Rice SNP-seek database update: new SNPs, indels, and queries

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ABSTRACT

We describe updates to the Rice SNP-Seek Database since its first release. We ran a new SNP-calling pipeline followed by filtering that resulted in complete, base, filtered and core SNP datasets. Besides the Nipponbare reference genome, the pipeline was run on genome assemblies of IR 64, 93-11, DJ 123 and Kasalath. New genotype query and display features are added for reference assemblies, SNP datasets and indels. JBrowse now displays BAM, VCF and other annotation tracks, the additional genome assemblies and an embedded VISTA genome comparison viewer. Middleware is redesigned for improved performance by using a hybrid of HDF5 and RDMS for genotype storage. Query modules for genotypes, varieties and genes are improved to handle various constraints. An integrated list manager allows the user to pass query parameters for further analysis. The SNP Annotator adds traits, ontology terms, effects and interactions to markers in a list. Web-service calls were implemented to access most data. These features enable seamless querying of SNP-Seek across various biological entities, a step toward semi-automated gene-trait association discovery. URL: http://snp-seek.irri.org.

INTRODUCTION

Genomic data play increasingly important roles in plant breeding by helping to discover new gene-trait associations and to understand how nucleotide variations are translated into phenotypic diversity of plants. While several other databases curate rice nucleotide variants, e.g. dbSNP at NCBI (1), Gramene (2), RiceVarMap (3), IC4R (4), RM-Breeding (5), the SNP-Seek database features interactive real-time visualization of millions of SNPs in thousands of rice varieties, making SNP-Seek a unique tool for allele mining (6).

Since the first release of Rice SNP-Seek Database (7), we have undertaken considerable development to incorporate new analysis results, datasets, viewers and query interfaces for multiple reference genomes and assemblies. We have also included features requested by users that will be useful to the broader rice research community.

NEW SNP SETS

We envision SNP-Seek to host data from not only our projects, but also as a repository for rice variant data in the public domain. Consequently, we redesigned the software architecture to handle multiple datasets and reference genomes. Further enhancement has occurred allowing the display and analysis of other variant types like indels and genetic markers and tools for analysis.

SNP and variety sets

The middleware of the SNP-Seek application was redesigned to handle multiple datasets and data formats. This enables use of the same user interface to display various SNP datasets and formats. Additional and updated analyses of the 3k varieties from the 3k Rice Genome Project (8) resulted in five SNP sets: All (32M positions), All biallelic (29M), Base (18M), Filtered (4.8M) and Core (404k). The details of these are described at the Download page (http://snp-seek.irri.org/download.zul). The Filtered SNP
set is the default dataset. We also imported the HDRA (9) data into SNP-Seek. With various SNP datasets available in a single interface, we can now efficiently query combined genotypes from these data. The current options are to query either/or both of the 3k or HDRA varieties; if both are selected, the SNP positions may be the union or the intersection of the two datasets.

We also have access to phenotype data from the International Rice Genebank Collection database (Genetic Resource Information Management System, GRIMS) (10) linked to SNP-Seek. When a new genotyping dataset is added to our database and the varieties can be traced to genetic stocks or source accessions in GRIMS, phenotype data for the genetic stocks or legacy data for the source accessions are immediately available, adding value to the dataset.

SNPs from multiple *O. sativa* assemblies

In addition to Nipponbare (japonica), the SNP-calling pipeline (11) was run on four sequenced rice genomes representing indica and aus, two of the major rice subpopulations: IR 64 (indica) and DJ 123 (aus) (12), Kasalath (aus) (13) and 93-11 (indica) (14). To avoid including redundant SNP positions, a custom pipeline was run on a path alignment of the five genomes computed from the pair-wise alignment between the five reference genomes. New SNPs for regions unique to each of the other genomes were sequentially added, with the union of SNP calls loaded into SNP-Seek. We used VISTA (15) for the genome comparison. Results of the pairwise genome alignments are viewable in the VISTA browser accessible through the SNP-Seek menu.

