Cytogenetic Diversity of Simple Sequences Repeats in Morphotypes of Brassica rapa ssp. chinensis

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A significant fraction of the nuclear DNA of all eukaryotes is comprised of simple sequence repeats (SSRs). Although these sequences are widely used for studying genetic variation, linkage mapping and evolution, little attention had been paid to the chromosomal distribution and cytogenetic diversity of these sequences. In this paper, we report the distribution characterization of mono-, di-, and tri-nucleotide SSRs in Brassica rapa ssp. chinensis. Fluorescence in situ hybridization was used to characterize the cytogenetic diversity of SSRs among morphotypes of B. rapa ssp. chinensis. The proportion of different SSR motifs varied among morphotypes of B. rapa ssp. chinensis, with tri-nucleotide SSRs being more prevalent in the genome of B. rapa ssp. chinensis.

We determined the chromosomal locations of mono-, di-, and tri-nucleotide repeat loci. The results showed that the chromosomal distribution of SSRs in the different morphotypes is non-random and motif-dependent, and allowed us to characterize the relative variability in terms of SSR numbers and similar chromosomal distributions in centromeric/peri-centromeric heterochromatin. The differences between SSR repeats with respect to abundance and distribution indicate that SSRs are a driving force in the genomic evolution of B. rapa species. Our results provide a comprehensive view of the SSR sequence distribution and evolution for comparison among morphotypes B. rapa ssp. chinensis.

Keywords: Brassica rapa, simple sequence repeats, fluorescence in situ hybridization, cytogenetic diversity, heterochromatin

INTRODUCTION

Simple sequence repeats (SSRs), also known as microsatellites, are composed of 1–6 nucleotide motifs that are repeated in tandem and are widely and non-randomly distributed in 100–1000s of copies in the genomes of both monocots and dicots (Tautz and Renz, 1984; Toth et al., 2000; Mortimer et al., 2005; Lawson and Zhang, 2006; Hong et al., 2007). Microsatellites are found predominantly in heterochromatin regions, such as centromeric, peri-centromeric, and sub-distal regions of eukaryotic chromosomes (Yang et al., 2005; Li et al., 2010), and sex chromosomes in animals (Beckmann and Weber, 1992; Cuadrado and Jouve, 2011), usually associated with
the constitutive heterochromatin (Lohe et al., 1993; Pedersen et al., 1996). Various non-random, taxon-specific patterns of SSR occurrence call for functional interpretations. Morgante et al. (2002) hypothesized that the relative frequency of microsatellites is higher in the single- or low-copy regions of the genome than in the repetitive regions. Mortimer et al. (2005) showed that SSRs are associated with and around transcribed sequences in Arabidopsis. Li et al. (2004) reviewed that SSRs in different positions in a gene can play important roles in regulating its expression and determining the function of its products. The accumulated evidence indicates that SSRs play an important role in chromatin organization, regulation of gene activity (Nagaki et al., 2004), recombination, DNA replication, the cell cycle, the mismatch DNA repair system (Li et al., 2002), and protein coding regions (Sonah et al., 2011).

The fluorescence in situ hybridization (FISH) technique has been used to localize one or more SSR loci on chromosomes. FISH has become a strategy for chromosome diagnosis and for investigating plant genome organization (Cuadrado and Schwarzacher, 1998; Cuadrado and Jouve, 2002). SSRs change rapidly during evolution, and thus display polymorphism at homologous sites between closely related species. Begum et al. (2009) showed that the rapid evolution of repetitive DNA sequences has resulted in species-specific repeat variants and the generation of novel repeat families. This characteristic has made SSRs useful as markers in comparative diversity analysis (Zhembetyayeva et al., 2003; Yu et al., 2010; Fang et al., 2013; Zhu et al., 2013) and genetics research (Nanda et al., 1991).

SSRs are highly abundant within genomes, they can be widely dispersed or be confined to certain chromosomal regions, and they display a high degree of length polymorphism (Katti et al., 2001; Lawson and Zhang, 2006). Carmona et al. (2013) defined cytotenic diversity in terms of the differences in abundance and distribution of microsatellites, and also found some specific and motif-dependent hybridization patterns. The repeats AG, AAG, ACT, and ATC presented different in situ hybridization patterns that provided cyogenetic landmarks for chromosome identification in barley, Hordeum vulgare ssp. vulgare (Carmona et al., 2013). Altogether, such variation could be used to determine evolutionary relationships between related species.

Brassica rapa belongs to the A genome species group in the Brassicaceae with 2n = 20 chromosomes (Nagaharu, 1935), which had a monophyletic origin (Lysak et al., 2005; Cheng et al., 2013). B. rapa comprises several sub-species, such as non-heading Chinese cabbage (B. rapa ssp. chinensis), Chinese cabbage (B. rapa ssp. pekinensis; Koo et al., 2004), and turnip (B. rapa L. ssp. Rapifera; Snowdon, 2007). B. rapa ssp. chinensis is one of the most important leafy vegetable forms in B. rapa, and consists of five morphotypes (Pak-choi, Wu ta cai, Cai xin, Fen nie cai, and Tai cai; Viehoever et al., 1920; Gladis and Hammer, 1992; Zheng et al., 2015). The phylogenetic relationships between some B. rapa ssp. chinensis morphotypes have been determined from morphological, ecological, and molecular data (Yu et al., 2010). In this paper, systematic research was performed to investigate the cytotenic diversity between intra-specific forms of B. rapa.

