

# Integrated Consensus Map of Cultivated Peanut and Wild Relatives Reveals Structures of the A and B Genomes of *Arachis* and Divergence of the Legume Genomes

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Edited by Kazuhiro Sato  
(Received 13 October 2012; accepted 21 December 2012)

## Abstract

**The complex, tetraploid genome structure of peanut (*Arachis hypogaea*) has obstructed advances in genetics and genomics in the species. The aim of this study is to understand the genome structure of *Arachis* by developing a high-density integrated consensus map. Three recombinant inbred line populations derived from crosses between the A genome diploid species, *Arachis duranensis* and *Arachis stenosperma*; the B genome diploid species, *Arachis ipaënsis* and *Arachis magna*; and between the AB genome tetraploids, *A. hypogaea* and an artificial amphidiploid (*A. ipaënsis* × *A. duranensis*)<sup>4×</sup>, were used to construct genetic linkage maps: 10 linkage groups (LGs) of 544 cM with 597 loci for the A genome; 10 LGs of 461 cM with 798 loci for the B genome; and 20 LGs of 1442 cM with 1469 loci for the AB genome. The resultant maps plus 13 published maps were integrated into a consensus map covering 2651 cM with 3693 marker loci which was anchored to 20 consensus LGs corresponding to the A and B genomes. The comparative genomics with genome sequences of *Cajanus cajan*, *Glycine max*, *Lotus japonicus*, and *Medicago truncatula* revealed that the *Arachis* genome has segmented synteny relationship to the other legumes. The comparative maps in legumes, integrated tetraploid consensus maps, and genome-specific diploid maps will increase the genetic and genomic understanding of *Arachis* and should facilitate molecular breeding.**

**Key words:** *Arachis* spp.; comparative genomics; genetic linkage map; integrated consensus map; legume genome

## 1. Introduction

Fabaceae, or Leguminosae, is composed of ~700 genera and 20 000 species.<sup>1</sup> The family is classified into three major subfamilies Mimosoideae, Caesalpinioideae, and Papilionoideae.<sup>2</sup> Papilionoideae is

divided into four clades Genistoids, Dalbergioids, Phaseoloids, and Galegoids. Most tropical and cool season legumes used as crops and studied as model plants for symbiosis of rhizobium are members of the Phaseoloids, e.g. genera *Cajanus*, *Glycine*, *Phaseolus*, and *Vigna*, or the Galegoids, e.g. genera *Lotus*,

*Medicago*, *Pisum*, *Trifolium*, and *Vicia*.<sup>2</sup> Therefore, legume genetics and genomics, including genome sequencing and comparative genomics, have greatly advanced in the Phaseoloids and Galegoids.<sup>3–9</sup>

The Dalbergioids, which include the genus, *Arachis*, are more basal in their divergence within the Papilionoideae than the Phaseoloids and Galegoids. The genus *Arachis* comprises 80 species ( $2n = 2x = 18$ ,  $2n = 2x = 20$ , and  $2n = 4x = 40$ ) possessing A, B, C, D, E, F, K, R, and T genomes.<sup>10–12</sup> As regarding the A genome diploids consisting of 15 wild species, they are morphologically similar, cross compatible, produce fertile hybrids with near normal chromosome pairing, and have a common karyotype structure.<sup>13</sup> The B-genome diploids also have similar morphological and chromosome feature, cross-compatibility, bivalent formation at meiosis in interspecific hybrids, and the same karyotype structure.<sup>12</sup> Peanut (*Arachis hypogaea*), or groundnut, is an autogamous allotetraploid legume ( $2n = 4x = 40$ ) harbouring homoeologous A and B genomes that are derived from two diploids, most likely *Arachis duranensis* (A genome) and *Arachis ipaënsis* (B genome).<sup>14,15</sup> While peanut is an important source of food and oil, the genetics and genomics of this legume lag far behind those of the Phaseoloids and Galegoids due to a complex genome structure with challenging features, such as tetraploidy, low genetic diversity, and a relatively large genome size of 2.8 Gb.<sup>16</sup>

Molecular genetic studies of *Arachis* initially progressed by using diploid rather than tetraploid peanut due to the greater simplicity of diploids as genetic models.<sup>17</sup> The first genetic linkage map in genus *Arachis* was developed based on an interspecific crossing between diploids *Arachis stenosperma* (A genome) and *Arachis cardenasii* (also A genome),<sup>17</sup> followed by a cross between *A. duranensis* and *A. stenosperma* (both A genomes)<sup>18,19</sup> and between *A. ipaënsis* and *Arachis magna* (both B genomes).<sup>20</sup> However, because of the small number of available DNA markers at the time, the number of mapped marker loci was limited to between 117 and 369. Recently, intraspecific maps for A-genome diploid (*A. duranensis*) and B-genome diploid (*Arachis batizocoi*) have been developed with 1724 and 449 marker loci, respectively, including single nucleotide polymorphisms and expressed sequence tag-simple sequence repeats (EST-SSRs) markers.<sup>21,22</sup>

Meanwhile, an artificial amphidiploid derived from a three-way cross between *A. batizocoi* (B genome) and a hybrid between *A. cardenasii* (A genome) and *A. diogeni* (A genome) has been crossed with *A. hypogaea* to introduce genetic diversity from wild diploid species into the tetraploid-cultivated peanut. The BC<sub>1</sub> progenies of the cross have contributed to the generation of a tetraploid genetic linkage map

with 370 restriction-fragment length polymorphisms.<sup>23</sup> In addition, another artificial amphidiploid line, derived from a hybrid between *A. duranensis* and *A. ipaënsis*, was used to establish a genetic linkage map made up of 298 SSR marker loci.<sup>24</sup> In the latter map, the linkage groups (LGs) were firstly anchored to the A and B genomes by identifying the genome origins of the mapped loci in the LGs based on the corresponding sizes of the DNA amplified from the original diploids. In the tetraploid-cultivated peanut (*A. hypogaea*), the population-specific linkage maps comprise 318 marker loci. Due to low genetic diversity,<sup>25–33</sup> this is the maximum number of marker loci for population-specific linkage maps, although >6000 SSR markers have been developed for mapping.<sup>34</sup> By integrating the 11 linkage maps of the artificial amphidiploids and cultivated tetraploid peanuts, a reference consensus map consisting of 897 marker loci was constructed and annotated with consensus nomenclature for the LGs (a01–a10 and b01–b10).<sup>35</sup>

