GAN: a platform of genomics and genetics analysis and application in Nicotiana

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Abstract

Nicotiana is an important Solanaceae genus, and plays a significant role in modern biological research. Massive Nicotiana biological data have emerged from in-depth genomics and genetics studies. From big data to big discovery, large-scale analysis and application with new platforms is critical. Based on data accumulation, a comprehensive platform of Genomics and Genetics Analysis and Application in Nicotiana (GAN) has been developed, and is publicly available at http://biodb.sdau.edu.cn/gan/. GAN consists of four main sections: (i) Sources, a total of 5267 germplasm lines, along with detailed descriptions of associated characteristics, are all available on the Germplasm page, which can be queried using eight different inquiry modes. Seven fully sequenced species with accompanying sequences and detailed genomic annotation are available on the Genomics page. (ii) Genetics, detailed descriptions of 10 genetic linkage maps, constructed by different parents, 2239 KEGG metabolic pathway maps and 209 945 gene families across all catalogued genes, along with two co-linearity maps combining N. tabacum with available tomato and potato linkage maps are available here. Furthermore, 3 963 119 genome-SSRs, 10 621 016 SNPs, 12 388 PIPs and 102 895 reverse transcription-polymerase chain reaction primers, are all available to be used and searched on the Markers page. (iii) Tools, the genome browser JBrowse and five useful online bioinformatics softwares, Blast, Primer3, SSR-detect, Nucl-Protein and E-PCR, are provided on the JBrowse and Tools pages. (iv) Auxiliary, all the datasets are shown on a Statistics page, and are available for download on a Download page. In addition, the user’s manual is provided on a Manual page in English and Chinese languages. GAN provides a user-friendly Web interface for searching, browsing and downloading the genomics and genetics datasets in Nicotiana. As far as we can ascertain, GAN is the most comprehensive...
source of bio-data available, and the most applicable resource for breeding, gene mapping, gene cloning, the study of the origin and evolution of polyploidy, and related studies in *Nicotiana*.

**Database URL:** http://biodb.sdau.edu.cn/gan/

### Introduction

The genus *Nicotiana* is the fourth largest genus in the family Solanaceae. Extensive research has accumulated a huge amount of data regarding genetics, evolution, genomics, taxonomy and breeding in the genus (1). Tobacco (*Nicotiana tabacum* L.) is cultivated worldwide as a plant of economic importance, as well as a model system in plant biotechnology (2). The collection and utilization of germplasm are the foundation of botanical research. A generally accepted international standard taxonomy for *Nicotiana* has existed for decades. It consists of 3 sub-genera, 14 sections and 66 species (3). However, most researchers now tend to classify *Nicotiana* into 13 sections and 76 species (4, 5). Regardless, several *Nicotiana* germplasm storehouses have been built over the years to collect and maintain *Nicotiana* germplasm resources. Presently, approximately 2152 *Nicotiana* accessions are maintained at the NPGS (National Plant Germplasm System, a world-famous genetic diversity warehouse for *N. tabacum*, http://www.ars-grin.gov/npgs/index.html) (6). In addition, a total of 1160 *Nicotiana* accessions are held at the CGRIS (Chinese Crop Germplasm Resources Information System, http://icgr.caas.net.cn/cgris_english.html) (7). Furthermore, the Tobacco Genetics and Breeding database (TGB, http://biodb.sdau.edu.cn/rgb/) also provides 1472 *Nicotiana* accessions with accompanying detailed annotations (8). Genus *Nicotiana* germplasm is generally understood to be available under >5000 accessions worldwide; however, no systematic database exists to preserve and catalogue all these germplasm resources.