The objective of the work presented here is to characterize the cytotenic diversity of SSRs between morphotypes of B. rapa ssp. chinensis that represent a broad range of cytotenic diversity. The available genome sequence of Chinese cabbage is an important and fundamental resource for understanding this species Wang et al., 2011, but the vast majority of heterochromatic regions remain essentially uncharacterized. The current estimates of SSR frequencies in many organisms differs from reality after comparisons with sequence databases (Hong et al., 2007; Cavagnaro et al., 2010; Gao et al., 2011). The distribution of SSRs in databases has been reported for the Chinese cabbage genome (Hong et al., 2007). In view of their ubiquity and functional importance, detailed information will be necessary to explore the comparative cyogenetics of SSRs in B. rapa.

Simple sequence repeats appear to be more abundant in non-coding regions than in coding regions of plant genomes (Hong et al., 2007; Cavagnaro et al., 2010). Tri-nucleotide repeats, the most abundant SSR types in many species (Gao et al., 2003; Shi et al., 2013), were found to be the most frequent in protein coding regions (Sonah et al., 2011). In addition to tri-nucleotide repeats, mono- and di-nucleotide repeats were also predominant in the B. rapa genome (Hong et al., 2007; Gao et al., 2011). In this study, we selected mono-, di-, and tri-nucleotide repeats for physical mapping on the chromosomes of B. rapa. FISH was performed with mono-, di-, and tri-nucleotides to detect the distributional profile of SSRs and to enhance our understanding of genome organization. The distributional characterization of SSRs revealed a range of cyogenetic diversity that could relate to genome organization, function, and evolutionary trends. We demonstrated that: (1) not all of the SSR-based probes produced FISH signals on all B. rapa ssp. chinensis chromosomes; (2) some SSR signal intensity did not show a relationship to the abundance in the genome database; (3) the distributional patterns of SSR signals depended on the SSR motif used and the species analyzed; and (4) differences in SSR abundance and density were shown within and between genomes.

MATERIALS AND METHODS

Plant Materials

Five morphotypes of B. rapa ssp. chinensis: Pak-choi (cv. NHCC002), Wu ta cai (cv. NHCC006), Cai xin (cv. NHCC008), Fen nie cai (cv. NHCC010), Tai cai (cv. NHCC015) were stored and cultivated at the Key Laboratory of Biology and Germplasm Enhancement of Horticultural Crops in East China, Ministry of Horticulture, China. All of these morphotypes have distinct phenotypic characteristics in terms of leaf shape and size, number of leaves, number of outgrowing axially buds, and flowering time (Figures 1a–e).

Chromosome Preparation

Mitotic metaphase chromosome preparation from root tips followed the procedure described by Zheng et al. (2014).

1http://www.plantnames.unimelb.edu.au/Sorting/Brassica_rapa.html#praecox

2http://brassica.bbsrc.ac.uk/brassica_genome_sequencing_concept.htm
Seeds of all morphotypes were allowed to germinate on moist filter paper in Petri dishes at 25°C until each root was approximately 1.5 cm long. To increase the number of cells at metaphase, seedlings were treated with 2.0 μM 8-oxoquinoline at room temperature for 1.5–2.0 h. After washing three-times for 5 min each in distilled water, the seedlings were fixed in a fresh 3:1 (v/v) mixture of 100% ethanol:glacial acetic acid for 24 h, and preserved in 70% (v/v) ethanol. Root tips were digested with 4% (w/v) cellulase plus 2% (w/v) pectinase for approximately 30 min at 37°C, after which they were squashed in a drop of 45% (v/v) acetic acid. After removing the cover slip by freezing, each slide was air-dried in preparation for FISH.

Fluorescence In situ Hybridization (FISH)

Table 1 shows all of the synthetic oligonucleotides of 10–20 bp that were used as SSR probes. For mono-nucleotides, the rate of A or C repeats was representative of itself and T and G. The AC/AG motifs represent both themselves and the complementary sequences TG/TC. Two di-nucleotides (AT and GC) were not be used in FISH for their self-complementary structure (Cuadrado et al., 2008). Hybridization with tri-nucleotides (Jurka and Pethiyagoda, 1995), together with mono- and di-nucleotides, were performed on metaphase chromosomes of five representative cultivars of B. rapa ssp. chinensis morphotypes. The SSR sequences were synthesized by Life Technologies (Nanjing, China), and were labeled with digoxigenin-11-dUTP by random primer labeling followed the manufacturer’s instructions (Roche). The reaction was performed following the method of Cuadrado and Jouve (1994). Detection of digoxigenin was performed by incubating the slides in anti-digoxigenin-rhodamine (Roche) at 37°C for 1 h. The chromosomes were then counterstained with 2 μg/μl DAPI (Sigma). Re-probing was performed following the method of Cuadrado and Jouve (1994).