Concurrent with the progress of the above genetic studies in *Arachis* spp., a high-density linkage map of cultivated peanut covering 2166 cM with 1114 marker loci has also been reported.<sup>36</sup> Because most of the markers on this map were published after the development of the other *Arachis* maps,<sup>36–38</sup> the markers do not appear on them.<sup>17–20,23–32</sup> In addition, due to an insufficient number of commonly mapped loci in the published diploid maps, the LGs have not been assigned to either A or B genome types, even though homoeologous LGs have been identified.<sup>36</sup> Anchoring the genetic LGs to the A and B genomes will enable enrichment of marker density on the consensus and diploid maps.<sup>19,20,35</sup> This would contribute to a better understanding of genome evolution within *Arachis* and among the Fabaceae. In addition, it would serve to increase the efficiency of introgression of desirable alleles from diploid wild species into the tetraploid-cultivated peanut through marker-assisted selection.

Comparative analysis of the A genome diploid *Arachis* species, e.g. *A. duranensis* and *A. stenosperma*, and model legumes has previously been performed. This comparative analysis revealed candidate synteny blocks between the genomes of *Lotus japonicus*, *Medicago truncatula*, and *Glycine max*.<sup>21,39</sup> To date, no comparative analysis of cultivated peanut and other legume species has been reported. Comparison of the tetraploid *Arachis* genome, represented by the high-density consensus map of the genomes of legumes *Cajanus cajan*, *G. max*, *L. japonicus*, and *M. truncatula*, would provide further insights into the legume genomes.

To address the above issues, it was considered that high-density linkage maps of the tetraploid and

diploid species, on which common makers were mapped, would be required. Therefore, we tried to map the large number of DNA markers reported for the two cultivated peanut maps published by Shirasawa *et al.*<sup>36</sup> onto the previously reported linkage maps of the A and B genomes,<sup>19,20</sup> along with that of an artificial amphidiploid derived from a hybrid of diploid species. The established linkage maps were integrated with 13 reported tetraploid *Arachis* maps to increase the number of mapped loci on the previously published consensus map.<sup>35</sup> Subsequently, the maps were subjected to comparative analysis with four legume genomes, *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula*, to clarify features of the genome structure of the genus *Arachis*.

## 2. Materials and methods

### 2.1. Plant materials

In previous studies, two F<sub>2</sub> mapping populations, derived from interspecific crosses between the two A genome diploid species, *A. duranensis* 'K7988' and *A. stenosperma* 'V10309', and between the two B genome diploid species, *A. ipaënsis* 'K30076' and *A. magna* 'K30097', were used to construct genetic linkage maps.<sup>19,20</sup> In the present study, the F<sub>5</sub> and F<sub>6</sub> progenies of the diploid A and B genome mapping populations, respectively, were generated by single seed descent, and used as recombinant inbred mapping populations. The A and B genome mapping populations were named as AF5 ( $n = 89$ ) and BF6 ( $n = 94$ ), respectively. In addition, other recombinant inbred lines were used to construct AB genome tetraploid ( $n = 91$ : population TF6) maps. This population consisted of F<sub>6</sub> lines derived from a cross between *A. hypogaea* 'Runner IAC 886' and an artificial amphidiploid (*A. ipaënsis* × *A. duranensis*)<sup>4×</sup>, which was developed by hybridizing *A. ipaënsis* 'K30076' and *A. duranensis* 'V14167'. Genomic DNA from each line was extracted from young leaflets essentially as described by Grattapaglia and Sederoff.<sup>40</sup>

### 2.2. Polymorphism analysis with DNA markers

A total of 3902 DNA markers comprising 1894 genomic SSRs *A. hypogaea* genomic SSR (AHGS),<sup>36</sup> 1571 EST-SSRs *A. hypogaea* EST-SSR (AHS),<sup>37</sup> and 437 transposon-based markers *A. hypogaea* transposable element (AhTE)<sup>36,38</sup> were screened against the parental lines of the populations AF5, BF6, and TF6 using an ABI-3730xl fluorescent fragment analyser (Applied Biosystems). Polymorphic markers were analysed on the three mapping populations using 10% polyacrylamide gels for markers exhibiting ≥10-bp allele size differences, or the fluorescent fragment analyser for markers exhibiting <10-bp allele

size differences between the parental lines. PCR, electrophoresis, and data scoring were performed as described previously.<sup>36–38</sup>

### 2.3. Construction of linkage maps and the integrated map

Linkage analysis of segregated genotypic data obtained in this study was performed together with linkage analysis of the previously investigated genotypes with mapped markers on the diploid maps<sup>19,20</sup> using JoinMap<sup>®</sup> version 4.<sup>41</sup> The marker loci were roughly classified using the JoinMap<sup>®</sup> Grouping Module with logarithm of odds (LOD) scores of 4.0–10.0. The JoinMap<sup>®</sup> Combine Groups for Map Integration Module was used to integrate the linkage maps developed in this study and the 13 previously published maps into a tetraploid map, i.e. the cultivated peanut map<sup>25–32,36</sup> and an artificial amphidiploid map.<sup>24</sup> Marker order and genetic distance were calculated using a regression mapping algorithm with the following parameters: Haldane's mapping function, recombination frequency ≤0.30, and LOD score ≥2.0. The graphical linkage maps were drawn using the MapChart program.<sup>42</sup>

### 2.4. Comparative analysis of the *Arachis* maps with other legume genomes

BLASTN<sup>43</sup> was used to conduct similarity searches (using the nucleotide sequences from which the mapped DNA markers were designed) against each pseudomolecule of the genome sequences of *C. cajan*,<sup>8</sup> *G. max* (Glyma1),<sup>6</sup> *L. japonicus* (build 2.5),<sup>5</sup> and *M. truncatula* (Mt3.5v4),<sup>7</sup> with a threshold *E*-value of 1e-20. The graphical comparative maps were drawn using the Circos program.<sup>44</sup>

