SBMDb: first whole genome putative microsatellite DNA marker database of sugarbeet for bioenergy and industrial applications

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Abstract

DNA marker plays important role as valuable tools to increase crop productivity by finding plausible answers to genetic variations and linking the Quantitative Trait Loci (QTL) of beneficial trait. Prior approaches in development of Short Tandem Repeats (STR) markers were time consuming and inefficient. Recent methods invoking the development of STR markers using whole genomic or transcriptomics data has gained wide importance with immense potential in developing breeding and cultivator improvement approaches. Availability of whole genome sequences and in silico approaches has revolutionized bulk marker discovery. We report world’s first sugarbeet whole genome marker discovery having 145 K markers along with 5 K functional domain markers unified in common platform using MySQL, Apache and PHP in SBMDb. Embedded markers and corresponding location information can be selected for desired chromosome, location/interval and primers can be generated using Primer3 core, integrated at backend. Our analyses revealed abundance of ‘mono’ repeat (76.82%) over ‘di’ repeats (13.68%). Highest density (671.05 markers/Mb) was found in chromosome 1 and lowest density (341.27 markers/Mb) in chromosome 6. Current investigation of sugarbeet genome marker density has direct implications in increasing mapping marker density. This will enable present linkage map having marker distance of ~2 cM, i.e. from 200 to 2.6 Kb, thus facilitating QTL/gene mapping. We also report e-PCR-based detection of 2027...
polymorphic markers in panel of five genotypes. These markers can be used for DUS test of variety identification and MAS/GAS in variety improvement program. The present database presents wide source of potential markers for developing and implementing new approaches for molecular breeding required to accelerate industrious use of this crop, especially for sugar, health care products, medicines and color dye. Identified markers will also help in improvement of bioenergy trait of bioethanol and biogas production along with reaping advantage of crop efficiency in terms of low water and carbon footprint especially in era of climate change.

**Database URL:** http://webapp.cabgrid.res.in/sbmdb/

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**Introduction**

Sugarbeet (*Beta vulgaris* L. *ssp. vulgaris*) is a biennial, dicotyledonous crop of temperate climate. It represents the world’s second highest source of sucrose with 15–20% sugar content (1) after sugarcane (*Saccharum officinarum* L.). It accounts for ~30% of the world’s annual sugar production and has also been considered as a potential biofuel crop (2) besides its potential as animal feed (3) and medicinal properties (4). With the ever increasing rise in the global population to be around 10 billion in 2050, finding sustainable solutions to the bioenergy research is becoming an important unanswered question. The use of potential food crops for biofuels will be one of the critical needs to support the global projected population. Its increasing importance in bioenergy has led to greater area for production of bioethanol and biogas (5).

Among the largest sugar beet producers, Europe and the United States share 75% of both, global area harvested and production. Among the main producers, France, Germany, the Russian Federation, Turkey and Ukraine, covers almost two thirds of the global production (6). Sugarbeet has been introduced in India in 1971 but its huge industrial potential has not been reaped so far. The demanding biofuel requirement in the country and globe as well, has necessitated the need of ethanol from sugarbeet. Very recently few cases of industrial level production in India, especially from the area of Punjab and Karnataka for sugar and alcohol production, respectively, has been started. If ensilage and anaerobic digestion approach is used, it has further potential of more energy per hectare than bioethanol (7).

Besides industrious use of sugarbeet crop in terms of sugar and bioenergy, it also possesses the additional multifold advantages like: it is tolerant to various climatic and soil conditions thus uncultivable land can also be used. In agriculture, it has three major importance namely, cash crop, soil amelioration/soil fertility improvement and use as by-products for cattle feed/mineral supplement during summer/drought, especially when there is scarcity of green fodder (8).

Beside agricultural importance, sugarbeet plays very important role in industrial area as sunless tanner dihydroxyacetone extracted from sugar beet (9). For human health, it has good medical potentials for anticancerous activity (10) and is a good source of antioxidant (11), antidepressant (12) and organic dyes (14).

Additionally, it is used in herbal therapy and hepatoprotective activity (4, 15). Furthermore, versatile industrial compounds like betaine (16), phenolics and betacyanins (17) obtained from the sugarbeet are also well documented in literature for their therapeutics. Betain is used in industry for PCR adjuvants as it improves amplification of GC-rich DNA sequences (18). Sugarbeet being, short season crop (6 months), offers advantage over sugarcane (12–18 months) along with its ability as most efficient crop in terms of water foot printing (19) and also for lowering ethanol’s carbon footprint (20).