Image Acquisition and Analysis

Fluorescence in situ hybridization signals and images of stained chromosomes were captured using a chilled charge-coupled device (CCD) camera (Axiocam HR, Carl Zeiss, Germany), and images were pseudo-colored and processed using Axiosvision software (Carl Zeiss). Detection signals and imaging acquisition were obtained by Zeiss Axio Imager A1 fluorescence microscope. For each SSR motif experiment, we analyzed at least 10 cells with distinct signals. The images from FITC and DAPI staining procedures were recorded separately using a cooled CCD camera. The exposure times depended on the intensity of the signals from each probe. The final images were prepared with Adobe Photoshop, version CS4.

RESULTS

We used 16 synthetic SSRs as probes for single- and double-target FISH. Differences were observed in the abundance and localization of motifs between the different B. rapa ssp. chinensis morphotypes, although a general distribution pattern emerged. In all five genomes, most motifs showed a higher density of signal at the centromeric or peri-centromeric regions.

Distribution of Mono- and Di-nucleotide SSRs in the Genomes of B. rapa ssp. chinensis Morphotypes

We did not detect visible signals for the mono-nucleotide repeats A and C on chromosomes of any target genome of B. rapa ssp. chinensis. The di-nucleotide probes, AC and AG, gave visible signals with different patterns among the five samples (Figure 2). Cai xin was the only morphotype in which we detected distinct signals on chromosomes from the two di-nucleotide probes. AG microsatellites showed weak hybridization signals with dispersed patterns in the genomes of Pak-choi and Fen nie cai (Figures 2f,g).
Physical Characterization of Tri-nucleotide Repeats in *B. rapa* ssp. *chinensis* Morphotypes

Chromosomal Localization Tri-nucleotide Repeats in Pak-choi

Metaphase chromosomes of five *B. rapa* ssp. *chinensis* morphotypes were hybridized by re-probing preparations with tri-nucleotide repeat probes ([Supplementary Figures S1–S3](#)). All microsatellite motif probes gave *in situ* hybridization signals on metaphase chromosomes of Pak-choi. Well-defined hybridization signals were produced by ATT, CAC, CAT, AGG, AGC, and ATC probes, and these sequences showed characteristic, motif-dependent distribution patterns ([Figure 3](#)). The ATT and CAC probes revealed weak hybridization signals restricted to the centromeres of chromosomes A4 and A5, and the CAT repeat probe gave no signal ([Figures 3a–c](#)). In addition, no signal was detected on chromosome A6 after hybridization with CAC, CAT, and ATT repeat probes. The AGC microsatellites co-localized with ATC repeats on Pak-choi metaphase chromosomes with similar intensity, but showed less intensity than AGG repeats ([Figures 3d–f](#)). AGG, AGC, and ATC probes showed obvious differences in signal intensity on chromosome A2, AGG being the most intense. These three SSR clusters differed extensively on chromosome A9. AGC and ATC repeats were confined to centromeric regions of this chromosome ([Figures 3e,f](#)); however, AGG repeats comprised nearly the entire length of the short arm of A9 ([Figure 3d](#)).

Chromosomal Localization Tri-nucleotide Repeats in Wu ta cai

Differences in the presence/absence and intensity of the hybridization signals were observed on chromosomes of Wu ta cai after hybridization with tri-nucleotide repeat probes ([Figures 4a,b](#)). All tri-nucleotide repeats were near the centromere on some chromosomes. Polymorphic intercalary signals were observed in two regions of chromosome A2 after hybridizing with AGC and ATC clusters ([Figures 4i,k](#)). Hybridization signals of differing intensity were obtained, depending on the SSR motif probe. ATC and AGC, which were present on all chromosomes, showed obvious differences in intensity, the former being more intense. A weak signal on the long arm of chromosome A5, after hybridization with and AGC repeat probe, was observed only after increasing the exposure time of the CCD (compare [Figures 4h and 4j](#)).

Chromosomal Localization Tri-nucleotide Repeats in Cai xin

Variation in intensity and location of the *in situ* signals were observed on metaphase chromosomes of Cai xin. The signals were confined to the centromere for all tri-nucleotide repeats with differing intensities. Polymorphism in terms of presence/absence was observed in the nucleolus organizing region (NOR). NOR signals were generated after hybridization with AG, CAC, ACT, GCC, and AGG repeats ([Figures 5b,c,e–g](#)). Visible centromeric signals were observed on chromosome A5 for AG, AGC, CAC, and ACT repeat probes ([Figures 5b–e](#)), but not for AC, GCC, and AGG repeats ([Figures 5a,f,g](#)).

Chromosomal Localization Tri-nucleotide Repeats in Fen nie cai

No specific clusters of SSR-specific signals were observed for tri-nucleotide repeats on chromosomes of Fen nie cai. Polymorphism in terms of presence/absence and intensity of the hybridization signals were observed for some SSRs. The absence signal was revealed from ACT and CAG clusters ([Supplementary Figures S1n,s](#)). However, signals of varying intensity were
observed on the homologous chromosomes after hybridization with AGG, AGC, and ATC repeat probes (Figures 4c–e).