The *N. tabacum* genome remained a challenge to sequence for quite a long time, because it is allotetraploid and very large. The allotetraploid (2n = 4x = 48) *N. tabacum* genome contains 24 pairs of chromosomes, derived from the two diploid species *N. sylvestris* and *N. tomentosiformis* (9, 10). High-throughput sequencing technologies have developed rapidly; comparative genomics can now provide insights into seven *Nicotiana* species: the allotetraploid *N. benthamiana* (11); three diploid species, *N. otophora*, *N. sylvestris* and *N. tomentosiformis* (10) and three common cultivated species, *N. tabacum* BX, *N. tabacum* K326 and *N. tabacum* TN90 (12). In the recent years, there was numerous sequences read archives available for other *Nicotiana* species like *N. obtusifolia*, *N. nudicaulis* and *N. repanda* and so on. Some of them were successfully used to identify allopolyploid species origin at a great accuracy (13, 14). So the high research value promoted to construct a genome information database which would provide convenient access to all the data.

The establishment of molecular markers and genetic linkage maps has greatly influenced gene mapping, gene cloning, quantitative trait loci and marker-assisted selection research (15, 16). As a model plant, genetic analyses in different *Nicotiana* genera employ a diversity of molecular markers (17–20). Molecular markers, restriction fragment length polymorphisms (RFLPs) (21), random amplified polymorphic DNAs (RAPDs) (22), amplified fragment length polymorphisms (AFLPs) (23), short sequence repeats (SSRs) (24), intron length polymorphisms (25), single nucleotide polymorphisms (SNPs) (26) and inter-simple sequence repeats (ISSRs) (27), were all excavated from previous research on a large scale. Ten *Nicotiana* genetic linkage maps were developed based on these markers, in our previous studies (28). However, these results were all derived from contig or scaffold sequences, not from whole genomes. Consequently, these marker linkage analyses needed to be redone on a large-scale using genome sequences.

An excellent resource for the online tobacco research community, the TGB database contains 1472 *Nicotiana* germplasms, with accompanying detailed annotations and 12 388 potential intron polymorphisms (PIPs), 10 351 EST-simple sequence repeats and 66 297 genomic-SSR markers (G-SSRs) (8). However, with the huge amount of new bio-data emerging in recent years, TGB can not satisfy all new research needs. Therefore, we found it necessary to establish a comprehensive platform of Genomics and Genetics Analysis and Application in *Nicotiana* (GAN, http://biodb.sdau.edu.cn/gan/). GAN contains a Source Section (Germplasm and Genomes), a Genetics Section (Markers, Genetics Maps and Genetics Annotations), a Tools Section [JBrowse (29) and Online Tools] and an Auxiliary Section (Datasets, Download and Manual) for *Nicotiana* research. GAN will greatly expedite genomics and genetics research in *Nicotiana* and related plants.

### Platform content and web interface

GAN provides information regarding the detailed annotation of 5267 *Nicotiana* accessions, 7 genomes, 10 genetic
linkage maps, 2239 KEGG metabolic pathway maps, 209 945 gene families, two co-linearity maps, 3 963 119 genome-SSRs, 10 621 016 SNPs, 12 388 PIPs, 102 895 reverse transcription-polymerase chain reaction (RT-PCR) primers, JBrowse and 5 useful online softwares. The GAN Web interface was designed to include the following components: Source, Genetics, Tools and Auxiliary (Figure 1).

JBrowse and the five online bioinformatics tools, Blast (30), Primer3 (31), SSR-detect (32), Nucl-Protein and E-PCR (33), are provided on the JBrowse and Tools pages. An Auxiliary Section contains all the datasets in GAN, displayed on a Statistics page and available for download on the Download page. In addition, the Auxiliary Section contains a user’s manual, provided on the Manual page in English and Chinese languages. GAN provides a user-friendly Web interface for searching, browsing and downloading genomics and genetics datasets for Nicotiana. As far as we know, GAN provides the most comprehensive Nicotiana bio-data available, and is an excellent resource for research in Nicotiana breeding, gene mapping, gene cloning, the origin and evolution of polyploidy and related studies.