Our comprehensive all-against-all pair-wise alignments showed that all five genomes are highly similar to one another. However, a significant fraction of each genome sequence was found to be unique, namely 8.59% of Nipponbare, 5.35% of 93-11, 4.11% of IR 64, 19.85% of Kasalath and 3.56% of DJ 123 (Supplementary Table S1). Reference genome-specific regions may be informative for the discovery of novel variants since variants in these unique genomic regions (which represent additional 12 Mb to 79 Mb, Supplementary Table S1) would not be detected in accessions aligned to reference genomes that are too distant. Over ~11 million additional SNPs and ~0.5 million indels were discovered from the additional reference genomes (Supplementary Table S2).

In a genotype query result, a position in the genotype table is based on the selected reference genome. The chromosome/contig and locus choices also depend on the selected reference. Selecting the ‘Show all reference alleles’ will display the alleles for all reference genomes. A gap in the allele means the position is not found, and possibly deleted for that genome (shown in Figure 1). The location of the queried region in the other references is reported in a message at the top of the results page.

**Indels**

In addition to SNPs, the variant calling pipeline (11) yielded short indel data. Unlike most variant databases where indels are presented as alleles, we optionally display them in the genotype matrix along with the SNPs, in a multiple sequence alignment-like format (shown in Figure 2). To accomplish this, the longest indel at each anchor point is determined for all 3024 varieties, call this length N. Then N columns are inserted to the right of the anchor point column. For insertion of length I less than N, the inserted nucleotides are filled in the first I columns, and gap(s) are padded from columns I + 1 to N. If the anchor point is at position P, the insertion region columns position are P,01, P,02, to P,N and the reference alleles are set to gap. For deletion D of length less than N, gap(s) are filled in the first D columns, and the reference is copied for columns D + 1 to N. If the anchor point is at position P, the deletion region column positions are P + 1, P + 2, to P + N and the reference alleles are set from the reference genome.

**Genomic features**

We also developed features to query genomic data. Although most of the genomic data were incorporated as provided by external data sources, we introduced some conventions to uniformly name and merge various gene models. First, we merged the Nipponbare MSUv7 (16) and RAP (17) gene models, plus in-house FGenesh++ (18) annotation into a single set of gene models and named them using our convention, of the form OsNippo{YY}g{NNNNNN}. Details about the merging and naming procedure are described in Supplementary Information Sections I and II, respectively. The advantage with this approach is that we were able to map and interconvert all Nipponbare genes between gene models based on location and overlaps instead of by using names. We also introduced a naming convention for gene loci for the other reference genomes, described in Supplementary Information Section II and summarized in Table I.

Where data are available, genes for any of the five reference assemblies can be queried using any of these constraints: functional annotation, gene name/symbols, accession number, Gene Ontology terms, traits or Trait Ontology terms, sequence or lists of SNP positions. The data sources are listed in Table 2 and were imported into SNP-Seek using the CHADO schema (19). Storing them with genotype and phenotype data allows complex queries to be performed for various analyses with display in SNP-Seek interface.

**NEW QUERY METHODS AND TOOLS**

Data alone will not have an impact unless it’s made available to those who need it the most and who have the best understanding of its biological significance. Large data sets require above average data handling and programming skills that may not be available to the average biologist. With this motivation, we implemented several query, data management and visualization features that use large variant and genomic data sets, but are relatively quick and intuitive for the general researcher.

**Ontology-driven queries**

Taking advantage of the CHADO data model, we use ontology terms to constrain gene queries, exploiting transitive closure. That is, selecting a term in the ontology will
Figure 1. Genotype query options (A) and results table (B) with multiple reference genomes alleles. The selected reference genome (Nipponbare) is displayed at the top header row just below the SNP positions. The alleles for the other genomes (Kasalath, DJ 123, IR 64, 93-11) are shown below in the table header. The corresponding positions in the other genomes are displayed in the message box.

Table 1. Gene loci names we used for rice reference genomes we are using

| Reference genome | Reference | Gene loci names |
|------------------|-----------|----------------|
| 93-11            | (14)      | Os9311[YY]g[NNNNNN], Os9311[XXXX]g[NNNNNN] |
| IR 64            | (12)      | OsIR64[XXXX]g[NNNNNN] |
| DJ 123           | (13)      | OsDJ123[XXXX]g[NNNNNN] |
| Kasalath         |           | OsKasal[YY]g[NNNNNN] |
| Nipponbare       | (16,17)   | OsNippo[YY]g[NNNNNN] |