Chromosomal Localization Tri-nucleotide Repeats in Tai cai
The most intense and rich patterns of in situ hybridization signals were produced by AGC, ACT, and ATT repeat probes on chromosomes of Tai cai, and were confined to the pericentromeric regions. No specific clustered sites were observed for GCC and AAC repeats (Supplementary Figures S1e and S3t). AAT and CAC repeats were confined around centromeric regions. The ATT repeat probes gave more dispersed and intense centromeric signals, clustered in centromeric and pericentromeric regions, than did the CAC repeats (Figures 4f,g). Variations in the number of signal sites were observed on chromosomes A2 and A6. An intercalary signal was observed on the long arms of chromosome A2 (compare Figures 4l with 4n) and A6 (compare Figures 4m with 4o) from hybridization with ATT repeats, in contrast to only one centromeric signal from CAC repeats (Figures 4n,o). No signals were observed from hybridization with a CAC repeat probe on chromosomes A4, A5, and the NOR, but the ATT repeat probe gave weak signals (Figures 4f,g).

DISCUSSION

Rogan et al. (2001) showed that base composition, probe length, and chromosomal location contribute to hybridization signal intensity in FISH. In the present study, all probes ranged from 15 to 20 bp in length. The hybridization patterns obtained depended on the base composition of the probes used and chromosomal location. The synthetic SSR probes were labeled by the random primer method, compared with end labeling, which improved the resolution of SSR loci in FISH (Bouilly et al., 2008). Different SSR probes of the same lengths gave different signal intensities, indirectly reflecting the influence of the target size and the copy numbers of the repeat sequences. The SSR probes gave specific hybridization patterns that provide cytogenetic landmarks for chromosome identification.

Different SSR Distribution Patterns on Chromosomes Depend on the Species Analyzed

Sonah et al. (2011) showed that mono-, di-, and tri-nucleotide repeats compose the major proportion of SSRs in plant genomes. For tri-nucleotides, A/T-rich repeats (e.g., AAC/GTT, AAG/CTT, and AAT/ATT) were predominant in dicot species. In the
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**FIGURE 4** | Photomicrographs showing the distribution of several tri-nucleotide SSRs (AGG, AGC, and ATC) on metaphase chromosomes of Wu ta cai, Fen nie cai, and Tai cai. (a,b) in Wu ta cai; (c–e) in Fen nie cai; (f,g) in Tai cai; (h–k) variation in the number of signal sites in Wu ta cai; (l–o) variation in the number of signal sites in Tai cai; lines indicate the number of fluorescent signals in Wu ta cai and Tai cai; arrows indicate different intensity fluorescent signals. Chromosomes were counterstained with DAPI.

monocot barley, however, repeated AAT SSR motifs gave poor hybridization signals (Cuadrado and Jouve, 2007b). Morgante et al. (2002) found that GCC repeats accounted for half of the tri-nucleotide repeats in rice, whereas they were rare in dicots. Lawson and Zhang (2006) examined the most common mono- (A/T) and di-nucleotide (AT and AG) repeats in the *Arabidopsis* genome, and found that polyA/T repeats were predominant, while polyC/G repeats were rare (Sonah et al., 2011). The distribution of AC and AG repeats were linked to the euchromatic and heterochromatic genomic regions, respectively (Cuadrado and Jouve, 2007a). The di-nucleotide AG repeats were located on all chromosomes in *Dendrobium aphyllum* and *D. aggregatum* (Begum et al., 2009), but were exclusively concentrated at the centromeres in *Triticum* (Carmona et al., 2013). The most abundant repeat motifs were A (28.8%), AG (15.4%), AT (13.7%), and AAG (13.3%) clusters, reflecting the A/T rich nature of the *B. rapa* genome (Hong et al., 2007). In this study, nearly all the di- and tri-nucleotide SSRs were detected on metaphase chromosomes of Cai xin and Pak-choi, inferring that these two morphotypes have more types and increased abundance of SSRs.
FIGURE 5 | The polymorphic characterization of the chromosome bearing the NOR in Cai xin following *in situ* hybridization with digoxigenin-labeled probes (detected with red rhodamine) and DAPI counterstaining. (a) (AC)$_8$; (b) (AG)$_{12}$; (c) (CAC)$_5$; (d) (ACG)$_5$; (e) (ACT)$_5$; (f) (GCC)$_5$; (g) (AGG)$_5$; arrows indicate chromosome A5; no signal was detected on the NOR in (a,d); fluorescent signal on the NOR in (c); weak centromeric signal in (d); signals on the NOR and the centromere in (b,e,g).

compared to the other three morphotypes (Wu ta cai, Fen nie cai, and Tai cai).

Various types of SSR motifs display taxon-specific patterns in the genomes of prokaryotes and eukaryotes (Toth et al., 2000). For example, the most intense hybridization signals were produced by the AGG, AAG, and AAC tri-nucleotide probes in barley (Cuadrado and Jouve, 2007b). The AAC clusters showed the same distribution patterns as AAG repeats in wheat (Cuadrado and Schwarzer, 1998). AAG repeat units are major contributors to the genomes of dicots (Sonah et al., 2011); they are preferentially associated with peri-centromeric heterochromatin in *Hordeum* species (Carmona et al., 2013), and generally reflect the distribution of heterochromatin and the C-banding pattern in wheat (Cuadrado and Jouve, 1994). AAC repeats are organized in a more dispersed manner, with centromeric regions being largely excluded in chickpea and tomato (Gortner et al., 1998; Gindullis et al., 2001). Some repeats, such as CAG, CAC, and ACG, have specific hybridization sites restricted to the centromeres of metaphase chromosomes in barley (Cuadrado and Jouve, 2007b).

