Interview with Michael Bevan: The Arabidopsis genome is sequenced, what next?

Abstract

Professor Michael Bevan is at the John Innes Centre, Norwich, UK, where he is Head of the Cell and Developmental Biology Department. He was the co-ordinator of the European Union Arabidopsis Genome Sequencing Consortium, which contributed to the sequence and analysis of the complete Arabidopsis genome that was published late last year. He is the co-ordinator of the EC funded EXOTIC program and is also a member of GARNet and UK CropNet. (For more background detail on these projects, please see our Featured Organism: Arabidopsis thaliana piece, also in this issue of CFG, or visit the websites listed below).

CFG: Your group at the John Innes Centre works on the development of tools, such as GUS reporter systems and in vivo footprinting techniques: Do you see ways that these can be adapted, or applied in a global sense due to the knowledge of the sequence?

MB: There are several potential applications of these methods to systematic analysis of the Arabidopsis genome. One is the use of the GUS reporter gene in gene trap constructs. For example, in Rob Martienssen’s and my lab we use a modified Ds transposon that makes in-frame fusions with GUS upon insertion in a gene (Sundaresan et al. 1995). Large populations of these gene traps are being generated to reveal the spatial and temporal patterns of gene expression. This will complement array analyses, which generally do not reveal the cellular specificity of gene expression. I don’t see in vivo footprinting being applied on a large scale, as it is very difficult. But chromatin immunoprecipitation, especially allied to array analysis, can reveal binding sites of transcription factors. Our ability to interpret the sequence information of promoters is very under-developed. But Arabidopsis, with essentially contiguous sequence, has nearly all of the potential regulatory sequences available for analysis. New bioinformatics techniques can be used to associate patterns of gene expression with promoter sequences, and establish likelihoods of particular sequences being associated with particular expression patterns. Here the availability of array data and gene trap data permits these approaches to be explored.

CFG: You have an interest in characterising transcription factors and their binding sites, the sequence offers a further tool for the study of upstream regions of genes, do you plan to capitalise on this? Are there any projects moving in this direction?

MB: There are opportunities to develop some prototypes for searching for promoter motifs in the “Exploiting Genomics” initiative from the BBSRC, but I don’t know of any work in this field in Arabidopsis apart from the work of Maleck et al. (2000) who defined the WRKY element as a conserved functional feature in SA regulated genes, using cluster analysis of microarray data. This is clearly a field ripe for further development.

CFG: The sequence also offers an unparalleled opportunity for comparative studies, which form part of your UK CropNet project, what are your plans in this area? How far do you think Arabidopsis data can be carried? [evidence for some synteny with tomato and soybean has been reported (Ku et al., 2000; Grant et al., 2000)]

MB: Comparative studies are very important at the John Innes Centre. The general goal is to identify crop plant orthologs of Arabidopsis genes, especially among the cereals and the Brassicas. Our strategy is
to use rice to model potential orthology within cereals, using conserved linkage as a clue to possible ancestry, and use Arabidopsis as a model for Brassicas. Arabidopsis will not do for cereals because of the evolutionary distance, and the problem of ploidy levels is very complex, with rape having three copies of most Arabidopsis genes. It's very difficult to generalise if you don’t have data on gene function in both compared species. With respect to transcriptional control, comparisons of promoter sequences between closely related species has been shown to reveal conserved sequences that may be regulatory. Comparisons between Arabidopsis ecotypes may reveal this, but one may have to go further, to Capsella (9–12 Myr distance) or Brassicas (15–20 Myr distance) to search for the appropriate level of sequence divergence needed to reveal conserved sequences of functional significance. Arabidopsis gene order is quite well conserved in Brassicas, and much less so in tomato.

CFG: In your role as coordinator of EXOTIC, you are studying the expression pattern of around 5000 Arabidopsis genes. Will you be using microarrays? What strategies are you going to be using (e.g. will you be exposing the plants to a range of conditions)? How will this data be made available to the public?

MB: In EXOTIC we will use gene trap expression data to assess gene function and expression in a wide range of conditions. At the JIC we are interested in a wide range of expression patterns, from those seen during specification of meristems, to stress responses. Ultimately, data generated here will probably be deposited in public databases, but that is a decision for the originators of the data.

CFG: As a service provider for GARNet you are involved in a project to produce a large collection of transposon insertion lines and also to sequence the sites of insertion in these lines. Can you describe your approach? How will the data be distributed?

