Construction of an ultrahigh-density linkage map and graphical representation of the arrangement of transcriptome-based unigene markers on the chromosomes of Allium cepa L.

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Abstract

Background

Genomic information for *Allium cepa* L. is limited as it is heterozygous and its genome is very large. To elucidate potential SNP markers obtained by NGS, we used a complete set of *A. fistulosum* L.-*A. cepa* monosomic addition lines (MALs) and doubled haploids (DHs). These were the parental lines of an *A. cepa* mapping population for transcriptome-based SNP genotyping.

Results

We mapped the transcriptome sequence reads from a series of *A. fistulosum*-*A. cepa* MALs onto the unigene sequence of the doubled haploid shallot *A. cepa* Aggregatum group (DHA) and compared the MAL genotype call for parental bunching onion and shallot transcriptome mapping data. We identified SNP sites with at least four reads on 25,462 unigenes. They were anchored on eight *A. cepa* chromosomes. A single SNP site was identified on 3,278 unigenes and multiple SNPs were identified on 22,184 unigenes. The chromosome marker information was made public via the web database Allium TDB (http://alliumtdb.kazusa.or.jp/). To map these markers, we gathered RNA sequence data from 96 lines of a DHA × doubled haploid bulb onion *A. cepa* common onion group (DHC) mapping population. After selecting co-dominant SNP sites, 16,872 SNPs were identified in 5,339 unigenes. Of these, at least two SNPs with identical genotypes were found in 1,435 unigenes. We developed a linkage map using genotype information from these unigenes. All unigene markers mapped onto the eight chromosomes and graphical genotyping was conducted based on the unigene order information. Another 2,963 unigenes were allocated onto the eight chromosomes. To confirm the accuracy of this transcriptome-based genetic linkage map, conventional PCR-based markers were used for linkage analysis. All SNP - and PCR-based markers were mapped onto the expected linkage groups and no inconsistency was found among these chromosomal locations.

Conclusions

Effective transcriptome analysis with unique *Allium* resources successfully associated numerous chromosome markers with unigene information and an ultrahigh-density *A. cepa* linkage map. The information on these unigene markers is valuable in genome sequencing and useful trait detection in *Allium*.

Background

The genus *Allium* comprises economically important vegetable crops such as bulb onion (*A. cepa* L.), garlic (*A. sativum* L.), bunching onion (*A. fistulosum* L.), leek (*A. porrum* L.), and numerous wild species (Hanelt, 1990) [1]. Bulb onion is a major vegetable crop worldwide. According to the FAOSTAT database, global bulb onion production was ~ 96 million t and ranked second after tomatoes in terms of vegetable crop cultivation in 2018 [2]. *Allium cepa* L. consists of the common onion (bulb onion) and the Aggregatum (shallot) groups (Jones and Mann, 1963) [3]. Shallot is also an important vegetable crop and is cultivated mainly in Europe, Southeast Asia, and Africa. Though it differs morphologically and ecologically from bulb onion, both are easily crossed (Astley et al., 1982) [4]. Shallot has a short growing period and is resistant to *Fusarium oxysporum* (Vu et al., 2012) [5]. Hence, analysis of its genome might generate valuable information applicable to bulb onion breeding. The
latter is time-consuming and labor-intensive as bulb onion is a biennial and heterogeneous because of severe inbreeding depression. To facilitate bulb onion breeding efforts, then, it is necessary to develop effective methods such as DNA marker-assisted selection.

Various DNA markers have been developed for *Allium* species. Simple sequence repeat (SSR) markers have been used to construct linkage maps as they are co-dominant and useful for PCR-based detection (Baldwin et al., 2012; Bradeen and Havey, 1995; Ipek et al., 2005; Martin et al., 2005; McCallum et al., 2012; van Heusden et al., 2000a, 2000b; Wilkie et al., 1993; Tsukazaki et al., 2006, 2007, 2008, 2011, 2015; Fischer and Bachman, 2000; Song et al., 2004) [6–20]. However, only ~500 SSR markers have been developed for bulb onion so far (Fischer and Bachman, 2000; Kuhl et al., 2004; McCallum et al., 2006; Tsukazaki et al., 2008, 2011; Martin et al., 2005) [9, 16, 17, 19, 21, 22]. These are inadequate for precise genetic analysis and DNA marker-assisted selection in bulb onions.

