1	Alien genes introgression and development of alien monosomic addition lines
2	from a threatened species, Allium roylei Stearn, to Allium cepa L.
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4	Hoa Q. Vu ¹ , Yasuyuki Yoshimatsu ² , Ludmila I. Khrustaleva ³ , Naoki Yamauchi ^{1, 2} , and
5	Masayoshi Shigyo ^{1, 2, *}
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7	¹ The United Graduate School of Agricultural Sciences, Tottori University, 4-101
8	Koyama-Minami, Tottori 680-8553, Japan; ² Faculty of Agriculture, Yamaguchi
9	University, 1677-1 Yoshida, Yamaguchi 753-8515, Japan; ³ Timiryazev Agricultural
10	Academy, Timiryazev Street 44, Moscow, Russia; *Corresponding author, E-mail:
11	shigyo@yamaguchi-u.ac.jp
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1 Abstract To produce alien monosomic addition lines (AMALs) of Allium cepa (genomes CC, 2n=2x=16) carrying extrachromosomes from Allium roylei (RR, 2n = 2x = 16), 2 3 reciprocal backcrossing of allotriploids (2n = 24, CCR) with diploids (2n = 16, CC) and 4 selfing of a single allotriploid were carried out. The chromosome numbers in the BC_2F_1 5 and BC₁F₂ progenies ranged from 16 to 32. Forty-eight plants were recorded to possess 6 2n = 17 among a total of 169 plants in observation. Through the analyses of isozymes, 7 expressed sequence tag (EST) markers, and karyotypes, all eight possible types of A. 8 cepa - A. roylei monosomic addition lines (CC+1R - CC+8R) could be identified. Seven 9 types of representative AMALs (without CC+2R) were used for the GISH analysis of 10 somatic chromosomes. Except for CC+6R, all AMALs showed an entire (unrecombined) 11 extrachromosome from A. roylei in the integral diploid background of A. cepa. A single 12 recombination between A. cepa and A. roylei was observed on the extrachromosome in 13 the remaining type. All alloplasmic AMALs possessing A. roylei cytoplasm showed high 14 or complete pollen sterility. Only the autoplasmic CC+4R with A. cepa cytoplasm 15 possessed relatively high pollen fertility. The bulbs of CC+4R displayed the distinct 16 ovoid shape that discriminates them from spherical or oval ones in other AMALs. Downy 17 mildew screening in the field showed higher resistance in A. roylei, a hypo-allotriploid (CCR-nR, 2n = 23), and an allotriploid (CCR, 2n = 24). Meanwhile, no complete 18 19 resistance was found in some AMALs examined. This was the first trial toward the 20 establishment of a complete set of A. cepa - A. roylei monosomic additions.

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1 Introduction

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3 Allium cepa L. is one of the most economically important species in the section Cepa of 4 the genus Allium (Brewster 2008). It contains two main groups: the Common onion group 5 and the Aggregatum group. The Common onion group includes the bulb onion, which is 6 the most important Allium vegetable crop grown worldwide, with a long cultivation 7 history dating back to approximately 3200 B.C. (McCollum 1976). Meanwhile, the 8 shallot is the most important vegetable and spice crop of the Aggregatum group (Fritsch 9 and Friesen 2002) and is cultivated mainly in low-latitude areas (Siemonsma and Piluek 10 1993). Up to now, onion breeding has focused on the F_1 hybrid seed production in most 11 parts of the world because of its many favorable characteristics, such as male sterility, 12 resistance to bolting and bulb splitting, high yield, and multiple types of bulb shapes and 13 skin colors (Dowker 1990). On the contrary, crossbreeding of the shallot has been poorly 14 examined because of its main propagation system based on division. Most probably, A. 15 cepa itself does not carry resistance genes against modern diseases and pests or other 16 interesting traits, e.g., a vigorous rooting system (Shigyo and Kik 2008). The exploitation 17 of wild species through interspecific hybridization has, therefore, been recommended 18 for introducing novel characteristics, especially those that are limited in the A. cepa gene 19 pool (Kik 2002).

Researchers are now interested in *Allium roylei* with a potential gene reservoir, as this wild species proved to be completely resistant to downy mildew (*Peronospora destructor*) (Kofoet et al. 1990) and partially resistant to leaf blight (*Botrytis squamosa*) (De Vries et al. 1992b) and *Fusarium* basal rot (Galván et al. 2008). This species is

1 considered as a species from the primary (Shigyo and Kik 2008) or secondary (Shigyo 2 2007) gene pool of A. cepa based on the gene pool concept proposed by Harlan and De 3 Wet (1971). Interspecific hybrids between A. roylei and A. cepa and its backcrosses to A. 4 cepa were first recorded by Van der Meer and De Vries (1990). In a study involving a 5 crossing experiment, only a slight isolation barrier was found between A. cepa and A. 6 roylei (Van Raamsdonk et al. 1992). A close relationship between these two species was 7 also demonstrated by meiotic observations of their F_1 hybrid (De Vries et al. 1992a). 8 Furthermore, A. roylei could be used as the bridging species to introduce the genes from 9 A. fistulosum into A. cepa genomes (Khrustaleva and Kik 2000). Using the tri-hybrid 10 population derived from this bridge-cross, Galván et al. (2011) recently found two 11 quantitative trait loci (QTLs) from A. roylei located on chromosomes 2 and 3 contributing 12 to mycorrhizal response. Vu et al. (2011) reported the substitution of A. roylei cytoplasm 13 for the production of novel alloplasmic male sterile lines in A. cepa via the use of their 14 amphidiploid.

15 An alien monosomic addition line (AMAL), which is a line with an extrachromosome 16 from a related species, can improve the process of genetic introgression from donor 17 species into recipient species through the production of chromosome substitution and 18 translocation lines (Singh 2003). In Allium, a combination of A. fistulosum and A. cepa 19 has been reported for AMAL production (Peffley et al. 1985; Shigyo et al. 1996; Hang et 20 al. 2004a). The AMALs of Allium crops could be used as the tools for the chromosomal 21 assignment of several morphological characteristics (Shigyo et al. 1997b) and 22 phytochemical phenotypes (Shigyo et al. 1997a; Hang et al. 2004b; Dissanayake et al. 23 2008; Yaguchi et al. 2008, 2009a, b; Masamura et al. 2011) as well as the assignment of

