Shanghai RAPESEED Database: a resource for functional genomics studies of seed development and fatty acid metabolism of *Brassica*

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**ABSTRACT**

The Shanghai RAPESEED Database (RAPESEED, http://rapeseed.plantsignal.cn/) was created to provide the solid platform for functional genomics studies of oilseed crops with the emphasis on seed development and fatty acid metabolism. The RAPESEED includes the resource of 8462 unique ESTs, of which 3526 clones are with full length cDNA; the expression profiles of 8095 genes and the Serial Analysis of Gene Expression (SAGE, 23 895 unique tags) and tag-to-gene data during seed development. In addition, a total of ~14 700 M3 mutant populations were generated by ethylmethanesulfonate (EMS) mutagenesis and related seed quality information was determined using the Foss NIR System. Further, the TILLING (Targeting Induced Local Lesions IN Genomes) platform was established based on the generated EMS mutant population. The relevant information was collected in RAPESEED database, which can be searched through keywords, nucleotide or protein sequences, or seed quality parameters, and downloaded.

**INTRODUCTION**

*Brassica* species, including *Brassica napus*, *B. oleracea* and *B. rapa*, are important vegetable and oilseed crops. Seed of *B. napus* contains abundant fatty acids such as oleic acid and linoleic acid, and serves as one of the main sources of plant oil for human daily life. As the second oilseed crop in the world, compared to soybean, numerous studies have been performed on *B. napus* with the focus on yield increase and quality improvement through genetic breeding (1–3), and development of genetics and molecular biology tools has significantly contributed to the relevant studies. However, the deficiency of genome sequences and the detailed genomic information counteract the functional genomics studies of rapeseed, especially the genetics approach through T-DNA tagged mutant, of which the gene sequences and expression information are required (4–6). Thus, the increased information of gene sequences and expression profiles will be beneficial for rapeseed functional genomics studies and further bioengineering.

Currently, several databases for *Brassica* species are available. Handa reported a database functions on *B. napus* mitochondrial genome and the comparative analysis to *Arabidopsis* (7). The EST and fatty acid metabolism sources of *B. napus* are available in KEGG database (8). Love et al. described a database (http://hornbill.espp.latrobe.edu.cn) incorporating the *Brassica* EST, Gene Ontology (GO) annotation and information of simple sequence repeat molecular markers, of which the EST, microarray (the expression profiles of 7000 uni-genes in vegetative tissues root and leaf), and MarkerQTL information were updated (9–11), providing a useful tool for *Brassica* research. However, there is still no resource providing the gene expression information and the SAGE data during the reproductive development of *B. napus* yet. In addition, there is also no database providing biosources including full-length cDNA or mutants.

Based on constructing cDNA libraries using seed materials at various developmental stages, large-scale sequencing, generation of glass-based cDNA microarray and hybridization, we are able to obtain the dataset of *B. napus* ESTs and the relevant expression profiles during seed development, which are incorporated into Shanghai RAPESEED (RAPESEED) Database, providing a solid platform for functional genomics studies of oilseed crops. In addition, the Serial Analysis of Gene Expression (SAGE) during seed development and mutant population by ethylmethanesulfonate (EMS) mutagenesis will further facilitate the relevant studies. These will significantly enrich the relevant resources and improve the studies of the area.

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SYSTEM ARCHITECTURE AND IMPLEMENTATION

RAPESEED was constructed on the platform of Sun Solaris 9 operating system and TomCat5.5 Web server. The database was implemented using a database management system MySQL5.0.20 (12). RAPESEED has been set up on a World Wide Web server allowing internet access with a web client.

SOURCES IN RAPESEED

The purpose of the RAPESEED is to provide helpful informations of EST, gene expression profiles and bioresources (full-length cDNA, TILLING population) to researchers and to promote the functional genomics studies and quality breeding of *Brassica* crops (Table 1).

ESTs, the annotation and Gene Ontology classification

RAPESEED contains 8462 unique ESTs of *B. napus*, of which 6892 were functionally annotated by sequence similarity to entries within GenBank (using BLAST with a cut-off value of *E*-value <10^-5. 13). GO classification was performed via mapping to UniProt to get the GO annotation. In addition, 3526 clones were full-length cDNA among the 8462 unique ESTs.

Gene expression profiles during seed development

Gene expression data for 8095 of 8462 genes during *B. napus* seed development are studied through cDNA microarray hybridization and available in RAPESEED, which include the expression profiles of relevant genes at 7, 9, 12, 17, 19, 21, 25, 31 day after pollination (DAP), normalized to 3 DAP (1.0).

Serial Analysis of Gene Expression (SAGE) data

A total of 23 895 unique tags from *B. napus* immature seeds (5 and 9 DAP) were obtained after sequencing and deposited into RAPESEED. Based on the virtual tags derived from GenBank and full-length cDNA from our cDNA library, the ‘tag-to-gene mapping database’ for *B. napus* and *Brassica* including *B. oleracea* and *B. rapa* were constructed. The whole SAGE dataset can be downloaded from RAPESEED.