## 3. Results

### 3.1. Polymorphism screening of the parental lines of the mapping populations and construction of the linkage maps

Prior to polymorphism analysis between the parental lines, a total of 3902 markers, i.e. 1894 AHGS, 1571 AHS, and 437 AhTE markers, were pre-selected from the 6680 AHGS, 3187 AHS, and 1039 AhTE markers that have been published.<sup>36–38</sup> The markers were pre-selected according to the following criteria: (i) AHGS markers were selected if they were expected to detect polymorphism among the cultivated peanut lines according to the results of *in silico* polymorphism analysis;<sup>36</sup> (ii) AHS markers were selected if they showed polymorphisms between cultivated lines and wild species;<sup>37</sup> (iii) AhTE markers were selected if they amplified single or double bands.<sup>36,38</sup>

**3.1.1. The AF5 population** A total of 582 markers (14.9% = 582/3902), including 450 AHGS (23.8% = 450/1894), 109 AHS (6.9% = 109/1571), and 23 AhTE (5.3% = 23/437) markers, were selected as polymorphism candidates between the parental lines of the AF5 population, *A. duranensis* 'K7988' and *A. stenosperma* 'V10309'. Out of the 582 polymorphic markers, 395 generated a total of 437 segregation loci, since several markers detected more than one locus. Specifically, 359, 31, 4, and 1 marker(s) detected 1, 2, 3, and 4 polymorphic loci, respectively, which were suffixed with '\_a1' and '\_a2', e.g. AHGS1647\_a1 and AHGS1647\_a2, to distinguish the loci each other. The 437 polymorphic loci, 284 codominant, and 153 dominant loci, were successfully mapped onto 10 LGs covering 544 cM, together with 160 previously mapped polymorphic loci from 158 markers (Fig. 1, Table 1, and Supplementary Table S1).<sup>19</sup> The average marker density of the map was 0.9 cM, and segregation distortions were observed in 65% (390 loci) of the mapped loci (Table 1).

**3.1.2. The BF6 population** A total of 862 markers (22.1% = 862/3902), including 513 AHGS (27.1% = 513/1894), 279 AHS (17.8% = 279/1571), and 70 AhTE (16.0% = 70/437) markers, were selected as polymorphism candidates between the parental lines of the BF6 population, *A. ipaënsis* 'K30076' and *A. magna* 'K30097'. Out of the 862 polymorphic markers, 624 generated a total of 680 segregation loci, since, as with the AF5 population, several of these markers detected more than one locus. Specifically, 579, 35, 9, and 1 marker(s) detected 1, 2, 3, and 4 polymorphic loci, respectively, which were suffixed with '\_b1' and '\_b2', e.g. AHGS1478\_b1 and AHGS1478\_b2, to distinguish the loci each other. The 680 polymorphic loci, 549 codominant, and 131 dominant, were successfully mapped onto 10 LGs covering 461 cM together with 118 previously mapped loci from 116 markers (Fig. 1, Table 1, and Supplementary Table S1).<sup>20</sup> The average marker density of the map was 0.6 cM, and segregation distortions were observed in 31% (250 loci) of the mapped loci (Table 1).

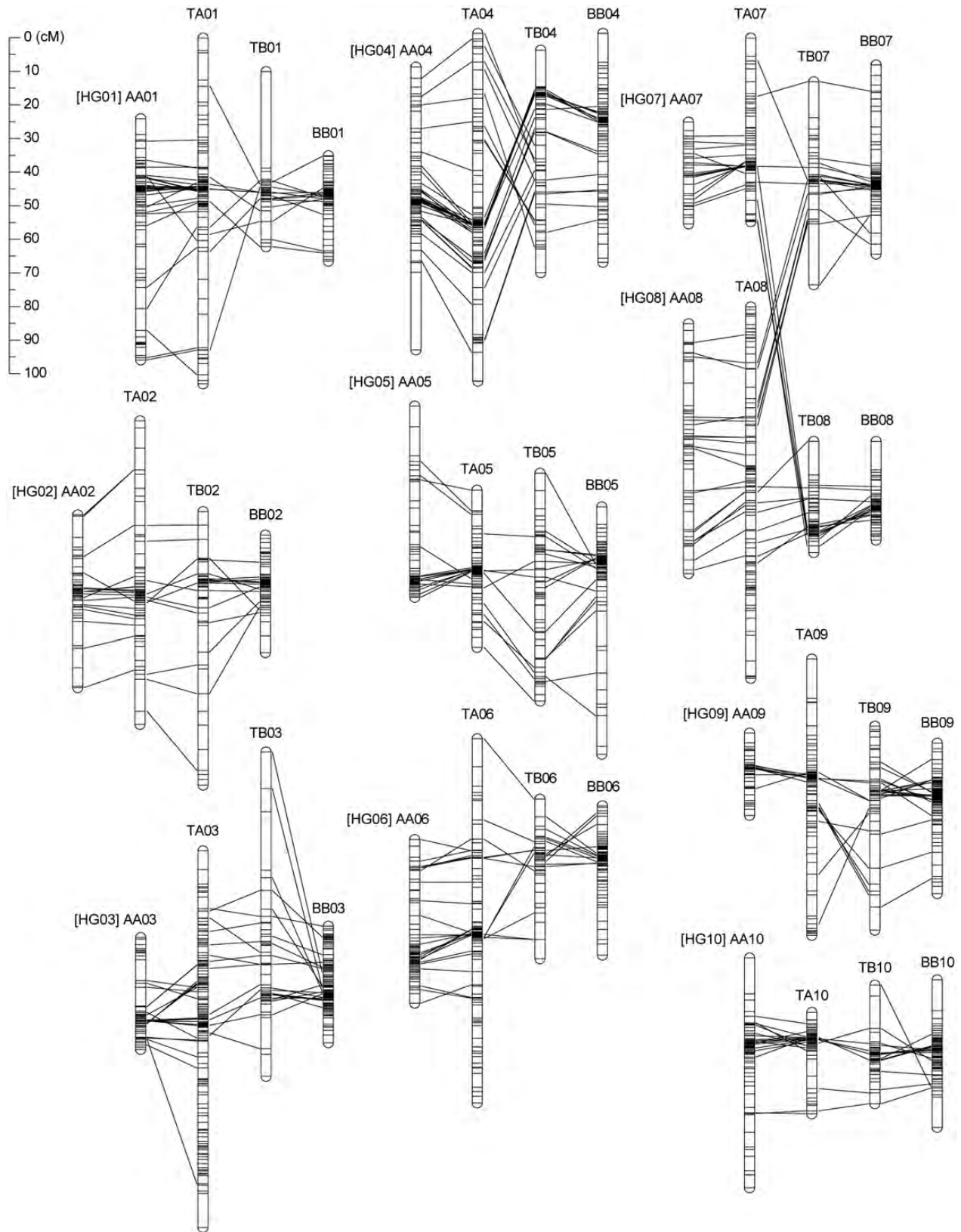
**3.1.3. The TF6 population** Together with the parental lines of TF6, the two original diploids of the artificial amphidiploid, *A. ipaënsis* 'K30076' and *A. duranensis* 'V14167', were also genotyped to identify the genome origin (A or B genome) of the polymorphic DNA fragments. A total of 1144 markers (29.3% = 1144/3902), including 664 AHGS (35.1% = 664/1894), 257 AHS (16.4% = 257/1571), and 223 AhTE (51.0% = 223/437) were selected as polymorphism candidates between the parental lines of the TF6 population, *A. hypogaea*