The GAN platform provides researchers all necessary data, and is composed of five main parts: Germplasm, Genomics, Genetics, Markers and Tools (Figure 2). (i) The Germplasm Section holds 5267 Nicotiana accessions, which can be queried using several different routes: specific ID, species, Nicotiana type, cultivar origin, agronomic characteristics and disease resistance characteristics. Furthermore, detailed germplasm information is also provided with high-quality images from the field or laboratory. (ii) The Genomics Section contains the seven available Nicotiana genome sequences, and all sources and annotations. (iii) The Genetics Section contains genetic maps, gene annotations, synteny analyses and gene family make up analyses for those seven Nicotiana species with complete genome data. (iv) The Markers Section contains all of the three widely used molecular markers, SSRs, SNPs and PIPs, that have been detected in those seven Nicotiana genes, and some of the exons, introns and UTRs. Furthermore, because RT-PCR is such a sensitive and widely used method for the detection of mRNA expression levels, RT-PCR primers are also available, predesigned using the coding sequences (CDSs) of current Nicotiana species. (v) The Tools Section contains the online bioinformatics tools Blast, SSR-detect, Primer3, Nucl-Protein and E-PCR, which have all been integrated into GAN, as well as the currently popular genome browser JBrowse for the display of detailed annotation and marker information overlaid on the genome.

Germplasm

We divide the 5267 GAN accessions into 35 Nicotiana species, of which ~92.8% are N. tabacum L., in accordance
The germplasm type is differentiated into seven tobacco categories: Aromatic, Burly, Cigar, Flue-cured, Wild, Rustica and Sun-cured. The 5267 Nicotiana accessions come from 45 different cultivar origins, with most cultivar originating in China and the next most from North America (Table 1). Detailed disease resistance information is provided for all accessions, including Black Shank, Granville Wilt, Root-knot Nematode, Brown Spot, Tobacco mosaic virus, Cucumber mosaic virus and Potato virus Y (Figure 3).

The Germplasm page can be searched using ID-Name, species, type, cultivar origin, agronomic characteristics and disease resistance keywords. Inputting your desired words and clicking the ‘Keywords Search’ button initiates a global keyword search. Users can also click the ‘Submit’ button in the Germplasm gray-box to browse all germplasms, displaying all 35 Nicotiana species, 7 Nicotiana types and 45 cultivar origins, which can then be chosen among. Finally, three different specific identifiers, the germplasm’s name, its GAN ID and its National Uniform Number, can be used to search for a specific germplasm. Thirty-five common field agronomic characteristics are provided including phenotype and physiology. Users can choose one or more parameters to generate exact results. Similar to agronomic characteristics, the disease resistance search box offers seven kinds of disease, including different levels of resistance, such that the user can obtain accurate germplasm identifications. Results in the Web interface
appear as simple information entries in tables in which users can click a ‘Detail’ hyperlink to generate detailed information about a specific germplasm. The Detail Web page displays four main attributes: basic information, agronomic characteristics, disease resistance and *Nicotiana* images for each germplasm. The basic information entry provides the *Nicotiana* species of the germplasm along with an introduction, common names, economic importance and references.

### Genomics

Seven sequenced *Nicotiana* genomes, including assembly and annotation information, genome size (~2.7 G through ~5.18 G), contigs and scaffolds can be found on the Genomics page.

### Genetics

Ten specific genetic maps were constructed in previous research using different RFLP, RAPD, ISSR and SSR molecular markers. Wild species, Burly and Flue-cured are all used as materials among the 24 chromosomes. Enzymes of 2115 relate to metabolic pathways found in 6 *Nicotiana* CDSs, which participate in 383 specific pathway maps. A total of 506 553 protein domains and/or families were identified using Pfam, Pkinase, WD domain and PPR family analyses in the 6 *Nicotiana* CDSs.

