Use of translational genomics to identify genes important for legume seed development

Mélanie Noguero, Christine Le Signor, Vanessa Vernoud, Isabelle d’Erfurth, Jérôme Verdier, Gregoire Aubert, Jerome Gouzy, Jean-Marie Prosperi, Thierry Huguet, Judith Burstin, et al.

To cite this version:
Mélanie Noguero, Christine Le Signor, Vanessa Vernoud, Isabelle d’Erfurth, Jérôme Verdier, et al.. Use of translational genomics to identify genes important for legume seed development. 6. International Food Legumes Research Conference (IFLRC VI), University of Saskatchewan. CAN., Jul 2014, Saskatoon, Canada. 225 p. hal-01268897

HAL Id: hal-01268897
https://hal.archives-ouvertes.fr/hal-01268897
Submitted on 2 Jun 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
PROGRAM & ABSTRACT BOOK

TCU Place
Saskatoon, Saskatchewan, Canada
July 7-11, 2014
Keynote Speaker: Seeds and Nutrition (ICLGG)

Richard D. Thompson. After a Ph.D. in Molecular Biology U. Edinburgh (1979) and periods at the PBI Cambridge UK, and the MPI for Plant Breeding Research, Köln, currently researcher in the INRA Agroecology Unit, Dijon, (2002-present), Grain Legumes Unit Director 2002-2010, focussing on the genetic control of seed development and composition in legumes, using the *Medicago truncatula* model, and more recently, Pea (*Pisum sativum*). By applying proteomics and transcriptomics approaches, gene expression profiles for isolated tissues of the seed have been obtained and clusters of co-regulated genes acting during discrete stages in seed filling identified. To narrow down candidate genes for the control of seed composition and yield, we have selected genes that co-locate with QTLs for these traits and are analysed the phenotypes of corresponding TILLING and Tnt1 insertion mutants. Also coordinating the EU project LEGATO (Legumes for the Agriculture of Tomorrow (2014-2017)), aimed at increasing grain legume cultivation in Europe.

Use of translational genomics to identify genes important for legume seed development

Noguero, M.1, Le Signor, C.1, Vernoud, V.1, D’Erfurth, I.2, Verdier, J.3, Aubert, G.4, Gouzy, J.5, Prosperi, J-M.6, Huguet, T.7, Burstin, J.8, Gallardo, K.8, and Thompson, R.D.8*. 1INRA, UMR Agroécologie, DIJON, FRANCE; 2INRA, UMR1347 Agroécologie, DIJON, FRANCE; 3Shanghai Center for Plant Stress Biology, Shanghai Institutes of Biological Sciences, Chinese Academy of Sciences, Shanghai, PR China; 4INRA, UMR1347 Agroécologie DIJON, FRANCE; 5INRA, Lab. Interactions Plantes - Microorganismes, TOULOUSE; 6UMR Diversité et adaptation des plantes cultivées, MONTPELLIER; 7ENSAT, TOULOUSE; 8INRA, UMR1347 Agroécologie, DIJON, FRANCE. *(thompson@dijon.inra.fr)

We have been using genomics approaches with *Medicago truncatula* as a tool to identify key genes determining seed yield and composition in Medicago and in closely related legumes. Analyses of the proteome and transcriptome of the component tissues of the developing seed revealed extensive compartmentalization of gene expression and metabolic activities (Gallardo et al, 2007). Using a TF (Transcription Factor) qRT-PCR platform (Verdier et al., 2008) the Affymetrix Gene Chip (Benedito et al, 2008), and more recently, Nimblegen arrays (Buitink et al. submitted), putative regulatory genes specific for each seed tissue were identified, along with putative target genes of transcription factors (TFs). These genes have been located on the *M. truncatula* genetic map and correlations between map positions of TF loci and QTLs for protein quantities and other seed phenotypes were detected. These correlations were confirmed in certain cases by the existence of similar QTLs at syntenic positions in pea. This approach has enabled us to attribute roles to two genes, both specifically expressed in the developing endosperm of *M. truncatula* and present in pea. One encodes a DOF class transcription factor, whose mutant phenotype severely affects endosperm development. The second gene encodes an endosperm-specific subtilase (*SBT1.1*), which affects final seed weight in both species (D’Erfurth et al, 2012). The importance of the endosperm in determining legume seed size and composition will be discussed.

The research leading to these results has received funding from the European Community FP7 grant agreements FP7 KBBE-613551, LEGATO, and FP7 KBBE-289562, ABSTRESS, from the ANR (QualityLegSeed and GenoPea), and from the Burgundy Regional Council.