The advent of next-generation sequencing (NGS) has realized the accumulation of large amounts of sequence information and the identification of numerous single-nucleotide-polymorphisms (SNPs) to develop markers in plants with large genomes (Takahagi et al., 2016) [23]. NGS has been used to generate SNP markers in bulb onions via transcriptomic and selected genomic region bases (Duangjit et al., 2013; Jo et al., 2017; Choi et al., 2020) [24–26]. Numerous SNPs were identified by these approaches. However, only 597, 202, and 319 SNP markers were anchored on each genetic map, respectively, because of parental line heterozygosity in the mapping population. For the effective use of NGS technology, plant materials with sufficient homozygosity must be applied to the parental lines of the mapping population.

Doubled haploid (DH) techniques use chromosome doubling of haploid plants to generate materials that are homozygous enough for genetic analysis (Alan et al., 2003; Ajisaka et al., 2001; Jia et al., 2005) [27–29]. For *Allium*, we developed shallot and bulb onion DH lines and their F₁ hybrids for use in genetic analysis (Abdelrahman et al., 2015; Wako, 2016) [30, 31]. We also developed several bunching onion (*Allium fistulosum* L.)-shallot monosomic addition lines (MALs) (Shigyo et al., 1996) [32]. These have been used to assign genetic linkage maps to *A. cepa* chromosomes by seeking shallot-type alleles among the eight MALs (van Heusden et al., 2000b; Martin et al., 2005) [9, 12]. The combination of these plant resources could enhance potential SNP genotyping by NGS.

Here, we performed a transcriptome analysis on MALs to generate information about chromosome-specific unigene markers. We conducted transcriptomics on the F₂ population derived from a cross between shallot and bulb onion DH lines. We also constructed a high-density genetic linkage map by elucidating the potential SNP sites generated by NGS.

### Results And Discussion

#### Unigene chromosome anchoring by SNP genotyping via MAL RNA sequencing

To create unigene markers, we collected MAL transcriptome data via frequent RNA sequencing (RNA-Seq). We identified candidate genes related to the physiological traits of each line (Abdelrahman et al., 2017) [33]. The transcriptome sequence reads obtained for each MAL were mapped onto a doubled haploid shallot (DHA)
unigene sequence. The unigene transcript levels were evaluated by RPKM (reads per kilobase exon per million mapped reads).

We performed SNP discovery and genotyping as advanced mapping data applications. SNP sites with alternative homozygous calls in bunching onion and reference homozygous calls in shallot were selected by comparing the genotype call of the transcriptome mapping data between the MAL parental lines (bunching onion and shallot). Among 56,161 DHA unigenes, sites with \( \geq 4 \) reads coverage in all eight MALs were identified on 25,462 unigenes (Table 1). Of these, one SNP was identified in 3,278 unigenes. Those whose chromosome assignments could be completed as heterozygous genotypes were identified in only one MAL in each case. In contrast, multiple SNP sites were identified in 22,184 unigenes. Of these, 21,996 could be allocated to single physical chromosomes. Extrachromosomal MALs with heterozygous genotypes are consistent with chromosomal unigene locations with multiple SNPs. For the remaining 188 unigenes, \( \geq 2 \) multiple SNPs were ambiguous. There were heterozygous genotypes in eight MAL types and/or parental homozygous genotype(s). The corresponding gene may have been downregulated and the shallot gene had partial homology. These unigenes were assigned to the chromosome based on other marker(s) with a single heterozygous genotype in MALs with the R indication and mapped by representative SNPs. A total of 25,462 unigenes were anchored on eight chromosomes. There were 4,513 unigenes on chromosome 2 and only 2,169 unigenes on chromosome 8.