To accelerate the rate of genetic gain for high sugar content, resistance towards biotic (disease causing pathogens) and abiotic stresses (high temperature and saline/alkaline conditions) molecular markers are imperative and have been developed in various crops. Apart from abiotic stresses, sugarbeet is susceptible to over 60 disease caused by pathogens like bacteria, fungi, nematodes, viruses, phytoplasmal, spiroplasmal pathogens, aphids etc. (21–23). Biotic stress can lead to loss even upto 50% of sugarbeet yield (24). Molecular markers play major roles in higher root yield, strong selection against premature bolting, annuality and winter hardiness which are the major problems in sugarbeet abiotic management (25).

Present linkage map of sugarbeet constitutes of nine groups with ~700 cM marker coverage (26, 27). Dohm *et al.* (28) reported an extended genetic map consisting of 983 markers, and Holtgrawe *et al.* (29) in 2014 further added 307 markers to the existing dataset. A sugar beet physical map based on 8361 EST-derived probes was also provided (28). Fugate *et al.* (30) has reported 7680 putative SSR markers.

In *vitro* methods of Short Tandem Repeats (STR) development is disadvantageous as it is time-consuming and
expensive. Availability of whole genome sequence and in silico approach has revolutionized the marker discovery. Recently, a new class of functionally relevant microsatellites called as simple sequence repeats functional domain markers (SSR-FDMs) (31–33) have gained wide importance. This is being widely applied in a number of crop species including the biofuel and energy crop species such as sugarcane (34). For molecular breeding program of sugarbeet, its recently available genome assembly (569 Mb) of KWS2320 genotype (3) needs in silico approach for bulk marker discovery. Further, there is a need of in silico discovery of polymorphism of these markers utilizing resequencing data of four additional genotypes namely, KWS230-DH1440, STR06A6001, SynMono and SynTilling. These markers should be in the form of ‘ready to use’ and readily available to the global community in form of freely accessible database.

Our present work aims at development of microsatellite marker database of sugarbeet whole genome-based STR mining. We further aimed, the user defined primer designing with precise selection from each chromosome, at defined location and equal interval along with evaluation of polymorphism. This work also aims at mining of SSR-FDM from various major sources which can be assessed for the genotyping for direct functional markers using genomic DNA primers.

**Material and methods**

**Data collection and search flexibilities**

For mining of markers, the recently sequenced sugar beet genome data of genotype KWS2320 was used. This haploid line genome was of 567 Mb of which 85% data assigned over its nine chromosomes (2n = 18) having an assembly coverage of 63% was used in our study.

This assembly is having more than 27,000 predicted genes (3). This de novo assembly was downloaded from http://www.ncbi.nlm.nih.gov/assembly/GCA_000511025.1/#/st in FASTA format. These were cleaved using in house PERL scripts and parsed for the identification of the microsatellite markers using the MISA tool (http://pgrc.ipk-gatersleben.de/misa/) with default parameter setting.

For the mining of the functional SSRs markers (SSR-FDMs), Expressed Sequence Tags (ESTs) were downloaded from NCBI (www.ncbi.nlm.nih.gov). Additionally, Putative Unique Transcripts (PUTs) for sugarbeet were systematically downloaded from PlantGDB (Version release 187) available at http://www.plantgdb.org/. All the ESTs and PUTs were first scanned for the presence of the homopolymers errors and sequence ambiguity was further removed using the est_trimmer available at http://pgrc.ipk-gatersleben.de/misa/download/est_trimmer.pl with the following settings: -amb=2,50 -tr5=T,5,50 -tr3=A,5,50 and were subsequently screened for the SSRs identification using MISA. For the identification of the functional domains, the PUTs were translated into all the coding frames and were searched against Interpro. PUTs having SSRs and Interpro assigned functional domain were classified as SSR-FDMs (31–33, 37). For genotyping of SSR-FDM, primers were designed on genomic DNA sequence.

Whole genome based markers were generated with descriptive information on motif size, motif type, repeat numbers with their length and size, repeat type, GC content, start and end position. Provision was made for locating markers on each chromosome at desired interval for mapping of Quantitative Trait Loci (QTL)/gene. Additionally, marker can be selected based on motif type, repeat kind, GC content, number of base pair and copy number of repeat unit as markers with more than eight repeat often exhibits polymorphism due to slippage event in DNA replication. An additional plug-in of primer generation was implemented for the markers, using the primer3 core executable with further flexibility of 500 bp upstream and downstream sequence extraction using PERL scripts targeting approximately 1000 bp as a template for primer designing. Figure 1 demonstrates the flow of analytical pipeline developed for the SBMDb.

For the identified markers, web-based application was created in the window web development environment, WAMP Server with Apache, PHP and MySQL Database.