Users can browse ten genetics maps in the Genetics Section. Tables store constitutive data for these genetic maps in the main Web page. A graphical ‘Go’ button is pressed to get the requested genetic map; the maps contain detailed marker information obtained by clicking on the marker. KEGG annotation can then be queried if users...
select a specific *Nicotiana* species and input a sequence ID or KEGG Orthology number. Gene family, *Nicotiana* species or type, Gene ID, PF(Plant Family) accession and name can all be used to start the search.

**Molecular markers**

A total of 3,429,801 G-SSRs, 326,253 gene-SSRs and 18,485 CDS-SSRs were catalogued in this research. Additionally, 267,699 primer pairs were designed for *Nicotiana* genes. SNP markers of 10,621,016 were searched against and located on the reference *N. benthamiana* genome (Table 2). Furthermore, 12,338 PIP makers were obtained from the TGB platform. Finally, 102,859 pairs of E-PCR verified primers were generated for scientific research.

Generally, all of the Markers search pages require a *Nicotiana* species to be chosen as a necessary option. Different markers have their own search options beyond that: The SSR page provides options for sequence type, Gene ID, SSR motif and repeat number; the SNP page allows for searching by genome sequence ID, reference base and query base; the PIP page contains options for PIP ID, PlantGDB ID (http://www.plantgdb.org/) and gene name; the RT-PCR page requires a gene sequence ID. Clicking the ‘Detail’ hyperlink in the Result Web interface for all markers displays detailed information on the marker.

**Tools**

Five frequently used software tools are provided in GAN (Blast, Primer3, SSR-detect, Nucl-Protein and E-PCR) and one genome browser (JBrowse). Generally speaking users can paste sequences in FASTA format directly into a text-area, or upload sequence files from a local computer, into the five software pages. Users select from five Blast variants (blastp, blastx, blastn, tblastx and tblastn) on the Blast page, and search against any of the seven *Nicotiana* species genome sequences (or corresponding translations) along with basic program parameters to obtain results. In the SSR-detector page, SSRs can be identified using the default parameters of MISA (34). The Nucl-Protein page provides a tool for the translation of nucleotide sequences to protein sequences; users only need to provide a nucleotide sequence and working name, similarly with SSR-detector. The Primer Design page requires users to input the Min-TM, Max-TM, Min-GC, Max-GC and return number to predict maximal Primer3 results. The primer verification tool E-PCR uses the Primer3 results and requires users to select one *Nicotiana* species as a template, followed by a few alternative parameters. Results are obtained by online perusal or can be downloaded to the local computer.

JBrowse is an open-access genome browser that allows for the storage and display of genome sequences, genes, mRNAs, CDSs, introns, exons, UTRs, SSRs and SNPs, providing detailed annotation for each. In our implementation, the left column lists options consisting of *Nicotiana* genomes, a GFF choice, SNP loci, SNP coverage and SSR locations. The right column will display the corresponding information when users click the square box before the options in the left column. The GFF option provides an example showing how to work with the browsier. By clicking the GFF square all genome information will graphically display, then a dialog box will pop up if you click a specific graphic. Users can browse and download detailed information from the dialog box. We have added several new and useful features to the basic JBrowse browser (Figure 2).

**Download and help**

Users can download all the data used in GAN on the Download page. A download tree is used to access all the data relative to the Germplasm, Genomics, Genetics and Markers pages.

**Conclusions and perspectives**

GAN is a platform for providing *Nicotiana* accessions while integrating useful genomic and genetic data, and supplying relevant annotations and molecular markers in *Nicotiana*. The 5,267 *Nicotiana* accessions are the main content of the platform; however, GAN also expedites scientific work by identifying *Nicotiana* gene annotations and markers and primers for genetics analysis and breeding. Furthermore, the incorporation of several useful tools and JBrowse make it easy for users unfamiliar with bioinformatics to perform big-data scientific research. The platform will be updated continuously as new *Nicotiana* accessions are collected, and new *Nicotiana* genome sequences are generated.

