might be required for normal PMC development in tomato. Additionally, EMS1/EXS acts
19	upstream of DYT1 and downstream of SPL/NZZ. The ems1/exs mutant produces anthers lacking

1	the tapetum but with extra PMCs in Arabidopsis (Zhao et al. 2002). This finding may suggest
2	that EMS1/EXS-like (Solyc09g098420) acts downstream of SISES and has similar functions as
3	its homologue in Arabidopsis.

The locule number and ovary size were significantly reduced in the Slses mutant 4 compared with the WT (Fig. 1I, 4E), indicating that SISES is partly involved in determining $\mathbf{5}$ carpel number and ovary development. Although SPL/NZZ is essential for ovule primordia and 6 nucellus development, integuments develop normally in Arabidopsis (Schiefthaler et al. 1999; $\overline{7}$ 8 Sieber et al. 2004a; Bencivenga et al. 2011). Interestingly, the Slses mutant formed a pistil 9 (carpel) with ovules whose development arrested at the 2 to 3 mm bud stages, which was associated with a failure in nucellus (MMC) and integument development (Fig. 3I, J), indicating 10 that SISES functions in nucellus development, similar to Arabidopsis and integument 11 12development in tomato. INO plays a key role in the formation of the outer integuments in 13Arabidopsis (Baker et al. 1997; Villanueva et al. 1999). INO expression is not affected in the spl/nzz mutant compared with WT in Arabidopsis; therefore, the spl/nzz mutant forms 14integuments normally (Balasubramanian and Schneitz 2002; Sieber et al. 2004a). In tomato, the 15Slses mutant did not exhibit any integument development (Fig. 3O), and SlINO expression in 16the mutant is minimally detectable in the pistils of the 4.5 mm bud and anthesis stages (Fig. 177D), suggesting that the molecular regulation between SPL/NZZ and INO in Arabidopsis and 1819SISES and SIINO in tomato differs. SISES function is necessary for *SlINO* expression in tomato.

1	Arabidopsis forms ovules from two integuments (inner and outer integuments and bitegmic
2	ovules), whereas tomato forms ovules from a single integument (unitegmic ovules) (Wang and
3	Ren 2008; Skinner et al. 2016). Our results indicate that the characteristics of the single
4	integument in tomato are similar to that of the outer integument in Arabidopsis.
5	It has been demonstrated that exclusive and timely WUS expression in the nucellus is
6	critical for normal ovule development in Arabidopsis (Groß-Hardt et al. 2002). In the Slses
7	mutant, SIWUS mRNA expression was considerably downregulated compared with WT (Fig.
8	7E). In Arabidopsis, WUS expression in the nucellus was reduced in the <i>spl/nzz</i> mutants same
9	as our study (Sieber et al. 2004a), and this information suggests that SISES expression is
10	required to induce SIWUS mRNA transcription in tomato. ANT-like and BEL1-like mRNA
11	expression was downregulated in the pistils of the 4.5 mm bud in the Slses mutant compared
12	with the WT (Fig. 7G, I). These genes also play key roles in the development of ovules and
13	integuments in Arabidopsis (Elliott et al. 1996; Schneitz et al. 1997; Brambilla et al. 2007).
14	ANT is expressed within the developing integuments and funiculus. BEL1 is strongly expressed
15	within the developing integuments, and the <i>spl/nzz</i> mutant exhibits ectopic expression of ANT
16	and BEL1 within the ovule primordia in Arabidopsis (Balasubramanian and Schneitz 2000;
17	Bencivenga et al. 2012). ANT and BEL1 expression are required for the ovule integument
18	development in Arabidopsis (Schneitz et al. 1997). Ovule development, especially the ovule
19	integument (Fig. 3O), in the Slses mutant was arrested, suggesting reduced expression of ANT-

