Separomics applied to the proteomics and peptidomics of low-abundance proteins: Choice of methods and challenges – A review

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Abstract

The enrichment and isolation of proteins are considered limiting steps in proteomic studies. Identification of proteins whose expression is transient, those that are of low-abundance, and of natural peptides not described in databases, is still a great challenge. Plant extracts are in general complex, and contaminants interfere with the identification of proteins involved in important physiological processes, such as plant defense against pathogens. This review discusses the challenges and strategies of separomics applied to the identification of low-abundance proteins and peptides in plants, especially in plants challenged by pathogens. Separomics is described as a group of methodological strategies for the separation of protein molecules for proteomics. Several tools have been used to remove highly abundant proteins from samples and also non-protein contaminants. The use of chromatographic techniques, the partition of the proteome into subproteomes, and an effort to isolate proteins in their native form have allowed the isolation and identification of rare proteins involved in different processes.

Key words: sample preparation, complex protein extract, subproteomes, low-abundance proteins, cell wall proteins.

Introduction

The separomics challenges in plants

Proteomics tools have been widely used in recent years. Proteomics and peptidomics involve sophisticated methodologies which accurately detect alterations respectively in protein and peptide synthesis, under different physiological situations. Two main tools are widely used to isolate proteins, especially so two-dimensional electrophoresis (2-DE) associated with mass spectrometry (MS), and liquid chromatography associated with MS (LC-MS). Yet both present limitations inherent to the techniques (Cho, 2007). Multi-dimensional liquid chromatography has been valued in recent years as a technique to obtain native samples, due to the need to validate biological events observed in proteomic studies. Obtaining native proteins is one of the biggest challenges in proteomics. Difficulties in isolating and identifying protein and peptide groups occur due to the high complexity of proteins present in the samples, the presence of non-protein contaminants that are difficult to remove and the occurrence of post-translational modifications. The difficulties are even greater for proteins and peptides present in low abundance (LAP) in the tissues. These, however, have attracted the attention of researchers since they are in general very effective and/or transiently present as components of finely controlled metabolic pathways. Thus, alternative methods to detect and identify these proteins are necessary.

Alternative strategies applied to the extraction, purification and biochemical and functional analyses of these molecules have been proposed, favoring access to structural and functional information of hard-to-reach proteins and peptides (Kolodziejek and van der Hoorn, 2010). Separomics is described as a group of methodological strategies aimed at separating protein molecules for proteomics, including fractionation and enrichment of specific molecules (Fang and Zhang, 2008). The use of separomic tools is especially important for peptidomics, which is described as the group of methodologies and procedures applied to the analysis of native peptides by means of proteomics tools, since they are small and non-abundant proteins (Jurgens and Schrader, 2002).

Especially for peptidomics, 2-DE is difficult to apply, due to the low concentration of the peptide molecules, their removal and the occurrence of post-translational modifications.
small sizes (up to 10 kDa), their partial hydrophobic character, and their ionic characteristics, as many peptides are strongly cationic. For the identification of these molecules, the greatest challenges are the small number of available specific databases and the low number of studied and posted molecules, which makes their identification through limited proteolysis techniques and MALDI-MS difficult. In addition, the partial hydrophobicity characteristics and surface charges facilitate peptide molecular associations, making them unavailable for analysis by any known proteomics tools.

For plant proteomics, the greatest challenges are to reduce sample complexity and remove contaminants which are incompatible with the isolation and identification tools (Kolodziejek and van der Hoorn, 2010). The correct use of separations is imperative, especially for the identification of proteins and peptides expressed in low concentrations. This said, the current review aimed at discussing difficulties and challenges in plant proteomics and peptidomics, and to point out methodologies and strategic tools capable of detecting low-abundance proteins, especially biological peptides, differentially expressed in soybean plants after biotic and abiotic stresses.