**Table 2. The statistics of SSRs and SNPs**

| Species                  | Genome-SSR | Gene-SSR | SNP     |
|--------------------------|------------|----------|---------|
| *Nicotiana benthamiana*  | 492 540    | 61 568   | -       |
| *Nicotiana otophora*     | 358 420    | -        | 785 204 |
| *Nicotiana sylvesteris*  | 420 177    | 63 159   | 1 872 257 |
| *Nicotiana tabacum BX*   | 594 364    | 47 536   | 2 391 165 |
| *Nicotiana tabacum K326* | 620 818    | 45 437   | 2 403 132 |
| *Nicotiana tabacum TN90* | 637 331    | 44 660   | 2 442 813 |
| *Nicotiana tomentosiformis* | 306 151   | 63 893   | 726 445 |
Materials and methods

Germplasm and sequences

Information on 51 traits, and plant and inflorescence images of 5267 *Nicotiana* accessions were exhaustively collected. The genome sequences of *N. benthamiana*, *N. otophora*, *N. tabacum* BX, *N. tabacum* K326 and *N. tabacum* TN90 were downloaded from SGN (35) (Sol Genomics Network, https://solgenomics.net/) and the genome sequences of *N. sylvestris* and *N. tomentosiformis* were obtained from NCBI (National Center for Biotechnology Information, https://www.ncbi.nlm.nih.gov/).

Genetic maps and annotation

Ten *Nicotiana* genetic linkage maps with associated molecular markers were collected from previous research. Colinearity maps with tomato and potato were also extracted from previous *Nicotiana* sequence research (12). All relevant information can be found on the appropriate GAN page.

The KAAS Web interface (36) (http://www.genome.jp/tools/kaas/) was used annotate data with the KEGG pathway assignments (37) (Kyoto Encyclopedia of Genes and Genomes). PfamScan software (38) (http://www.ebi.ac.uk/Tools/pfa/pfamscan/) was used to scan gene families from Pfam (39) (http://pfam.xfam.org/).

Marker identification

From different type sequences of *Nicotiana* species, the SSRs (Genome-SSR, Gene-SSR, Cds-SSR, Exon-SSR and Intron-SSR) were detected by scanning monomer, dimer, trimer, tetramer, pentamer and hexamer nucleotide motifs with at least 10, 6, 5, 5 and 5 repeats by using perl-based MISA program (http://pgrc.ipk-gatersleben.de/misa/), respectively. For complex SSRs, the maximum difference between two SSRs had to be <100 bp. Primer3 was used for SSR primers design (31). Using *N. benthamiana* sequences as reference, and the other six *Nicotiana* species sequences as queries, SNPs were extracted and filtered using the BWA (Burrows-Wheeler Aligner) (40), Samtools (41) and Bcftools. Among them, BWA is responsible for database building and comparison. Samtools is responsible for processing and converting formats. Finally, Bcftools was used for SNP calling. All parameters were used at default settings.

Platform implementation

The LAMP framework (Linux, Apache, MySQL and PHP/Perl) was used to construct the GAN platform. All standardized data were imported into MySQL for storage and management. Commands are submitted in the Web-friendly interface using HTML on the Apache Web server, generating PHP scripts, which extract corresponding data from MySQL. Perl and Java scripts are used to enhance the appearance of the Web interface and to improve user query efficiency.

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Conflict of interest

None declared.