MB: In GARNet we are making a population of 30,000 gene trap lines and will sequence the insertion sites of the transposons in all these lines. This sequence will be sent to Genbank and also integrated with the MIPS genome database. We are also making a new database called ATIS, which integrates genome sequence with insertion sites and expression patterns.

CFG: Is there any follow up planned on this project in terms of phenotypic analyses, or will this be left to the wider Arabidopsis community? Are there any other experiments you would like to do with these lines?

MB: The lines made in GARNet will be bulked and sent to the stock centres for distribution. We have no intention to screen systematically for phenotypes, but if funding permits, we would like to screen for GUS expression patterns, to extend the scope of work in the EXOTIC EC project.

CFG: The Arabidopsis genome has a bigger proportion of genes in large gene families than C. elegans, or Drosophila, which could imply increased redundancy. Do you expect that most Arabidopsis genes will have a phenotype, or that it will be more similar to the case in yeast, in which many genes have no discernible phenotype under a range of conditions?

MB: Previously we have been involved in a project to define the functions of MYB transcription factors in Arabidopsis, and identified disruptions in 32 genes (Meissner et al., 1999). While phenotype screening is not yet completed, it seems that the closer you look, and the more you think about what to look for, coupled to good luck, will generally give you a phenotype. Well-established cases of highly overlapping gene functions in Arabidopsis are few, with the Shatterproof genes being the best example (Liljegren et al., 2000).

CFG: As coordinator of the European Union Arabidopsis Genome Sequencing Consortium you will have seen at first hand the excitement and motivation the genome sequence has given to the Arabidopsis community, would you like to comment on that and what has been achieved?

MB: I think the Arabidopsis, and indeed the wider plant community, were pleased to have the sequence finished relatively quickly, with an unprecedented completeness and accuracy (The Arabidopsis Genome Initiative, 2000). The sequence has been thoroughly analysed by MIPS and TIGR and made available to the public in comprehensive databases. The bioinformaticians, sequencers and mappers did a fantastically thorough job against tight deadlines, and it was both a privilege and
enormously exciting to work with these groups. Although there was a parallel industrial effort to sequence Arabidopsis, we were able to publish jointly with them, and their data is a potent source of polymorphisms for mapping. Like most scientific communities studying model organisms, the Arabidopsis genome project was generally characterised by openness and mutual support, and all but a few scientists respected the unspoken right of the sequencers to publish initial analyses of the whole genome sequence they had produced. The genome sequence can be used in a wide range of applications, and its final value won’t be realized for many years. One particularly exciting development is the power of comparison between genomes – Arabidopsis and Synechocystis have many genes in common, and so do humans and Arabidopsis.

CFG: You were also involved in organising the recent Arabidopsis Functional Genomics workshop at Cold Spring Harbor. What progress was made there towards defining the goals for Arabidopsis Functional Genomics studies? Do you think there will be similar international initiatives and consortia as for the sequencing? Was there any discussion of data standards or ontology issues?

MB: The sequencing work was relatively closely organised, and I don’t think at this stage functional genomics activities in Arabidopsis need to be as highly regulated. Gene disruptions are made by random insertions, so the first goal is to just get enough lines (primarily T-DNA) to yield a high probability of finding a disruption, then making these lines available through screening and ultimately sequencing the insertion sites, and integrating the data into public databases. This is all ongoing. Coordination is required in the integration of array analysis data from different labs, and this is also in hand. Beefing up the stock centres to handle the vast number of lines is a very high priority, and is needed if the gene disruption lines are to be readily available to users.

Various approaches are being taken to derive general descriptions of gene function. Presently the Arabidopsis functional catalogue consists of yeast functional categories, which have proved to be remarkably adaptable. Nevertheless, ontologies based on Drosophila gene descriptions are being developed, and there are also more novel approaches being considered. This work will continue to require a good deal of input from the community to devise standards and capture the varieties of descriptions of gene function.

Related Websites

Arabidopsis Transposon Insertion Service (ATIS)  
http://www.jic.bbsrc.ac.uk/STAFF/michael-bevan/ATIS/

Cold Spring Harbor Arabidopsis Genetrap Database  
http://spot.cshl.org/genetrap_database/mainframe.html

Genomic Arabidopsis Resource Network (GARNet)  
http://garnet.arabidopsis.org.uk

MIPS Arabidopsis thaliana database (MATDB)  
http://websvr.mips.biochem.mpg.de/proj/thal/

The TIGR Arabidopsis thaliana Database  
http://www.tigr.org/tdb/at/at.html

UK CropNet (UK Crop Plant Bioinformatics Network)  
http://ukcrop.net

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