DHA unigene information has been made public through the web database ‘Allium Transcriptome DataBase’ (TDB) at http://alliumtdb.kazusa.or.jp. We integrated the chromosome marker information onto each page of the corresponding unigene. These anchoring markers are useful in genome sequencing projects.

**SNP detection in *Allium cepa* doubled haploids**

To link the eight chromosome-specific unigene sets to genetic linkage map information, we accumulated transcriptome data from the F\(_2\) mapping population derived from a cross between the *A. cepa* DH lines (DHA for shallot \( \times \) DHC for bulb onion). As the parental lines were doubled haploid, genotyping the mapping population should be classified as reference (DHA) homozygous, alternative (DHC) homozygous, and heterozygous. RNA sequence data were collected from 96 F\(_2\) lines (population A) of the mapping population and from DHC. The intraspecific SNPs identified by mapping DHC reads with \( \geq 2 \) reads coverage on all 96 lines were selected for genotyping. Selecting co-dominant SNP sites with heterozygous genotypes among the 96 lines identified 16,872 SNP sites in 5,339 unigenes. One SNP site was identified on 2,109 unigenes. These genotypes were used for map calculation with an O indication meaning that one SNP site was supported. Of the 3,230 unigenes with multiple SNP sites, \( \geq 2 \) SNP sites with identical genotype patterns on the 96 lines were identified on 1,435 unigenes. These patterns were selected as the solid genotype (S) of the corresponding unigenes. For the remaining 1,795 unigenes, inconsistencies between the homozygous and heterozygous calls were identified among the 96 lines. The representative genotype (R) was created by selecting the most abundant genotype in each of the 96 lines.

**Genetic linkage map construction and physical chromosome assignment**

We used the solid co-dominant genotype information obtained from 1,435 unigenes in population A to plot a genetic linkage map with JoinMap v. 4.0 (Kyazma BV, Wageningen, The Netherlands). By applying the LOD 5
cutoff, all tested markers were assigned to eight linkage groups. Based on the unigenes with anchored chromosome information, all of these could be anchored to each of the eight bulb onion chromosomes. No inconsistency was detected between each linkage group and assigned chromosome. Hence, this linkage map was reliable.

A graphical genotype list was constructed according to the unigene order information. A total of 610 genotype blocks were assigned based on the patterns of the tested 96 lines (Table S1). The remaining unigenes with O- and R-coded genotypes were allocated to the most probable genotype block and permitted genotype inconsistencies for ≤5 lines. A total of 1,537 O-marked and 1,426 R-marked unigenes were allocated onto the genotype blocks (Tables 2).

To confirm transcriptome-based genetic linkage map accuracy, we applied conventional PCR-based markers to the same F2 population (A). The PCR-based SSR and InDel markers were previously reported (Fischer and Bachmann, 2000; Kuhl et al., 2004; Martin et al., 2005; McCallum et al., 2012; Tsukazaki et al., 2006, 2007, 2008, 2011; Wako, 2016) [9,10,14,15,16,17,19,21,31] and used in the present study. Thirty-three markers were polymorphic between DHA and DHC. Fourteen InDel polymorphisms were detected for the sequence comparisons between DHA and DHC in Allium TDB. We designed primer sets that included these polymorphism sites and amplified them by PCR. We used 47 PCR-based markers in a linkage analysis on another F2 population (B). All linkage groups were assigned to eight physical chromosomes in MALs confirmed by amplification. These marker locations matched those in previous reports (Tsukazaki et al., 2008, 2011, 2015; Masuzaki et al., 2006a, 2006b; Wako, 2016) [16,17,18,31,34,35]. We selected 14 reliable PCR-based markers covering all eight chromosomes, applied them to population A, and integrated them onto the transcriptome-based genetic linkage map. The reconstructed map consisted of eight linkage groups with all SNP solid and PCR-based markers covering 936.6 cM. The average marker interval was 0.65 cM. All PCR-based markers were integrated onto positions corresponding to those on population B. The latter was based on a PCR marker-based linkage map. No contradiction in marker location was caused by using common markers between these maps and another linkage map previously constructed with a gynogenic population (C) derived from the same F1 hybrid between DHA and DHC (Fig. 1) (Wako, 2016) [31]. We also compared the genetic maps against a published transcriptome-based SNP marker analysis (Duangjit et al., 2013) [24]. Comparison of the positions of 137 SNP markers on sequences overlapping in both analyses revealed that the anchored chromosomes and relative positions were consistent for all SNP markers (Table S2). Therefore, our transcriptome-based genetic linkage map is reliable.

