Introduction

Advances in next-generation sequencing and genotyping technologies have paved the way for the application of genomics-assisted breeding (Varshney et al. 2005) to various major crops, such as maize, wheat, and rice. In the last several years, minor (or orphan) crops, which are those crops that have received less attention from researchers compared with major crops, have also started to garner attention as targets for genomics-assisted breeding (Armstead et al. 2009, Varshney et al. 2012). Minor crops are of great importance in the marginal environments of Africa, Asia and South America because they are often more nutritious and more adapted to the harsh environments of these areas than many of the major crop varieties. Because they play an important role in regional food security, the development of genomic resources has rapidly progressed for various minor crops (e.g., Ly et al. 2013, Varshney et al. 2013).

Common buckwheat, *Fagopyrum esculentum* Moench (2n = 2x = 16), is an example of a minor yet important crop. Common buckwheat is easy to grow and can be adapted to almost all types of soil, except for sandy, heavy crusted, or wet soils. Genus *Fagopyrum* has great potential as a crop for forage, human consumption, and medicine because of its high-quality protein content (Campbell 1997). For common buckwheat in particular, its rapid rate of growth and ability to grow at high altitudes make it an especially important crop in the mountainous regions of South Asia (e.g., Baniya 1990, Joshi 1999). Despite its importance, the genetic improvement of common buckwheat has achieved only limited success, mainly due to buckwheat’s heteromorphic self-incompatibility (Campbell 1997). Common buckwheat is a heteromorphic, self-incompatible species that has heterostyous flowers controlled by the S-locus (reviewed in Lewis and Jones 1992). Genomics-assisted breeding is expected to accelerate the genetic improvement of common buckwheat; however, the genomic resources currently available remain incomplete. Furthermore, the development of genomic resources for use in common buckwheat breeding has been hampered because buckwheat is the only crop species within the Polygonaceae family, and therefore is phylogenetically distant to other major crops and model plants. While some linkage map has been published (e.g., Konishi
and Ohnishi 2006, Pan and Chen 2010, Yasui et al. 2004), the utility of the map has been limited due to buckwheat’s large genetic diversity.

The potential of genomics-assisted breeding has been demonstrated in various plant species (e.g. Varshney et al. 2013). Yabe et al. (2014) expect genomic selection (Meuwissen et al. 2001), which is a promising method in genomics-assisted breeding, to accelerate the genetic improvement of allogamous crops such as common buckwheat. However, to fully utilize genomic selection, a rapid, high-throughput genotyping technology is necessary because a large number of genome-wide markers need to be genotyped for all of the plants in a breeding population before the plants mature. To widen the scope for the application of genomics-assisted breeding technologies, genotyping technologies should be able to conduct genotyping or constructing linkage map even without any a priori genomic information of target species.

In the present study, we evaluated the potential of a novel marker system (Enoki et al. 2012, Iehisa et al. 2014) for use as a rapid, high-throughput genotyping technology for genomics-assisted breeding. We used common buckwheat as the test plant, and conducted a genome-wide linkage analysis without using any a priori genomic information (e.g., DNA sequences) or any existing genomic tools (e.g., DNA markers). Moreover, to verify the utility of the linkage map we constructed, we applied it to a quantitative trait loci (QTL) analysis of the same population. We selected two plants from a breeding population and crossed them to generate a segregating population. This approach can be generalized to linkage and QTL mapping in other crop species, thus the potential of the novel marker system demonstrated in this study may be feasible in various ‘orphan’ or ‘less-studied’ crops. Since common buckwheat is a completely allogamous species, breeding populations have large effective population sizes; common buckwheat is therefore expected to have a low level of linkage disequilibrium (Sved 1971). Linkage map containing a high number of markers (i.e., a high-density linkage map) could be highly beneficial for the success of genomics-assisted breeding.

Materials and Methods

Development of mapping population and agronomic characterization

To construct a linkage map and detect QTL related to agronomic characteristics, we employed 92FE1-F4, a population produced by bulk crossing among ‘Tempest’, ‘Kitawasesoba’, ‘Natsusoba’, and ‘Shinanonatsusoba’. These cultivars are classified into a single agroecotype: summer type. We developed a mapping population consisting of 178 F1 progeny derived from a cross between P1 and P2, which were selected from 92FE1-F4. In the present study, P1 had thrum-type flowers and P2 had pin-type flowers. The genotype of the S-locus was heterozygous (i.e., S/s) in P1, and recessive homozygous (s/s) in P2.