**Database development**

Sugarbeet MicroSatellite Database (SBMDb) has been developed using PHP and MySQL database under the web development environment, WAMP Server. This relational database was developed based on ‘three tier architecture’ having client tier, middle tier and database tier. Provision to store all in silico mined STRs was made at the backend in MySQL database. PHP scripts were written to properly query and execute the search made by users. The primer3 core was integrated to compute primers of the selected STRs. Primer call for specific locus, i.e. output of primer designing is with list of five primers with their respective melting temperature, GC content, start position and estimated PCR product size are available in the database. Functional domains linked with the simple sequence repeat patterns as an add-on utility to search for the simple sequence repeats functional domain markers (SSR-FDMs) has also been made. To identify the functional domains, all the sequences were translated into all the six reading frames and Interproscan tool was used to analyse and
predict the protein domains using the default settings (31–33). Sequences harboring the functional domains and the simple sequence repeats along with the primer pairs were classified as the functional markers. The database has been designed to cater the needs of the plant biologist and breeders thus making it very flexible to access with user defined options. The choice of motif type, namely, mono, di, tri, tetra, penta and hexa, repeat type and repeat kind (simple and composite) over all the nine chromosomes will be useful to breeding researchers and QTL placements to select desired type of STR markers.

**In silico discovery of polymorphic markers**

A total of five genotypes namely, KWS2320, KWS230 DH1440 (KDHBv), STR06A6001 (UMSBv), SynMono (YMoBv) and SynTilling (YTiBv) (http://bvseq.molgen.mpg.de/) were used for *in silico* discovery of polymorphic markers using selected SSRs. Since polymorphism is exhibited by SSR having greater than or equal to eight repeat unit (35), these were selected and all simple repeats except ‘mono-nucleotide’ repeats were selected for discovery of polymorphic markers. For this, in house perl scripts were written accordingly. Further, selected primers were put in e-PCR (36) among five genotypes. Locus having difference in PCR product size were considered as polymorphic.

**Results and discussion**

**Analysis of sugarbeet genome and relative abundance**

The overall analysis of available sugarbeet genome gives the association of the distribution of the microsatellite

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**Figure 1. Flow of the database search.**
markers to the genomic attributes. A total of 145 K STR markers were successfully mined and populated in database as user friendly application. The distribution of simple and compound repeat types were 88 and 12%, respectively. Among simple type, ‘mono’ repeat type were more prevalent with 76.82%, followed by ‘di’ repeats, which was 13.68%. Although ‘di-nucleotide’ repeat type are observed abundantly in eukaryotes (38), on the contrast, our analysis reports ‘mono’ repeat patterns as the most abundant type (Figure 2). Since MISA parameters were not set for any threshold for mono-repeats, thus this prominence might be due to the inherent limitation of the NGS technology used which causes more mono nucleotide stretches as sequencing error (39).

STR markers being ubiquitously distributed, proportionately higher repeat content for longer chromosomes are expected (40), which is also observed in the present analysis. The most abundant STRs were distributed in Chromosome 1, followed by Chromosome 6 and 5, while Chromosome 3 contains the least abundant STRs (Table 1). The proportion of STRs with size less than (<10 bp) was maximum (57.05 %) followed by the ones between the size range of 11–13 bp (28.38%) and size range 14–25 bp (13.32%). Only 1.26% of the total STRs belonged to the size more than 25bp (Figure 3). Chromosome 1 showed highest density (671.05 markers/Mb) of markers and chromosome 6 reports minimum density of markers (341.27 markers/Mb), while the relative density of the sugarbeet whole genome is 378.54 markers/Mb, showing that these markers are ubiquitously distributed with homogeneity in terms of distance, which is inherent attribute of microsatellite to be used as marker of choice. Remaining all seven chromosomes were having the marker density of 341.27 to 384.94 marker/Mb.

The relative density of the sugarbeet whole genome reported in the present study is 379 markers per Mb, which is more than the range in Arabidopsis (157 markers per Mb). The other crops having similar number of markers are, cucumber (367 markers per Mb), rice (370–490 markers per Mb), poplar (485 markers per Mb) and grape (487 markers per Mb).

The initial linkage map of sugarbeet was having nine groups, with 700 cM coverage with just 500 STR markers (26, 27). An extended genetic map of sugar beet (Beta vulgaris L.) was achieved with 177 segregating markers on nine linkage groups (26). The linkage map comprises 1057.3 cM. Marker density calculations of present genetic map reveal a distance of ~2 cM between markers. The bulk set of markers (145 K), identified in the present study were assigned to the projected physical map and showed 430-fold higher marker density i.e. segregating two markers with a distance of 2.6 Kb. Since the average size of any eukaryotic gene falls within this distance between markers. Thus, these set of markers can ensure mapping of almost all genes.