MALs have been used extensively to assign DNA markers to physical chromosomes (van Heusden et al., 2000b; Martin et al., 2005; Tsukazaki et al., 2008) [9,12,16]. Here, we identified chromosome-specific SNPs by comparing transcriptome data with MALs. For the first time, we used F2 populations from Allium DH parental lines. The parental line has each homozygous allele. SNPs between the parental lines DHA and DHC are easily detected. Transcriptome data from the DH lines efficiently found SNPs (Baldwin et al., 2012) [6] and we obtained abundant and reliable SNP information here. We constructed a reliable genetic map based on S-marked SNP markers. No inconsistency was found between the physical chromosome assignments and S-labeled markers in the linkage group. The genetic map comprised 1,435 SNP markers, one bulb onion SSR marker, and 13 InDel markers and covered 936.6 cM. To our knowledge, this map has the highest number of markers to date. Integrated linkage maps include markers associated with phenotypic characteristics for the
nuclear male fertility restoration loci of cytoplasmic male sterility (Chr. 2) and bulb color (Chr. 7) (Wako, 2016) [31]. Shallot is a genetic breeding resource for bulb onion as it produces certain distinctive chemical compounds such as saponins conferring pathogen resistance (Shigyo et al., 1997; Abdelrahman et al., 2017) [33,36]. By combining these DH lines with linkage map information, progress is anticipated in Allium molecular breeding by marker-assisted selection for several agronomic bulb onion traits.

**Conclusions**

In the present study, we constructed an ultrahigh-density linkage map in Allium cepa using numerous SNP markers obtained from the transcriptome information of the Allium DH lines and the MALs. As DH techniques depress inbreeding, they are useful for making homozygous pure lines that resemble inbred lines (Abdelrahman et al., 2015) [30]. Though bulb onion and shallot have different characteristics, they both belong to A. cepa and are easy to cross (Astley et al., 1982) [4]. The MALs have all A. fistulosum chromosomes and one A. cepa chromosome (Shigyo et al., 1996) [32]. We performed a transcriptome analysis to identify unigenes and assign them to physical chromosomes. To this end, we compared shallot DH and MAL transcriptome data. We then used the F2 mapping population between bulb onion DH and shallot DH to detect SNP sites. A total of 16,872 SNP sites were identified on 5,339 unigenes. Of these, 1,435 were selected as the solid genotype of the corresponding unigenes. By constructing a linkage map with SNP solid markers, all markers were mapped and the locations between the physical chromosomes and linkage groups were consistent. The number of SNPs located on the linkage map was much higher than those previously reported. Thus, the linkage map resolution was high. Furthermore, linkage maps integrated with PCR-based markers are now available. Shallots produce chemical compounds conferring resistance to certain bulb onion diseases (Abdelrahman et al., 2017) [33]. Hence, connecting phenotype and genotype information is a holistic approach towards Allium gene expression analysis for plant breeding and an effective, low-cost method of developing novel disease-resistant Allium varieties.

**Methods**

**Plant materials and genetic cross**

F2 plants generated by a cross between shallot and bulb onion doubled haploid (DH) lines (DHA and DHC) were used to construct a linkage map. DHA was derived from ‘Chiang Mai’ shallot in Thailand while DHC was derived from long-day ‘Sapporo-ki’ onion cultivar in Japan (Abdelrahman et al., 2015) [30]. For the transcriptome and conventional PCR-based marker analyses, populations A (96 individuals) and B (186 individuals) were created. They were raised in the greenhouses of Yamaguchi University (34°11” N; 131°28” E) and the Institute of Vegetable and Floriculture Science of NARO (34°61” N; 136°25” E) in Japan.

