Physiological and molecular aspects of pod shattering resistance in crops

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Abstract: Pod shattering resistance is a trait acquired by crops in the process of evolution. Manipulation of physiological and molecular processes is fundamental for the improvement of shattering resistance in crops. In this review we discuss several enzymes, key hormones and their possible roles or relationships involved in pod shattering, and highlight responsible genes, quantitative traits loci (QTLs) and their implications for increased pod shattering resistance. Cell wall degrading enzymes, particularly β -glucanases and endopolygalacturonases play an important role in the process of pod dehiscence. It is not clear how and to what extent a specific hormone regulates the dehiscence zone differentiation and the dehiscence process is not clear. Resistance to shattering is highly heritable and is not controlled by a single gene. Several QTLs associated to dehiscence have been identified in crops, while the underlying genetic functions of these QTLs deserve further investigation. Further physiological analyses of the pod wall will help to understand better the pod dehiscence.

Keywords: enzymes; gene identification; hormones; QTLs

Crops have been selected for acquired resistance to pod shattering or pod dehiscence to avoid yield losses in the process of domestication (Fuller & Allaby 2009; Bennett et al. 2011; Funatsuki et al. 2014). Still, susceptibility to pod shattering remains one of major traits resulting in yield losses in many seed crops including soybean (*Glycine max* (L.) Merr.), sesame (*Sesamum indicum* L.), oilseed rape (*Brassica napus* L.), birdsfoot trefoil (*Lotus corniculatus* L.) and other pulse crops (ABD EL-Moneim 1993; Grant 1996; Langham & Wiemers 2002; Weeden et al. 2002; Child et al. 2003; Romkaew et al. 2008).

In soybean, the contribution to total yield loss by pod shattering in the South Eastern USA was around

37% (Philbrook & Oplinger 1989), and respective yield loss insusceptible and intermediately susceptible cultivars was 57–175 and 0–186 kg/ha respectively (Tukamuhabwa et al. 2002). As much as 40 to 60% shattering rate of matured pods was reported in some common vetch (*Vici sativa* L.) cultivars (Sattell et al. 1998; Dong et al. 2016). For birdsfoot trefoil, possible yields could be as high as 400 to 600 kg/ha with best management, but as low as 50 kg/ha under poor weather conditions in which seed yield losses by shattering were 3 to 5.3 kg/ha/day (Garcia-Diaz & Steiner 2000; Řepková & Hofbauer 2009). Seed yield losses in big trefoil (*L. uliginosus* Schkuhr.) ranged from 7 to 88% pods with a rate of 10% per

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day due to pod shattering (HARE & LUCAS 1984). Yield loss was 11–25% in the pods of many *Brassica* species once they reached maturity (PRICE et al. 1996), and up to 50% in adverse seasons (MACLEOD 1981). Besides the economic importance from the yield loss, pod shattering can also lead to the emergence of volunteer weeds in the subsequent growing season, thus impeding future crop rotations. Differences in pod anatomy and morphology in soybean may cause pod dehiscence (ZHANG et al. 2018), while the physiological and molecular nature factors are pivotal for the improvement of shattering resistance in crops. This paper intends to discuss an overview of the control of pod dehiscence from the aspects of enzymes, phytohormones, genes and QTLs participation and functioning, with an aim to provide insight and further understanding of the pod dehiscence mechanism and to devise strategies in manipulating and reducing the problem.

Enzymes involved in pod shattering process

Dehiscence zone (DZ) is the element involved in pod dehiscence (Ferrándiz 2002). The pod shattering and seed release are mostly due to the loss of cellular cohesion and degradation of the middle lamella in DZ (Carlson & Lersten 2004; Bennett et al. 2011). The degradation of pectin, especially homogalacturonan-rich pectin in the middle lamella has been commonly found in fruit shattering, and pectin integrity is suggested to be important for cell adhesion (Ridley et al. 2001; Dong & Wang 2015).

An increased β -glucanase activity in the cells of the DZ and accompanied degradation of the cell wall at the site of fruit dehiscence, were found in *Brassica* (Kemmerer & Tucker 1994). Result with antibodies against β -glucanase indicated that the break strength in the abscission zone was significantly reduced by antibodies injection, while the changes of this enzyme and its distribution were completely involved in abscission processes, indicating that β -glucanase is actually required for abscission (Sexton *et al.*1980).