1	like and BEL1-like genes, which was actually in accordance with our results. PIN1 is an auxin
2	efflux facilitator, and the pin1-5 mutant exhibits abnormal ovule development. Therefore,
3	SPL/NZZ expression in the ovule is required to induce PIN1 transcription in Arabidopsis
4	(Bencivenga et al. 2012). Moreover, the class III homeodomain leucine zipper (HD-ZIPIII)
5	genes CNA, PHB, PHV and REV are expressed within the chalaza and restrict WUS and INO
6	expression to the nucellus and gynobasal side of the ovule in Arabidopsis (Kelley et al. 2009ab;
7	Yamada et al. 2016). PIN1, CNA, PHB, PHV and REV are required for precise spatiotemporal
8	expression for correct ovule development. In tomato, PIN1-like, CNA-like, PHB/PHV-like and
9	SIREV expression was reduced in the pistils of the 4.5 mm bud and/or anthesis flower, but the
10	reduction was comparatively small. Nevertheless, ovule development was arrested in the Slses
11	mutant (Fig. 7H, J-L). Given that the precise spatial expression of these genes is essential to
12	successfully form ovules, the control of the spatial expression and expression level of these
13	genes were collapsed in the Slses mutant.
14	SPL/NZZ expression is positively regulated by AG in Arabidopsis (Ito et al. 2004). On
15	the other hand, the highest mRNA expression point of SISES and TAG1 was in the 1 mm bud
16	and the pistils of 4.5 mm bud, respectively (Fig. 6, 7D). This may represent that TAG1 doesn't
17	regulate <i>SlSES</i> expression or they need other factor to regulate the mRNA expression of <i>SlSES</i> .
18	In any case, we could not judge there is such a relationship between SISES and TAG1 in our
19	study. In order to clarify, we need to do further experiments.

1	In this study, we identified a key gene, SISES, that is responsible for both male and
2	female gametogenesis in tomato. SISES is a homologue of SPL/NZZ in Arabidopsis, and the
3	loss-of-function SISES mutant exhibits sexual sterility caused by a failure to form pollen and
4	ovules. Possible roles of SISES in ovule and anther development are proposed in Fig. 8 and are
5	compared with SPL/NZZ of Arabidopsis. In ovule development, SPL/NZZ positively regulates
6	PIN1 and WUS expression and plays a key role in nucellus development in Arabidopsis (Sieber
7	et al. 2004a; Bencivenga et al. 2012). On the other hand, SISES is required to express SIWUS
8	and SlINO. Moreover, we demonstrate that SlSES has a role in nucellus and integument
9	development via controlling SlINO expression in tomato (Fig. 8). In addition, in anther
10	development, SPL/NZZ positively regulates BAM1/2 and EMS1/EXS expression and is essential
11	for archesporial cell development in Arabidopsis (Yang et al. 1999; Zhao et al, 2002; Hord et
12	al. 2006; Ma et al. 2012). SISES positively regulates EMS1/EXS-like expression and plays a key
13	role in archesporial cell development in tomato (Fig. 8). Isolation of the SISES gene may
14	positively impact gametogenesis research in tomato.
15	

16 Materials and Methods:

17 Plant materials and growth conditions

18 The miniature tomato 'Micro-Tom' and the cultivated species 'Ailsa Craig' were used 19 in this study. The *Slses* mutant was obtained from gamma ray-irradiated Micro-Tom mutant populations produced by the National BioResource Project (NBRP, http://www.nbrp.jp/) (Saito et al. 2011). All lines were grown under a regimen of 16-h days and 8-h nights at a constant temperature of 25°C under fluorescent lights at 100 to 150 μ Em⁻²s⁻¹.

