TNAU PDB - Tamil Nadu Agricultural University Proteome DataBase - Black Gram Proteome

S. Swathi¹, N. Senthil¹*, V. Vinod Kumar¹, S. Sathish², N. Jagadeesh Selvam¹ and M. Raveendran¹

¹Department of Plant Molecular Biology & Bioinformatics, Tamil Nadu Agricultural University, Coimbatore, Tamil Nadu, India.
²Department of Seed Science and Technology, Seed Centre, Tamil Nadu Agricultural University, Coimbatore, Tamil Nadu, India.

Authors’ contributions

This work was carried out by the first author SS in collaboration between all authors. Guided by the expertise advice of the second author NS and last author MR, the corresponding author NS designed the study, first author SS and third author VVK designed the web page and wrote the first draft of the manuscript. Fourth author SS and fifth author NJS did the proteomic analysis. All the authors managed the analyses of the study. However, all authors read and approved the final manuscript.

ABSTRACT

Tamil Nadu Agricultural University Proteome DataBase (TNAU PDB) – Black gram proteome is an open accessible database that focuses on proteome of Black gram (Vigna mungo L.). Currently, the database contains reference maps of Two-Dimensional Polyacrylamide Gel Electrophoresis (2D-PAGE) of proteins obtained from artificially aged black gram seeds of variety TNAU blackgram CO 6, which is compared to that of fresh seeds. The database provides information about experimentally identified properties, such as molecular weight, pl value, of the differentially expressed protein due to accelerated ageing and protein sequences obtained using MALDI-TOF mass spectrometry. This database runs on WAMP server with HTML as the front end and MySQL as the backend using PHP as interface and it is hosted in TNAU genomics domain. The basic intention of this database is to provide the detailed information about proteome of black gram. This will help us to understand adaptive and general protective mechanism related to seed aging and effect of ageing on germination.

*Corresponding author: Email: senthil_natesan@tnau.ac.in;
Keywords: Accelerated ageing; black gram; database; proteomics; proteome database; TNAU PDB.

1. INTRODUCTION

Black gram (Vigna mungo L) is a protein rich food, containing about 26 per cent protein, which is almost three times that of cereals. It ranks fourth among the major pulses cultivated in India. Black gram supplies a major share of protein requirement of vegetarian population of the country. It is consumed in the form of split pulse as well as whole pulse, which is an essential supplement of cereal based diet. In India, black gram occupies 12.7 per cent of total area under pulses and contribute 8.4 per cent of total pulses production. However, area and production of black gram has declined from 3.01 million ha and 1.30 million tonnes in 2000-01 to 2.97 million ha and 1.23 million tonnes, respectively in 2009-10 [1]. In the current scenario of flourishing “omics” segment that provides deep insight into the mechanism of crop physiology and assist in further crop improvement, proteomics of blackgram is not yet completely exploited as that of cereals like paddy [2]. Since analysis of the proteome provides a direct link of genome sequence with biological activity, data on proteomes are slowly mounting [3]. Currently, in our lab, proteome study on blackgram crop were set in motion. Since seed is the prime source of crop production [4] and storage is inevitable for carryover seeds [5], knowledge on molecular basis of seed ageing is warranted for effective maintenance of seed quality during storage. Accelerated ageing has been widely used to study the pattern of seed deterioration in various crops [6-9]. As a initial phase of proteome study in blackgram, proteome of seed ageing was analysed through comparative study of accelerated aged and fresh seeds and differentially expressed proteins due to ageing were identified. This growing data on proteome of blackgram and unavailability of no separate database concentrating on proteomes of black gram had encouraged us to develope a database to integrate the essential data derived in our lab which would help the researchers to understand the biological process and thereby assists in further research. The main objective of this database is to promote quantitative and qualitative proteome research in black gram that helps in understanding the mechanism of various biotic and abiotic stress related proteins which in turn helps in annotating their function. Currently detailed proteome information about the blackgram variety TNAU blackgram CO 6 seeds are made available in the database.