Proteomic analysis applied to the response of soybean plants to pathogens

Proteomic analysis has become the most powerful tool for the functional characterization of plants. Information on soybean sequencing (Kim et al., 2010; Schmutz et al., 2010) and the soybean genome database (Phytozome v7.0: Glycine max) are available. Soybean was the first legume species to have its complete genome sequenced, becoming therefore a key reference for the more than 20,000 legume species (Schmutz et al., 2010). Several genomes are emerging as model for plants, including that from soybeans. Komatsu and Ahsan (2009) have discussed the advantages and limitations of different proteomics tools applied to the study of soybean defense. In the last two years, different research groups have also discussed the difficulties and alternatives concerning proteomic analyses of plants in general (Sudaric et al., 2010; Yamaguchi and Sharp, 2010; Bindschedler and Cramer, 2011). Low protein concentration, difficulties in protein extraction, genome ploidy, interference of highly abundant proteins in green tissue are some of the main challenges in plant proteomics (Bindschedler and Cramer, 2011). In general, there are many additional challenges for plant proteomics, such as the identification of proteins expressed transiently and in relatively low concentration, or the fact that the natural peptide has not yet been deposited in any of the databases. In silico information is essential for proteomic analysis of important physiological processes, such as plant defense. The subcellular proteome analysis of soybean plants submitted to stress conditions aims to identify proteins and peptides that are differentially expressed and potentially involved in plant defense or pathogen resistance induction. Biotechnological methods may then be developed in order to interrupt the pathogen action before infection, or to produce defense agents to boost the plant’s defense system. Organelle proteins and specific soybean tissues have been studied, such as membrane (Komatsu and Ahsan, 2009; Bindschedler and Cramer, 2011), primary roots (Nouri and Komatsu, 2010) and cell wall (Yamaguchi and Sharp, 2010).

Searching For Native Low-Abundance Proteins and Peptides By Proteomics and Peptidomics

Techniques with different sensitivities and accuracies should be used in the analysis of the proteome. Due to the physical, chemical and biological diversity of proteins, proteomics tools present limitations that make it unfeasible to analyze the entire proteome with only a single separation strategy, even if it is orthogonal, such as the 2-DE or multi-dimensional liquid chromatography. Furthermore, the cellular protein concentration may vary from mg mL$^{-1}$ to pg mL$^{-1}$ (Fang and Zhang, 2008; Jorrín-Novó et al., 2009).

Different proteomic analysis platforms were developed in recent decades based on biochemical tools already available for protein isolation and identification. By means of these strategies, thousands of proteins have been identified, especially high-abundance proteins (HAP), whereas the identification of low-abundance proteins (LAP) is still limited. The search for LAP is growing, for they can represent important biological markers (Lescuyer et al., 2007), be responsible for eliciting important cellular responses, and may even correspond to low molecular weight transcription factors. Not only is their synthesis transient but they can also be easily lost (Corthals et al., 2000). Different procedures have been used to reduce HAP levels in extracts and to improve LAP detection, such as: (1) partial removal of RUBISCO by increasing dithiothreitol concentration in extracts of rice leaves (Cho et al., 2008), or by fractionating extracts of soybean leaves with calcium chloride and sodium phyate (Krishnan and Natarajan, 2009), (2) addition of solvents such as the isopropanol to remove storage proteins from extracts of soybean seeds (Natarajan et al., 2009), (3) division of the proteome into subproteomes, as done for banana leaf membranes (Vertommen et al., 2011), or (4) the use of affinity and immunoaffinity columns for protein purification (Azarkan et al., 2007; Fang and Zhang, 2008).

Due to the need for improvement and/or development of protein separation methods for proteomic studies, separomics makes its appearance as the science for proteomic separation. The main goal of separomics is to obtain a specific set of proteins in a given biological system for proteome composition analysis, for protein-protein interaction studies, or for analysis of alterations in protein synthesis in biological materials submitted to different physiological conditions. The concept of Separomics (originally called Seppromics) was proposed to describe and define technolo-
gies, processes, requirements, patterns and applications on proteomic separation, including fractionation and enrichment (Huang et al., 2005). The greatest challenge was to reduce the complexity of a given proteome, aiming to increase the effectiveness of proteomic analysis as well as the identification of new proteins through mass spectrometry (Fang and Zhang, 2008; Kosová et al., 2011).

Specific structures in plants, such as the cell wall and vacuoles, contain substances responsible for the inferior and non-reproductive outcomes during the separation of proteins due to proteolytic breakdown, streaking and charge heterogeneity. In most plant tissues, the proteins are part of complex structures, requiring special care for their isolation in a soluble form, as a native or non-native protein. The most commonly found interfering molecules during protein isolation are phenolic compounds, proteolytic and oxidative enzymes, terpenes, pigments, organic acids, inhibiting ions, and carbohydrates (Carpentier et al., 2005).