1 linkage groups in the donor species on its specific chromosomes (Van Heusden et al. 2 2000b; Martin et al. 2005; Tsukazaki et al. 2008, 2011). On the other hand, an increment 3 of the combination number for AMAL development seems to provide effective 4 alternatives to extend the genetic variability of cultivated Allium species. In a previous 5 study (Vu et al. 2011), a successful result was reported in the backcrossing of A. cepa 6 diploid (genomes CC) with A. cepa - A. roylei allotriploid (CCR, 2n=3x=24). Plenty of 7 hyper-diploid plants with 2n = 17, which may possess an alien extrachromosome from A. 8 roylei, could be observed in the BC₂ progeny derived from this cross combination. The 9 aims of the present study were to select a complete set of A. cepa - A. roylei monosomic 10 addition lines from the BC₂ progeny via the use of molecular marker and cytogenetic 11 techniques and to find some favorable agronomic traits such as male sterility or downy 12 mildew resistance in this set. 13 14 Materials and methods 15 16 Crossing procedure for production of monosomic addition lines 17

The plant materials were *A. roylei* (RR, 2n = 2x = 16) and two groups of *A. cepa* (CC, 2n = 2x = 16): shallot (Aggregatum group) and bulb onion (Common onion group). In a previous study, a capital letter "A" was used to indicate the shallot chromosome (Shigyo et al. 1996). However, the "C" used for the onion chromosomes can also be used for those of shallot (De Vries 1990). Therefore, a capital "C" was used for both shallot and bulb onion chromosomes in this study. To produce *A. cepa - A. roylei* monosomic

1 addition lines, an interspecific F₁ hybrid was obtained from a cross between A. roylei '97175' and shallot '86208' at first (Fig. 1). After doubling the chromosomes of the 2 3 clones derived from this single F_1 hybrid, two amphidiploids (plant codes: '28-1' and '33', CCRR, 2n = 4x = 32) were obtained. These two amphidiploids (as seed parents) 4 5 were backcrossed with a late bolting bulb onion unknown variety, 'Banchusei', and an 6 asexually propagated shallot clone, 'Chiang Mai' (as pollen parents), to produce BC₁ 7 progenies as allotriploids (CCR, 2n = 3x = 24), i.e., 'CM26' and 'CM23', respectively. 8 Backcrosses with A. cepa or selfings were made to obtain BC₂ or BC₁F₂ populations that 9 were named as BR, SR, RB, and SY (see Fig. 1). The parts surrounded by the frame in 10 this figure were carried out in this study. All processes of the crossings and raising 11 seedlings were carried out according to Vu et al. (2011). 12 13 Chromosome observation and karyotype analysis

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All plants of BR, SR, RB, and SY lines were used to determine chromosome number.
Somatic chromosomes of root tips were observed by Feulgen nuclear staining followed
by the squash method. The karyotype analyses were according to the standard
nomenclature system for the chromosome of *Allium* (Kalkman 1984), which was
generally agreed upon at the Eucarpia 4th *Allium* Symposium (De Vries 1990).

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- 1 Characterization of alien chromosome based on isozyme and EST markers
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3 Chromosome constitutions of the plants in BC₂ and BC₁F₂ progenies, which possess the 4 chromosome number 2n = 17, namely alien monosomic addition lines (AMALs), were 5 further characterized using isozyme and EST markers. Van Heusden et al. (2000b) 6 reported that isozyme loci Lap-1, Pgm-1, and Pgi-1 are respectively located on 1R, 4R, 7 and 5R in A. roylei. The characterization of extrachromosomes from A. roylei was, 8 therefore, based on the analyses of these three isozyme loci. The extraction of enzymes, 9 electrophoresis, and staining were carried out following the method of Shigyo et al. 10 (1995a, b) and Van Heusden et al. (2000b).

11 Because there is a close genetic relationship between A. roylei and shallot, we 12 assumed the chromosomal locations of ten EST markers in A. roylei from a 13 homoeologous chromosome of the shallot chromosome to which the shallot allele 14 corresponding to each of the markers had been assigned (Table 1). To find the 15 chromosomal locations in A. cepa of the three specific markers CA4H, TC2418, and 16 TC1088, the analyses of these markers were carried out in a complete set of A. fistulosum 17 - shallot monosomic addition lines produced by Shigyo et al. (1996). For EST marker 18 analyses of the A. cepa - A. roylei monosomic additions as well as the complete set of A. 19 fistulosum - shallot monosomic additions, the total genomic DNA of the parental plants 20 and AMALs was isolated from fresh leaf tissue using a mini-prep DNA-isolation method 21 (Van Heusden et al. 2000a). The reaction mixture contains 20 ng template DNA, 1 x Ex 22 buffer, 0.2 mM dNTPs, 0.5 µM forward primer, 0.5 µM reverse primer, and 0.25 units Ex 23 Taq in a total volume of 10 μ l. All PCRs were performed as follows: initial denaturation

1 for 3 min at 94°C and 35-40 cycles of PCR amplification (1 min denaturation at 94°C, 1 2 min annealing at 60-79°C, and 1 min primer extension at 72°C) on a Program Thermal Cycler iCycler[™] (Bio-Rad, Hercules, CA, USA). The number of cycles in PCR 3 4 amplification and the annealing temperature were optimized for each EST marker. The 5 ramp times were carried out under default conditions with adjusted temperatures at the 6 maximum ramp rate with minimum ramp time. The PCR products were separated by 2%7 agarose electrophoresis or 5% polyacrylamide gel electrophoresis (PAGE) according to 8 the method of Yaguchi et al. (2009a). If they were monomorphic after denaturing PAGE, 9 they were subjected to restriction analysis. Five microliters of PCR products was 10 incubated for 24 h at 37°C in a total volume of 10 µl using 2 U of restriction enzyme and 11 subsequently resolved by 2% agarose gel electrophoresis or PAGE. Restriction enzymes 12 AluI (for the markers TC131, ACAHN07, and ACAHM36), AfaI (for CA4H and 13 TC1088), MboI (for MSPS), HinfI (for TC2418 and DFR), and TaqI (for FLS) (Toyobo, 14 Osaka, Japan) were used in an attempt to generate polymorphism between A. cepa and A. 15 roylei. Three restriction enzymes XspI (for CA4H), MboI (for TC2418), KpnI (for 16 TC1088) were used to reveal polymorphism between shallot and A. fistulosum.

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18 Bulb morphology assessment

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After harvesting, the bulb morphologies of some lines in each of eight AMALs were evaluated. The maximum horizontal and vertical diameters of the bulbs were measured, bulb weights were recorded, and bulb colors were noted.

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1 GISH analysis

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The procedure of GISH analysis was according to the method of Khrustaleva and Kik
(2000) with minor modifications.

Seven individual plants of *A. cepa - A. roylei* AMALs (CC+1R, +3R, +4R, +5R, +6R,
+7R, and +8R) were used for GISH analysis. Total genomic DNA was extracted from 4 g
of young leaves from shallot 'Chiang Mai' and *A. roylei* '97175'. The extracted DNA of *A. roylei* was labeled with digoxigenin-11-dUTP by a standard nick-translation protocol
(Roche Diagnostics GmbH, Mannheim, Germany). The extracted DNA of shallot was
used as a blocking DNA.