Table 2. Data sources for genomics data

| Data             | Source     | URL                                      | Reference |
|------------------|------------|------------------------------------------|-----------|
| Gene model       | MSU v7     | http://rice.plantbiology.msu.edu        | (16)      |
| Gene model       | RAP        | http://rapdb.dna.affrc.go.jp             | (17)      |
| Gene names/symbols | Oryzabase | http://shigen.nig.ac.jp/rice/oryzabase  | (20)      |
| Gene ontology    | MSU v7     | http://rice.plantbiology.msu.edu        | (16)      |
| Trait genes      | OGRO       | http://qtaro.abr.affrc.go.jp/ogro/table | (21)      |
| Trait ontology-genes | Oryzabase | http://shigen.nig.ac.jp/rice/oryzabase  | (20)      |
| Plant ontology-genes | Oryzabase | http://shigen.nig.ac.jp/rice/oryzabase  | (20)      |
| QTL              | Q-TARO     | http://qtaro.abr.affrc.go.jp             | (26)      |
| Sequence         | MSU v7     | http://rice.plantbiology.msu.edu        | (16)      |
Figure 2. Genotype matrix with short indels. The table displays deletions (positions in blue) at anchor positions (region): 27698 (27699), 27791 (27792–27794), 27836 (27837–27841). Insertion regions (positions in green) are at 27722.01 27722.04 and 27797.01. For deletion regions, the reference is copied from the reference genome, while for insertions the reference genome is set to gaps.

return entities related to the term and its descendants as defined in the ontology. In the gene locus query, Gene Ontology terms can be used to constrain gene locus queries. Trait Ontology terms are also used to query genes where gene–trait association data are available from Oryzabase (20) or OGRO (21). In variety queries, phenotypes are mapped to Trait Ontology (http://browser.planteome.org/amigo/term/TO:0000387) and Crop Ontology (Rice) terms (http://www.cropontology.org/terms/CO320:ROOT/).

Allele frequency display
The allele frequency chart (shown in Figure 3) displays the frequency or count of major and minor alleles for the queried region or positions, for all varieties or by subpopulation. The chart can be useful for detecting haplotype blocks in the queried region, since adjacent SNPs in a block tend to have the same allele frequencies. If the major frequency of a subpopulation is 1, but less for the other groups, it means the subpopulation has no variant at those positions, which may be an important discriminator for the group. It can also show genotype instead of allele statistics.

New JBrowse tracks and reference genomes
This is a significant update to the JBrowse genome browser (22) since our first release. First, new sets of tracks for Nipponbare are organized by categories. We added tracks for trait associated genes and the BAM and VCF analysis results for each of the 3024 varieties stored in Amazon S3. All tracks for Nipponbare are listed in Table 3.

The second update is a separate JBrowse instance for each of the other four reference assemblies. The tracks in JBrowse are loaded with the same data as Nipponbare using gff format. Each instance has the option to display the sequence and gene models as provided by the source using original locus names and another track using our locus naming convention described in Supplement II. The alignments with the other four genomes can also be viewed as tracks. These instances are accessible through the main menu.

Large SNP queries
In the prior version of SNP-Seek, querying SNPs for large regions or using a list of many positions was prohibited by server timeout where the client application needed to wait for a query to finish before the user could proceed with other tasks. To accommodate large queries, we implemented an asynchronous query engine, utilizing Spring (https://spring.io) @Async annotation to manage parallel processes. This allowed us to extend the genotype query limit to a 5Mb region, 500kb SNPs or 1000 gene loci; however, the results are not displayed but are available for download. The user need not wait for the task to finish since a dynamic link is given to monitor the progress and download the results when ready.

Alternate sequence download
A common task is to reconstruct the alternate sequence for a list of regions and varieties, by substituting the SNPs, and integrating the insertions and deletions into the reference genome. This feature uses tabix (23) to query the VCF files from Amazon S3 and process the VCF using the FastaAlternateReferenceMaker tool in GATK (11). Results are downloadable as compressed Fasta files in gzip format (*.fasta.gz).
**Figure 3.** Allele frequency chart with major/minor allele/genotype frequency/count at each SNP position in the queried region for all or each subpopulation.