4

5 Histological and microscopic analysis

6 Whole anthesis flowers and flower buds were collected and fixed in FAA (formalin : acetic acid: 70% EtOH = 1:1:18), vacuum-infiltrated twice for 10 min each, and left overnight. $\overline{7}$ 8 The fixed tissues were dehydrated with a t-butyl alcohol series from No. 1 to 6, which contained *t*-butyl alcohol, ethanol and DW (Supplementary Table S4). Each step required a greater than 9 6-hour incubation at room temperature, and the No. 6 solution was used twice. The tissues in 10 the No. 6 solution were incubated at 60°C for 2 hours. Then, melted paraffin was added, and 11 the tissues were stored at 60°C overnight. The embedded tissues were sliced into 10-µm 12sections using a microtome (MICROM HM325, Leica, http://www2.leicabiosystems.com/). 13The sections were stained with 0.05% toluidine blue (pH 7.0). Paraffin was removed with 14xylene, and the sections were mounted with entellan new (Merck Millipore). Then, sections 15were observed under an optical microscope (Olympus BX53, http://www.olympus-16lifescience.com/). 17

18

19 Map-based cloning using DNA markers

1	To conduct SolCAP analysis (http://solcap.msu.edu/index.shtml) for rough mapping,
2	an F ₂ mapping population was constructed from a cross between S. lycopersicum 'Ailsa Craig'
3	and the Slses heterozygous mutant in the 'Micro-Tom' background. Then, we used 28 F2 plants
4	that exhibited the Slses mutation for this analysis. In total, 7,720 markers were used for this
5	experiment, and 2,098 markers showed polymorphism to distinguish between 'Micro-Tom' and
6	'Ailsa Craig'. Then, the region of the responsible gene was narrowed down.
7	
8	Genome re-sequencing and identification of responsible gene
9	The genome sequence of the Slses mutant was obtained by Illumina HiSeq 2000. The
10	sequences of 'Heinz 1706' and 'Micro-Tom' Japan were used as reference genomes (Ariizumi
11	et al. 2014; Kobayashi et al. 2014). Linkage analysis was conducted with F2 plants which
12	obtained crossed with the ses mutant and 'Micro-Tom', WT or 'Ailsa Craig'. The marker (SISES
13	marker-1, Supplementary Table S2) was designed to amplify the 13-bp deletion region. The
14	amplicon sizes were different in the ses mutant and WT.
15	
16	Characterization of the identified gene
17	The genome sequences and amino acid sequences were obtained from the Sol
18	Genomics Network (SGN) (https://solgenomics.net/). The estimation of the SISES protein
19	domains was conducted using the Conserved Domain Database (CDD)

1	(http://www.ncbi.nlm.nih.gov/cdd). The homologous genes from Arabidopsis and Tomato were
2	searched using The Arabidopsis Information Resource (TAIR) (https://www.arabidopsis.org/)
3	and SGN, respectively. We selected the homologous gene of which e-value is under '0.1'. The
4	phylogenetic tree was constructed with multiple sequence alignment using CLUSTALW
5	(http://www.genome.jp/tools/clustalw/). The amino acid homology of SISES and SPL/NZZ was
6	analyzed using the EMBOSS Needle (http://www.ebi.ac.uk/Tools/psa/emboss_needle/) in the
7	European Bioinformatics Institute (<u>http://www.ebi.ac.uk/</u>).
8	
9	Creation of transgenic plants
10	The binary vector pIG121Hm was used for the creation of transgenic plants. The
11	complete SISES gene, which included 2563 bp of upstream sequence and 763 bp of downstream
12	sequence, was amplified with gene-specific primers (whole SISES gene primer, Supplementary
13	table S2) using the In-Fusion technique. The amplified fragments of the complete genomic
14	region were inserted into pIG121Hm digested by HindIII and SalI using the In-Fusion
15	technique (TAKARA) (Supplementary Fig. S3). Then, the construct was transformed into
16	Agrobacterium tumefaciens GV2260 by electroporation, and transformants were selected on
17	LB plates containing kanamycin (50 mg L ⁻¹). The transformation construct was transformed
18	into 'Micro-Tom' WT and plants heterozygous or homozygous for the Slses mutation using an
19	Agrobacterium-mediated transformation method (Sun et al. 2006). The transgenic plants were

1	selected on MS plates containing kanamycin (100 mg L ⁻¹). Ploidy and transgene copy number
2	transgene were assessed by flow cytometry and Southern blotting analysis, respectively, in the
3	T0 generation. Plants that were both diploid and single-copy transgenic were selected, and their
4	T1 or T2 generations that harbored the Slses homozygous mutation were used for further
5	analysis. The Slses homozygous mutants were selected using "SlSES marker-2" and the
6	restriction enzyme Bsl I, we could distinguish among homozygous, heterozygous and WT by
7	electrophoresis band pattern (Supplementary Table S2; Supplementary Fig. S4). The presence
8	of transgene was confirmed by detecting "NPT II marker" using PCR (Supplementary Table
9	S2).