The hybridization mixture contained 50% (v/v) deionized formamide, 10% (w/v) sodium dextran sulfate, 10% (v/v) 20x SSC, 0.25% (w/v) SDS, 1 ng/µl digoxigeninlabeled DNA of *A. roylei*, and 0.06-0.08 µg/µl blocking DNA of shallot. Digoxigeninlabeled DNA was detected with anti-digoxigenin-FITC raised in sheep (Roche, Mannheim, Germany), then amplified with anti-sheep-FITC raised in rabbit (Vector Laboratories, Burlingame, USA), and finally amplified with anti-rabbit-FITC raised in goat (Vector Laboratories, Burlingame, USA).

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19 Meiotic observation and pollen fertility test

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Acetocarmine smears of pollen mother cells (PMCs) were used for meiotic studies. Fresh pollen grains were stained with 1% acetocarmine to observe their viability. The evaluation of pollen fertility was performed according to Shigyo et al. (1999). The pollen fertility was checked during the first or second week after blooming of the first floret in
 each flower umbel.

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4 Downy mildew screening in field

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6 A test on downy mildew resistance for some AMALs and two multiple alien addition 7 lines from RB lines was carried out in the experimental field of Takii & CO., LTD., in 8 Konan City, Shiga Prefecture, Japan (N35°00['], E136°05'). The bulbs were sown in pots, 9 and germinated plants were then transplanted to the infected field in the beginning of October 2009. Spores were collected from the surface leaves of artificially inoculated 10 11 plants with a fine paintbrush and suspended in distilled water with a final concentration of 10³ spores/ml. A spore suspension was sprayed over the plants for six times in 2010 12 (on 5, 17, and 23 March and on 5, 8, and 20 April). The first day of sporulation for each 13 14 inoculated plant was recorded. Symptoms on plant leaves were observed in the field 15 during the growing period.

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- 17 **Results**
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19 Production of alien monosomic addition lin	nes
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In backcrossing and selfing of allotriploids (CCR, 2n = 24) to produce BC₂ (RB, SY, and BR lines) and BC₁F₂ (SR line), seed setting could be obtained and most of the seedlings survived (Table 2). All plants of BC₂ and BC₁F₂ progenies were used for the 1 chromosome count. Their chromosome numbers are shown in Table 3. The chromosome 2 number of BC₂ plants varied from 2n = 16 to 24. On the other hand, in the BC₁F₂ 3 progeny, the chromosome number ranged from 2n = 16 to 32. In total, the highest 4 frequency of plants was found with chromosome number 2n = 16 (110 plants), followed 5 by 2n = 17 (48 plants). Other chromosome numbers, 2n = 18, 19, 22, 23, 24, 28, and 32, 6 were recorded with a low number of plants (one to three in each case). Finally, all 48 7 plants recorded as 2n = 17 were used for further characterization of their chromosome 8 constitution.

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10 Identification of alien monosomic additions

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12 The analyses of karyotypes, isozymes, and EST genetic markers were carried out to identify the extrachromosomes of A. roylei in plants with 2n = 17 that had been produced. 13 14 As the chromosome 6 of A. roylei showed the most distinctive sub-telocentric shape in its 15 basikarvotype (Sharma and Gohil 2008), the AMALs possessing this chromosome could 16 easily be recognized by conventional karyotype analysis. However, the identification of 17 the other AMALs by karyotyping was not always easy. Therefore, the chromosome 18 constitutions of all AMALs were confirmed further by molecular markers. Every plant of 19 48 AMALs was analyzed with at least one chromosome-specific marker (isozyme or EST 20 markers) for each of eight different chromosomes derived from A. roylei (1R-8R).

The three isozymes could show the clear polymorphism between the parental lines [*A*. *cepa* (shallot and bulb onion) and *A. roylei*]. Therefore, they could be adopted for determining the presence of specific chromosomes of *A. roylei* in the BC₂ and BC₁F₂ progenies. The F₁ hybrid, amphidiploid, and allotriploid plants showed both bands of *A*. *roylei* and *A. cepa* for the locus *Lap-1* (Fig. 2a). An additional band at intermediate
mobility between the parental bands was found for the locus *Pgi-1* (Fig. 2b). The
summary of isozyme analysis in AMALs was included in Table 4. With the isozyme
markers *Lap-1* and *Pgi-1* the AMALs CC+1R and +5R were characterized.

6 The three EST markers CA4H, TC2418, and TC1088 were, respectively, determined 7 to be located on the chromosomes 1, 5, and 6 of *A. cepa* via the use of the complete set of 8 *A. fistulosum - A. cepa* monosomic additions (Fig. 3). These three markers were, 9 therefore, included in EST marker analyses for identification of extrachromosome in *A.* 10 *cepa - A. roylei* monosomic additions in this study.

11 All EST markers, except for the SiR marker on chromosome 3, appeared as one 12 single PCR product on 2% agarose gel in both A. roylei and A. cepa, and no 13 polymorphism was detected. For the SiR marker, two PCR products were observed in A. 14 roylei, from which the size of one product is distinguishable from one product of A. cepa. 15 For the monomorphic markers, an interspecific polymorphism was generated using 16 restriction enzymes to show polymorphisms between A. cepa and A. roylei. The DNA fragments derived from A. roylei were used to confirm the presence of A. roylei 17 18 respective chromosomes (Fig. 4). The results of EST-marker analyses in AMALs were 19 summarized in Table 4. All EST markers, except for TC1088, were present in one or 20 more AMALs. Seven AMALs (CC+1R-5R and CC+7R-8R) were identified by EST 21 markers. Although the marker TC1088 for chromosome 6 was not detected in any AMAL, 22 two AMALs, RB82 and RB120, were determined to possess three sub-telocentric 23 chromosomes (homoeologous chromosome group 6) by karyotype analysis. RB82,

however, also showed the presence of the marker for chromosome 7 of *A. roylei*, and its
 chromosome constitution was determined as unidentified. RB120 was confirmed by the
 following GISH analysis. Figure 5 shows representative somatic metaphase
 chromosomes of a complete set of AMALs.

5 Finally, this study demonstrated that the production of a complete set of AMALs was 6 not impossible in the species combination of A. cepa (recipient) and A. roylei (donor). 7 The chromosome constitutions of 13 plants could not be identified due to the absence of 8 all markers (11 plants) or the presence of more than one different chromosome-specific 9 markers (two plants). Among AMALs of both RB and SY lines, the highest frequency of 10 plants was found in CC+8R (RB: 34%, SY: 38%), followed by CC+7R (13%, 25%) and 11 CC+3R (8%, 13%). The other AMALs, CC+1R, +2R, +5R, and +6R, appeared with 12 lower percentages. All eight possible AMALs, except for CC+4R, were obtained in RB 13 and SY lines. However, one plant of CC+4R appeared in BR lines when the allotriploid 14 was used as the pollen parent.