'Runner IAC 886' and an artificial amphidiploid (*A. ipaënsis* 'K30076' × *A. duranensis* 'V14167')<sup>4×</sup>. Out of the 1144 polymorphic markers, 1055 generated a total of 1261 segregation loci, since, as with the AF5 and BF6 populations, several of these markers detected more than one locus. Specifically, 879, 150, 22, and 4 markers detected 1, 2, 3, and 4 polymorphic loci, respectively, which were suffixed with '\_t1' and '\_t2', e.g. AHGS1991\_t1 and AHGS1991\_t2, to distinguish the loci each other. The 1261 polymorphic loci, 970 codominant, and 291 dominant, were successfully mapped onto 20 LGs covering 1442 cM together with 208 loci from 180 markers (Moretzsohn *et al.* unpublished data). The 20 LGs were classified into 10 pairs of A and B genomes, according to the DNA fragment size of mapped loci corresponding to 'K30076' or 'V14167' (Fig. 1, Table 1, and Supplementary Table S1). The average marker density of the map was 1.0 cM, and segregation distortions were observed in 19% (285 loci) of the mapped loci (Table 1).

In total, the number of non-redundant AHGS, AHS, and AhTE markers mapped on the three maps were 890, 446, and 211, respectively. Out of the total 1547 markers, 73 (67 AHGSs and six AHSs) and 381 (298 AHGSs, 59 AHSs, and 24 AhTEs) were mapped on three and two maps, respectively, while 1093 (525 AHGSs, 381 AHSs, and 187 AhTEs) were mapped on just one map; the AF5, BF6, or TF6 map (Supplementary Table S1). Comparing the three maps, 203 of the 597 mapped loci on the AF5 map, and 157 out of 798 loci on the BF6 map, were also located on the TF6 map (Supplementary Table S1).

The LGs were designated AA, BB, and TA or TB in the AF5, BF6 and TF6 maps, respectively. The nomenclature of the LGs of the diploid maps was tentatively determined according to the commonly mapped marker loci between the present and the previously constructed maps.<sup>19,20</sup> Eight LGs of the AF5 map (AA01 and AA03–AA09) and all 10 of the LGs of the BF6 map (BB01–BB10) were assigned to the corresponding LGs on the previous maps (Supplementary Table S2). The two LGs of the AF5 map were assigned to 'Group 02' in the previous map,<sup>19</sup> and they were tentatively named AA02 and AA10 (Table 1 and Supplementary Table S2). 'Group 10' in the previous map<sup>19</sup> was disassembled on the AF5 map. On the other hand, in the TF6 map, TA01, TA03–TA09, and TB01–TB10 were assigned to the corresponding LGs of the AF5 and BF6 maps, and TA02 and TA10 were assigned to the AA02 and AA10 LGs, respectively (Table 1 and Supplementary Table S2). After integrating all of these data, 10 homoeologous groups (HGs) were identified on the TF6 map.

The marker order was almost completely conserved between the HGs except for HG04, HG07, and HG08



**Figure 1.** Genetic linkage maps obtained for AF5, BF6, and TF6 populations. The linkage groups in the AF5, BF6, and TF6 maps are indicated by AA, BB, and TA or TB, respectively, and the homoeologous groups are prefixed with HGs. Homologous or homoeologous marker loci are connected by lines.

**Table 1.** Descriptions of the genetic linkage maps AF5, BF6, and TF6

LGs	AF5 map				LGs	BF6 map				LGs	TF6 map			
	Locus no.	cM	cM/locus	Percentages of loci with segregation distortion <sup>a</sup>		Locus no.	cM	cM/locus	Percentages of loci with segregation distortion <sup>a</sup>		Locus no.	cM	cM/locus	Percentages of loci with segregation distortion <sup>a</sup>
AA01	84	71.8	0.9	69	—	—	—	—	TA01	108	102.9	1.0	36	
—	—	—	—	—	BB01	70	31.4	0.5	46	TB01	47	52.1	1.1	17
AA02	55	51.3	1.0	20	—	—	—	—	TA02	73	30.2	0.4	33	
—	—	—	—	—	BB02	62	34.9	0.6	92	TB02	47	35.3	0.8	36
AA03	63	33.3	0.5	78	—	—	—	—	TA03	154	112.1	0.7	4	
—	—	—	—	—	BB03	113	34.7	0.3	8	TB03	50	96.5	2.0	26
AA04	75	84.2	1.1	97	—	—	—	—	TA04	105	103.5	1.0	7	
—	—	—	—	—	BB04	85	68.1	0.8	48	TB04	62	66.2	1.1	23
AA05	56	56.9	1.0	82	—	—	—	—	TA05	81	46.7	0.6	23	
—	—	—	—	—	BB05	75	73.5	1.0	13	TB05	60	67.7	1.1	8
AA06	63	48.7	0.8	65	—	—	—	—	TA06	67	90.2	1.4	7	
—	—	—	—	—	BB06	84	44.4	0.5	35	TB06	49	81.3	1.7	55
AA07	31	30.3	1.0	16	—	—	—	—	TA07	70	54.6	0.8	20	
—	—	—	—	—	BB07	78	56.3	0.7	31	TB07	40	60.4	1.5	20
AA08	54	74.4	1.4	28	—	—	—	—	TA08	98	110.4	1.1	16	
—	—	—	—	—	BB08	61	29.4	0.5	11	TB08	50	33.1	0.7	14
AA09	48	24.5	0.5	100	—	—	—	—	TA09	96	82.2	0.9	3	
—	—	—	—	—	BB09	91	44.8	0.5	35	TB09	56	60.7	1.1	18
AA10	68	68.4	1.0	65	—	—	—	—	TA10	109	108.4	1.0	39	
—	—	—	—	—	BB10	79	43.9	0.6	11	TB10	47	47.5	1.0	2
Total	597	543.8	0.9	65	Total	798	461.4	0.6	31	Total	1469	1441.8	1.0	19

<sup>a</sup>Percentages of loci that showed segregation distortion ( $P < 0.01$ ).