The F1 progeny were sowed at a density of one seed per plastic pot (diameter, 24 cm; height, 24 cm) on 6 August 2012 and cultivated under natural conditions in an isolation chamber (L × W × H, 630 × 540 × 230 cm) at the University of Tsukuba (36°06′N, 140°05′E). We examined main stem length as the trait for QTL mapping after harvesting on 9 October 2012.

DNA extraction

Total genomic DNA from both parental individuals and the F1 progeny was extracted as follows: About 100 mg milled leaf tissue was mixed with 500 μl lysis buffer (0.3% sodium dodecyl sulfate, 20 mM Tris-HCl [pH 8.0 at 25°C], 5 mM EDTA, 400 mM NaCl) and 4 μl RNase A (100 mg/ml; Qiagen Inc., CA, USA), and incubated at 65°C for 10 min. The mixture was then centrifuged (14,000 rpm, 20°C) for 2 min. Buffer AP2 (130 μl; Qiagen Inc.) was added to the cleared lysate and the samples were mixed, incubated on ice for 5 min, and centrifuged for 2 min (14,000 rpm, 20°C). Then, 500 μl isopropanol (Wako Pure Chemical Industries, Ltd., Osaka, Japan) was added to the supernatant, and the samples were mixed by inversion, incubated for 5 min at room temperature, and centrifuged for 10 min. The DNA pellet was washed twice with 500 μl of 70% ethanol (Wako Pure Chemical Industries, Ltd.). After each wash, the sample was centrifuged for 5 min, and 500 μl of 99.5% ethanol (Wako Pure Chemical Industries, Ltd.) was added to the pellet. After incubation for 5 min at room temperature, the sample was centrifuged for 5 min and resuspended in 100 μl Tris-EDTA buffer.

Sequence library construction and next-generation sequencing analysis for marker preparation

Total genomic DNA was prepared from the mapping parents (P1 and P2) and from 40 plants of the 92FE1-F4 population (hereafter referred to as the 40-mix population). Total genomic DNA was digested with Psfl (New England Biolabs, Inc., Ipswich, MA) at 37°C for 1 h and then with BstNI (New England Biolabs) at 60°C for 1 h. Genomic DNA fragments obtained were ligated with Psfl-sequence adapters (5′-CACGATGGATCCAGTGCA-3′, 5′-CTGGATTCCATCGTGCA-3′) by using T4 DNA Ligase (New England Biolabs, Inc., Ipswich, MA) at 65°C for 1 h. Polymerase chain reaction (PCR) amplifications were performed with Promega DNA polymerase (Takara Bio, Inc., Otsu, Japan). The PCR primer was designed based on the adapter sequences (5′-GATGGATCCAGTGCA-3′). The PCR conditions were as follows: 30 cycles at 98°C for 10 s, 55°C for 15 s, and 72°C for 1 min, followed by 1 cycle at 72°C for 3 min. These steps amplify only the Psfr–Psfl DNA fragment but not Psfl–BstNI fragments. After sonication, DNA fragments from 200 to 500 bp were recovered by electrophoresis on a 1.5% agarose gel. DNA fragments from P1 and P2 were recovered separately, and two sequence libraries (LIB-P1 and LIB-P2) were constructed by using a Paired-End DNA Sample Preparation Kit (Illumina, Inc.,
San Diego, CA). PCR products from each plant in the 40-mix population were equally mixed before sonication and sizing, and a sequence library (LIB-40mix) was constructed by using a Paired-End DNA Sample Preparation Kit (Illumina). LIB-P1 and LIB-P2 were analyzed by using an Illumina GAII with a paired-read of 300 nt, and LIB-40mix was analyzed by using an Illumina HiSeq2000 with a paired-read of 200 nt. The sequences were submitted to the DNA Data Bank of Japan (submission no. DRA001140).