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16 GISH analysis of alien monosomic additions

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The GISH analyses included seven individual plants, RB34, SY45, BR1, SY7, RB120, SY24, and RB66, which were determined previously through karyotype and molecular markers as AMALs CC+1R, +3R, +4R, +5R, +6R, +7R, and +8R, respectively (Fig. 6a). All types of the examined AMALs, except for CC+6R, showed an intact chromosome from *A. roylei* as a single chromosome addition with a complete set of 16 chromosomes from *A. cepa* without any translocation. This result confirmed the integrity of the additional chromosome from *A. roylei* in each of these six AMALs. One exception,
RB120, showed a single recombinant chromosome 6R' with segments derived from both *A. roylei* and *A. cepa* in an integral diploid back ground of *A. cepa*. The green
fluorescence segment derived from *A. roylei* occupied the terminal half of the long arm of
the chromosome 6R'.

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7 Meiosis in alien monosomic additions

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9 Five different AMALs derived from five individual plants comprising four plants of SY7 10 (CC+5R), RB120 (CC+6R'), SY24 (CC+7R), and RB66 (CC+8R) that were previously 11 used for GISH analysis and plant SY42 (CC+1R) were further investigated for their 12 meiotic behavior during flowering time of 2011 (Fig. 6b). Meiosis of the AMALs CC+2R, +3R, and +4R was not observed because some plants of these lines did not develop 13 14 flower stalks in this year, and the collection time of flower buds from the other plants was 15 late to obtain PMCs at metaphase-I (MI). In the four AMALs, CC+1R, +5R, +7R, and 16 +8R, a high percentage of PMCs at the MI of meiosis formed eight bivalents and one 17 univalent; and the cells with seven bivalents and one trivalent also appeared, but with 18 limited numbers ranging from 6.7 to 24.3% (Table 5). However, CC+6R' showed eight 19 bivalents and one univalent or seven bivalents and one trivalent of chromosome 6 (Fig. 20 6b), with the possibility of around 50% for each case (Table 5).

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A total of the 24 plants that came into bloom in the spring of 2007-2010 were checked for
their pollen fertility (Table 6). All plants except BR1 (CC+4R) showed very low pollen
fertility or completely sterility (0-4.3%). The CC+4R showed a 42.6% of the pollen
fertility. This plant is an autoplasmic line derived from the cross when a bulb onion was
used as a seed parent. Meanwhile, all other plants are alloplasmic lines carrying
cytoplasm from *A. roylei*.
Bulb morphology of alien monosomic additions

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12 Bulb characteristics including shape, color, and weight were investigated with several 13 types of AMALs. Shallot showed spherical bulbs with purple outer scale, while A. roylei 14 displayed ovoid bulbs and brown outer scale. Therefore, the introgression of single 15 chromosome from A. roylei might actually alter the morphological features on bulbs in 16 the recipient species. The bulb of BR1 (CC+4R) was distinct from the others and had an 17 ovoid shape and brown skin color (Table 7, Fig. 6c). The bulbs of CC+2R, +3R, +5R, and +8R were spherical, whereas those of CC+1R, +6R', and +7R, were oval (Fig. 6c). 18 19 The red purple color of the outer bulb skin was observed in CC+2R, +3R, +5R, and +7R. The other three AMALs, CC+1R, +6R', and +8R, were light purple. The heaviest bulb 20 21 was found in CC+4R (Table 7).

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3 Fourteen plants of six AMALs [CC+1R (one plant), +3R (two), +4R (one), +5R (one), 4 +7R (four), and +8R (five)], one plant of hypo-allotriploid, RB35 (2n = 3x-1 = 23, CCR-5 8R), and one plant of allotriploid, RB51 (2n=3x=24, CCR), were used to screen for 6 resistance to downy mildew in the infected field (Table 8). The screening test for CC+2R 7 and +6R were not carried out due to insufficient number of plant materials. Seven plants 8 (one from CC+3R, four from CC+7R, and two from CC+8R) showed an early sporulation 9 after the first inoculation of spores. Six other plants (one plant from CC+4R, one from 10 CC+5R, two from CC+8R, and two from the allotriploids) showed a late sporulation after 11 four cycles of inoculation. Furthermore, late sporulation (after six inoculation cycles) was 12 observed in three remaining plants, each from the CC+1R, +3R, and +8R. However, after 13 six cycles of inoculation, all plants of the AMALs had severe disease symptoms and died 14 at the end of the growing period. On the other hand, light and moderate symptoms 15 appeared in the hypo-allotriploid, RB35, and the allotriploid, RB51. These two lines 16 could survive until a final phase.

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18 Discussion

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In diploid recipient species, AMALs had thus far been obtained in 20 different species combinations, including that of the present study (Singh 2003). However, all the possible types of AMALs were found in only eight cases, while, in others, a complete set was lacking. The largest holders were *Beta* with three cases, *Allium* with two, and the
 remaining three species (*Lycopersicon*, *Nicotiana*, and *Oryza*) with one.

3 In Allium, Peffley et al. (1985) reported four types out of the A. cepa - A. fistulosum 4 monosomic additions derived from a cross between allotriploid (seed parent) and bulb 5 onion. They assumed that hyperploid zygotes or seedlings may possess a low survival 6 rate, which resulted in few progeny plants with extrachromosomes. It was also difficult to 7 complete eight possible shallot (A. cepa Aggregatum group) - A. fistulosum monosomic 8 additions (Hang et al. 2004a). They suggested that the possible existence of deleterious 9 alleles that are maintained in the heterozygous state remaining in shallot genomes may 10 reduce the survival rate of progeny plants homozygous for these alleles. In this study, we 11 produced A. cepa - A. roylei monosomic additions through three different cross 12 combinations. In 2005, a pilot crossing, allotriploid CCR (\bigcirc) x bulb onion, had been 13 carried out at first in an experimental field at Yamaguchi University (lat. 34°N, long. 14 131°E), being less than successful due to an insufficient number of the obtained progeny. 15 A sufficient number of the BC₂ progenies (RB and SY) could be surprisingly obtained in 16 the same way for crossing at Hokkaido (44°N, 142°E) in 2006. This could be interpreted 17 assuming that the seed and pollen fertilities of A. roylei and its derivatives vary 18 considerably with latitude. Furthermore, AMALs appeared with obviously high 19 frequency (28.8%) in these progenies. This result shows a good agreement with that of 20 Peffley et al. (1985). Multiple crossings at high latitude regions seem to be a valid and 21 feasible way to produce monosomic addition lines of A. cepa - A. roylei although those of 22 A. fistulosum - shallot were not the case (Shigyo et al. 1996). On the other hand, all types 23 of AMALs, excepting CC+4R, were developed from RB and SY. One possibility for the