(Fig. 1). In HG04, the lower part of TA04 corresponded to the upper part of TB04 (Fig. 1). In HG07, the loci mapped onto TB07 corresponded to those of TA07 and the upper part of TA08. Moreover, corresponding loci on TB08 were observed on the lower parts of TA08 and TA07 (Fig. 1), indicating a translocation between the A and B genomes as previously suggested.<sup>20,22,24</sup>

### 3.2. Integration of the genetic linkage maps into a consensus map

An integrated consensus map was constructed based on the segregation genotypes of 16 populations: three from this study and 13 from previous studies (Supplementary Table S2).<sup>19,20,24–32,36</sup> The integrated consensus map was 2651 cM in total length and comprised 20 LGs, on which 3693 loci, including 1564 AHGS, 569 AHS, 450 AhTE, and 1110 other loci, were mapped (Table 2, Fig. 2, and Supplementary Table S3), out of which markers mapped on more than two loci were suffixed with ‘\_c1’ and ‘\_c2’, e.g. AHGS1403\_c1 and AHGS1403\_c2, to distinguish each other. The average marker density of this map

was 0.7 cM (Table 2). On the consensus map, 391 marker loci were commonly mapped onto pairs of LGs in each HG, and the order of the markers was roughly conserved (Fig. 2).

In the present study, locus clusters were defined as regions in which  $\geq 10$  loci were mapped in 5 cM windows. They were observed in all of the LGs (Fig. 2, Table 2, and Supplementary Table S3). The total length of the locus clusters was 505 cM (101 clusters), varying from 15 cM (three clusters) in A06 and B05 to 40 cM (eight clusters) in A03 and B03. A total of 2319 loci, representing 62.8% of the mapped loci, were located in 101 clusters, which varied in loci content from 53 (A08) to 175 (A03) loci.

### 3.3. Comparative mapping between the *Arachis* spp. and the four comparison legumes, *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula*

Corresponding sequences were available for 3473 of the 3693 mapped loci (Supplementary Table S3). Of these 3473 sequences, 869 showed significant similarity to one of the four legume genome

**Table 2.** Descriptions of the integrated consensus map and integrated LGs of the population-specific maps

LGs	Locus no.	cM	cM/locus	Length of the locus cluster <sup>a</sup>	Locus no. in the marker cluster
A01	238	183.8	0.8	25	158
B01	168	84.7	0.5	25	129
A02	126	78.6	0.6	20	86
B02	154	112.0	0.7	25	104
A03	272	151.6	0.6	40	175
B03	234	144.8	0.6	40	167
A04	213	106.8	0.5	20	106
B04	208	132.1	0.6	30	122
A05	189	126.0	0.7	30	135
B05	169	112.7	0.7	15	77
A06	187	136.4	0.7	15	85
B06	176	152.5	0.9	20	109
A07	145	167.0	1.2	20	76
B07	162	179.1	1.1	25	105
A08	167	126.3	0.8	20	53
B08	176	107.3	0.6	30	138
A09	194	126.5	0.7	30	141
B09	179	170.5	1.0	30	121
A10	172	131.1	0.8	20	117
B10	164	121.3	0.7	25	115
Total	3693	2651.1	0.7	505	2319

<sup>a</sup>Regions in which  $\geq 10$  loci were mapped in 5 cM windows.

sequences (*C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula*). For each legume species, 515, 781, 436, and 419 sequences showed similarities to sequences in the *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula* genomes, respectively, and 170 sequences showed similarity to sequences in all four legume genomes.

Segmental syntenic regions were observed between the *Arachis* HGs and the genomes of the four comparison legumes (Fig. 3, Table 3, and Supplementary Table S3). Most of the LG pairs in each HG showed similar synteny relationships to those in the four comparison legumes. Between the four comparison legume species, the genome of *G. max* showed the densest synteny of LGs, corresponding to those of *Arachis*. In addition, most of the LGs in *Arachis* were identified as shared syntenies with LGs on each of two chromosomes of *G. max*. This might reflect the palaeopolyploid nature of the genomes of these species. Comparing *Arachis* with the other three legume genomes, clearly syntenic segments were observed as follows: *Arachis* HG01 shared synteny with *C. cajan* chromosome 11 (Cc11) and *M. truncatula* chromosome 4 (Mt04); *Arachis* HG04 shared synteny with *L. japonicus* chromosome 4 (Lj04);

*Arachis* HG05 shared synteny with Cc06, Lj02, and Mt05; *Arachis* HG06 shared synteny with Lj01 and Mt07; *Arachis* HG07 shared synteny with Cc02, Lj02, and Mt05; *Arachis* HG08 shared synteny with Cc03; *Arachis* HG09 shared synteny with Cc02, Lj05, and Mt01; and *Arachis* HG10 shared synteny with Lj01. On *Arachis* HG03, segmental blocks were identified that showed shared synteny between the comparison legume genomes, while no obvious synteny was observed between *Arachis* HG02 and the genomes of *C. cajan*, *L. japonicus*, and *M. truncatula*.