2. METHODOLOGIES

2.1 Database Content and Source

The black gram proteome database consists of proteome information about pods, leaves, stem and root. Currently, only proteome data of pods alone is made available. Possible changes in proteins due to artificial ageing on dry seeds of blackgram variety TNAU blackgram CO 6 were analyzed and the quantitative and qualitative proteome changes were recorded. Artificial ageing was done for 6 days by packing seeds in perforated butter paper bags and placed in an ageing jar containing 100 ml of distilled water to maintain 98 ± 2% relative humidity and incubated at a temperature of 40 ± 1°C [10]. In the first dimension, 150µg of protein was loaded on a 17 cm IPG strip with a linear gradient of pH 4-7. In the second dimension, 12% SDS-PAGE gels were used with molecular weight standards. Proteins were visualized by silver staining as suggested by Blum et al. [11]. Their properties such as molecular weight, pI values and their expression were calculated using image
analyzer software. Significant spots were cut, digested in trypsin and analysed in MALDI-TOF for sequencing [12]. The generated data were uploaded in the database.

2.2 Languages and Software’s Used

The front end of the web application is developed on HTML 5.0 (Hyper Text Markup Language) and the validations are done using javascript. The server side scripting was done on PHP 5.3.0 (Hypertext Pre Processor) and the application was connected to the database using MySQL 5.0.8. Web application was created by using wamp server 2.2 and each spot in the gel image was linked to the corresponding protein information with the help of Macromedia dreamweaver.

2.3 Database Architecture

The application is built on a three tier architecture model consisting of presentation layer, logic layer and the database layer. The presentation layer is the front end of the application created using HTML with which the user interacts. The middle layer is the application server or logical layer created using PHP, which serves the application with data and accepts the requests from the user. The data layer contains the relational database which contains the data to be fetched by the application.

3. RESULTS AND DISCUSSION

3.1 Flow of Data

Proteome database of blackgram has been added as a sub link under TNAU genomics web page which has home page under the link http://www.tnaugenomics.com. From the home page the blackgram database can be reached through the drop down link named TNAU PDB under the tab databases. The flow of data and the procedure to access the database were meticulously depicted in Fig. 1. Similar flow of data can also be seen in a database developed for maize [13].
3.2 Database Schema and its Features

The data is classified based on the tissue such as pods, leaf, stem, root, etc. from which it is identified. Proteomes obtained using the 2D PAGE and MALDI-TOF methods are displayed using image analyzer software. The description of each spot is stored in the database and they are mapped to the corresponding spots. Based upon the request from the user, the application queries the database and fetches the spot information.

Currently, database contains the proteome information obtained from pods of black gram. The reference 2D-PAGE gel shows the position of each identified protein in that crop. The entire list of protein can also be obtained by selecting the crop name displayed inside the table. By selecting the spot in the gel image the entire information about that protein can be obtained. The results are displayed in such a manner that they are also compared with their control (Fig. 2). All the reference maps are also displayed under 2D gel section for a quick reference. The experiment protocols are listed under the protocol section. Some of the major proteomics tools like Mascot, Compute pI/Mw tool in ExPASy and ExPASy Proteomics tools are displayed under proteome tools section. Similar advanced proteome databases were developed for Arabidopsis and maize by Cornell university [14] and Dyna Prot 2D for dynamic online access to proteomes and two-dimensional electrophoresis gels [15].
3.3 Utility

The primary users of this database will be plant breeders, seed technologist, students and other researchers concentrating on blackgram improvement. It shows the proteome changes that could take place at the dry state of aged seeds. Determination of the expression patterns in response to stress, and an understanding of their functions in stress adaptation will provide the researchers with the basis for effective genetic engineering strategies for improving the tolerance of crops to various stresses.

3.4 Future Developments

The database content will be updated routinely. In the near future proteome data for whole plant parts such as root nodules, stem, leaves and expression of protein under different environmental stress conditions will be made available. With the availability of ample proteome data, architecture of the web page might be modified to suit to the excess flow of information.