1 absence of CC+4R is that a relatively low number of AMALs (46 plants) in RB and SY 2 lines were produced. Actually, we could find the presence of chromosome 4R via Pgm-13 marker in some multiple addition lines (2n=19, 22, 23, 24) of BC₂ plants in RB and SY 4 (data not shown). In addition, it is well possible that the apperance frequency differs 5 between chromosomes. Shigyo et al. (1996) reported that the appearance frequencies 6 varied largely among eight types of A. fistulosum - shallot monosomic additions. The 7 authors also suggested that the extrachromosomes might affect the survival of AMALs. 8 While, one plant of CC+4R was obtained via a special cross combination between the 9 bulb onion 'Sapporo-Ki' (seed parent) and the allotriploid. Therefore, the other 10 possibility might be that the transmission rate of chromosome 4R through male gametes 11 was higher than that through female gametes. The maternal transmission of alien 12 chromosomes from a donor into a recipient is generally higher than the paternal one in 13 many crops (Multani et al. 1994; Singh et al. 1998; Akaba et al. 2009). However, in some 14 special cases, the preferential transmission through male gametes has been reported in 15 some specific chromosomes of wild species, such as chromosome 5H^tL in the 16 transmission from Elymus trachycaulus to common wheat (Jiang and Gill 1998) or 17 chromosome 5R from rye to wheat (Efremova et al. 1996). To explain the phenomenon in 18 our study, it is necessary to investigate the occurrence rate of CC+4R in a larger number 19 of BC₂ plants from reciprocal backcrosses.

The modifications of morphological characteristics in AMALs are mostly due to the interaction between genes of the recipient and donor parents (Singh 2003). It has been reported that the traits of color, shape, and size of bulbs in *A. cepa* are quantitatively inherited (El-Shafie and Davis 1967; McCollum 1971). McCollum (1966) suggested the

1 genetic determiners of the bulb height and shape index (the ratio of bulb height to bulb 2 diameter), which might be viewed as two measures of the same genetic traits. On the 3 other hand, bulb diameter and bulb weight were affected by the environmental conditions. 4 The distinct ovoid bulbs of CC+4R, resembling those of A. roylei, may suggest that some 5 major genes related to this bulb feature are located on the chromosome 4 of A. roylei and interact with the genes in A. cepa genomes to define the final trait of this AMAL. Either 6 7 way, it is possible to identify at least CC+4R morphologically. In this study, isozyme and 8 EST markers, the chromosomal locations of which had been reported in A. cepa (Shigyo 9 et al. 1995a, b; Martin et al. 2005; Masuzaki et al. 2006; McCallum et al. 2007; Yaguchi 10 et al. 2008, 2009b) and A. fistulosum (Yaguchi et al. 2009a), were applied for the 11 characterization of extrachromosomes from A. roylei in the AMALs. The result of marker 12 analyses corresponded well with that of the karyotypic observation, except for TC1088 13 which was not present in the translocation line CC+6R'. Three of the seven isozyme 14 markers employed for the correspondence of paternal (A. roylei) and maternal (A. cepa) 15 linkage groups in the previous study (Van Heusden et al. 2000b) were presently used to 16 identify A. roylei alien chromosomes in an A. cepa diploid background. The universal 17 chromosome-specific markers in Allium could accurately detect the homoeologous 18 chromosomes of A. roylei as well as the two cultivated species in alien addition lines. The 19 six lines possessing each extrachromosome of 1R, 3R, 4R, 5R, 7R, and 8R were proven 20 to be true AMALs through GISH. In these AMALs, the majority of PMCs showed 21 preferential parings between homologous chromosomes of A. cepa, leaving the 22 chromosomes of A. roylei as univalent. On the other hand, the presence of trivalents 23 between chromosomes of the two species in these lines showed a different phenomenon

1 from A. fistulosum - A. cepa monosomic additions reported by Shigyo (1997c), in which 2 only bivalents and univalents were observed. This result suggested that A. roylei 3 possesses a higher degree of homoeology with A. cepa than A. fistulosum. The 4 translocation was observed in only one AMAL with a 6C-6R recombinant chromosome 5 in the integral diploid background of A. cepa. The recombination point might have been located at the interstitial region. This translocation line probably resulted from chiasma 6 7 formation among chromosome 6 of A. cepa and A. roylei during meiosis of the 8 allotriploid 'CM23' used in backcrossing. This line showed nearly an equal rate of PMCs 9 with the univalent and trivalent, suggesting that the recombinant chromosome 6C-6R has 10 high homology with the chromosome 6C of A. cepa. In the present study, the integrated 11 applications of the three different methodologies enabled us to clarify chromosomal 12 constitutions of our AMALs.

13 In our previous study, a complete male sterility was recorded in diploid A. cepa 14 harboring cytoplasm from A. roylei or alloplasmic lines (Vu et al. 2011). It was suggested 15 that the sterility would be caused by incompatibility between the nuclear genomes of A. 16 cepa and the cytoplasm of A. roylei. The data of this study would, therefore, indicate the 17 effect of A. roylei cytoplasm on the pollen fertility of the AMALs. Tsutsui et al. (2011) 18 newly found stable male sterility in alloplasmic AMALs of *Brassica rapa* carrying the 19 cytoplasm and an extrachromosome from Moricandia arvenis. The probable presence of 20 its fertility restoring gene could not be proved since all of the alloplasmic AMALs 21 carrying cytoplasm from A. roylei showed an adequate level of male sterility. The 22 restoring genes of pollen fertility were found in some specific types of alloplasmic 23 Moricandia arvenis - Raphanus sativus monosomic additions (Bang et al. 2002) or in the syntenic group 6 of alloplasmic *Brassica campestris - B. oxyrrhina* monosomic additions
(Srinivasan et al. 1998). Relatively high pollen fertility was only found in the autoplasmic
CC+4R, likely because of the lack of effect of the *A. roylei* cytoplasm. The direct
introduction of the 4R chromosome to the alloplasmic line might be possible through a
crossing with the CC+4R as a pollen parent. If a fertility-restoring gene existed on this
chromosome, the restoration of pollen fertility would be clearly observed in the
alloplasmic CC+4R.