## 4. Discussion

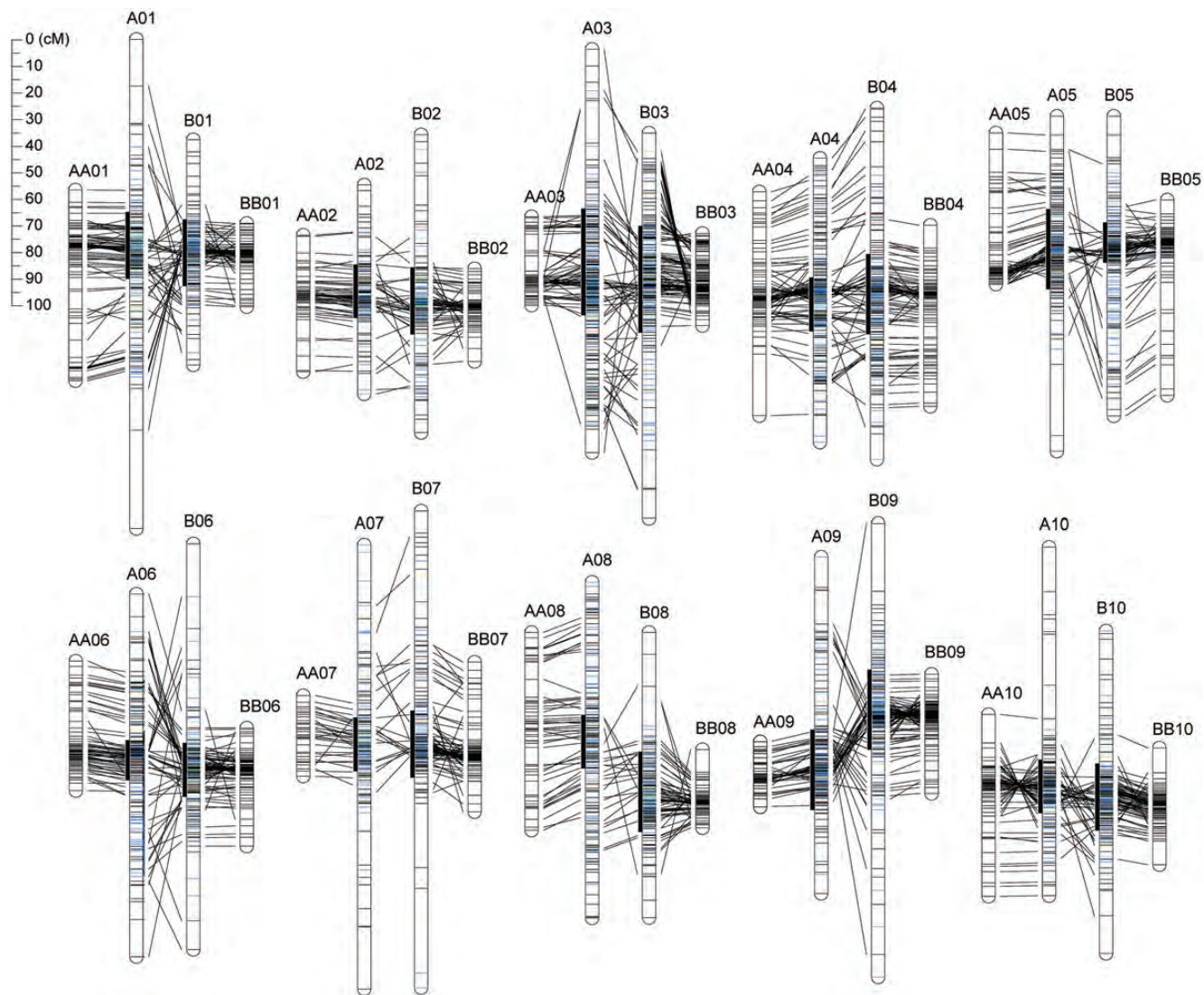
### 4.1. Polymorphism potential of the DNA markers mapped on the genetic linkage map

The polymorphism potential of DNA markers in *Arachis* is usually evaluated on the basis of the polymorphism information content (PIC) value.<sup>34</sup> While the PIC values of the AHS were estimated from genotyping data from 16 *Arachis* lines,<sup>37</sup> those of the AHGS and AhTE have not been investigated thoroughly.<sup>36,38</sup> In the present study and in previous studies,<sup>36</sup> 968 (416 AHGS, 395 AHS, and 157 AhTE), 666 (392 AHGS, 65 AHS, and 209 AhTE), 245 (197 AHGS, 10 AHS, and 38 AhTE), 73 (72 AHGS and 1 AhTE), and 3 (AHGS only) markers were mapped on single, double, triple, quadruple, and quintuple maps, respectively. Of these, the PIC values of the 395, 65, and 10 AHS markers were estimated to be, on average, 0.23, 0.28, and 0.36, respectively, from the results of Koilkonda *et al.*<sup>37</sup> A positive correlation was observed between the PIC values and the number of maps on which the AHS was mapped. Therefore, if this estimation can be generalized, the AHGS and AhTE markers mapped on multiple maps can be considered as potential sources for obtaining polymorphic markers in other mapping populations.

In the screening of polymorphic markers between the parental lines of the mapping populations, 582, 862, and 1144 markers were selected as polymorphism candidates, but 187, 238, and 89 were excluded from the mapping analysis. The excluded markers might amplify DNAs from not the target locus but non-allelic repetitive and putative paralogous sequences in the *Arachis* genome,<sup>38,45,46</sup> which are not suitable for segregation and linkage analysis.

### 4.2. Genetic linkage maps for three mapping populations

Comparison of the AF5, BF6, and TF6 maps revealed that the marker order was almost perfectly conserved between the HGs, with the exception of HG04, HG07, and HG08 (Fig. 1 and Supplementary Fig. S1). This result indicated that the A and B genomes of *Arachis*



**Figure 2.** Comparison of the integrated consensus tetraploid *Arachis* map with diploid maps. The abbreviations AA and BB indicate LGs on the AF5 and BF6 maps, respectively, and the prefixes A and B show LGs on the integrated consensus map. Horizontal lines on the LGs indicate the positions of the mapped loci; the loci derived from one, two, three, four, five, and more than five genetic linkage maps are shown in black, blue, light blue, green, yellow, and red. Vertical bars on the left side of the LGs indicate the locus clusters. Homologous and homoeologous loci are connected by black lines.