Despite the stable chromosome number and similarities in chromosome size and morphology, differences in numbers and distribution of SSR blocks have been observed among *B. rapa* ssp. *chinensis* morphotypes. The ATT repeat probe gave two signals with different intensities on chromosomes A2 and A6 of Tai cai, indicating that this SSR experienced chromosome-specific accumulation during evolution, as did AGG and ATC repeats in Wu ta cai. In addition, several SSRs (ATT, CAC, AGG, AGC, and ATC) also showed chromosome-specific signals on marker chromosomes of Pak-choi. These three morphotypes may have resulted from chromosomal recombination during speciation and development.

The Different Distribution Patterns on Chromosomes Depend on the SSR Probes Used

A negative correlation has been observed between repeat numbers and the total length of repeat units in both monocots and dicots (Cavagnaro et al., 2010; Gao et al., 2011; Sonah et al., 2011). Mono-nucleotide repeats of A are the most abundant repeats among all SSRs analyzed (Gao et al., 2011); however, no visible signal was generated, partly implying their dispersed distribution along chromosomes, or may be related to the centromeric function (Cuadrado and Jouve, 2007a). In this work, the di-nucleotide repeats AC and AG only showed distinct *in situ* hybridization signals on chromosomes of Cai xin, although there were weak signals in Pak-choi and Fen nie cai for AG repeats. In addition, three tri-nucleotide repeat probes (AAG, AGG, and AGC) gave distinct signals on all analyzed genomes. Di-nucleotide AG and tri-nucleotide AAG repeats are
relatively abundant in the *B. rapa* genome (Hong et al., 2007). AG-rich microsatellites might be the most prevalent SSRs that cluster around centromeric regions in *B. rapa ssp. chinensis* morphotypes.

We found that some tri-nucleotide repeats distinctly co-localized within the same genome, although the signals differed in intensity, which could be explained by their intermixed structure at the given loci. Some SSR probes gave weak or dispersed signals at the same physical position, which could possibly be due to closely linked blocks of repeated sequences. Here, we show that individual SSRs vary widely in their relative proportions at the chromosome level, and that the distribution of SSRs along the chromosomes is non-random. The characteristic patterns of SSR distribution show that centromeric regions are more densely populated than the central regions (Rogan et al., 2001). The similar distribution patterns of some SSR motifs indicate that long stretches of different SSRs are of functional importance, and could possibly represent an ancient component of plant genomes (Cuadrado et al., 2000; Cuadrado and Jouve, 2007b). These results are supported by evidence showing that microsatellites display relatively uniform coverage in the genome, and that there are taxon-specific distribution patterns among *B. rapa ssp. chinensis* morphotypes. Our results suggest that SSR repeats are the major component of the satellite DNA fraction, and show evolutionary conservation among *B. rapa ssp. chinensis* morphotypes.

**Relationship between SSRs and Centromeres**

In most higher eukaryotic organisms, chromosomal centromeres are composed of long arrays of satellite repeat sequences and retro-transposons (Henikoff et al., 2001; Jiang et al., 2003). The centromeric and distal regions play important roles during mitosis and meiosis (Li et al., 2002; Hong et al., 2006), and highly divergent sequences are present in the peri-centromeric regions (Wang et al., 2012). The repetitive DNA sequences frequently form clusters within heterochromatin blocks, which are predominantly concentrated at peri-centromeric regions and have been detected in plants with small and compact genomes (Cuadrado and Jouve, 2010; Falistocco and Marconi, 2013). In the present work, most *in situ* SSR signals were confined to centromeric or adjacent regions on *B. rapa ssp. chinensis* chromosomes. This distributional characterization may be related to their effect on DNA replication, chromatin organization, and the cell cycle.

**Evolutionary Trends of SSRs among *B. rapa ssp. chinensis* Genomes**

The relationship between microsatellites and chromosomal evolution has not been clearly documented. The frequency of repeats decreases exponentially with their length, type, and number of SSR motif repeats. This characterization appears to be more conservative in coding than non-coding sequences (Cavagnaro et al., 2010; Sonah et al., 2011; Shi et al., 2013), and less pronounced for di-nucleotides compared to longer repeat types (Cavagnaro et al., 2010). The trends for various repeat types are similar between different chromosomes within the same genome, but the density of repeats may vary between different chromosomes in the same species (Schafer et al., 1986; Katti et al., 2001). Sonah et al. (2011) found species-specific accumulation of particular motif repeats. Variation is present between related species in terms of the abundance and chromosomal distribution of SSR clusters among morphotypes. Schmidt and Heslop-Harrison (1996) demonstrated that microsatellites, representing a substantial fraction of the genome, showed chromosome-specific amplification in plants. High levels of polymorphism and heterozygosity between homologs, in terms of the distribution of AAG and AAC repeats, was shown for out-breeding species in the *Secale strictum* species complex (Cuadrado and Jouve, 2002).

Our results suggest that SSR sequences are more predisposed to being amplified or deleted as a result of independent events. The balance among SSR sequences generated by strand-slipage replication, or recombination and repair mechanisms, cannot be the only explanation for the observed differences in their chromosomal distributions. Another possible explanation would involve selection pressure or mutation (Li et al., 2002). The different chromosomal positions of SSRs involved in the regulation of gene expression (Lawson and Zhang, 2006; Gao et al., 2011), could indicate their underestimated roles in genome evolution (Cuadrado et al., 2008).