8 The resistance of A. roylei to downy mildew (P. destructor) in greenhouse and field 9 conditions is well-known (De Vries et al. 1992a). Recently, its resistance gene allocated 10 on the chromosome 3 was transferred successfully into a bulb onion, and a disease-11 resistance test was carried out in a trial field in Europe (Scholten et al. 2007). In this 12 study, we carried out the disease-resistance test under field conditions in Japan with A. 13 roylei and the A. cepa - A. roylei chromosome addition lines. From the results of downy 14 mildew screening, no complete resistance was found in any of the tested plants. In this 15 study, even A. roylei showed the symptoms with sporulation on the leaves. This finding 16 was irreconcilable with the results of the above two articles which reported a complete 17 resistance in A. roylei. It has been reported that disease resistance in various plants is 18 affected by environmental conditions as well as pathogen populations. In maize, the 19 resistance inheritance to downy mildew (Peronospora sorghi) was affected not only by 20 certain conditions (Williams 1984) but also by regions and pathogenic differences 21 (Singburaudom and Renfro 1982). As mentioned in a previous report (Schwartz 2008), a 22 high-humidity environment, as in Japan, seems to be more favorable for downy mildew 23 development than a relatively dry environment, as in Europe. One possible hypothesis for

1 this phenomenon is that the susceptibility was due to the severe condition of the disease-2 resistance test under high humidity at the field in Japan. In addition, Japanese fungal 3 strains might be different from European ones. All the AMALs including those of CC+3R were completely susceptible. This result is in contrast to the result of Scholten et 4 5 al. (2007) who used GISH to show that a single introgression of A. roylei on chromosome 6 3 was enough for resistance to downy mildew. However, the light-to-moderate symptoms 7 and the survival of A. roylei, the single deletion, and the multiple addition lines indicated 8 that they had higher levels of resistance than those of all the tested AMALs. Among the 9 AMALs, the plants of CC+1R (RB34), CC+3R (RB86), and CC+8R (RB85) may show 10 higher resistance than those of the others, because they showed the latest sporulation. 11 This result may indicate the existence of some genes responsible for disease resistance 12 located on some chromosomes of A. roylei, which is in the contrast to the conclusion of 13 previous studies that the resistance is based on one, dominant gene. Sabry et al. (2006) 14 found that one locus on chromosome 2 of maize had a major effect and was associated 15 with downy mildew resistance in all test environments. Moreover, they found two 16 additional QTLs on chromosomes 3 and 9 of maize showing minor additive effects on 17 resistance, but only in a specific environment. The present study would raise the question 18 about whether the other genes exist on the other chromosomes of A. roylei, which might 19 show additive effects with the resistance gene previously found on chromosome 3 under 20 the field condition of Japan. Another issue that needs to be considered is the plants of the 21 same AMAL type, such as CC+3R or CC+8R, showed different appearance of 22 sporulation. One possibility might be derived from the discrete segregations of the several genetic factors related to their resistibility in the diploid background of these two
 AMAL types.

3 In this study, a complete set of seven A. roylei whole chromosomes and one 4 recombinant chromosome, 6C-6R, were added to A. cepa. Even when the completion of 5 possible eight AMALs proves impossible, assignment of DNA markers to A. roylei 6 chromosomes can be done using the DNA fingerprints of our incomplete set. This set 7 would be helpful in the construction of a genetic map in A. roylei, which serves as the 8 potential tool for its genetic studies. Furthermore, the complete set of AMALs is an 9 invaluable material that makes it easier to localize loci responsible for the interested traits 10 in breeding program, as the whole donor genome is divided into individual 11 extrachromosomes adding to the genetic background of the recipient species. It, therefore, 12 would enable rapid introgression of desired alien genes from A. roylei, a currently 13 threatened species, into A. cepa.

14 At present, the set of AMALs has been maintained vegetatively in our laboratory. To 15 avoid high risk of losing materials via vegetative propagation, it is essential to find a way 16 of the permanent maintenance of our AMALs for future verification in several other 17 institutes. Shigyo et al. (1998, 2003) reported that another useful method for maintaining 18 a complete set of A. fistulosum - shallot monosomic additions was seed propagation via a 19 combination of selfings and backcrossings AMALs (seed parent) x A. fistulosum. While, 20 the transmission rates of extrachromosomes of A. cepa - A. roylei monosomic additions 21 will show different tends from those of A. fistulosum - shallot monosomic additions. The 22 detailed analyses of the transmission rates are underway to find the best maintenance

1	procedure of the novel Allium AMALs. In near future, it will be possible for us to provide
2	the seed of the AMALs for scientists as well as breeders all around the world.
3	
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Primer set	GenBank	Forward and reverse primers	Chromosome	Reported	Restriction
	accession No.				enzyme
CA4H	AY541032	5'-TCGGCAACTGGCTCCAAGTC-3'	1	This study	Afa I
		5'-TGATTGACCAGTTCCGCTATGCC-3'			
TC131	AY541031	5'-CCTGCAGAATGAGCTCATC-3'	2	Yaguchi et al.	Alu I
		5'-GAAGTTTCCGCCATGCAC-3'		(2009b)	
ACAHN07	CF443350	Outside 5'-TTGATAACTCCAATGGGTGTAAATGTCC-3'	3	Kuhl et al. (2004)	Alu I
		5'-ATCCTGCTTGGTATCAAGCGGCATGT-3'			
		Inside 5'-GCAAAGAAAGGACTGCTTGTCAATGCT-3'			
~		5'-ATTGCTCGGATCTCTGAGTCCATGTAG-3'			9
S1R	CF434863	5'-TGCAGCTCTTTCTCAAGTTGG-3'	3	McCallum et al.	_"
		5'-CAGAGCAGGACATGCCATAG-3'		(2007)	
FLS	AY221247	5'-TTAAGGACGACCACTGGTT-3'	4	Masuzaki et al.	Taq I
		5'-CCACGACATCCGTGACT-3'		(2006)	
TC2418	CF438524,	5'-CATCAGGAGAAGATATCAATGCTC-3'	5	This study	Hinf I
	CF438854	5'-GCCTGTCTTCACTCCTAGTC-3'			
TC1088	CF451692,	5'-TAACGCCGTCAAACTCTACC-3'	6	This study	Afa I
	CF448264	5'-CTCGGTGGACAACCTTACC-3'			
ACAHM36	CF443252	Outside 5'-TTCCTTGCTCAAGGATACGAGGATG-3'	7	Martin et al. (2005)	Alu I
		5'-GATCATTTCATCGTCATTTGCCTCG-3'			
		Inside 5'-GAAGAGAAGATGCTGGGGAATCTCG-3'			
		5'-GAGCAGATAGCTCTGCAGCACTCTG-3'			
DFR	AY221250	5'-ACAGATGTACTTTGTGTCCA-3'	7	Masuzaki et al.	Hinf I
		5'-GCTTCATCGAACATACTTTCC-3'		(2006)	
MSPS	EU164758	5'-GAAGGCTGATATTGTTGGTGAAG-3'	8	Yaguchi et al. (2008)	Mbo I
		5'-TGTGTCGTAGGAGCCTGATG-3'			