A further limiting factor in proteomic studies in plants is the loss of protein solubility caused by either the enrichment of protein extracts or by the properties of the solvents. Solubility of proteins is usually dependent on their concentration and amino acid composition, type and solvent dielectric constant, ionic strength, and on the presence of contaminants (Jorrín-Novo et al., 2009). The optimum solubility conditions will be empirically defined for each system.

Preparing and Isolating Low-Abundance Proteins and Peptides From Plants

Plant extracts require specific sample preparation procedures especially adjusted for proteomic and peptidomic studies. These procedures should take into account the solubility, the physico-chemical characteristics and the cellular localization of the proteins, as well as the presence of interfering molecules (Chen and Harmon, 2006; Matros et al., 2010), which correspond to the most challenging and critical aspects (Komatsu and Ahsan, 2009).

Subproteomes

Since the preparation protocols do not allow evaluation of the complete proteome and peptidome, the protocol definition should take into consideration protein subgroups and source material of interest, whether it is an organelle, a cell or a tissue, in order to reduce sample complexity, enrich and increase the possibility of identifying LAP of interest involved in different cellular mechanisms and different locations. Hence, the information obtained for each subproteome of the same source material contributes to achieve greater coverage of the proteome being studied. This strategy has attracted the attention of several researchers (Cánovas et al., 2004; Zhang et al., 2004; Natarajan et al., 2005, 2009; Oehrle et al., 2008; Jorrin-Novo et al., 2009; Krishnan and Natarajan, 2009; Agrawal et al., 2010; Matros et al., 2010; Kota and Goshe, 2011).

Plant cell wall proteins

For the study of plant proteomes, the extracts may be successfully fractionated into soluble proteins and cell wall (CW) proteins (Figure 1). CW proteins represent a subproteome of great importance, since many of these molecules are involved in the maintenance of the cellular structure and in processes of plant defense, for instance in responses to abiotic and biotic stresses (Jamet et al., 2006; Kong et al., 2010), or as a constitutive barrier against pathogenic microorganisms. Studies in our laboratory evidenced high constitutive antimicrobial activity against two plant-pathogenic bacteria by the peptide fraction obtained from CW extracts from leaves of bell pepper (Figure 2, Teixeira et al., 2006) and 60-day-old eggplant (Almeida et al., 2008). Also for eggplants, the highest inhibition levels were obtained with CW extracts from 5-cm-tall plants, while soluble extracts promoted the highest inhibition rate when fully expanded leaves were analyzed (Table 1). These results suggested that young plants exhibit an innate defense mechanism, very likely to minimize plant microbial invasion, whereas expanded leaves produce soluble defense molecules (Almeida et al., 2007).
Plant CWs are highly dynamic and contain chemically active compounds secreted by the cells, which are essentially polysaccharides and proteins, the latter comprising approximately 10% of the CW mass (Jamet et al., 2008). These proteins, which are difficult to isolate from these complex matrices, require specific extraction methods (Teixeira et al., 2006; Zhu et al., 2006; Almeida et al., 2007, 2008; Kong et al., 2010), due to their low abundance. Differential extraction enriches the extract, allows access to the CW LAP, and facilitates the comparison of the expression profiles under different stress conditions (Watson and Summer, 2006; Negri et al., 2008).

Protein extraction from CWs can be achieved by methods that may involve or not cell disruption. For each procedure there are advantages and disadvantages, especially concerning contamination and experimental procedures. Methods commonly used involve calcium chloride and lithium chloride solutions (Feiz et al., 2006; Kong et al., 2010). Calcium chloride does not disrupt the cell wall but is capable of releasing ionic molecules externally anchored to the CW, whereas lithium chloride solution is used to extract proteins intrinsic to the CW, such as glycoproteins (Feiz et al., 2006). When comparatively evaluating the CW subproteomes of soybean leaves submitted or not to biotic stress using reversed-phase high performance liquid chromatography we obtained a group of differential protein peaks in the comparative chromatograms, demonstrating that sample complexity was reduced. This procedure allows the identification of the isolated proteins.