The microsatellite sequences analyzed here showed similar chromosome distribution polymorphism patterns, inferring that these SSR loci may result from convergent evolution. However, they differed in intensity or position, indicating that microsatellite repeats can contract or expand over a very short evolutionary time frame (Iwata et al., 2013). The wide distribution of SSRs, and the fact that their positions are restricted to chromosomal centromeres, as revealed by FISH, suggested a general model for the parallel chromosome evolution of repeat-rich heterochromatin in *B. rapa ssp. chinensis*.

Carmona et al. (2013) suggested that changes in the amount and distribution of tandem repetitive DNA sequences are major driving forces of genome evolution and speciation. The different regions are thought to undergo different selection pressures (Morgante et al., 2002), which might account for different motif preferences and frequencies among chromosomes. The evolutionary dynamics of microsatellites is generally consistent with plant divergence and evolution (Shi et al., 2013), and the distribution of microsatellites is related to the history of genome evolution and selective constraints (Morgante et al., 2002). The variation in SSRs at the chromosome level may be the result of adaptive divergence, or selection resulting from the stress response among species and populations (Cavagnaro et al., 2010; Gao et al., 2011; Carmona et al., 2013). Whether SSRs are under selection or are neutral as has been reported (Ellegren, 2004), and can be used for exploring the dynamics of the evolutionary process (Santos et al., 2010), will require study.

**AUTHOR CONTRIBUTIONS**

S-nZ and J-sZ: Designed the experiment. X-lH and GB: Provided the materials. J-sZ and C-zS: Performed the experiment. J-sZ: Analyzed the data and wrote the manuscript.
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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2016.01049

REFERENCES

Beckmann, J. S., and Weber, J. L. (1992). Survey of human and rat microsatellites. Genomics 12, 627–631. doi: 10.1016/0888-7543(92)90285-Z

Begum, R., Alam, S. S., Menzel, G., and Schmidt, T. (2009). Comparative molecular cytogenetics of major repetitive sequence families of three Dendrobiunm species (Orchidaceae) from Bangladesh. Ann. Bot. 104, 863–872. doi: 10.1093/aob/mcp178

Bouilly, K., Chaves, R., Leitao, A., Benabdelmouna, A., and Guedes-Pinto, H. (2015). Chromosomal organization of simple sequence repeats and the centromeric DNA of Pelargonium sidoides. Bot. Bull. Acad. Sin. 56, 541–551. doi: 10.1093/aob/mcs133

Bouilly, K., Chaves, R., Leitao, A., Benabdelmouna, A., and Guedes-Pinto, H. (2016). Chromosomal organization of simple sequence repeats (SSRs) and centromeric DNA of Pelargonium sidoides (Geraniaceae). Plant Cell Tissue Organ Cult. 125, 37–49. doi: 10.1007/s11240-015-0946-3

Carmona, A., Friero, E., de Bustos, A., Jouve, N., and Cuadrado, A. (2013). Genome-wide characterization of simple sequence repeats in cucumber (Cucumis sativus L.). BMC Genomics 14:569. doi: 10.1186/1471-2164-11-S69

Cheng, F., Mandáková, T., Wu, J., Xie, Q., Lysak, M. A., and Wang, X. W. (2013). Comparative molecular cytogenetics of major repetitive sequence families of three Dendrobiunm species (Orchidaceae) from Bangladesh. Ann. Bot. 104, 863–872. doi: 10.1093/aob/mcp178

Cuadrado, A., and Jouve, N. (2009). Chromosomal organization of simple sequence repeats in wheat and rye genomes. Chromosoma 117, 387–394. doi: 10.1007/s00412-009-0254-x

Cuadrado, A., and Schwarzacher, T. (1998). The chromosomal organization of simple sequence repeats in wheat and yye genomes. Chromosoma 107, 587–594. doi: 10.1007/s004120050345

Cuadrado, A., Schwarzacher, T., and Jouve, N. (2007). Identification of different chromatin classes in wheat using in situ hybridization with simple sequence repeat oligonucleotides. Theor. Appl. Genet. 101, 711–717. doi: 10.1007/s001220051535

Ellegren, H. (2004). Microsatellites: simple sequences with complex evolution. Nat. Rev. Genet. 5, 435–445. doi: 10.1038/nrg1348

Ellegren, H. (2004). Microsatellites: simple sequences with complex evolution. Nat. Rev. Genet. 5, 435–445. doi: 10.1038/nrg1348

Falistocco, E., and Marconi, G. (2013). Cytogenetic characterization by in situ hybridization techniques and molecular analysis of SS rRNA genes of the European hazelnut (Corylus avellana). Genome 56, 155–159. doi: 10.1139/gen-2013-0045

Fang, P., Chen, F. B., Yao, Q. L., Yang, K. C., Zhou, G. F., Fan, Y. H., et al. (2013). Analysis of genetic diversity in the tuber mustard (Brassica juncea var. tumida Tsen et Lee) in the Yangtze river basin of China. Genet. Resour. Crop Evol. 60, 129–143. doi: 10.1007/s10722-012-9821-y