References

1. Lewis,R.S. (2011) *Nicotiana*. In: Kole,C. (ed). Wild Crop Relatives Genomic and Breeding Resources, Plantation and ornamental crops. Springer, Berlin, pp. 185–208.
2. Nekrasov,V., Staskawicz,B., Weigel,D. et al. (2013) Targeted mutagenesis in the model plant *Nicotiana benthamiana* using Cas9 RNA-guided endonuclease. *Nat. Biotechnol.*, 31, 691.
3. Harper,G.T. (1955) *The Genus Nicotiana*. Soil Science, 250.
4. Lewis,R.S. and Nicholson,J.S. (2007) Aspects of the evolution of *Nicotiana tabacum* L. and the status of the United States *Nicotiana* Germplasm Collection. *Genetic Resources Crop Evol.*, 54, 727–740.
5. Knapp,S., Chase,M.W. and Clarkson,J.J. (2004) Nomenclatural changes and a new sectional classification in *Nicotiana* (Solanaceae). *Taxon*, 53, 73–82.
6. Smale,M., Hodgkin,T., Zohrabian,A. et al. (2001) The demand for crop genetic resources: international use of the U.S. National Plant Germplasm System. *RePEc*, 30,1639–1655.
7. Fang,W. and Cao,Y. (2012) Chinese crop germplasm resources information system. *E-Sci. Technol. Appl.*, 3, 66–73.
8. Cao,H., Wang,Y., Xie,Z. et al. (2013) TGB: the tobacco genetics and breeding database. *Mol. Breed.*, 31, 655–663.
9. Gerstel,D.U. (1960) Segregation in new allopolyploids of *nicotiana*. I. Comparison of 6x (N. tabacum x tomentosiformis) and 6x (N. tabacum x otophora). *Genetics*, 45, 1723–1734.
10. Sierro,N., Battey,J.N., Ouadi,S. et al. (2013) Reference genomes and transcriptomes of *Nicotiana sylvestris* and *Nicotiana tomentosiformis*. *Genome Biol.*, 14, R60.
11. Bombarely,A., Rosli,H.G., Vrebalov,J. et al. (2012) A draft genome sequence of *Nicotiana benthamiana* to enhance molecular plant-microbe biology research. *Mol. Plant-Microbe Interactions: MPMI*, 25, 1523.
12. Sierro,N., Battey,J.N., Ouadi,S. et al. (2014) The tobacco genome sequence and its comparison with those of tomato and potato. *Nat. Commun.*, 5, 5833.
13. Renny-Byfield,S., Chester,M., Kovarik,A. et al. (2011) Next generation sequencing reveals genome downsizing in allotetraploid *Nicotiana tabacum*, predominantly through the elimination of paternally derived repetitive DNAs. *Mol. Biol. Evol.*, 28, 2843.
14. Rennybyfield, S., Kovarik, A., Kelly, L.J. et al. (2013) Diploidization and genome size change in allopolyploids is associated with differential dynamics of low- and high-copy sequences. *Plant J.*, 74, 829–839.
15. Prospero,S. and Rigling,D. (2016) Using molecular markers to assess the establishment and spread of a mycovirus applied as a
biological control agent against chestnut blight. *Biocontrol*, 61, 313–323.
16. Ott,J., Wang,J. and Leal,S.M. (2015) Genetic linkage analysis in the age of whole-genome sequencing. *Nat. Rev. Genetics*, 16, 275.
17. Yuan,X., Feng,C., Zhang,Z. et al. (2017) Complete mitochondrial genome of *phytophthora nicotianae* and identification of molecular markers for the oomycetes. *Front. Microbiol.*, 8, 1484.
18. Huang,L., Cao,H., Yang,L. et al. (2017) Large-scale development of SSR markers and their complementary applied in *Nicotiana*. *Russian J. Genetics*, 49, 827–838.
19. Bindler,G., Plieske,J., Bakaher,N. et al. (2011) A high density genetic map of tobacco (*Nicotiana tabacum L.*) obtained from large scale microsatellite marker development. *Theor. Appl. Genetics*, 123, 219.