Processing of raw reads and design of probes for microarray analysis

The procedure for the processing of row reads and the design of probes was similar to the one provided in Iehisa et al. (2014), in which the same genotyping system was applied to the linkage mapping of common wheat. First, we performed trimming of adaptor and primer sequences and low-quality 3′ ends (quality score below 2) from raw reads using in-house developed software. Discarding pairs of reads when either of pairwise reads had length of <70 nt, we obtained three trimmed sequence libraries, TRIM-P1, TRIM-P2 and TRIM-40mix. CONT-P1, CONT-P2 and CONT-40mix were then assembled, respectively, from TRIM-P1, TRIM-P2 and TRIM-40mix using Velvet ver. 1.1.05 (Zerbino and Birney 2008). For the assemblies, velveth was run with a k-mer size of 65, and velvetg was run with expected coverage: ‘auto’, coverage cut-off: ‘auto’, insert length of 220 bp, minimum contig length of 100 bp, and allowing scaffolding. From assembled contigs and reads unused in the assemblies, tiling probes were designed with a Tm of around 76°C and the length of 50–75 bp. The probe design fundamentals were described in the NimbleGen technical note (http://www.roche-biochem.jp/products/2010/08/array/pdf/probe_design_2008_06_04.pdf). To construct a mapping array, designed probes were first aligned to two trimmed libraries, TRIM-P1 and TRIM-P2, by using Bowtie ver. 0.12.7 (Langmead et al. 2009), where no more than 3 mismatches were allowed and up to 30 alignments were reported (-v 3, -k 30). P1-derived probes that met all the following five conditions were selected: (1) the number of aligned P1 reads with no mismatch was larger than 3, (2) the ratio of aligned P1 reads with no mismatch over aligned P1 reads with one or two mismatches was larger than 4, (3) the ratio of aligned P1 reads with no mismatch over aligned P2 reads with less than three mismatch was larger than 10. Similar selection was performed for P2 probes. All candidate probes from CONT-40mix were mapped to TRIM-40mix by using the Bowtie software, and the probes that the number of aligned reads with no mismatch was larger than 30 were selected. Then, the selected probes were aligned to the trimmed sequence library TRIM-166, which were obtained from PsrI–PsrI DNA fragments that did not harbor BsrNI sites, was obtained from 166 F1 plants derived from the 40-mix population. The number of aligned reads with no mismatch to the sequences of each plant was counted. Probes that showed identical patterns of absence/presence of probe hits were merged. All probes from P1, P2 and 40mix and control probes were synthesized in triplicate on a NimbleGen HD-2 135K × 12plex microarray (Roche Diagnostics, Madison, WI). Although the company has terminated services for the microarrays, microarrays provided by other companies, such as Agilent Technologies, Inc., will also be usable for the genotyping assay of the marker system.

Hybridization of microarray

Genotypes of the parental plants, P1 and P2, and their 178 F1 progeny were determined by using HD-2 135K × 12plex microarrays as described. After digestion of total DNA with PsrI and BsrNI, adapter ligation, PCR amplification and purification of PCR products, as described in library preparation section, they were labelled with Cy3 using NimbleGen One-Color DNA Labeling Kit (Roche Diagnostics) according to the manufacturer’s instructions. Hybridization was performed at 42°C during 24 hours on NimbleGen Hybridization System (Roche Diagnostics). The microarrays were scanned with a NimbleGen MS200 Microarray Scanner (Roche Diagnostics) and genotype calling was performed based on the signal intensity. Present/absent calls of the probe-sets reflected the present/absent DNA fragments hybridized to the probe-sets. Thus, DNA polymorphisms genotyped with this system were used as dominant markers.

Linkage map construction

Linkage maps of P1 and P2 were constructed by using the pseudo-testcross strategy (Grattapaglia and Sederoff 1994). First, the deviation from the Mendelian segregation ratio was tested for each marker by using the Chi-square test (p < 0.01; statistically significance). Markers segregating in a 1 : 1 ratio were used to construct the linkage map of P1 and P2. Markers were assigned to linkage groups by setting the recombination rate threshold at 0.3 and the threshold for the minimum number of markers at 3. Locus ordering was performed by using AntMap software (Iwata and Ninomiya 2006) with a 50-run of an optimization process (i.e., maximization of the log-likelihood). The Kosambi mapping function (Kosambi 1943) was used to calculate map distances. To connect linkage groups constructed in the P1 and P2 linkage map, recombination rates between markers segregating in a 3 : 1 ratio and markers represented in the P1 and P2 map were calculated. Among the markers segregating in a 3 : 1 ratio, those that had low recombination rates (<0.02) with both markers on the P1 and P2 map were used as “bridges” between the P1 and P2 map. When multiple markers on the P1 or P2 map had recombination rates <0.02 and were segregated in a 3 : 1 ratio, the centers of gravity of these multiple markers were connected by a bridge.