Table 1. EST markers for identification of extrachromosomes from A. roylei

^a No restriction enzyme was used

		Number	Number of	Number of
Cross combination	Line	of seeds	seeds that	seedlings
		produced	germinated	that
				survived
CM23 x 'Kitami Kohai 39 go'	RB	178	130	127
CM23 x 'Sapporo-Ki'	SY	_ ^a	52	52
'Sapporo-Ki' x CM26	BR	3	3	3
CM26 selfed	SR	11	6	6

Table 2. Seed set, seed germination, and number of seedlings survival of allotriploids CM23 and CM26 (CCR) in backcrosses with bulb onion and selfing

Not counted

Line	Number of plants observed	Freque Chron	Frequency of plants Chromosome number									
2		16	17	18	19	22	23	24	28	32		
RB	108	68	38	_a	-	-	1	1	-	-		
BR	3	2	1	-	-	-	-	-	-	-		
SR	6	2	1	-	-	1	-	-	1	1		
SY	52	38	8	1	1	1	1	2	-	-		
Total	169	110	48	1	1	2	2	3	1	1		
a	· 1											

Table 3. Frequency of chromosome numbers in BC_2 (BR, RB, and SY) and BC_1F_2 (SR) progenies

^a Not determined

Line	Group	Frequency of	A. royle	i chromos	ome-speci	fic genetic mar	kers		/								Extrachromo-
		plants	$(1R)^{a}$		(2R)	(3R)		(4R)		(5R)		$(6R)^{b}$		(7R)		(8R)	some
			Lap-1	CA4H	TC131	ACAHN07	SiR	Pgm-1	FLS	Pgi-1	TC2418	Karyotype	TC1088	ACAHM36	DFR	MSPS	_
	1	1	1 ^c	_e	0	0	-	0	-	0	-	0	0	0	-	0	1R
	2	1	0^{d}	-	1	0	-	0	-	0	-	0	0	0	-	0	2R
	3	3	0	-	0	1	-	0	-	0	-	0	0	0	-	0	3R
	4	1	0	-	0	0	-	0	-	1	-	0	0	0	-	0	5R
	5	1	0	-	0	0	-	0	-	0	-	1	0	0	-	0	6R
RB	6	5	0	-	0	0	-	0	-	0	-	0	0	1	-	0	7R
	7	13	0	-	0	0	-	0	-	0	-	0	0	0	-	1	8R
	8	1	0	-	0	0	-	0	-	1	-	0	0	0	-	1	Unidentified
	9	1	0	-	0	0	-	0	-	0	-	1	0	1	-	0	Unidentified
	10	11	0	-	0	0	-	0	-	0	-	0	0	0	-	0	Unidentified
	1	1	-	1	0	-	0	-	0	-	0	-	0	-	0	0	1R
	2	1	-	0	0	-	1	-	0	-	0	-	0	-	0	0	3R
SY	3	1	0	0	0	-	0	0	0	1	1	-	0	-	0	0	5R
	4	2	0	0	0	-	0	0	0	0	0	-	0	-	1	0	7R
	5	1	-	0	0	-	0	-	0	-	0	-	0	-	0	1	8R
	6	2	0	0	0	-	0	0	0	0	0	-	0	-	0	1	8R
BR	1	1		0	0	-	0	-	1	-	0	-	0		0	0	4R
SR	1	1	-	0	0	-	0	-	0	-	1	-	0	-	0	0	5R

Table 4. Identification of extrachromosomes in *A. cepa - A. roylei* monosomic addition lines via chromosome-specific isozyme markers (*Lap-1*, *Pgm-1*, and *Pgi-1*) and EST markers (the others)

^a The symbols inside parentheses show chromosome numbers of *A. roylei* on which the genetic markers are located ^b Chromosome 6 was characterized by karyotype analysis

^c Presence

^d Absence

^e Not conducted

AMALs	Plant code	No. of PMCs	Frequency of PMCs Chromosome pairing				
		observed	8II + 1I	7II + 1III			
CC+1R	SY42	40	33 (82.5) ^a	7 (17.5)			
CC+5R	SY7	37	28 (75.7)	9 (24.3)			
CC+6R'	RB120	41	21 (51.2)	20 (48.8)			
CC+7R	SY24	30	28 (93.3)	2 (6.7)			
CC+8R	RB66	30	26 (86.7)	4 (13.3)			

Table 5. Chromosome configurations in PMCs at MI of *A. cepa - A. roylei* monosomic addition lines

^aNumbers in parentheses indicate percentages

Plant material	CC +	Plant code	Pollen	fertility	(%)						
	extrachromosome		2007		2008		2009		2010		Average
			1^{st}	2^{nd}	1^{st}	2^{nd}	1^{st}	2^{nd}	1^{st}	2^{nd}	
AMALs	CC+1R	SY42	<u>-</u> a	-	-	-	-	-	0.0	0.0	0.0 ± 0.0
	CC+2R	RB54	-	-	2.6	2.6	0.0	-	-	-	1.7 ± 0.9
	CC+3R	RB62	-	-	0.0	0.0	-	-	-	-	0.0 ± 0.0
		RB86	-	-	0.0	-	-	-	-	-	0.0 ± 0.0
	CC+4R	BR1	-	-	42.6	-	-	-	-	-	42.6
	CC+5R	RB109	-	-	0.0	0.0	-	-	-	-	0.0 ± 0.0
	CC+6R'	RB120	-	-	0.0	0.0	-	-	-	-	0.0 ± 0.0
	CC+7R	RB20	-	-	0.0	0.0	0.0	0.0	-	-	0.0 ± 0.0
		RB58	-	-	15.8	1.0	0.1	0.3	-	-	4.3 ± 3.8
		RB78	-	-	0.0	0.0	0.0	-	-	-	0.0 ± 0.0
		RB81	6.2	-	0.4	4.3	0.0	0.0	-	-	2.2 ± 1.3
		RB94	-	-	0.0	0.0	-	-	-	-	0.0 ± 0.0
	CC+8R	RB4	-	-	0.2	0.0	0.3	0.0	-	-	0.1 ± 0.1
		RB18	-	-	-	-	0.8	0.1	-	-	0.5 ± 0.4
		RB42	-	-	0.0	0.0	0.5	0.0	-	-	0.1 ± 0.1
		RB61	-	-	-	-	0.0	-	-	-	0.0 ± 0.0
		RB65	-	-	-	-	0.0	0.0	-	-	0.0 ± 0.0
		RB66	-	-	0.0	0.0	2.7	0.0	-	-	0.7 ± 0.7
		RB68	-	-	0.0	1.5	0.5	0.0	-	-	0.5 ± 0.3
		RB85	-	-	0.0	0.0	0.0	-	-	-	0.0 ± 0.0
		RB96	-	-	-	-	0.0	0.0	-	-	0.0 ± 0.0
		RB100	-	-	0.0	0.0	-	-	-	-	0.0 ± 0.0
		RB114	-	-	0.0	0.3	0.0	-	-	-	0.1 ± 0.1
		RB117	1.3	3.5	0.0	0.0	0.0	-	-	-	1.0 ± 0.7
A. roylei '97175'			-	-	23.0	-	-	-	-	-	23.0
Shallot 'Chiang Mai'			-	-	93.7	-	-	-	-	-	93.7
F_1 hybrid			-	-	0.2	-	-	-	-	-	0.2
Allotriploid 'CM23'			-	-	0.3	-	-	-	-	-	0.3
Allotriploid 'CM26'			-	-	11.5	-	-	-	-	-	11.5