20. Tong,Z., Yang,Z., Chen,X. et al. (2012) Large-scale development of microsatellite markers in *Nicotiana tabacum* and construction of a genetic map of flue-cured tobacco. *Plant Breed.*, 131, 674–680.
21. Lin,T.Y., Kao,Y.Y., Lin,R.F. et al. (2001) A genetic linkage map of *Nicotiana plumbaginifolia*/*Nicotiana longiflora* based on RFLP and RAPD markers. *Theor. Appl. Genetics*, 103, 905–911.
22. Linos,A., Nikoloudakis,N., Katsiotis,A. et al. (2014) Genetic structure of the Greek olive germplasm revealed by RAPD, ISSR and SSR markers. *Scientia Horticulturae*, 175, 33–43.
23. Xie,L., Wang,X., Peng,M. et al. (2014) Isolation and detection of differential genes in hot pepper (*Capsicum annuum L.*) after space flight using AFLP markers. *Biochem. Syst. Ecol.*, 57, 27–32.
24. Mudalkar,S., Golla,R., Ghatty,S. et al. (2014) De novo transcriptome analysis of an imminent biofuel crop, Camelina sativa L. using Illumina GAIIX sequencing platform and identification of SSR markers. *Plant Mol. Biol.*, 84, 159–171.
25. Yang,L., Jin,G., Zhao,X. et al. (2007) PIP: a database of potential intron polymorphism markers. *Bioinformatics*, 23, 2174–2177.
26. Gong,D., Huang,L., Xu,X. et al. (2016) Construction of a high-density SNP genetic map in flue-cured tobacco based on SLAF-seq. *Mol. Breed.*, 36, 100.
27. Hassani,T.F., Samizadeh,L.H. and Shoarei,D.M. (2014) Study of genetic diversity among and within types of tobacco (*Nicotiana Tabacum L.*) using ISSR markers. *Modern Genetics*, 9, 1–22.
28. Tong,Z., Xiao,B., Jiao,F. et al. (2016) Large-scale development of SSR markers in tobacco and construction of a linkage map in flue-cured tobacco. *Breed. Sci.*, 66, 381–390.
29. Skinner,M.E., Uzilov,A.V., Stein,L.D. et al. (2009) JBrowse: a next-generation genome browser. *Genome Res.*, 19, 1630.
30. Altschul,S.F., Gish,W., Miller,W. et al. (1990) Basic local alignment search tool. *J. Mol. Biol.*, 215, 403–410.
31. Untergasser,A., Cutcutache,I., Koressaar,T. et al. (2012) Primer3–new capabilities and interfaces. *Nucleic Acids Res.*, 40, e115.
32. Beier,S., Thiel,T., Munch,T. et al. (2017) MISA-web: a web server for microsatellite prediction. *Bioinformatics*, 33, 2583–2585.
33. Schuler,G.D. (1997) Sequence mapping by electronic PCR. *Genome Res.*, 7, 541–550.
34. Martins,W., de Sousa,D., Proite,K. et al. (2006) New softwares for automated microsatellite marker development. *Nucleic Acids Res.*, 34, e31.
35. Fernandez-Pozo,N., Menda,N., Edwards,J.D. et al. (2015) The Sol Genomics Network (SGN)–from genotype to phenotype to breeding. *Nucleic Acids Res.*, 43, D1036.
36. Moriya,Y., Itoh,M., Okuda,S. et al. (2007) KAAS: an automatic genome annotation and pathway reconstruction server. *Nucleic Acids Res.*, 35, 182–185.
37. Ogata,H., Goto,S., Sato,K. et al. (2000) KEGG: kyoto encyclopedia of genes and genomes. *Nucleic Acids Res.*, 28, 27–30.
38. Finn,R.D., Coggill,P., Eberhardt,R.Y. et al. (2016) The pfam protein families database: towards a more sustainable future. *Nucleic Acids Res.*, 44, D279.
39. Finn,R.D., Coggill,P., Eberhardt,R.Y. et al. (2016) The pfam protein families database: towards a more sustainable future. *Nucleic Acids Res.*, 44, D279.
40. Li,H. and Durbin,R. (2009) Fast and accurate short read alignment with Burrows–Wheeler transform. *Oxford Univ. Press*, 25, 1754–1760.
41. Li,H., Handsaker,B., Wysoker,A. et al. (2009) The sequence alignment-map format and SAMtools. *Bioinformatics*, 25, 2078–2079.