Gao, C. H., Tang, Z. L., Yin, J. M., An, Z. S., Fu, D. H., and Li, J. N. (2011). Characterization and comparison of gene-based simple sequence repeats across Brassica species. Mol. Genom. Genet. 286, 161–170. doi: 10.1007/s00438-011-0636-x

Gao, L. F., Tang, J. F., Li, H. W., and Jia, J. Z. (2003). Analysis of microsatellites in major crops assessed by computational and experimental approaches. Mol. Breed. 12, 245–261. doi: 10.1023/A:1023646121217

Gindullis, F., Desel, C., Galasso, I., and Schmidt, T. (2001). The large-scale organization of the centromeric region in Beta species. Genome Res. 11, 253–265. doi: 10.1101/gr.162301

Glads, T., and Hammer, K. (1992). The Gatersleben Brassica collection—Brassica juncea, B. rapa, B. nigra and B. rapa (Germ., Engl. summary). Feddes Repert. 103, 469–507. doi: 10.1002/fedr.19921030704

Gortner, G., Nenno, M., Weising, K., Zink, D., Nagl, W., and Kahl, G. (2001). The large-scale genome by BAC-End sequence analysis of Citrus (Citrus sinensis L.) chromosomes. Genom. Res. 11, 1773–1777. doi: 10.1101/gr.1131001

Gosálvez, J. A., and Jouve, N. (2007). Chromosomal organization of simple sequence repeats and the centromeric DNA of Pelargonium sidoides. Bot. Bull. Acad. Sin. 56, 541–551. doi: 10.1093/aob/mcs133

Hedreck, S., Ahmad, K., and Malik, H. S. (2001). The centromere paradox: stable inheritance with rapidly evolving DNA. Science 293, 1098–1102. doi: 10.1126/science.1062939

Heslop-Harrison, J. S. (1991). The molecular cytogenetics of palnts. J. Cell Sci. 100, 15–21.

Hong, C. P., Piao, Z. Y., Kang, T. W., Batley, J., Yang, T. J., Hur, Y. K., et al. (2007). Genomic distribution of simple sequence repeats in Brassica rapa. Mol. Cells 23, 349–356.

Hong, C. P., Plaha, P., Koo, D. H., Yang, T. J., Choi, S. R., Lee, Y. K., et al. (2006). A survey of the Brassica rapa genome by BAC-End sequence analysis and comparison with Arabidopsis thaliana. Mol. Cells 22, 300–307.

Iwata, A., Tek, A. L., Richard, M. M. S., Abernathy, B., Fonseca, A., Schmutz, J., et al. (2013). Identification and characterization of functional centromeres of the common bean. Plant J. 76, 47–60. doi: 10.1111/tpj.12269
Schafer, R., Ali, S., and Epplen, J. T. (1986). The organization of the evolutionarily
distinct patterns of SSR distribution in the Arabidopsis thaliana and rice genomes. Genome Biol. 7:R14. doi: 10.1186/gb-2006-7-2-r14

Li, C., Huang, B., Yan, X. H., Wang, I. J., Yang, Q., and Wei, W. H. (2010). Cloning, identification and characterization of a repetitive sequence flanking telomere and homologous to canrcep in Brassica napus. Bot. Stud. 51, 421–430.

Li, Y., Korol, A. B., Fahima, T., Beiles, A., and Nevo, E. (2004). Microsatellites within genes: structure, function and evolution. Mol. Biol. Evol. 21, 991–1007. doi: 10.1093/molbev/msh073

Li, Y. C., Korol, A. B., Fahima, T., Beiles, A., and Nevo, E. (2002). Microsatellites: genomic distribution, putative functions and mutational mechanisms: a review. Mol. Ecol. 11, 2453–2465. doi: 10.1046/j.1365-294X.2002.01643.x

Lobe, A. R., Hilliker, A. J., and Robert, P. A. (1993). Mapping simple repeated DNA sequences in heterochromatin of Drosophila melanogaster. Genetics 134, 1149–1174. doi: 10.1006/geno.1999.3455

Lyak, M. A., Koch, M. A., Pecinka, A., and Schubert, I. (2005). Chromosome triplication found across the tribe Brassicaceae. Genome Res. 15, 516–525. doi: 10.1101/gr.3531105

Morgante, M., Hanafey, M., and Powell, W. (2002). Microsatellites are preferentially associated with nonrepetitive DNA in plant genomes. Nat. Genet. 30, 194–200. doi: 10.1038/ng822

Mortimer, J. C., Batley, J., Love, C. G., Logan, E., and Edwards, D. (2005). Simple sequence repeat (SSR) and GC distribution in the Arabidopsis thaliana genome. J. Plant Biotechnol. 7, 17–25.

Nagaharu, U. (1935). Genome analysis in Brassica with special reference to the experimental formation of B. napus and peculiar mode of fertilization. Ipn. J. Bot. 7, 389–452.