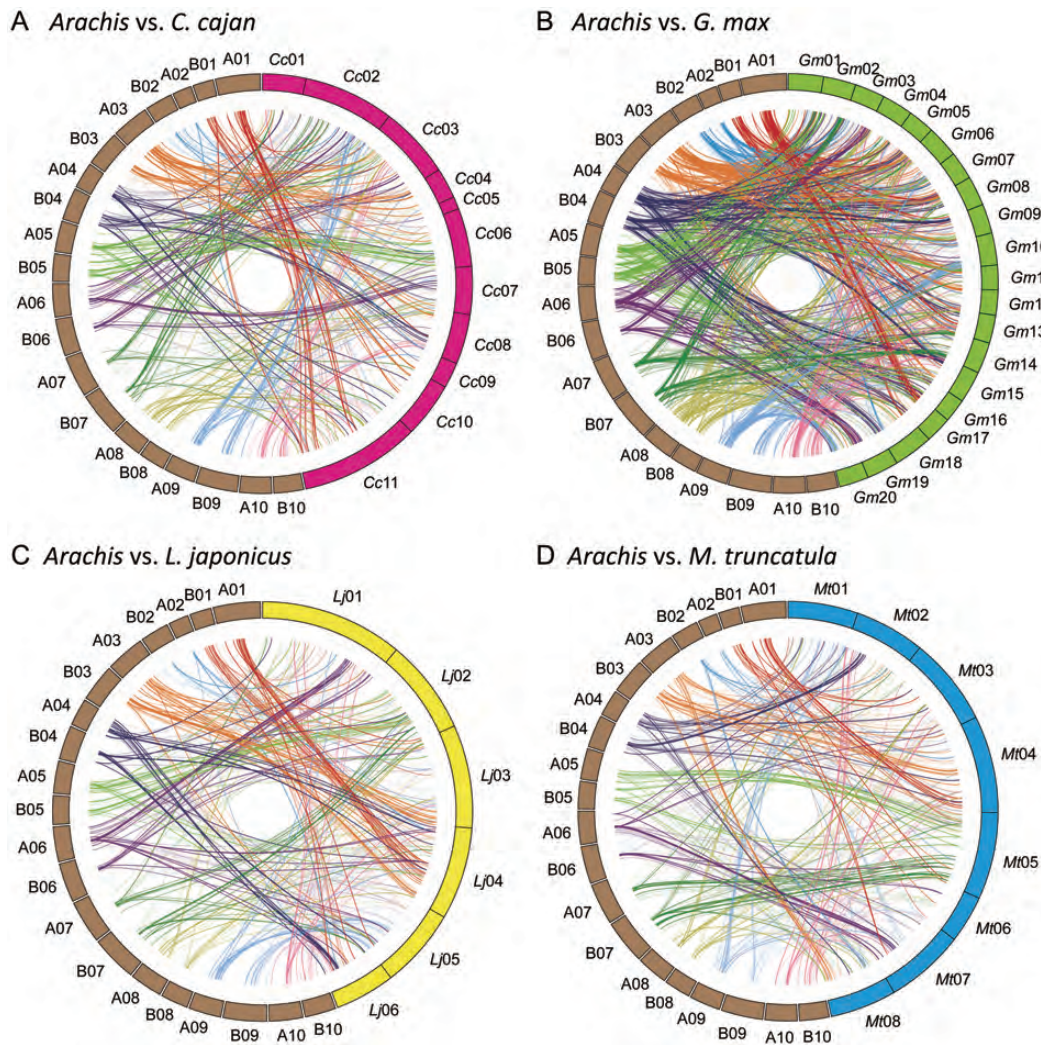
evolved from a common ancestor species. Exceptions to this shared marker order were found within HG04, and between HG07 and HG08. The former was observed for the first time in this study, while the latter confirms previous observations.<sup>20,24</sup> These findings suggest that the chromosome translocations might have occurred at, or after, the divergence of the A and B genomes in *Arachis*.

The AF5 and BF6 linkage maps have been improved by subsequent mapping of the newly developed markers after they were originally published.<sup>19,20</sup> However, despite the increase in the number of mapped loci from 170 to 597 in the AF5 map and from 149 to 798 in the BF6 map, the total map length decreased from 1231 to 544 cM in the AF5

map and from 1294 to 461 cM in the BF6 map. In the maps generated in the present study, the marker loci that mapped to the ends of LGs in previous studies<sup>19,20</sup> were excluded, and the intervals between the markers were shorter than those in previous maps. Mapping telomere-based makers will clarify whether the maps from this study have been saturated or not.

In the linkage map for the TF6 population, the total length of the map was 1442 cM, which was shorter than that of the maps reported for *Arachis* tetraploids (2210 cM with 370 loci,<sup>23</sup> 2166 cM with 1114 loci,<sup>36</sup> 1844 cM with 298 loci;<sup>24</sup> 1785 cM with 191 loci<sup>29</sup>). Nagy *et al.*<sup>21</sup> reported that the length of the genetic linkage map in *A. duranensis* ranged from





**Figure 3.** Comparative maps of *Arachis* and four other legumes, *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula*. Circled bars in brown, magenta, green, yellow, and blue indicate the genomes of *Arachis*, *C. cajan* (A), *G. max* (B), *L. japonicus* (C), and *M. truncatula* (D), respectively. Homologous loci are connected by lines.

**Table 3.** Similarity of other legume chromosomes with that of *Arachis* homoeologous groups

<i>Arachis</i>	<i>C. cajan</i>	<i>G. max</i>	<i>L. japonicus</i>	<i>M. truncatula</i>
HG01 <sup>a</sup>	11	5, 7, 13, 17	4, 6	4
HG02	—	3, 7	—	—
HG03	4, 5, 8, 9, 10	4, 5, 6, 8, 9, 11, 12, 18	3, 4	4, 8
HG04	7, 8	3, 7, 8, 9, 12, 13, 15, 18, 20	6	2, 7
HG05	6	1, 9, 11, 16	2	5
HG06	2, 3, 7	2, 3, 16, 19	1	7
HG07	2	2, 6, 13	2	5
HG08	3	4, 6, 8, 12, 13	1, 3	2, 3
HG09	2	7, 10, 20	5	1
HG10	1, 8	10, 15, 19	1	2, 3

<sup>a</sup>HG indicates homologue group.

1081.1 to 2056.5 cM (depending on the mapping program used) when 1054 segregation loci were used for the linkage analysis. Therefore, the different programs, algorithms, and functions used for mapping may explain the differences in map length reported by the present and previous studies. Our preliminary analysis, in which MapMaker program<sup>47</sup> gave longer map lengths than the JoinMap, also supported these explanations. Alternatively, as Sim *et al.*<sup>48</sup> suggested (based on linkage analysis in tomato), different ratios between the number of markers and population size in each population might have resulted in the length differences.