EMS mutant population and seed quality measurement

The mutant population mutagenesized with ethyl-methanesulfonate (EMS) was generated using *B. napus* (huyou-15). A population of ~14 700 M3 individuals was generated and relevant seed qualities were measured and analyzed using the Foss NIR System. The seed quality parameters including the content of glucosinolates, protein, total fat, erucic acid, oleic acid, linoleic acid and eicosatetraenoic acid in seeds (Table 2), providing a useful source for rapeseed quality studies.

Based on the EMS mutant population, a TILLING (Targeting Induced Local Lesions IN Genomes) platform was established.

DATABASE QUERY AND USER INTERFACE

BLAST and SEARCH

The genes of interest or EST information can be identified through the BLAST program (13). Three different BLAST programs including BLASTn, tBLASTn and tBLASTx were provided, of which the BLAST parameters can be freely chosen by users including *E*-value or number of alignments. The typical result page contains the collected information of the available genes in RAPESEED, including the clone ID, GenBank Accession, expression profiles during seed development, gene annotation and related *E*-value and score, full-length information and DNA sequences (Figure 1).

In addition, RAPESEED provides a key word SEARCH program to facilitate the gene identification. Clone ID, GenBank Accession or key word can be selected for search. The complex SEARCH of two search items combination is also supported through and/or option.

Resource download and order from RAPESEED

All the dataset including those for whole EST sequences and corresponding annotations against GenBank, SAGE data and related tag-to-gene information, raw data of the microarray hybridization and seed qualities of EMS population (Table 2) can be downloaded.

### Table 1. Available resource of RAPESEED

| Resource | Description |
|----------|-------------|
| Unique EST | 8462; relevant expression profiles during seed development |
| Full-length cDNA | 3526 |
| Unique SAGE tag | 23 895 |
| Tag-to-gene | 502 tags to *B. napus* |
| M3 EMS mutant individuals | ~14 700 |

### Table 2. The summary of the seed quality measurement of EMS mutagenesized population through NIR analysis

| Parameter | Glucosinolates | Protein | Lipid content | Erucic acid | Oleic acid | Linoleic acid | Eicosatetraenoic acid |
|-----------|----------------|---------|---------------|-------------|------------|--------------|----------------------|
| WT (Average) | 24.02 ± 2.17 | 24.74 ± 0.25 | 39.39 ± 0.42 | 7.03 ± 1.23 | 56.28 ± 1.66 | 19.62 ± 0.39 | 1.27 ± 0.77 |
| WT (Maximum) | 31.37 | 25.47 | 42.43 | 9.42 | 59.91 | 21.14 | 2.97 |
| WT (Minimum) | 18.99 | 21.72 | 38.26 | 3.71 | 52.86 | 18.60 | -0.49 |
| Criteria | >40 | <10 | >30 | <20 | >44 | >20 | <1 | >60 | >25 | <10 | >5 |
| Numbers | 1101 138 56 19 38 3553 125 1254 432 117 365 |

Parameters of seed qualities are indicated as the content (%) in seeds. Data are presented as mean ± SD (n= 77 for wild type).
Furthermore, all listed sources of RAPESEED were available, especially the full-length cDNA clones and EMS lines with altered seed qualities, which can be ordered under the accordance of ‘Biological Material Transfer Agreement’.

AVAILABILITY

RAPESEED can be freely accessed at http://rapeseed.plantsignal.cn via the World Wide Web. We have developed a mature data management system, and all newly released information will be announced on the website. Besides ordering the interested clones or lines, users can contact us for any suggestions or questions through internet or send the comments to rapeseed@sibs.ac.cn.

FUTURE DEVELOPMENTS

RAPESEED is a database developed for B. napus functional genomics studies, and we will do our best to continuously optimize the system and supply new records, to make it an integrated resource (Figure 2) and provide a base for Brassica research and breeding. Besides the gene identification and expression profiles, we will focus on the following in the future.

TILLING population

TILLING is a useful reverse genetic technique that is a powerful, non-transgenic and unbiased targeted mutagenesis method. It can help to identify single point mutations in genes of interest (14), and is suitable for mutation site identification of the EMS lines, which is appropriate and helpful for further crop breeding. Based on the developed TILLING population of M3 EMS lines of B. napus, an amphidiploid (AACC) consisting of the genomes of B. rapa (AA) and B. oleracea (CC), we will further develop a TILLING population (>4000 M2) of diploid (B. rapa) for Brassica breeding studies. In addition, a new B. napus EMS population (M2) has already been developed this year.

We will develop pooled DNAs of ~3000 M2 plants every year and relevant seed qualities will be measured. The screening data and seed qualities information of them will be deposited into RAPESEED annually.

Figure 1. Typical result page of query. The information of a gene including the clone ID, GenBank Accession, expression profiles during seed development, gene annotation and related E-value and score, full-length information and EST sequences will be shown after search.
Comparative gene expression profiles of *B. napus* to *B. oleracea* and *B. rapa*

*Brassica napus* (~1200 mb) consists of that from *B. rapa* (700 mb) and *B. oleracea* (500 mb), and differences between them are obvious in many aspect such as plant morphology, seed oil content and self-productivity. This suggested that the gene expression profiles were changed dramatically during breeding processes, and investigations on these differences will surely provide the information on the regulatory network of the gene expression and the relevant cross-talk. The related information will be available soon.

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Conflict of interest statement. None declared.

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