For the A. cepa unigene chromosome assignments via interspecific SNP detection, a complete set of MALs of Allium fistulosum L. with eight single shallot extra-chromosomes (Shigyo et al., 1996) [32] was used (Abdelrahman et al., 2017) [33].

**Transcriptome sequencing**
Total RNA was isolated with the RNeasy plant mini kit (QIAGEN Sciences, Germantown, MD, USA). RNA quality was checked with an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA, USA). Samples with RNA integrity number (RIN) > 8.0 were selected for further use. The cDNA library was generated with a TruSeq™ RNA sample preparation kit (Illumina, San Diego, CA, USA) in accordance with the manufacturer’s instructions. Sequencing was performed on the Illumina HiSeq 2500 platform (Illumina, San Diego, CA, USA).

**SNP detection and selection**

RNA sequence reads were filtered with PRINSEQ v. 0.20.4 (Schmieder and Edwards, 2011) [37] and fastx_clipper in FASTX-toolkit (http://hannonlab.cshl.edu/fastx_toolkit/). The filtered single-end reads were mapped onto DHA unigene sequences in end-to-end mode with Bowtie v. 22.1.0 (Langmead et al., 2009) [38]. The sequence alignment/map (SAM) format files were converted into BAM format with SAMtools v. 0.1.19 (Li et al., 2009) [39]. The BAM files were subjected to SNP calling with the mpileup option of SAMtools31 v. 0.1.19 and the mpileup2snp option of VarScan v. 2.3 to obtain a variant call format (VCF) file containing the SNP information.

**PCR-based marker analysis**

Total DNA was prepared from individual plants according to the method of Song et al. (2004) [20]. To detect polymorphisms between DHA and DHC, SSR markers derived from the *A. cepa* genome (Fischer and Bachmann, 2000) [19], the *A. fistulosum* genome (Ohara et al., 2005; Song et al., 2004; Martin et al., 2005; Tsukazaki et al., 2006, 2007, 2008, 2011; Wako, 2016) [9,14,15,16,17,20,31,40], and *A. cepa* expressed sequence tags (ESTs) (Jakse et al., 2005; Kuhl et al., 2004; Martin et al., 2005; Tsukazaki et al., 2008; 2011) [9,16,17,21,41] and other CAPS and SCAR markers (Kuhl et al., 2004; Masuzaki et al., 2006a, 2006b; McCallum et al., 2006; Yaguchi et al., 2008) [21,22,34,35,42] were screened. Moreover, polymorphisms were detected between DHA and DHC nucleotide sequences obtained from transcriptome information in Allium Transcriptome DataBase (AlliumTDB) (Abdelrahman et al., 2017) [33] by sequence comparison with BLASTn. InDels > 6 bp were selected and primer sets including the polymorphism site were designed with Primer 3 (http://bioinfo.ut.ee/primer3/) (IDTDBxxx). The markers were applied to the F$_2$ population and parental lines. PCR was performed in a 10-μL reaction mixture containing 10 ng template DNA, 0.2 μM of each primer, 0.2 mM of each dNTP, and 5 μL GoTaq Master Mix (Promega Corp., Madison, WI, USA). Amplification was performed for 35 cycles after initial denaturation at 94 °C for 4 min. Each cycle consisted of 15 s at 94 °C, 30 s at 55 °C, 1 min at 72 °C, and 4 min at 72 °C. The PCR products were evaluated by 2% (w/v) agarose gel electrophoresis. Certain forward primers were fluorescently labeled with 6-FAM, NED, PET, or VIC dyes (Applied Biosystems, Foster City, CA, USA) before PCR. The PCR products were loaded onto a capillary DNA sequencer (ABI3730; Applied Biosystems, Foster City, CA, USA) and analyzed with GeneMapper v. 3.0 (Applied Biosystems, Foster City, CA, USA).