In evaluation of 15 513 repeats by e-PCR, we found 2027 polymorphic markers in panel of five genotypes. Chromosome-wise distribution is summarized in Table 2 and details are given in supplementary table (Supplementary Table 1).

**Utility of the database**

Previously, several attempts have been made for increasing the markers based species delineation and genus identification events in *Beta vulgaris*. Earlier attempts have been made using the morphological descriptor and isozyme markers to differentiate *Beta vulgaris* and *B. webbiana* (41). Earlier attempts have been made to delineate the approaches for the varieties/lines differentiation within the *B. vulgaris* species using both STR and SNP markers.

Varieties/line differentiation within the species of *B. vulgaris* has been attempted by both STR and SNP
markers, e.g. a limited 677 SNP markers have been used for differentiation of 924 lines of sugarbeet (42).

However, there is limited use of STR markers in sugarbeet variety identification as reported earlier (43). Additionally, the number of informative morphological characters is limited in sugarbeet that often leads to some problems in variety registration (43). Previously there have been reports on the varietal differentiation using 12 STR markers in this species (43). However, the amount of the markers used were very few, which is a bottle-neck in this species. In the present report, the identified 145 K markers, can serve as a good reference resource for the development of the varietal identification markers. These whole genome markers have also played a role in the mapping and variety identification supplementing Distinctness, Uniformity and Stability (DUS) test and product trace ability (44). Use of STR in plant variety identification is well reported in other crops like barley varieties (45), S. tuberosum ssp. tuberosum (46), sugarcane (47), capsicum (48) and identification of Basmati rice from that of non-Basmati rice (49) etc.

STR markers have known to play an important role in regulating the gene expression. The observed markers in the present study are ubiquitously distributed can help in deciphering the gene level regulation. For example, length changes of microsatellites within promoters and other cis-regulatory regions can also change gene expression quickly, between generations. Such mechanism is already reported in large number of genomes. For example, in case of human genome, more than 16 000 STRs in regulatory regions are working as ‘tuning knobs’ for gene expression (50). Additionally, STR markers distributed across the intronic regions are reported to influence phenotype (51, 52). Such relation of STR with phenotype has not reported in Beta vulgaris. Our markers can be used for exploration of similar association. Transposable element contributes in plant gene regulation (53). Such transposable elements are present in sugarbeet also (54). It is probable that short sequence repeats in those locations are also involved in the regulation of gene expression (55). The repeat sequence mined in our database can be used for such studies where transposable elements play role in gene regulation.

STR has been used to trace hybridization and introgression events with wild beet to monitor feral or wild beet characters in GM beets (56), genetic diversity and root traits (57). Similarly, SNP markers have also been used for diversity analysis of sugarbeet (57). Our in silico discovered 2027 polymorphic markers can also be used for
diversity studies and phylogenetic studies of varieties or species.

A deep review of the previously published literature illustrates that identification of candidate genes for marker-assisted selection can improve the efficiency of breeding for increased drought tolerance (58). Need of markers to improve crop efficiency ratio in alkali soil for alcohol production is reported by Garg and Khanduja (59). Molecular markers are needed for mapping of disease resistance genes by linkage analysis in sugarbeet (60). Use of molecular markers are reported for construction of linkage map and identification of commercially valuable CMS in sugarbeet (61). Genes for various economically and commercially relevant trait of sugarbeet has been reported, e.g. seedling vigour (62), FLC-like gene BvFL1 associated with annuality and winter hardiness (25), root traits (37), non-restoring allele for Owen-type cytoplasmic male sterility, for development of molecular markers for the maintainer genotype (61), aphid resistance (63), nematode resistance (64), QTL for leaf spot (65), hardiness and bolting (66) and drought and salt tolerance (67). To increase the crop efficiency for bioenergy mapping of bioenergy traits are imperative (68). Markers can be used as a genomic resources to increase the biofuel potential from sugarbeet (30). We believe that the genome-wide STR makers developed and displayed graphically in our database with the options to synthesize the primers directly for the desired regions of the chromosome will serve the ease of developing markers for screening the mapping-based population for the genes involved in the several key domestication and biofuel traits.