Proteins with different ranges of molecular masses

When the aim is to identify proteins of a specific group of molecular mass present in a complex sample, methodologies based on ultrafiltration have an advantage, since they are capable of fractionating, concentrating and exchanging solutions, and thus recovering native samples with molecular masses of interest. Ultrafiltration is a low-cost process requiring simple equipment only, such as filtering membranes of different cut-off values, which allow the fractionation of different protein groups. Soluble (SE) and CW extracts (CWE) of soybean leaves submitted (Figure 1) or not to stress were successfully fractionated in our laboratory through ultrafiltration. Fractions for peptidomics and proteomics, with proteins smaller than 30 kDa and greater than 30 kDa, respectively, were recovered. After separating the fractions by reversed-phase chromatography (RPC) and recovering the proteins and peptides for mass spectrometry analysis, differentially expressed molecules were successfully identified.

Molecular exclusion chromatography (MEC) is also an alternative used by our group for this purpose. This method favors isolating a specific protein group according to the selected resin. In general, MEC presents low accuracy in the separation of small proteins and peptides, yet it is a fast procedure that generates clean samples with low salt concentration, which is important for later stages of purification, for instance during RPC. MEC has been successfully used in our proteomic analyses using multi-dimensional liquid chromatography. A constitutive antimicrobial peptide was identified in CW extracts from tomato plant after MEC following RPC while an induced defense peptide was detected in plants after abiotic stress (submitted for publication).

Proteins from different cellular compartments

Ultra-centrifugation enables the separation of cellular organelles according to their sizes and densities. Zonal centrifugation or centrifugation by differential velocity may be used for fractionation, the latter being a technique which generates greater sedimentation strength, enabling the separation of organelles of similar characteristics (Komatsu, 2006), such as nuclei, mitochondria or plastids.
while maintaining their integrity (Wijk, 2004; Kosová et al., 2011).

**Contaminant removal and use of additives**

A limiting factor for proteomic studies is the presence of protein and non-protein contaminants in the samples. Contaminants may interfere with the detection, isolation or identification of proteins of interest, acting by aggregation with the proteins, by affecting the signal/noise relationship of the detection equipment, by promoting protein degradation or reducing the activity of enzymes, among other factors. Under native conditions, the removal of these contaminants may be achieved by dialysis, salt fractionation, or with organic cold solvents, through ultrafiltration or preparative chromatography. Under denaturing conditions, steps including selective heating or addition of concentrated acids may favor sample bleaching and the enrichment of the protein group of interest. Also, anionic detergents like sodium dodecyl sulfate (SDS) or nonionic detergents such as Triton X-100 and Nonidet P-40 can be added in different concentrations. The concentration to be used must be empirically defined.

Fractionation by ultrafiltration allows the simultaneous desalting and concentration of the samples in the native form (Kong et al., 2010). However, during the ultrafiltration procedure certain caution is required so as to maintain the sample in its native form, as well as to avoid protein agglomeration. Some of the factors to be controlled are hydrophobicity, pH and ionic strength of the buffer or solution being used. The choice of salts and pH of the buffer solution for ultrafiltration depends on the subproteome under study. For instance, as pH values around 5.5 are close to the pH of the cell wall, the fractionation of CW proteins should be done at pH values close to 5.5 to avoid precipitation of these proteins (Watson and Summer, 2006). On the other hand, many storage proteins, which are usually contaminants, precipitate between pH 4.5 and 4.8 (Speroni et al., 2010), and extraction at these pH values produces less complex extracts. For ultrafiltration of membrane proteins and partially hydrophobic proteins, detergents should be used, as these become soluble in the presence of amphipathic compounds, such as 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate (CHAPS), SDS and the nonionic surfactant Triton X-100, among others.

Pressure is considered an important factor which frequently is not controlled during ultrafiltration. There are not sufficient studies reporting protein agglomeration, solubility and protein conformation under pressure (Chalikian and MacGregor, 2009; Speroni et al., 2010). Iwabuchi and Yamauchi (1987), fractionating glycine (360 kDa) and β-conglycinin (180 kDa), found low molecular mass proteins in the protein fraction containing these storage proteins, suggesting the occurrence of agglomeration of low molecular mass proteins together with high molecular mass proteins.