QTL analysis

For the QTL analysis, 171 plants for which both marker data and phenotypic data (i.e., scores of main stem length
Composite interval mapping (Zeng 1993, 1994) was performed by using the QTL Cartographer software ver. 1.17 (Basten et al. 2003). To analyze our pseudo-testcross data, we employed a model for inbred backcross design. In the analysis, we adopted the linkage phase estimated at the step of linkage map construction. A permutation test with 100 replicates was performed for each trait to estimate the empirical threshold value corresponding to the 5% significance level. The proportion of total (i.e., phenotypic) variance explained by each of detected QTL was calculated as ([residual variance under the null hypothesis]–[residual variance under the alternative hypothesis])/[phenotypic variance].

Results

Construction of microarray

To design probes for developing a microarray, we sequenced complexity-reduced genomic DNA samples from P1, P2 and 40mix-population. After trimming and filtering of raw reads, libraries of 13,055,602, 11,576,694 and 267,438,466 reads were generated (TRIM-P1, TRIM-P2 and TRIM-40mix; Fig. 1). All libraries were assembled independently by using the Velvet, yielding 9,054 contigs and 5,535,673 unused reads for TRIM-P1, 8,187 contigs and 4,684,155 unused reads for TRIM-P2, 63,558 contigs and 128,957,022 unused reads for TRIM-40mix. From assembled contigs and reads unused in the assemblies, tiling probes were designed. The probes developed from P1 and P2 were aligned to the TRIM-P1 and TRIM-P2, and the probes that showed presence/absence polymorphisms were selected. The probes developed from 40mix were aligned to the TRIM-40mix, and the probes that showed more than 30 hits were selected. The selected probes were then mapped to TRIM-166, and probes that showed identical patterns of absence/presence of probe hits were merged. Finally, we obtained 16,583 probes from P1, 16,584 from P2, and 11,480 from 40mix population.

Preparing the data for linkage map construction

DNA microarray genotyping was performed for 44,836 markers (i.e., 44,647 probes mentioned above and 189 control probes). For marker genotypes, we treated three signal types: present call, absent call and missing. The present/
absent calls were determined according to the bimodal microarray signals. The ambiguous signals were classified as missing values. From those markers, we used 16,841 markers that were heterozygous in one or both parents for further analysis. In the F1 population of 178 plants, the numbers of markers segregating in 1 : 1 and 3 : 1 were 14,442 (P1: 6,875 markers; P2: 7,567 markers) and 2,399, respectively. A segregation ratio of 1 : 1 is expected when one parent has a heterozygous genotype and the other has a recessive homozygous genotype. A segregation ratio of 3 : 1 is expected when both parents have a heterozygous genotype. Of the markers with segregation ratio of 1 : 1 and 3 : 1, 9,112 (P1: 4,325 markers; P2: 4,787 markers) and 1,701 markers, respectively, showed a clear distinction between the two genotypes. Of all 10,813 markers, 1,339 markers contained one or more missing values. Of these, 824 contained missing values for less than four plants, and 106 contained missing values for more than 17 (10%) plants. Markers derived from an identical contig with the same segregation type (i.e., P1 or P2 markers with a 1 : 1 segregation ratio or bridging markers with a 3 : 1 segregation ratio) were expected to reflect genotypes at the location of the contig because of the close linkage between the markers. We therefore collected markers from a single contig to make consensus genotypes. The missing genotypes of the collected markers were imputed on the basis of the consensus genotypes. In a few cases, two or three patterns of marker genotypes were observed in one contig. In those cases, we treated the groups of markers having different genotypes as separate contigs (sep-contigs). When sep-contigs derived from a single contig belonged to different segregation types, we excluded them from the subsequent analysis. Sep-contigs from a single contig showing a low level of consensus (i.e., genotypes were discordant in more than 30% of plants) were also excluded. We used flower morphology (i.e., pin or thrum) controlled by the S-locus as the phenotypic marker. The marker was located on the P1 map because P1 was heterozygous (i.e., S/s) at the locus. In total, we had 1,455, 1,631, and 869 contigs for P1, P2, and bridging markers, respectively. Next, to analyze the pseudo-testcross data, we inverted the genotype data by duplicating and converting all of the contigs (i.e., homozygous genotypes were converted into heterozygous genotypes and vice versa). We grouped contigs that had an identical genotype into a single marker group and then separated those marker groups into three marker types—P1 markers, P2 markers, and bridging markers. Marker groups were grouped into two and four clusters for P1 and P2 type, respectively, via a single-linkage cluster analysis based on Manhattan distances among marker groups. The distances were calculated based on the genotypes of marker groups, which were scored with 0 and 1. We used single-linkage clustering because the method is similar to the one used in the linkage grouping. In total, we attained 346, 410, and 360 marker groups, respectively, for P1, P2, and bridging markers. The genotypes of these marker groups were used to construct the linkage map.