Endopolygalacturonase (endo-PG) termed RD-PG1e, specifically expressed in the DZ, catalyzes the main chain of the homogalacturonan region of pectin, i.e. the hydrolysis of α-1,4-glycosidic bonds in polygalacturonic acid. This hydrolysis process was considered to facilitate the breakdown of the middle lamella (Christiansen *et al.* 2002). The endo-PG transcript analysis in soybean further indicated that the endo-PG was involved in the breakdown of the middle lamella before the occurrence of dehiscence,

and thus played an active role in the process of pod shattering. β -glucuronidase (GUS) staining of the transgenic Arabidopsis lines and inhibited formation of abscission layers without PG all demonstrated and supported this idea (Ogawa *et al.* 2009).

Although no correlation was observed between pod dehiscence and the PG activity either temporally or spatially (Meakin & Roberts 1990), and very few direct genetic evidence was found for the physiological importance of individual PGs, there was a report showing that increasing PG activity was correlated with cell separation in the shedding of fruit (Kalaitzis *et al.* 1997). The increased activities of endo-1,4-β-glucanases and endo-PG, but a significant decrease of the protein content in the DZ throughout maturation and senescence were also found (Christiansen *et al.* 2002). These findings indicated that endo-1,4-glucanases and endo-PG may disintegrate the middle lamella in the separation layer, and the outcome is a decreased cell to cell adhesion.

Therefore, cell wall-modifying enzymes affect the breakdown of DZ, which should be closely involved in the processes of pod dehiscence through cell wall disassembly, reconstruction and adjacent cell bindings required for dehisced pods (Hadfield et al. 1998; Del Campillo 1999; Rose & Bennett 1999). Further examination of their expression and activity in the pod tissues and/or contributions to the breakdown of the cell wall is required.

Hormones in pod shattering

As a biologically active and readily diffusible phytohormone, ethylene has been identified in association with the processes of fruit ripening and abscission (Abeles et al. 1992). The onset of pod dehiscence was correlated with a burst in seed ethylene production, and a transient peak production of ethylene was also temporally correlated with the increase in β -glucanase activity, the enzyme responsible for the degradation of the cell wall leading to shattering (Oeller et al. 1991). It should be noted that exposure to the gas does not accelerate pod shattering while the peak in ethylene production might only act as a signal factor for pod dehiscence since the climax usually precedes dehiscence. Actually, the fleshy and dehiscent fruits become more competent in response to ethylene ripening signals as they age (JOAQUIN et al. 2007). Therefore, peak production of the ethylene might be only responsible for the acceleration of the senescence onset but not dehiscence promotion (John et al. 1995).

In the abscission zone, the inhibition of β -1,4-glucanase mRNA accumulation by auxin has been reported, and the decreased auxin content in the DZ was correlated with increased β -glucanase activity (Tucker *et al.*1988). Pods treated with 4-CPA (auxin mimic 2-methy l-4-chlorophenoxyacetic acid) delayed β -1,4-glucanase activity approximately 10 d and concomitant cell separation in the DZ (Chauvaux *et al.* 1997). Dehiscence zone differentiation required auxin accumulation in early stages, while auxin depletion at later stages could be important for triggering cell separation (Ballester & Ferrándiz 2017). These findings suggest that the regulation of auxin and its concentrations on pod dehiscence is partially associated to the activation of the β -1,4-glucanase activity in the DZ.

Gibberellic acid (GA) was shown to be important for the continued pod elongation in oil seed rape (Bouttier & Morgan 1992) and is completely required, for the correct functioning of INDEHISCENT protein in Arabidopsis fruit (Arnaud *et al.* 2010). However, functions of GA on pod dehiscence have not been fully examined. Furthermore, less research has been conducted on the involvement of the two phytohormones, abscisic acid and cytokinins on pod dehiscence in crops, although their roles in other aspects of plant development have been highlighted (Finkelstein 2013; Humplik *et al.* 2017).