Additionally, SBMDb provides access to the first ever comprehensive catalogue of the SSR-FDMs along with the markers from the genome wide coverage. Previously, SSR-FDMs have been widely used for the fluorescent based markers with an average of 7.42 alleles per locus in sugarcane (34). Utility of these markers also established the structure–function relationship for the beta-amylase and protein kinase encoding unigenes, which harbors the functional repeats in the catalytic domains (34). It is worthwhile to mention that a high robust amplification efficiency (96.5%) and high intra-specific polymorphic potential (34%) has been recently been exploited for the genotyping and trait association mapping in Chickpea using the 1108 transcription factor gene-derived microsatellite (TFGMS) and 161 transcription factor functional domain-associated microsatellite (TFFDMS) markers (69). Based on the above observations and later on application of the SSR-FDMs in several crop species such as Ocimum basilicum (70), Seasme indicum (71), Elaeis guineensis (72) and Camellia sinensis (73), suggests that linking the identified markers to the possible functional domains extends the evaluation of these markers from genotypic arrays and possibly can help us to elucidate the possible linkage of the strand slippage mechanism to the functional relevance.

If STR markers from our database are used for mapping of these genes or markers of flanking regions of these specific genes are selected, then they can directly be used in molecular breeding program for introgression of these genes/traits.

Conclusion

Using a computationally intensive in silico approaches, we mined and catalog the 145 K STR markers and built the first whole genome based STR database, which is freely accessible to the public domain at http://webapp.cabgrid.res.in/sbmdb/. With the marker information present in the SBMDb, the linkage map’s marker density can be increased which will facilitate in QTL and gene mapping. In order to facilitate the use of these markers in various molecular breeding and QTL programs, we have implemented several plug-in to generate primers at user defined chromosomal locations, which can be directly exported for genotyping assays. Additionally, the identified polymorphic markers can also be used for the DUS test for variety identification and improvement, MAS/GAS, QTL and gene mapping and germplasm improvement and management through marker genotyping. The present database will overcome the need of the marker portal of the sugarbeet genomics and the user friendly design will also help in the easy to access the marker information for molecular breeding required to accelerate industrious use of this crop, especially for sugar, biofuel/bioenergy, health care products, medicines, color dye. These markers need widest utilization across globe for best industrious use of sugarbeet by improvement of bioenergy trait of bioethanol and biogas production. This will not only improve the crop efficiency, but will also be a model industrial crop in the endeavour of water and carbon footprint in the challenging climate change regime.

Availability and requirement

SBMDb, the sugarbeet microsatellite marker database is freely accessible for research purposes for non-profit and academic organizations at http://webapp.cabgrid.res.in/sbmdb/.

Supplementary Data

Supplementary data are available at Database Online.
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References

1. Marlander, B., Hoffmann, C., Koch, H.J., et al. (2003) Environmental situation and yield performance of sugar beet crop in Germany: heading for sustainable development. J. Agron. Crop Sci., 189, 201–226.

2. Zhou, G.Q., Zhang, G.F. and Qi, D.M. (2011) A new method of producing bio-energy by using sugar-beets. Energy Procedia., 12, 873–877.

3. Dohm, J.C., Minoche, A.E., Holtgrawe, D. et al. (2014) The genome of the recently domesticated crop plant sugar beet (Beta vulgaris). Nature, 505, 546.

4. Pal, R., Girgenrunde, K., Gevraya, H. et al. (2010) Hepatoprotective activity of Beta vulgaris against CCl4 induced acute hepatotoxicity in rats. Arch. Appl. Sci. Res., 2, 14–18.

5. Klieffmann. (2011). http://www.klieffmann.com/openncms/openncms/de/mi德center/Pressemitteilungen/Zuckereruebe/ December 2014, date last accessed

6. FAO Statistical Yearbook, World Food and Agriculture. (2013) Food and Agriculture Organization of United Nations, ISSN 2225-7373, Rome.

7. Kumar, S., Singh, P.K., Swapna, M. et al. Souvenir: IISR-Sugarcane Technology Interface on Research and Development Initiatives for Sugarbeet in India. 2013; Association of Sugarcane Technologists of India, Indian Institute of Sugarcane Research, Lucknow, India.

8. Bhattacharya, A.N., Khan, T.M. and Uwayjan, M. (1975) Dried beet pulp as a sole source of energy in beef and sheep. J. Anim. Sci., 41, 616–621.

9. Burkhardt, C.G. and Burkhardt, C.N. (2009) Dihydroxyacetone and methods to improve its performance as artificial tanner. T. O. Derm. J., 3, 42–43.

10. Concha, J., Weinstein, C. and Zuniga, M.E. (2013) Production of pectic extracts from sugar beet pulp with anti-proliferative activity on a breast cancer cell line. Front. Chem. Sci. Eng., 7, 482–489.

11. Sakac, M.B., Pericin, D.M., Mandic, A.I. et al. (2004) Antioxidant properties of ethanolic extract of sugar beet pulp. APTEFF, 35, 1–280.