**Table 3.** JBrowse tracks for Nipponbare

| Category          | Track names (count)                                                                 | Reference          |
|-------------------|------------------------------------------------------------------------------------|--------------------|
| Gene model        | MSU7 RAP representative RAP predicted FGenesh++ Merged MSU7, RAP, FGenesh++ all trait genes | (16,17)            |
| Trait Genes       | 28 OGRO trait track OGRO all traits genes Oryzabase                                 | (20,21)            |
| QTL               | 28 QTARO QTL tracks QTARO all QTL                                                  | (26)               |
| BAM               | 3024 varieties                                                                      | https://aws.amazon.com/public-data-sets/3000-rice-genome/     |
| BAM Coverage      | 3024 varieties                                                                      | https://aws.amazon.com/public-data-sets/3000-rice-genome/     |
| Alignment         | Nipponbare versus 9311 Nipponbare versus IR64-21 Nipponbare versus DJ123 Nipponbare versus Kasalath | This project       |
| Variants          | SNPs v2 INDELS v2 SNPs v1                                                          | This project       |

**List management**

We want SNP-Seek to be used by researchers with minimal data interface/conversion issues. The List Manager is designed for this purpose. There are currently three types of lists implemented: variety, SNP and locus lists. The user can create a list, use it to constrain a query, generate a list from the query results and then download or submit it to other analysis tools or queries. The flow of data and queries available in SNP-Seek is illustrated in Figure 4. Arrows show the possible data from an initial set of information. Along with the available set operations, the system can be used for gene-trait association discovery. We extended the SNP List functionality to perform SNP-Marker annotation and genotype matching:

**SNP/Marker annotator**

The user can create a list of SNP positions and this feature annotates the markers with evidence collected from various other databases and analyses. This list may be significant markers from gene expression or GWAS studies. The annotations can include gene models (RAP (17), MSUv7 (16) or FGenesh++ (18)) or promoter regions (FGenesh++, PlantPromDB (24)) if SNPs are located within these loci. The effects of SNP variants were also added using results from SNPEff (25). For SNPs within gene models, additional evidence about the gene are included using Gene Ontology terms, Plant and Trait Ontology terms and gene names collected from Oryzabase (20), trait genes from OGRO (21), and QTL from Q-TARO (26), interacting genes from RiceNet v2 (27) and rice proteins from PRIN (28). The list of annotations and references are in Supplementary Table S3.

**Genotype match**

A common use case is to find the most related genotypes among the 3k/HDRA varieties when given a particular genotype. We created a query where the constraint is a list of SNP positions with allele values, and the result is a genotype table where the varieties are sorted based on the number of matching alleles between the query and each variety in the selected dataset. The genotype table can also display values for a selected phenotype allowing quick evaluation of the effect of the queried genotype on the phenotype.

**SOFTWARE AND DATABASE UPDATES**

To provide the data and query requirements described in the previous sections, several modifications were made in the underlying software architecture. Our major objectives
Figure 4. Query capabilities of SNP-Seek. The blocks at the left (rounded) are possible query constraints to the query modules (rectangles). The query results may be stored as a list (parallelograms), and used as constraints in further queries. Lists may also be created by the user as initial constraints. The marker annotator accepts a list of SNP positions, which may be the result from experiments or GWAS studies, to generate constraints for further queries or loop back to the initial constraints, increasing the confidence of the association.

for these updates were to improve query performance, to increase query capacity, to easily integrate new datasets and to serve data to other software systems. Query speed was improved by using hybrid storage wherein genotype matrices are located on the web server as HDF5 (https://www.hdfgroup.org/HDF5) formatted files while other data and the indices for HDF5 files are located in the RDMS. The middleware was re-designed into Service and Data Access layers using Java interfaces, further allowing multiple genotypic data sets to be accommodated. The details of these updates are described in Supplementary Information III.

Web services
We defined a set of RESTful web-service calls for internal use and shared to collaborators through our development site (http://snp-seek.irri.org/dev). The calls are focused on Germplasm, Phenotypes and Genotypes. Another set of calls were implemented for SNP-Seek for compliance to the Breeding API (BrAPI, http://docs.brapi.apiary.io) for Germplasm, Phenotypes, Maps, Studies and Genotypes. The web services documentation can be accessed from the Help menu. Most calls are open but some require login and password. Interested collaborators may contact the authors to use the protected calls.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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