In our experience, when a low salt concentration buffer (5 mM Tris buffer, pH 7.0) was used after the ultra-filtration procedure, the occurrence of agglomeration in extracts from soybean leaves increased. Agglomeration was also increased under cold temperatures, and an insoluble precipitate was observed in fractions containing low and high molecular masses. Replacing this buffer with 20 mM Tris-HCl, pH 7.0, containing 20 mM ammonium acetate, agglomeration was no longer observed, suggesting that agglomeration was an ionic behavior. In an attempt to disaggregate the precipitate formed in the high molecular fraction, different procedures were used. Disaggregation was not achieved in the presence of 0.5% (v/v) Triton X-100 or by dilution (up to 10 times) with 100 mM Tris-HCl, pH 7.0, but the sequential addition of 0.4, 1.0 and 2.0% (w/v) SDS partially reduced the agglomeration, thus corroborating the hypothesis of ionic behavior.

Denaturing one-dimensional electrophoresis (16.5% T) done with extracts from soybean leaves enriched for low molecular mass proteins (up to 30 kDa) yielded protein bands higher than 30 kDa, suggesting the occurrence of agglomeration after ultrafiltration. This was confirmed when a 60-kDa band was eluted from the gel and again separated under the same conditions, this time producing 6-kDa and 30-kDa bands. Similarly, the elution and new separation of a 6-kDa band has also produced the same bands, confirming that heating the sample for 10 min in the sample buffer (Teixeira et al., 2006) was unable to solubilize proteins in the sample.

Fractionation with ammonium sulfate has been used by our group since 2000 for bleaching plant samples, to remove contaminants such as carbohydrates, phenolic compounds and pigments, among others. Plant extracts obtained in the absence of this compound presented high noise/signal relationship in the spectrometry analyses and protein agglomeration was observed. Recently, Park et al. (2008) stressed the importance of this procedure to remove interfering molecules. Nevertheless, caution should be taken, as the presence of salt may interfere with electrophoretic properties and the MS procedure.

Specific stages of the proteomic analysis require samples with specific characteristics. 2-DE, for instance, is highly affected by salts, as these may accumulate in the tips of the strips during the first-dimension separation. They can cause heat generation and the formation of zones with different conductivities and degrees of hydration. During MS analysis, salts may increase noise that interferes with the ionization of the molecules. SDS interferes with the RFC procedure by causing the degradation of the resins. Samples with high protein concentration may precipitate and those with low protein concentration may prevent the detection of LAP. Hence, samples should be prepared in buffers with low ionic strength and in the presence of protease.
inhibitors with different specificities (unpublished date). The use of non-interfering additives is an empirical decision, which should be carefully evaluated because these compounds may have to be removed in a subsequent purification step.

Refined methods for isolation

Separation by two-dimensional electrophoresis

Two-dimensional electrophoresis (2-DE) is the most commonly used separation technique in proteomics for comparative and global protein analyses under differential conditions. A high number of proteins can be identified in a single gel. Considering that in general, plant tissues do not present a high protein concentration, and that the presence of proteases and interfering molecules can drastically affect proteomic analyses, an efficient protein extraction protocol must be used. This protocol should be able to eliminate secondary metabolites, remove additives which are not compatible with the different stages of purification, and enrich the LAP. Extraction methods using phenol in conjunction with ammonium acetate/methanol precipitation have proved to be highly efficient (Isaacson et al., 2006). Using this procedure Xu et al. (2006) were able to obtain a large number of intense and well-resolved spots from soybean leaves in 2-D gels. Similar results were obtained in our laboratory. Methods based on precipitation with trichloroacetic acid (TCA) have also been used in the literature (Chen and Harmon, 2006) and in our laboratory. Using a method based on an initial TCA precipitation step followed by extraction by the dense phenol/SDS method and a final precipitation with ammonium acetate/methanol, Wang et al. (2003) obtained a high number of soluble and membrane proteins from soybean leaves. To optimize this method, we introduced some modifications, such as the use of protease inhibitors (1 mM phenylmethanesulfonyl fluoride), reducing agents (different concentrations of β-mercaptoethanol and dithiothreitol) and 1% polyvinylpolypyrrolidone, which eliminate or prevent the action of proteases and compounds that interfere with the subsequent steps in protein separation and identification.

Liquid chromatography (LC) can be used as an alternative to 2-DE when the proteins of interest present extremes of molecular mass and pI, when they are highly hydrophobic or LAP (Natarajan et al., 2009; Gilmore and Washburn, 2010; Kolodziejec and van der Hoorn, 2010).