Table 6. Pollen fertility in A. cepa - A. roylei monosomic addition lines

^aNot observed

AMALs	Number of	Number of	Bulb	Bulb height	Bulb weight	Bulb	Bulb shape	Outer scale
	lines	plants	diameter	(mm)	(grams)	diameter/		color
	observed		(mm)			bulb height		
CC+1R	1	2	21.6	33.0	7.3	0.65	Oval	Light purple
CC+2R	1	2	34.9	33.0	20.9	1.05	Sphere	Red purple
CC+3R	3	9	29.2±3.8	28.4±1.5	5.3±0.5	1.04 ± 0.03	Sphere	Red purple
CC+4R	1	2	38.6	60.8	43.4	0.63	Ovoid	Brown
CC+5R	1	4	29.7±1.9	28.2±0.9	13.0±2.3	1.05 ± 0.04	Sphere	Red purple
CC+6R'	1	2	34.1	31.6	17.4	1.08	Sphere	Light purple
CC+7R	5	25	17.4±0.5	25.5±0.6	3.4±0.2	0.69 ± 0.02	Oval	Red purple
CC+8R	10	36	27.0 ± 0.8	29.5±0.7	9.5±0.6	0.92 ± 0.03	Sphere	Light purple

Table 7. Morphological characteristics of bulbs in *A. cepa* - *A. roylei* monosomic addition lines

Plant materials	Chromosome constitution	Plant code	Days of first appearance of sporulation after first inoculation	No. of inoculation cycles before sporulation	Degree of symptom on leaves	Final evaluation
AMALs	CC+1R	RB34	47	6	$++^{a}$	S^d
	CC+3R	RB62	11	1	++	S
		RB86	47	6	++	S
	CC+4R	BR1	32	4	++	S
	CC+5R	RB109	32	4	++	S
	CC+7R	RB58	11	1	++	S
		RB78	4	1	++	S
		RB81	11	1	++	S
		RB94	11	1	++	S
	CC+8R	RB42	32	4	++	S
		$RB65(2)^{f}$	32	4	++	S
		RB68	11	1	++	S
		RB85	47	6	++	S
		RB96	4	1	++	S
Hypo-allotriploid	CCR-8R	RB35 (2)	32	4	$+^{b}$	R ^e
Allotriploid	CCR	RB51	32	4	+_ ^c	R
A. roylei	RR	95001-3 (5) 95001-6 (5)				
		95001-10 (5) 95001-21 (5) 95001-24 (5) 97175-1 (2)	47	6	+- or +	R

Table 8. Downy mildew screening of *A. cepa - A. roylei* monosomic addition lines, hypo-allotriploid, allotriploid, and *A. roylei*

^a Severe symptom ^b Moderate symptom ^c Light symptom ^d Susceptible ^e Resistant ^f Number of bulbs used



Fig. 1 Method for producing alien monosomic addition lines of *A. cepa* with extrachromosomes from *A. roylei*



Fig. 2 *Lap-1* (**a**) and *Pgi-1* (**b**) zymograms in *A. roylei* '97175' (RR), shallot '86208' (CC1), shallot 'Chiang Mai' (CC2), bulb onion 'Kitami Kohai 39 go'(CC3), F₁ hybrid (CR), amphidiploid '33' (CCRR), allotriploid 'CM23' (CCR), AMAL 'RB34' (CC+1R) and AMAL 'RB109' (CC+5R)



Fig. 3 Representative PCR amplification profiles of EST markers CA4H (**a**), TC2418 (**b**) and TC1088 (**c**) after digestion by restriction enzymes on 2% agarose in *A. fistulosum* (FF), shallot 'Chiang Mai' (CC) and a complete set of *A. fistulosum* - shallot monosomic addition lines (FF+1C-FF+8C). *Arrows* indicate chromosome-specific markers of shallot. *M* molecular size marker (100bp ladder)



Fig. 4 Representative PCR amplification profiles of the chromosome-specific EST markers ACAHN07 (3R) (**a**), FLS (4R) (**b**), DFR (7R) (**c**), and MSPS (8R) (**d**) after digestion by restriction enzymes on 2% agarose (**a-c**) and 5% polyacrylamide (**d**) in *A. roylei* '97175' (RR), shallot '86208' (CC1), shallot 'Chiang Mai' (CC2), bulb onion 'Kitami Kohai 39 go'(CC3), BC₂ (RB, SY and BR), and BC₁F₂ (SR).. *Arrows* indicate chromosome-specific markers of *A. roylei*. *M* molecular size marker (100bp ladder)

KNIKNIIK/
)))///////////////////////////////////
2R
X)XXXXXXXX
1C 2C 3C 4C 5C 6C 7C 8C 8R

Fig. 5 Somatic metaphase chromosomes of eight different *A. cepa - A. roylei* monosomic addition lines (CC+1R, RB34; CC+2R, RB54; CC+3R, RB62; CC+4R, BR1; CC+5R, RB109; CC+6R', RB120; CC+7R, RB81; CC+8R, RB68)



Fig. 6 a Somatic metaphase cells of seven types of *A. cepa - A. roylei* monosomic addition lines (1R, 3R, 4R, 5R, 6R', 7R, and 8R) after genomic in situ hybridization. *Arrow* indicates recombinant chromosome 6. **b** Chromosome pairings at MI in PMCs of five types of monosomic addition lines (1R, 5R, 6R', 7R, and 8R). *Black* and *white arrows* indicate univalent and trivalent, respectively. **c** Dormant bulbs of shallot (P1), *A. roylei* (P2) and a set of monosomic addition lines (1-5R, 6R', 7-8R)