11 Gene expression analysis

mRNA expression levels were analyzed by quantitative RT-PCR (qRT-PCR) using 12SYBR Premix Ex Taq II (TAKARA). The template cDNAs were synthesized from total RNA 13 extracted from 1 mm whole buds or pistils of unopened buds classified by bud length (3 mm, 144.5 mm, 6 mm, 1 day before anthesis) and anthesis flowers. Total RNA was extracted with an 15RNeasy Mini Kit (Qiagen, http://www.qiagen.com/), and cDNA was synthesized with a 16SuperScript VILO cDNA Synthesis Kit (Invitrogen, http://www.lifetechnologies.com/). SGN-17U316474 (SAND) mRNA was used as an internal control (Expósito-Rodríguez et al. 2008). The 18expression level was calculated according to Pfaffl 2001. The primers were designed with 19

[Primer3Plus] (<u>http://primer3plus.com/cgi-bin/dev/primer3plus.cgi</u>) (Supplementary Table S5).
 2

3 Statistical	analysis
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4	We used Student's t test (P = 0.05 and 0.01) to show the significance between two
5	values, WT and the <i>Slses</i> mutant. Then, Chi-squared (χ^2) test was performed to examine the
6	correctness of fit between the expected Mendelian ratios and segregation data for the Slses
7	mutation. We calculated χ^2 value on below conditions; degrees of freedom, 1; the expected
8	Mendelian ratios, WT : $Slses = 3 : 1$.
9	
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13	Micro-Tom (TOMJPF00001), Ailsa Craig (TOMJPF0004), and Slses were obtained from the
14	National BioResource Project, Ministry of Education, Culture, Sports, Science and Technology
15	(MEXT), Japan.
16	
17	Disclosures:
18	The authors have no conflicts of interest to declare.

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19	

Tables:

Table 1. Comparison of the numbers of flower organs between the *Slses* mutant and WT.

	Flower organ			
_	Sepal	Petal	Anther	Pistil
WT	5.20 ± 0.09	5.05 ± 0.09	5.15 ± 0.08	1.00 ± 0.00
Slses	5.30 ± 0.11	5.25 ± 0.10	5.30 ± 0.11	1.00 ± 0.00

Values are means \pm standard error. Statistical significance was analyzed using the Student's *t* test.

Table 2. Comparison of the phenotypes of F_1 and F_2 plants obtained by crossing the *Slses* mutant (heterozygous) and WT plants.

Cross	Generation –	Phenotype		w ² walua	Devalue
Cross		WT	Slses	χ^2 value	P value
<i>Slses</i> (Heterozygous) × WT	F_1	17	0	-	-
	F ₂ (A)	118	40	0.008	0.927
	F ₂ (B)	149	0	-	-

Two patterns of F_2 population were observed: segregated (A) and non-segregated (B).

See supplementary Fig. S1.

The χ^2 value was calculated from the F_2 populations (A)

1 Figure legends

2 Figure 1.

3	The floral morphology of the <i>Slses</i> mutant at the anthesis stage. (A, B) The flower, (C, D) the
4	anther and pollen grains, (E-H) the pistil. (A, C, E, G) WT, (B, D, F, H) Slses mutant. Bar = 1
5	mm. (I) The diameter and length of the ovary. Statistical analysis was performed using the
6	Student's <i>t</i> -test. **P<0.01; *P<0.05, n = 10, error bar = SE.
7	
8	Figure 2.
9	Cross-sections of developing anthers in WT and the Slses mutant. (A, C) Cross sections of