Separation by chromatographic methods

Chromatographic methods using a combination of different separation principles (multi-dimensional) in on-line or off-line processes, or the selective separation by affinity have been used and improved to resolve the complexity of the extracts and/or to enrich them. Commercially available on-line chromatographic systems generally use a strong ionic exchange column (IEC) followed by a reverse phase. Molecular exclusion chromatography (MEC) followed by reverse phase chromatography (RPC) is also used (Issaq, 2001), though less often. Our group has worked with off-line chromatography with success, especially associating MEC and RPC for prospection of proteins with different ranges of molecular masses, when two antimicrobial peptides from tomato leaves were purified under native conditions and subsequently identified (data submitted for publication). These procedures are orthogonal, creating a large number of fractions to be analyzed, making proteomic analyses by MDLC difficult and lengthy. However, it allows a greater proteome and peptidome prospection, with little restriction for very large or very small and electrically charged membrane proteins. And in particular, it allows the isolation of native purified samples for evaluation of the proteome functionality.

Although these methods are very efficient, complementary techniques are required for obtaining further information regarding the proteome. Thus, the development and application of technologies and methods for separation by affinity and enrichment have become a high priority for defining separomics protocols to be used for different organisms.

A great variety of affinity and immuno-affinity columns are already available and allow the separation of complex biological extracts into different protein classes, or even to fractionate LAP. These commercially available columns are important for the identification of protein-protein interaction networks (Azarkan et al., 2007) and to remove unwanted HAP proteins, peptides and nucleotides, metals, etc. (Keshishian et al., 2007; Cellar et al., 2008; Fang and Zang, 2008; Huang and Fang, 2008; Miernyk and Hajduch, 2011).

Identifying Soybean Proteins and Peptides By Proteomics Tools

For comparative proteomics and peptidomics, 2-DE is the proteomic platform giving highest-throughput (Brandão et al., 2010). More sophisticated procedures, such as 2-D DIGE and isotopic labeling can be used. In most soybean proteomic studies using 2-D procedures, the proteins undergo electrophoresis, are trypsinized in gel and are then analyzed by MALDI-MS/MS. The PMF technique (Peptide Mass Fingerprint) following MS correlates the peptide masses after proteolysis of the unknown protein with the peak list of theoretical proteins deposited in public databases (Chamrad et al., 2004; Elias et al., 2005), which are assessed by search software such as MASCOT (Thiede et al., 2005). Similarly, PFF (Peptide Fragment Fingerprint) correlates mass spectra of fragments of these peptides (Chamrad et al., 2004; Elias et al., 2005; Nielsen et al., 2005). Fragmentation spectra can also be analyzed by “de novo” peptide sequencing using MS, which gives the amino acid sequences (Roepstorff and Fohlman, 1984). The development of a plethora of bioinformatics resources has
allowed the identification and characterization of proteins with accuracy, thanks to genomic and protein sequence information deposited in public databases (Schmutz et al., 2010).

Conclusions

Proteomics is an invaluable tool for functional genomic analysis of plants. Techniques with different sensitivities and accuracies should be used in the analysis of the respective proteome. Highly sensitive and accurate identification equipment is available; however, better protein and peptide extraction procedures and methods for enrichment and isolation of low-abundance proteins with transient expression and of rare peptides are needed. Some alternatives to overcome or minimize these limitations are: (1) the decrease of extract complexity to allow evaluation of subproteomes, (2) use of methodologies and/or additives to remove highly abundant proteins known as contaminants, (3) use of chromatographic techniques which allow the enrichment of specific protein fractions, and (4) isolation of proteins in their native form to aid their identification and validation of metabolic pathways in which they are involved. The use of basic biochemistry tools which separate protein and peptides based on their physico-chemical characteristics can further the enrichment process. Knowledge about the chemical and structural characteristics of non-protein contaminants is helpful in the choice of methods and tools for their removal or inactivation. This information will also aid in defining methods to prevent solubilization of these contaminants and their consequent contact with the proteins of interest.

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Internet Resources

Phytozome ver. 7.0: Glycine max. Center for Integrative Genomics (CIG), Joint Genome Institute (JGI), University of California Regents, http://www.phytozome.net/soybean (January 24, 2011).

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