12. Satyanand, V., Vati, S.M., Krishna, B.P. et al. (2014) A study of beet root derived dietary nitrate efficacy on performance of Runners. Ind. J. Basic Appl. Med. Res., 3, 690–695.

13. Carlezon, Jr. W.A., Mague, S.D., Parow, A.M. et al. (2005) Antidepressant-like effects of uridine and omega-3 fatty acids are potentiated by combined treatment in rats. Biol. Psychiat., 57, 343–350.

14. Aksu, Z. and Isoglu, I.A. (2006) Use of agricultural waste sugar beet pulp for the removal of Gemazol turquoise blue-G reactive dye from aqueous solution. J Hazard. Mater., 137, 418–430.

15. Jain, N.K. and Singhai, A.K. (2012) Protective role of Beta vulgaris L. leaves extract and fractions on ethanol-mediated hepatic toxicity. Acta Pol. Pharm. Drug Res., 69, 945–950.

16. Craig, S.A.S. (2004) Betaine in human nutrition. Am. J. Clin. Nutr., 80, 539–549.

17. Kujala, T.S., Loponen, J.M., Klika, K.D. et al. (2000) Phenolics and betacyanins in red beetroot (Beta vulgaris) root: distribution and effect of cold storage on the content of total phenolics and three individual compounds. J. Agr. Food Chem., 48, 5338–5342.

18. Henke, W., Herdel, K., Jung, K., et al. (1997) Betaine improves the PCR amplification of GC-rich DNA sequences. Nucleic Acids Res., 25, 3957–3958.

19. Gerbens-Leenesa, W., Hoekstra, A.Y. and van der Meer, T.H. (2009) The water footprint of bioenergy. Proc Natl. Acad. Sci. USA, 106, 10219–10223.

20. de Figueiredo, E.B., Panoso, A.R., Romao, R. et al. (2010) Greenhouse gas emission associated with sugar production in southern Brazil. Carbon Bal Manage., 5, 3–10.

21. Biancardi, E., McGrath, J.M., Panella, L.W. et al. (2010) SugarBeet. In: Bradshaw, J.E. (ed.), Root and Tuber Crops, Handbook of Plant Breeding. New York, Springer, 7, 173–219.

22. Kirchhoff, M., Svirchchevskaya, A., Hoffmann, C. et al. (2012) High degree of genetic variation of winter hardness in a panel of Beta vulgaris L. Crop Sci., 52, 179–188.

23. Ober, E.S., Clark, C.J.A., Bloa, M.L. et al. (2004) Assessing the genetic resources to improve drought tolerance in sugar beet: agronomic traits of diverse genotypes under droughted and irrigated conditions. Field Crop. Res., 90, 213–234.

24. Shane, W.W. and Teng, P.S. (1992) Impact of Cercospora leaf spot on root weight sugar yield and purity of Beta vulgaris. Plant Dis., 76, 812–820.

25. Frerichmann, S.M., Kirchhoff, M., Muller, A.E. et al. (2013) EcoTILLING in Beta vulgaris reveals polymorphisms in the FLC-like gene BvFL1 that are associated with annuality and winter hardness. BMC Plant Biol., 13, 52.

26. Grimmer, M.M.K., Trybush, S., Hanley, S. et al. (2007) An anchored linkage map for sugar beet based on AFLP, SNP and RAPD markers and QTL mapping of a new source of resistance to beet necrotic yellow vein virus. Theor. Appl. Genet., 114, 1151–1160.

27. Paesold, S., Borchardt, D., Schmidt, T. et al. (2012) A sugar beet (Beta vulgaris L.) reference FISH karyotype for chromosome and chromosome-arm identification, integration of genetic linkage groups and analysis of major repeat family distribution. Plant J., 72, 600–611.

28. Dohm, J.C., Lange, C., Holtgrawe, D. et al. (2012) Palaeoexaploid ancestry for Caryophyllales inferred from extensive gene-based physical and genetic mapping of the sugar beet genome (Beta vulgaris). Plant J., 70, 528–540.

29. Holtgrawe, D., Sorensen, T.R., Viehover, P. et al. (2014) Reliable in silico identification of sequence polymorphisms and their application for extending the genetic map of sugar beet (Beta vulgaris). PLoS One, 9, e110113.

30. Fugate, K.K., Fajardo, D., Schlautman, B. et al. (2014) Generation and characterization of a sugarbeet transcriptome and transcript-based SSR markers. Plant Gen., 7, 1–13.
31. Yu,J.K., Paik,H., Choi,J.P. et al. (2010) Functional domain marker (FDM): An in silico demonstration in Solanacea using simple sequence repeats (SSRs). Plant Mol. Biol. Rep., 28, 352–356.