11 I) 1 to 2 mm bud, (F, J) 2 to 3 mm bud, (G, K) 4 to 5 mm bud, (H, L) 6 to 7 mm bud. (A, B, E-

anthers at the anthesis stage, (B, D) enlarged view of a piece of anther at the anthesis stage, (E,

- 12 H) WT, (C, D, I-L) the *Slses* mutant. En, endothecium; Ep, epidermis; MC, meiotic cell; ML,
- 13 middle layer; Msp, microspore; P, pollen; PMC, pollen mother cell; S, style; T, tapetum; VB,
- 14 vascular bundle. Bar = $500 \,\mu m$.
- 15

10

16	Figure	3.
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17 Sections of developing pistils in WT and the *Slses* mutant. (A, C) Cross-section, (B, D) 18 longitudinal section at the anthesis stage. (E, I) 1 to 2 mm bud, (F, J) 2 to 3 mm bud, (G, K) 4 19 to 5 mm bud, (H, L) 6 to 7 mm bud. Bar = $500 \,\mu$ m. (M) Ovule primordia of 2 to 3 mm bud, (N,

1	O) ovule primordia of 4 to 5 mm bud. Bar = 50 μ m. (A, B, E-H, M, N) WT, (C, D, I-L, O) the
2	Slses mutant. MMC, megaspore mother cell; O, ovule; OI, ovule integument; OP, ovule
3	primordia; P, placenta.
4	
5	Figure 4.
6	Morphology of ripe red fruits in the WT and the Slses mutant. (A) Appearance of red ripe fruits,
7	(B) cross-section of the red fruits. Bar = 1 cm. (C) Fruit weight, (D) fruit size and (E) the number
8	of locules per fruit in the WT and the <i>Slses</i> mutant. $n = 20$, error bar = SE, statistical analysis
9	was performed using the Student's <i>t</i> -test, $**P < 0.01$.

10

11 Figure 5.

Gene structure of Solyc07g063670. (A) The location and gene structure of Solyc07g063670. 12The blue bar indicates chromosome 7, the gray box indicates the UTR, white boxes indicate 13 exons, and regions between the boxes indicate introns. (B) Genome and amino acid sequences 14of WT and the Slses mutant at the mutation point. The upper black letters indicate genome 15sequences, and the lower red letters indicate putative amino acids. Hyphens indicate the 13-bp 16deletion. (C) The putative amino acid length and domain of the SISES protein. The putative 17amino acid length and domain region were exhibited under and upper of SISES protein models 18respectively. The domain is indicated with yellow and red boxes. * = stop codon. (D) 19

1	Phylogenetic tree of <i>SlSES</i> and its homologous proteins in Arabidopsis. (E) The EAR motif of
2	SISES and SPL/NZZ at the C-terminal region. The red frame indicates the EAR motif.
3	
4	Figure 6.
5	Expression analysis of <i>SlSES</i> in the WT by qRT-PCR. <i>SlSES</i> expression level in the WT in the
6	1 mm bud and in pistils from the 3 mm, 4.5 mm, 6 mm, 1 day before anthesis (DBA) and
7	anthesis stages. $n = 3$.
8	
9	Figure 7.
10	Expression analysis of gametogenesis genes in the WT and the <i>Slses</i> mutant by qRT-PCR. (A)
11	Male Sterile 10 ³⁵ (MS10 ³⁵), (B) BARELY ANY MERISTEM1 and 2-like (BAM1/2-like), (C)
12	EXCESS MICROSPOROCYTES1/EXTRA SPOROGENOUS CELLS-like (EMS1/EXS-like), (D)
13	Tomato AGAMOUS1 (TAG1), (E) WUSCHEL (SIWUS), (F) INNER NO OUTER (SIINO), (G)
14	AINTEGUMENTA-like (ANT-like), (H) PIN-FORMED1-like (PIN1-like), (I) BELL1-like
15	(BEL1-like), (J) REVOLUTA (SIREV), (K) CORONA-like (CNA-like), (L)
16	PHABULOSA/PHAVOLUTA-like (PHB/PHV-like), (M) GOBLET (GOB). mRNA expression
17	was analyzed at the 1 mm bud stage and in pistils from the 4.5 mm bud and anthesis stages. n
18	= 3, error bar = SE. Statistical analysis was performed using the Student's <i>t</i> -test. **P< 0.01 ;

19 *P<0.05.