**Linkage map construction**

By using a pseudo-testcross strategy, we constructed linkage map for P1 and P2, and connected them with bridging markers (Fig. 2; Supplemental Tables 1, 2). We mapped 346 loci on the P1 map, and 410 loci on the P2 map (Table 1; Supplemental Table 1). We used 283 groups segregating 3 : 1 (bridging markers), which represented small

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**Fig. 2.** Linkage map for common buckwheat. Dashed lines represent “bridge” markers that connect marker groups between the linkage maps of P1 and P2. Phenotypic marker position (i.e., S-locus) is indicated by “S”.

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recombination rates (i.e., <0.02) with loci in both the P1 and P2 map, to combine the loci in the P1 and P2 linkage map (Supplemental Table 2). We used markers segregating 3:1 only for the P1 and P2 map bridging, not for mapping on the linkage map. Because the precision of the estimation of recombination rates between markers segregating 3:1 and between markers segregating 1:1 and 3:1 was low (Ritter et al. 1990), the inclusion of 3:1 markers in the linkage map was thought to affect the estimation of the linkage map positions of 1:1 markers. Thus, as shown in Fig. 2, we used bridging markers only to connect the linkage groups between the P1 and P2 linkage map. The phenotypic marker, (flower morphology) was located on linkage group P1_3 (“S” in Fig. 2). After connecting the P1 and P2 linkage map, the number of linkage groups converged to eight, which is the basic chromosome number of common buckwheat. The eighth linkage group was divided into two groups of short length in the P1 map. The P1 and P2 linkage map covered 773.8 and 800.4 cM, and contained 1,455 and 1,631 contigs, consisting of 4,227 and 4,657 markers, respectively (Table 1; Supplemental Table 1). The means of the intervals between adjacent positions were 2.30 and 1.99 cM (Table 1) and the medians were 1.68 and 1.15 cM in the P1 and P2 linkage map, respectively. Most (90%) adjacent positions had intervals shorter than 5.07 cM.

On the linkage map, one position (i.e., a single marker group) harbored a number of contigs, and one contig consisted of a number of markers. Fig. 3 shows the number of contigs per loci (A) and the number of markers per contig (B). Among the 756 loci on the map, 555 loci consisted of more than one contig, and 492 loci consisted of less than 10 contigs. Among the 3,086 contigs, 1,140 contigs consisted of more than one marker, and among those, 1,036 contigs consisted of less than 15 markers and 13 contigs consisted of more than 35 markers.

**QTL analysis**

Phenotypic values for main stem length observed in the mapping population had a unimodal, continuous distribution (Fig. 4), suggesting that main stem length is controlled by multiple QTL and is influenced by environmental effects. For main stem length, significant QTL were detected at map positions 9.3 cM on the P1-1 group, 49.0 cM on the P1-2 group, 9.0 cM on the P1-5 group, and 16.9 cM on the P2-4 group (Fig. 5; Table 2). The four QTL accounted for 5.64% to 8.51% of the phenotypic variance observed in main stem length (Table 2).