32. Sablok,G. and Shekhawat,N.S. (2008) Bioinformatics analysis of distribution of microsatellite markers (SSRs)/single nucleotide polymorphism (SNPs) in expressed transcripts of Prosopis Juliflora: frequency and distribution. J. Comput. Sci. Syst. Biol., 1, 87–91.

33. Sablok,G., Luo,C., Lee,W.S. et al. (2010) A comprehensive database of distribution of microsatellite markers (SSRs)/single nucleotide polymorphism (SNPs) in expressed transcripts of Prosopis Juliflora: frequency and distribution. J. Comput. Sci. Syst. Biol., 1, 87–91.

34. Parida,S.K., Pandit,A., Gaikwad,K. et al. (2010) Functionally relevant microsatellites in sugarcane unigenes. BMC Plant Biol., 10, 251.

35. Kelkar,Y.D., Strubczewski,N., Hile,S.E. et al. (2010) What is a microsatellite: a computational and experimental definition based upon repeat mutational behavior at A/T and GT/AC repeats. Genome Biol. Evol., 2, 620–635.

36. Schuler,G.D. (1997) Sequence mapping by electronic PCR. Genome Res., 7, 541–550.

37. Quevillon,E., Silventoinen,V., Pillai,S. et al. (2005) InterProScan: protein domains identifier. Nucleic Acids Res., 33, 116–120.

38. Shioiri,C. and Takahata,N. (2001) Skew of mononucleotide frequencies, relative abundance of dinucleotides and DNA strand asymmetry. J. Mol. Evol., 53, 364–376.

39. Haseneyer,G., Schmutzer,T. and Seidel,M. (2011) From RNA-seq to large-scale genotyping-genomics resources for rye (Secale cereale L.). BMC Plant Biol., 11, 131–143.

40. Zietkiewicz,E., Rafalski,A. and Labuda,D. (1994) Genome fingerprinting by simple sequence repeat (SSR)-anchored polymerase chain reaction amplification. Genomics, 20, 176–218.

41. Reamond-Ramos,S.M. and Wricke,G. (1992) A full set of monosomic addition lines in Beta vulgaris from Beta Webbiana: morphology and isozyme markers. Theor. Appl. Genet., 84, 411–418.

42. Wurschum,T., Reif,J.C., Kraft,T. et al. (2013) Genomic selection in sugar beet breeding populations. BMC Genet., 14, 85–92.

43. Smulders,M.J.M, Esselink,G.D., Everaert,I. et al. (2010) Characterisation of sugar beet (Beta vulgaris L. ssp. vulgaris) varieties using microsatellite markers. BMC Genet., 11, 41.

44. Iqubal,M.A., Sarika, Vasu,A. et al. (2013) First whole genome based microsatellite DNA marker database of tomato for mapping and variety identification. BMC Plant Biol., 13, 41.

45. Karakousis,A., Barr,A.R., Chalmers,K.J. et al. (2003) Potential of SSR markers for plant breeding and variety identification in Australian barley germplasm. Aust. J. Agr. Res., 54, 1197–1210.

46. Kawchuk,L.M., Martin,R.F. and McPherson,J. (1990) Resistance in transgenic potato expressing the potato leafroll virus coat protein gene. Mol. Plant Microbe In, 3, 301–307.

47. Manigbas,N.L. and Villegas,L.C. (2004) Microsatellite markers in hybridity tests to identify true hybrids of sugarcane. Philipp J. Crop. Sci., 29, 23–32.

48. Shirasawa,K., Ishii,K., Kim,C. et al. (2013) Development of Capsicum EST-SSR markers for species identification and in silico mapping onto the tomato genome sequence. Mol. Breed. 31, 101–110.

49. Archak,S., Lakshminarayanareddy,V. and Nagaraju,J. (2007) High-throughput multiplex microsatellite marker assay for detection and quantification of adulteration in Basmati rice (Oryza sativa). Electrophoresis, 28, 2396–2405.

50. Rockman,M.V. and Wray,G.A. (2002) Abundant raw material for cis-regulatory evolution in humans. Mol. Biol. Evol., 19, 1991–2004.

51. Akagi,T., Yin,D., Kawamata,N. et al. (2009) Functional analysis of a novel DNA polymorphism of a tandem repeated sequence in the asparagine synthetase gene in acute lymphoblastic leukemia cells. Leuk. Res., 33, 991–996.

52. Kersting,C., Angelopoulos,K., Schmidt,H. et al. (2008) Biological importance of a polymorphic CA sequence within intron I of the epidermal growth factor receptor gene (EGFR) in high grade central osteosarcomas. Gene Chromosomes Cancer, 47, 657–664.