2 Figure 8.

3	A model for ovule (left) and anther (right) development regulated by SISES (SPL/NZZ) in
4	tomato and Arabidopsis. Solid arrows represent positive regulation, and dotted arrows indicate
5	gene function (red = tomato, blue = $Arabidopsis$). The gray box shows the model of integument
6	development in Arabidopsis (left) and tomato (right). I = integument, II = inner integument, OI
7	= outer integument, PMC = pollen mother cells, PPC = primary parietal cells, PSC = primary
8	sporogenous cells, SPC = secondary parietal cells.
9	

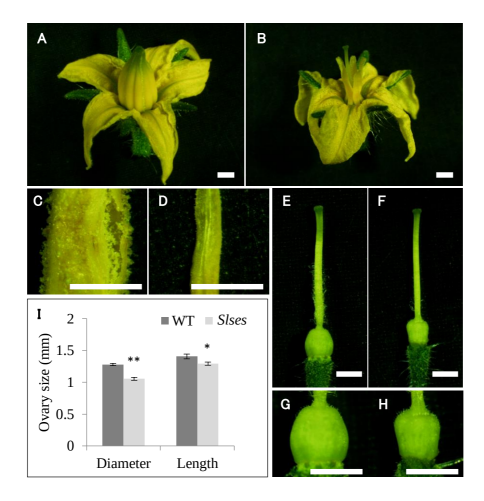
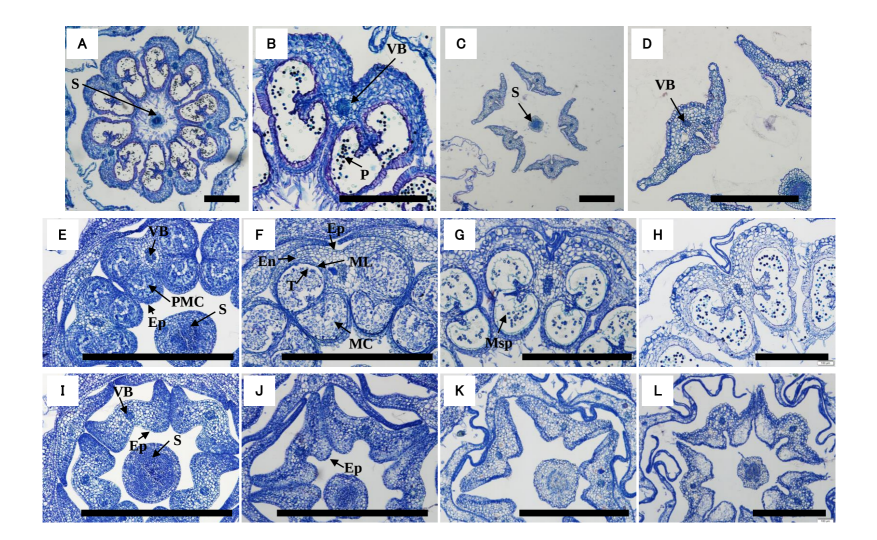
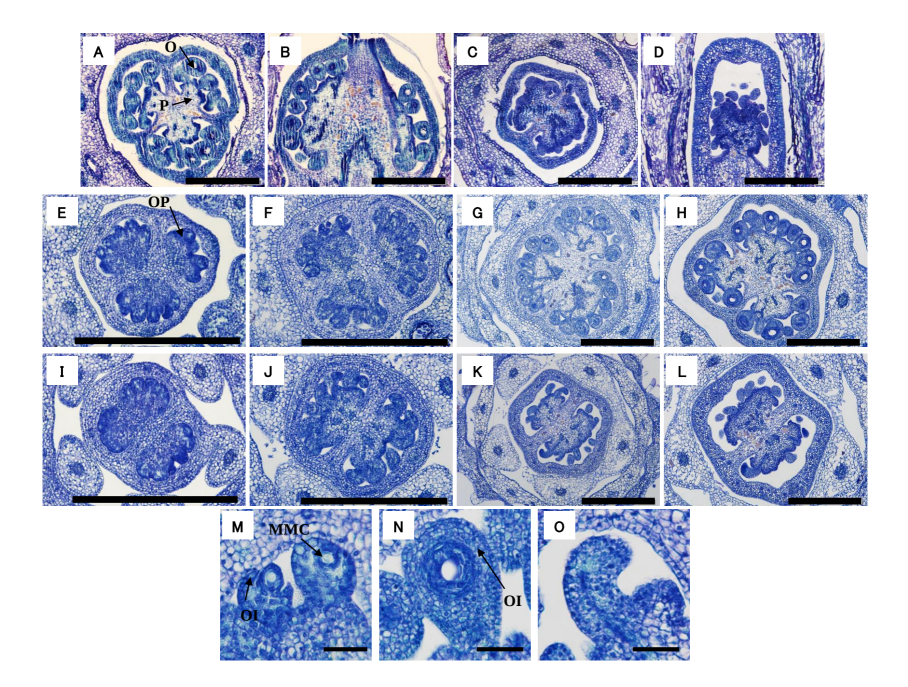
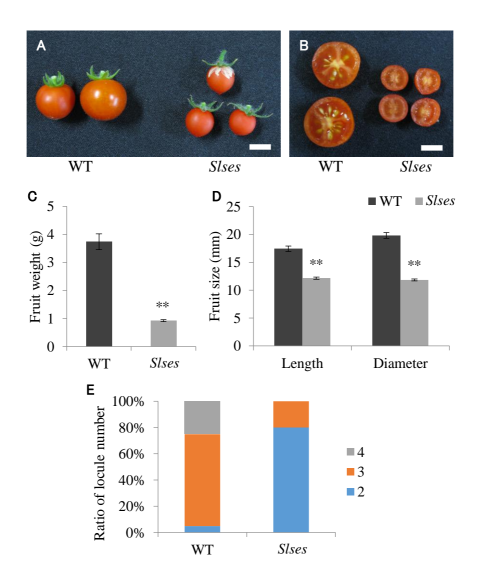
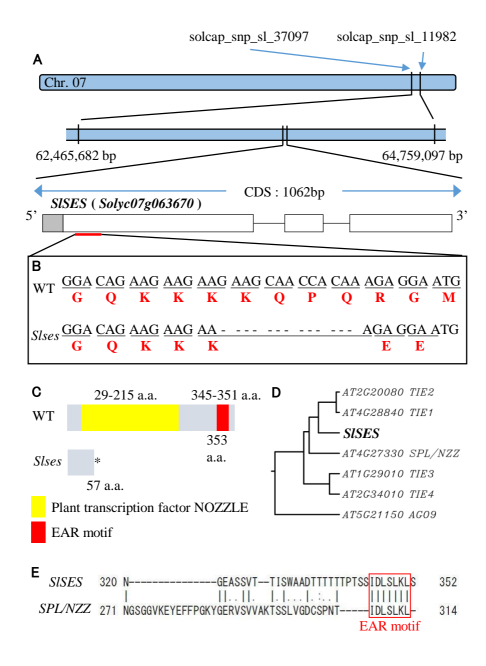