Segregation distortions were observed in the three maps, the AF5, BF6, and TF6, and the ratios were different among the populations and LGs (Table 1). Similar observations were found in our previous studies on tomato on which high and low ratios of

segregation distortion were observed in inter- and intraspecific maps, respectively.<sup>49,50</sup> Segregation distortion is generally caused by the chromosomal structural differences or the presence of transmission distortion factors on some chromosome.<sup>51</sup> Our findings will contribute to identify factors for the segregation distortion.

#### 4.3. Characterization of the cultivated peanut genome using the *Arachis* consensus map

The consensus map was constructed using segregation data from 16 populations, in which five species, *A. duranensis* (A), *A. stenosperma* (A), *A. ipaënsis* (B), *A. magna* (B), and *A. hypogaea* (AB), were involved (Supplementary Table S2). A total of 889 loci derived from 421 markers were commonly mapped onto all of the paired LGs as HGs; however, the locus orders were not always conserved. The regions in which mapped loci showed non-corresponding orders between the paired LGs in each HG were considered as candidate regions having A or B genome-specific structures. Most of the locus orders were conserved between AA (diploid) and A (tetraploid) LGs, or between BB (diploid) and B (tetraploid) LGs (Fig. 2). However, several regions showed evidence of rearrangement, such as between BB03 and B03, BB09 and B09, and AA10 and A10. *Arachis duranensis* and *A. ipaënsis* are considered to be the most probable ancestors of *A. hypogaea*, because of similar karyotypes of *A. duranensis* and *A. ipaënsis* to that of *A. hypogaea*.<sup>12–15</sup> By contrast, even though the structures of A genome of different species are well conserved, the *A. stenosperma* genome is distinguishable from that of *A. duranensis* on the basis of the variability observed in the heterochromatin and 18S–26S rRNA loci.<sup>13</sup> It is considered the possibility that such differences in the genome structure of the parental lines of the mapping populations might disrupt the marker order between diploid and tetraploid consensus maps.

The average marker density of the consensus map was 0.7 cM/locus (Table 2). Because the genome size of *A. hypogaea* is estimated to be ~2.8 Gb,<sup>16</sup> the DNA markers were located at <760 kb intervals on average. In other words, a map-based cloning strategy has become a realistic approach for molecular genetics in *Arachis* spp., because a candidate region for the location of a target gene can, on average, be identified to within 760 kb. Furthermore, this high-density map should aid in genome sequencing analysis in *Arachis* spp., because the locus order can work as a reference for the orientation of sequence contigs and scaffolds. Moreover, the marker clusters were observed in all of the LGs (Fig. 2 and Supplementary Table S3). Because chromosomal

crossover, and the resulting genetic recombination, are severely suppressed in heterochromatin,<sup>6,49</sup> the clustered regions were speculated to be located in heterochromatin, which makes up 10–20% of the *Arachis* chromosomes in somatic metaphase.<sup>13,15</sup> In the case of soybean, broad pericentromeric regions show markedly diminished recombination (4.2 Mb/cM), while the euchromatic regions at the chromosome ends maintain consistent levels of recombination (193 kb/cM).<sup>6</sup> This pattern is also observed in tomato, with 6042 and 172 kb/cM in heterochromatic and euchromatic regions, respectively.<sup>49</sup> As with soybean and tomato, it was predicted that the physical marker-to-marker distance intervals in the peanut genome were different between heterochromatic and euchromatic regions.

#### 4.4. Comparative genomics of *Arachis* spp. with *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula*

All of the *Arachis* HGs showed similarities to the genomes of the four legumes (Fig. 3, Table 3, and Supplementary Table S2). The genetic composition of *Arachis* corresponded to that of the *C. cajan*, *L. japonicus*, and *M. truncatula* genomes on a one-on-one basis, but was doubled when compared with the *G. max* genome. These observations support the estimation that a whole-genome duplication occurred in legumes ~58 million years ago (mya), at the time of the Papilionoid origin,<sup>52</sup> whereas the soybean-specific duplication is estimated to have occurred <13 mya.<sup>6</sup> These insights are based on a comparison of the A genome species between *Arachis* and *L. japonicus* and *M. truncatula*.<sup>39</sup>

At the chromosome level, each chromosome of *Arachis*, which belongs to the Dalbergioids, might have evolved differently and show different rearrangements from those in the ancestral chromosomes of the legumes after isolation from the other subfamilies, the Phaseoloids and Galegoids, ~55 mya.<sup>52</sup> Of the 10 *Arachis* HGs, HG01, 05, 07 and 09 showed microsynteny with the chromosomes of *C. cajan*, *L. japonicus*, and *M. truncatula*. This suggested the possibility that the major chromosome rearrangements did not occur after the time of divergence of subfamilies in the Papilionoideae. By contrast, HG02 showed the most disparate synteny with the genomes of *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula* (Table 3), which suggested frequent chromosome rearrangement.

## 5. Conclusion

In this study, we developed three linkage maps for *Arachis* spp., and anchored the LGs to the A and B genomes. Integration of the tetraploid maps

developed in this study with the 13 previously published studies generated a high-density consensus map of tetraploid *Arachis*. The developed maps identified structural features within the *Arachis* genome using comparative genomic analysis, and also identified differences between the *Arachis* genome and that of other legumes, *C. cajan*, *G. max*, *L. japonicus*, and *M. truncatula*. The results obtained in this study will bridge the gaps in our knowledge regarding the genomes of *Arachis* and other legumes, and will further the genetic/genomic study and molecular breeding of *Arachis*.

### 5.1. Availability

Information for the genetic linkage maps and the integrated consensus map is available at <http://marker.kazusa.or.jp>.

**Acknowledgements:** We are grateful to S. Sasamoto, S. Nakayama, A. Watanabe, T. Fujishiro, M. Kato, Y. Kishida, M. Kohara, C. Minami, K. Nanri, C. Takahashi, and H. Tsuruoka in the Kazusa DNA Research Institute for their technical assistance.

### Funding

This work was supported by the Kazusa DNA Research Institute Foundation.

**Supplementary data:** Supplementary data are available at [www.dnaresearch.oxfordjournals.org](http://www.dnaresearch.oxfordjournals.org).

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