53. Bennetzen,J.L. (2000) Transposable element contributions to plant genome evolution and genome evolution. Plant Mol. Biol., 42, 251–269.

54. Menzel,G., Krebs,C., Diez,M. et al. (2012) Survey of sugar beet (Beta vulgaris L.) hAT transposons and MITE-like hATpin derivatives. Plant Mol. Biol., 78, 393–405.

55. Tomilin,N.V. (2008) Regulation of mammalian gene expression by retro elements and non-coding tandem repeats. BioEssays, 30, 338–348.

56. Stark,C., Liepelt,S., Dieckvoss,M. et al. (2006). Fast and simple monitoring of introgressive gene flow from wild beet into sugar-beet. J Sugar Beet Res., 43, 145–154.

57. Stevanato,P., Daniele,B., Bullet,T. et al. (2013) Evaluation of genetic diversity and root traits of sea beet accesses of the Adriatic Sea coast. Euphytica, 189,135–146.

58. Hajheidari,M., Abdollahian-Noghabi,M., Askari,H. et al. (2005) Proteome analysis of sugar beet leaves under drought stress. Proteomics, 5, 950–960.

59. Garg,V.K. and Khanduja,S.D. (1986) Energy efficiency ratio of sugar beet grown on alkali soils in Northern India. Biomass, 11, 147–151.

60. Hunger,S., Di Gaspero,G., Mohring,S. et al. (2003) Isolation and linkage analysis of expressed disease-resistance gene analogues of sugar beet (Beta vulgaris L.). Genome, 46, 70–82.

61. Moritani,M., Taguchi,K., Kitazaki,K. et al. (2013) Identification of the predominant nonrestoring allele for Owen-type cytoplasmic male sterility in sugar beet (Beta vulgaris L.): development of molecular markers for the maintainer genotype. Mol. Breed., 32, 91–100.

62. de los Reyes,B.G. and McGrath,J.M. (2003) Cultivar-specific seedling vigor and expression of a putative oxalate oxidase germ-line-like protein in sugar beet (Beta vulgaris L.). Theor. Appl. Genet. 107, 54–61.

63. Leijman,E. (2011) Fine mapping of resistance against root aphids in sugar beet. The Faculty of Landscape Planning, Horticulture and Agriculture Science, SLU, Alnarp, Master thesis in Biology, Horticulture Programme, Alnarp.
nematode resistance from *Beta procumbens*. Mol. Anal. Translocat. Sugarbeet. 31, 27–42.

65. Taguchi,K., Kubo,T., Takahashi,H. et al. (2011) Identification and precise mapping of resistant QTLs of cercospora leaf spot resistance in sugar beet (*Beta vulgaris* L.). *Genes Genom. Genet.*, 1, 283–291.

66. Pfeiffer,N., Trankner,C., Lemnian,I. et al. (2014) Genetic analysis of bolting after winter in sugar beet (*Beta vulgaris* L.). *Theor. Appl. Genet*.

67. Monteiro,F., Romeiras,M.M., Batista,D. et al. (2013) Biodiversity assessment of sugar beet species and its wild relatives: linking ecological data with new genetic approaches. *Am. J. Plant Sci.* 4, 21–34.

68. Feltus,F.A. and Vandenbrink,J.P. (2012) Bioenergy grass feedstock: current options and prospects for trait improvement using emerging genetic, genomic, and systems biology toolkits. *Biotech. Biofuel.*, 5, 80.

69. Kujur,A., Bajaj,D., Saxena,M.S., et al. (2013) Functionally relevant microsatellite markers from chickpea transcription factor genes for efficient genotyping applications and trait association mapping. *DNA Res.*, 20, 355–373.

70. Gupta,S., Shukla,R., Roy,S. et al. (2010) *In silico* SSR and FDM analysis through EST sequences in *Ocimum basilicum*. *POJ.*, 3, 121–128.

71. Bhattacharyya,U., Pandey,S.K. and Dasgupta,T. (2014) Identification of EST-SSRs and FDM in sesame (*Sesamum indicum* L.) through data mining. *Sch. J. Agric. Sci.*, 4, 60–69.

72. Tranbarger,T.J., Kluabmongkol,W., Sangsraekru,D. et al. (2012) SSR markers in transcripts of genes linked to post-transcriptional and transcriptional regulatory functions during vegetative and reproductive development of *Elaeis guineensis*. *BMC Plant Biol.*, 12, 1.

73. Sahu,J., Sarmah,R., Dehury,B. et al. (2012) Mining for SSRs and FDMs from expressed sequence tags of *Camellia sinensis*. *Bioinformatics*, 8, 260–266.