Nagaki, K., Cheng, Z. K., Ouyang, S., Talbert, P. B., Kim, M., Jones, K. M., et al. (2004). Sequencing of a rice centromere uncovers active genes. Nat. Genet. 36, 138–145. doi: 10.1038/ng1289

Nanda, I., Zischler, H., Epplen, C., Guttenbach, M., and Schmid, M. (1991). Chromosomal organization of simple repeated DNA-sequences used for DNA fingerprinting. Electrophoresis 12, 193–203. doi: 10.1002/elps.1150120216

Pedersen, C., Rasmussen, S. K., and Linde-Laursen, I. (1996). Genome and chromosomal identification in cultivated barley and related species of the Triticaceae (Poaceae) by in situ hybridization with the GAA-satellite sequence. Genome 39, 93–104. doi: 10.1139/g96-013

Rogan, P. K., Cazzarco, P. M., and Knoll, J. H. M. (2001). Sequence-based design of single-copy genomic DNA probes for fluorescence in situ hybridization. Genome Res. 11, 1086–1094. doi: 10.1101/gr.171701

Santos, J., Serra, L., Sole, E., and Pascual, M. (2010). FISH mapping of microsatellite loci from Drosophila subobscura and its comparison to related species. Chromosome Res. 18, 213–226. doi: 10.1007/s10577-010-9112-4

Schafer, R., Ali, S., and Pascual, M. (1996). The organization of the evolutionarily conserved GATA GACA repeats in the mouse genome. Chromosoma 93, 502–510. doi: 10.1007/BF00386791

Schmidt, T., and Heslop-Harrison, J. S. (1996). The physical and genomic organization of microsatellites in sugar beet. Proc. Natl. Acad. Sci. U.S.A. 93, 8761–8765. doi: 10.1073/pnas.93.16.8761

Shi, J., Q., Huang, S. M., Fu, D. H., Yu, J. Y., Wang, X. F., Hua, W., et al. (2013). Evolutionary dynamics of microsatellite distribution in plants: insight from the comparison of sequenced Brassica, Arabidopsis and other angiosperm species. PLoS ONE 8:e59988. doi: 10.1371/journal.pone.0059988

Snowdon, R. J. (2007). Cytogenetics and genome analysis in Brassica crops. Chromosome Res. 15, 85–95. doi: 10.1007/s10577-006-1109-y

Sonah, H., Deshmukh, R. K., Sharma, A., Singh, V. P., Gupta, D. K., Gacche, R. N., et al. (2011). Genome-wide distribution and organization of microsatellites in plants: an insight into marker development in Brachypodium. PLoS ONE 6:e21298. doi: 10.1371/journal.pone.0021298

Tautz, D., and Renz, M. (1984). Simple sequences are ubiquitous repetitive components of eukaryotic genomes. Nucleic Acids Res. 12, 4127–4138. doi: 10.1093/nar/12.10.4127

Toth, G., Gaspari, Z., and Jurka, J. (2000). Microsatellites in different eukaryotic genomes: survey and analysis. Genome Res. 10, 967–981. doi: 10.1101/gr.10.7.967

Viehoever, A. (1920). Studies in mustard seeds and substrates: I. Chinese colza (Brassica campestris chinoilefera Viehoever). J. Agric. Res. 20, 117–140.

Wang, K., Zhang, W. P., Gao, Y. J., Zhang, Z. X., Zheng, D. W., Zhou, B. L., et al. (2012). Localization of high level of sequence conservation and divergence regions in cotton. Theor. Appl. Genet. 124, 1173–1182. doi: 10.1007/s00122-011-1777-3

Wang, X., Wang, H., Wang, J., Sun, R., Wu, J., Liu, S., et al. (2011). The genome of the mesopolyploid crop species Brassica rapa. Nat. Genet. 43, 1035–1039. doi: 10.1038/ng.919

Yang, T. J., Lee, S., Chang, S. B., Yu, Y. J., Hwang, R. A. (2005). In-depth sequence analysis of the tomato chromosome 12 centromeric region: identification of a large CAA block and characterization of pericentromere retrotransposons. Chromosoma 114, 103–117. doi: 10.1007/s00412-005-0342-8

Yu, S. C., Zhang, F. L., Wang, X. Y., Zhao, X. Y., Zheng, D. S., Yu, Y. J., et al. (2010). Genetic diversity and marker-trait associations in a collection of Pak-choi (Brassica rapa L. ssp Chinensis Makino) accessions. Genes Genomics 32, 419–428. doi: 10.1186/gene-1003-6

Zhebentyayeva, T. N., Reighard, G. L., Gorina, V. M., and Abbott, A. G. (2003). Simple sequence repeat (SSR) analysis for assessment of genetic variability in apricot germplasm. Theor. Appl. Genet. 106, 435–444.

Zheng, J. S., Sun, C. Z., Xiao, D., Zhang, S. N., Bonnema G., and Hou, X. L. (2015). Karyotype variation and conservation in morphotypes of non-heading Chinese cabbage. Plant Syst. Evol. 301, 1781–1791. doi: 10.1007/s00606-014-1177-7

Zheng, J. S., Sun, C. Z., Zhang, S. N., and Hou, X. L. (2014). Karyotype of mitotic metaphase and meiotic diakinesis in non-heading Chinese cabbage. Plant Syst. Evol. 300, 295–302. doi: 10.1007/s00606-013-0882-y

Zhu, X. C., Raman, H., Wu, H. W., Lemerle, D., Burrows, G. E., and Wing, R. A. (2005). Intron recombination analysis of rice leaf nighthshade (Solanae elaegnifolium) and related species. Plant Mol. Biol. Rep. 31, 248–254. doi: 10.1101/g3.003105-12473-z

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