4. CONCLUSION

Tamil Nadu Agricultural University Proteome DataBase (TNAU PDB) – Black gram proteome contains 2D reference map of black gram seeds developed through comparative proteome analysis of artificially-aged and fresh seeds and shows the identified differentially expressed proteins due to seed ageing. The database content will be updated routinely. It will be useful for plant breeders, seed scientist and other researchers working in black gram.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCE

1. ASSOCHAM. Study paper on “Emerging pulses scenerio in 2015 - productivity, awareness and affordability to drive pulses economy in India”. The Associated Chambers of Commerce and Industry of India, New Delhi, India. 2012;1–18.
2. Setsuko Komatsu, Keiichi Kojima, Kouji Suzuki, Kazuo Ozaki, Kenichi Higo. Rice Proteome Database based on two-dimensional polyacrylamide gel electrophoresis: its status in 2003. Nucleic Acids Research. 2004;32(Database issue):D388 - D392.
3. Montpetit B. Proteomics in the post-genomics era: weighing in the data. BioTeach Journal. 2003;1:9–12.
4. Ajeigbe HA, Abdoulaye T, Chikoye D. Legume and cereal seed production for improved crop yields in Nigeria. Proceedings of the Training Workshop on Production of Legume and Cereal Seeds, 2008. Accessed 4 June 2013. Available: http://www.iita.org/c/document_library/get_file?uuid=e2d9972d-41f1-4e25-a38a-fb0309716105&groupId=25357.
5. Justice OL, Bass LN. Principles and practices of seed storage. 1st ed. Science and Education Administration's Federal Research; 1978.
6. Jatoi SA, Afzal M, Nasim S, Anwar R. Seed deterioration study in pea, using accelerated ageing techniques. Pakistan Journal of Biological Sciences. 2001;4(12):1490–1494.
7. Scialabba A, Bellani LM, Dell’Aquila A. Effects of ageing on peroxidase activity and localization in radish (Raphanus sativus L.) seeds. Eur. J. Histochem. 2002;46:351-358.

8. Rajjou L, Lovigny Y, Groot SPC, Belghazi M, Job C, Job D. Proteome-wide characterization of seed aging in arabidopsis: a comparison between artificial and natural aging protocols. Plant Physiology. 2008;148:620–641.

9. Xin, X, Lina XH, Zhoub YC, Chena XL, Liua X, Lua XX. Proteome analysis of maize seeds: the effect of artificial ageing. Physiologia Plantarum. 2011;143:126-138.

10. Delouche JC, Baskin CC. Accelerated ageing techniques for predicting the relative storability of seed lots. Seed Sci. & Technol. 1973;1:427-452.

11. Blum H, Beier H, Gross HJ. Improved silver staining of plant proteins, RNA and DNA in polyacrylamide gels. Electrophoresis. 1987;8:93-99.

12. Shevchenko A, Wilm M, Vorm O, Mann M. Mass spectrometric sequencing of proteins silver-stained polyacrylamide gels. Anal. Chem. 1996;68:850-858.

13. Murukarthick J, Senthil N, Raveendran M, Prabhakaran P, Sreedevi G, Sumanth Kumar M, Shobhana VG, Dhanya S, Khushboo R, Arumugasamy S, Ravikesavan R, Jagadish SVK. Biogen Base – An Interactive Maize Database for Phenomics Platform. International Journal of Computer Application. 2011;2:56–61.

14. Qi Sun, Zybaïlov B, Majeran W, Frisco G, Oliñares PDB, Wijk KJV. PPDB, the Plant Proteomics Database at Cornell. Nucleic Acids Research Database issue. 2009;37:D969–D974.

15. Drews O, Gorg A. Dyna Prot 2D: an advanced proteomic database for dynamic online access to proteomes and two-dimensional electrophoresis gels. Nucleic Acids Research Database issue. 2005;33:D583–D587.

© 2013 Swathi et al.; This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Peer-review history:
The peer review history for this paper can be accessed here:
http://www.sciencedomain.org/review-history.php?id=236&id=2&aid=1883