**Table 1.** Summary information for the linkage map of common buckwheat

| Linkage group | No. of loci | Genetic length (cM) | Average interval of adjacent loci (cM) | No. of contigs | No. of markers |
|---------------|-------------|---------------------|----------------------------------------|----------------|---------------|
| P1_1          | 45          | 107.6               | 2.45                                   | 198            | 674           |
| P1_2          | 51          | 109.1               | 2.18                                   | 211            | 549           |
| P1_3          | 51          | 114.3               | 2.29                                   | 223            | 691           |
| P1_4          | 42          | 80.2                | 1.96                                   | 193            | 511           |
| P1_5          | 47          | 125.8               | 2.73                                   | 203            | 626           |
| P1_6          | 49          | 118.0               | 2.46                                   | 196            | 560           |
| P1_7          | 41          | 87.1                | 2.18                                   | 161            | 453           |
| P1_8.1        | 3           | 1.8                 | 0.88                                   | 10             | 18            |
| P1_8.2        | 17          | 30.0                | 1.88                                   | 60             | 145           |
| P2_1          | 37          | 83.3                | 2.32                                   | 119            | 235           |
| P2_2          | 54          | 104.4               | 1.97                                   | 206            | 621           |
| P2_3          | 70          | 116.1               | 1.68                                   | 273            | 886           |
| P2_4          | 49          | 71.1                | 1.48                                   | 184            | 498           |
| P2_5          | 57          | 120.8               | 2.16                                   | 217            | 598           |
| P2_6          | 57          | 134.5               | 2.40                                   | 224            | 695           |
| P2_7          | 43          | 83.6                | 1.99                                   | 189            | 544           |
| P2_8          | 43          | 86.7                | 2.06                                   | 219            | 580           |
| P1            | 346         | 773.8               | 2.30                                   | 1,455          | 4,227         |
| P2            | 410         | 800.4               | 1.99                                   | 1,631          | 4,657         |
| All           | 756         | 1,574.3             | 2.13                                   | 3,086          | 8,884         |

![Fig. 3.](image-url) Distribution of the number of contigs per map position (A) and the number of markers per contig (B).
By using a novel marker system, we successfully constructed a high-density linkage map for common buckwheat. The number of linkage groups converged to eight, which is the basic number of chromosomes in common buckwheat. The size of the linkage groups in the P1 and P2 linkage map were 773.8 and 800.4 cM, respectively. Some linkage maps have previously been developed for common buckwheat. Yasui et al. (2004) constructed a linkage map with eight linkage groups covering a total of 508.3 cM that contained 223 amplified fragment-length polymorphism (AFLP) markers. However, some of these markers were not included in the eight linkage groups. Konishi and Ohnishi (2006) conducted linkage mapping with the pseudo-testcross strategy. Their female map had 12 linkage groups covering 911.3 cM that contained 54 microsatellite and 77 AFLP markers; their male map had 12 linkage groups covering 909.0 cM that contained 37 microsatellite and 34 AFLP markers. The female and male map converged to eight paired linkage groups, although three of the linkage groups in the female map did not converge to the eight groups. Pan and Chen (2010) constructed a linkage map with 10 linkage groups covering a total of 692.4 cM that included 87 randomly amplified polymorphic DNA markers, 12 sequencetagged sites, four seed protein subunit markers, and three morphological character alleles. Hara et al. (2011) constructed a linkage map with nine linkage groups covering a total of 311.6 cM that contained two photoperiod-sensitivity candidate genes and 63 expressed sequence tag markers.

Compared with the previously constructed map, the map constructed in the present study had the largest number of markers on one side (i.e., P1 map or P2 map), and the average interval of adjacent markers was 2.13 cM, which is the smallest interval reported to date. Moreover, the map constructed in the present study converged to eight linkage groups including all markers used in the mapping analysis, whereas the others did not. Together, these results show that the map constructed in the present study will be suitable for use as a basic linkage map in the future studies of common buckwheat.

The linkage map constructed in the present study had 756 independent loci and 8,884 markers (Table 1; Fig. 3). Thus, the map harbored multiple markers at a single position. This characteristic may contribute to the versatility of the linkage map. Common buckwheat has a high level of genetic variation; therefore, genetic composition differs by population. This high level of genetic variation makes it difficult to apply markers detected in one population to another population. However, if a single position has multiple markers, at least one of the markers may be polymorphic in a different target population. This allows the linkage-mapping step to be skipped for new target populations because the positions of the co-located polymorphic markers are already known.