**Allium cepa linkage map construction by two methods and map comparison**

Linkage analysis was performed with JoinMap v. 4.0 (van Ooijen, 2006) [43]. The Kosambi function was used to obtain the cM values (Kosambi, 1944) [44]. The DNA markers used to construct the *A. cepa* linkage map are listed in Table S3. Eight different *A. fistulosum-A. cepa* MALs were used to determine the corresponding physical chromosomes for the linkage groups. The linkage maps were compared according to the SNP and PCR-based markers by using the 14 anchor markers on both maps.
Abbreviations

DH: doubled haploid

NGS: next-generation sequencing

PCR: polymerase chain reaction

MAL: monosomic addition lines

SSR: simple sequence repeat

InDel: insertion-deletion

EST: expressed sequence tag

SNP: single-nucleotide polymorphism

CAPS: Cleaved amplified polymorphic sequence

SCAR: Sequence Characterized Amplified Region

Declarations

Ethics approval and consent to participate

Not applicable.

Consent to participate

Not applicable.

Availability of data and materials

RNA-Seq data for the MALs and F₂ mapping population (DHA × DHC) are available in the DDBJ sequence read archive under Accessions No. DRA00509 and DRA009194, respectively. The DHA unigene information has been made public through the web database ‘Allium Transcriptome DataBase’ (AlliumTDB) located at http://alliumtdb.kazusa.or.jp.

Competing interests

The authors declare that they have no competing interests.

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Author contributions
All authors have read and approved the submitted version of the manuscript, and have agreed both to be personally accountable for the author’s own contributions and to ensure that questions related to the accuracy or integrity of any part of the work, even ones in which the author was not personally involved, are appropriately investigated, resolved, and the resolution documented in the literature. Authors’ contribution are, 1. Conceptualization: TW, SS, MS; 2. Methodology: HH, YM, SS, MS.; 3. Investigation: SF, TYA, TM, KY, HT, KT; 4. Writing-Original Draft: SF, TA, SS, MS; 5. Writing-Review and Editing: SF, TYA, TM, TW, KY, HT, HH, YM, SS, MS.

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**Tables**

**Table 1.** DHA unigenes with SNP(s) detected on MALs

| Chr. 1 | Chr. 2 | Chr. 3 | Chr. 4 | Chr. 5 | Chr. 6 | Chr. 7 | Chr. 8 | Total |
|--------|--------|--------|--------|--------|--------|--------|--------|-------|
| Multiple SNPs | 3,404 | 3,900 | 3,148 | 2,538 | 2,588 | 2,546 | 2,032 | 1,840 | 21,996 |
| allocated to single chromosome | | | | | | | | | |
| including representative SNP | 30 | 35 | 33 | 21 | 16 | 16 | 18 | 19 | 188 |
| One SNP | 486 | 578 | 455 | 397 | 357 | 376 | 319 | 310 | 3,278 |
| Total | 3920 | 4,513 | 3,636 | 2,956 | 2,961 | 2,938 | 2,369 | 2,169 | 25,462 |

**Table 2.** Genetic markers developed for each chromosome
|       | Chr. 1 | Chr. 2 | Chr. 3 | Chr. 4 | Chr. 5 | Chr. 6 | Chr. 7 | Chr. 8 | Total |
|-------|--------|--------|--------|--------|--------|--------|--------|--------|-------|
| Solid (S) | 270    | 126    | 200    | 170    | 199    | 198    | 170    | 102    | 1,435 |
| Representative(R) | 175    | 416    | 195    | 163    | 148    | 123    | 110    | 96     | 1,426 |
| One (O)   | 239    | 278    | 229    | 192    | 135    | 186    | 148    | 130    | 1,537 |
| Total     | 684    | 820    | 624    | 525    | 482    | 507    | 428    | 328    | 4,398 |

**Figures**

![Figure 1](image1)

*Figure 1*
Linkage maps. Left, SNP-based map including PCR-based markers (F2 population A). Middle, PCR-based map (F2 population B). Right, linkage map constructed using a gynogenic population derived from F1 (population C) (Wako, 2016).

**Supplementary Files**

This is a list of supplementary files associated with this preprint. Click to download.

- SupplementalTableS1colormapDHAxDHCF2population.xlsx
- SupplementalTableS2SNPcomparison.xlsx
- SupplementalTableS3201108.xlsx