Figure 1









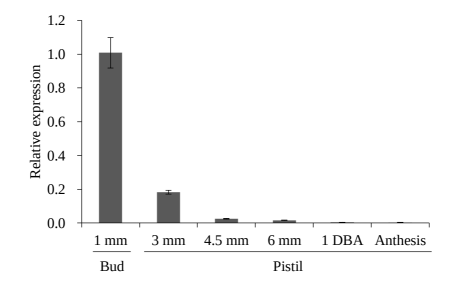


Figure 6

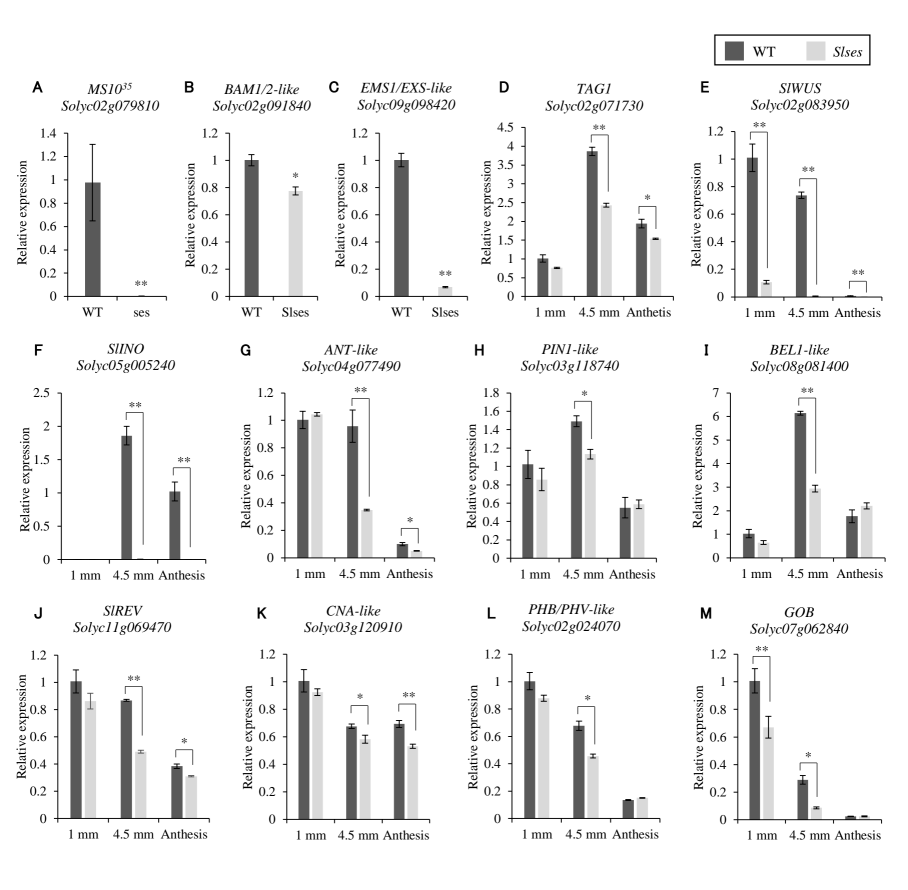
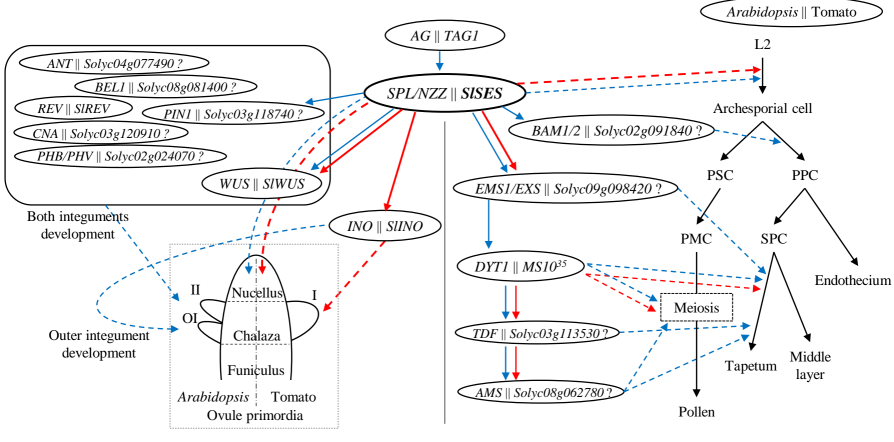


Figure 7



Ovule development

Anther development