QTL were detected for main stem length (Fig. 5; Table 2). The linkage map consisted of 756 loci, and marker density was high. QTL detected on the P1 map were not detected on the P2 map, suggesting that the QTL detected in one population (i.e., the loci heterozygous in one or both parents) may not be detected in another population. That is, QTL efficient in one population may not totally explain the genetic variation of the target trait in another population. This suggests that marker-assisted selection may not be a suitable method for use in common buckwheat populations. Because it is necessary to map QTL on a population-by-
population basis, genome-wide association studies (GWAS) may instead be better to detect QTL that contribute to the genetic variation of a target trait in multiple populations. When undertaking GWAS or multi-family (i.e., population-by-population) QTL mapping, a high-throughput genotyping system is essential. Therefore, the genotyping system proposed in the present study may be useful for GWAS and multi-family QTL mapping.

The results of the present study suggest that genomics-assisted breeding can be used in crop plants for which genomic resources are poor. We conducted linkage mapping and QTL mapping in a breeding population of common buckwheat without using any a priori genomic information or any existing genomic tools. It took eight months to construct the microarray markers in this study. Because we did not use any a priori genomic information in common buckwheat, our results show the timescale in which the genotyping system can be used for the analysis of other crops for which no genomic information is available. When a genotyping system is used for genomic selection (Meuwissen et al. 2001), the genotyping of plants needs to be conducted at every selection cycle. Yabe et al. (2014) conducted genomic selection simulations with open-pollinated plants, and the results suggest that it is important to control pollination among parents in allogamous species. To select parent plants before pollination, it is necessary to obtain their marker genotypes prior to flowering. In this study, it took 37 days (5 days for DNA extraction, 7 days for the Cy3 labeling, 17 days for microarray analysis, 6 days for data analysis) to process 178 samples from DNA extraction to genotyping, suggesting the genotyping system allows us to obtain genotypes of genome-wide markers of a few hundreds of common buckwheat plants before the plants mature.

The marker system used in this study was a microarray-based method. Diversity arrays technology (DArT) is a well-known microarray-based method for genotyping DNA polymorphisms, which can detect a single base pair change in a restriction site. Although DArT has potential to detect insertion, deletion, and rearrangement type DNA polymorphisms (Jacquard et al. 2001), DArT mainly detected restriction site polymorphisms (Wittenberg et al. 2005). The method used in this study can detect insertion, deletion, and rearrangement type DNA polymorphisms as well as restriction site polymorphisms, because a mutation at a single to several nucleotides can be detected accurately by a probe whose length is shorter than a genomic DNA fragment (Enoki et al. 2012). In the present study, we obtained a large number of markers (10,813) showing clear segregation patterns, suggesting that the system has high capability to detect DNA polymorphisms in a common buckwheat population.

Currently, genotyping systems using next generation sequencing, such as restriction site-associated DNA sequencing (RAD-Seq; Baird et al. 2008) and genotyping-by-sequencing (GBS; Elshire et al. 2011), are used in various crops to score a massive number of SNPs for a large number of genotypes. Genotyping using next generation sequencing has several advantages over genotyping using microarrays: (i) it does not require SNP discovery and array design (Davve et al. 2011), (ii) it is free from ascertainment bias attributable to SNP discovery (Heslot et al. 2013), and (iii) it is lower in cost (Elshire et al. 2011). Genotyping using next generation sequencing, however, shows higher rates of missing data than genotyping using microarrays in most cases, and requires a special procedure to estimate markers with missing data (e.g., Poland et al. 2012). The genotyping system used in this study showed a low rate (0.4%) of missing data. Low rate of missing data is necessary to estimate recombination fraction accurately in the linkage mapping. Iehisa et al. (2014) used the same genotyping system to construct a high-density linkage map of common wheat and reported low levels of missing data. The result also suggests a high yield of data from the genotyping system.

Here, we demonstrated the potential of a novel marker system for the linkage and QTL mapping of common buckwheat. Our results suggest that this system can be used as a rapid, high-throughput genotyping technology for genomics-assisted breeding, and that it will be particularly useful for use in crop plants for which no prior genomic information is available. Because of their high marker density, the linkage map constructed in this study will be useful for genomic studies based on other common buckwheat populations. By using the novel marker system, marker genotypes of a few hundreds of plants can be acquired in ca. 40 days, suggesting that the system can be used for genomic selection that requires marker genotypes to be obtained prior to flowering.

### Acknowledgments

This research was supported by cooperative research funds from the TOYOTA MOTOR CORPORATION and partly by JSPS KAKENHI Grant Number 25·6921. We express our deepest gratitude to Yasuko Shirai and Atsuko Ushitani for their technical assistance.

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