Therefore, how and to what extent a specific hormone regulates DZ differentiation and the dehiscence process is not clear. Since phytohormones could change during pod development, and can either act synergistically or antagonistically towards one other, it is possible that not a single hormone but phytohormones interactions are actively involved in pod shattering. Changes of individual hormone during pod development particularly hormone balance or interplay in relation to pod dehiscence need to be fully identified, which may add more knowledge and shed insight for their roles in pod and seed development.

Genes involved in pod dehiscence

In soybean, the first gene, *Pdh1* (*Pod Dehiscence1*), was identified as a candidate for leading protein involved in pod dehiscence (RALPH *et al.* 2007; FUNATSUKI *et al.* 2008). This gene is highly expressed in the lignin-rich inner sclerenchyma of pod walls at the stage of initiation in lignin deposition, which promotes pod dehiscence by increasing the torsion of dried pod walls, and thus serves as a driving force for pod dehiscence (FUNATSUKI *et al.* 2014).

Another gene, NAC (The NAM, ATAF1/2 and CUC2 domain protein) *SHATTERING1-5* (*SHAT1-5*) activates secondary wall biosynthesis and enhances pod-wall binding strength through the thickening of fibre cap cells in secondary walls of the pod ventral suture (Dong *et al.* 2014). In *Arabidopsis*, evidences indicated that a subset of closely related NAC domain proteins, including NST1/ANAC043, NST2/ANAC066, and NST3/SND1(Secondary Wall-associated NAC Domain Protein)/ANAC012 act as master transcriptional switches governing secondary cell wall biosynthesis in a partially redundant manner (MITSUDA *et al.* 2005; ZHONG *et al.* 2007).

A recessive allele was responsible for non-shattering characteristics in wild types of common vetch, whereas dominant alleles were identified for shattering in cultivated types (ABD EL-MONEIM 1993). In pea, a main candidate gene for pod dehiscence was localized on LGIII, and identified as proline-rich extensin-like protein or a homolog of peptidoglycan-binding domain protein (PGDB) of Medicago truncatula (TAYEH et al. 2015). As extensins are structural cell-wall proteins specific to plants, and can substantially regulate mechanical cell wall properties through linkages to other cell wall component (LAMPORT et al. 2011). The MACE-P015 gene is supposed to have a general peptidoglycan binding function and thought to play a critical role in pod dehiscence of pea. In Arabidopsis thaliana, not only SHATERPROOF1 and SHATER-PROOF2 genes dominantly regulate DZ formation, but are also two partially redundant genes that control the formation of secondary cell walls of pods (LILJEGREN et al. 2000; Dinneny & Yanofsky 2005; Mitsuda & Онме-Така 2008). This suggested that multiple genes with minor effects fundamentally determine the resistance to pod shattering in crops (LIU et al. 2007).

Therefore, genes as the main regulators of lignified layer establishment in pod wall development, cause pod dehiscence. Since current literature shows that it is not a single candidate gene underlying pod dehiscence for any crop, the identification and differentiation of homologous or novel genes defining dehiscence zone from the pod suture as well as venture tissue are still a great challenge in understanding the molecular mechanism of the pod dehiscence resistance in crops.

QTL involved in pod dehiscence

Common quantitative trait loci (QTLs) related to shattering resistance have already been identified from several crops (PATERSON 2001). A single locus

controlling pod dehiscence (PD) was found in lentil, while two loci, one controlled the number of twists along the length of the shattered pod, and the other one controlled the percentage of shattered pods, were identified in mung bean (ISEMURA *et al.* 2007). Two similar loci were found in pea (WEEDEN *et al.* 2002; WEEDEN 2007), and common bean (KOINANGE *et al.* 1996). *Dpo* locus responsible for loss of PD in pea was localized on LGIII (BORDAT *et al.* 2011).

Considerable progress has been achieved in QTL identification in soybean. Restriction fragment length polymorphism (RFLP) loci linked to QTL conditioning resistance to PD in soybean were identified in 1997. The consistency of QTL across locations indicated that PD was highly heritable and conditioned by one major and a few minor QTLs. Their findings provided a basis for comparative mapping of the trait in other legumes. QTL for Pod Dehiscence1 (qPDH1) was the major QTL controlling pod dehiscence in soybean (BAILEY et al. 1997), which most likely affected the process of lignin biosynthesis or composition of the lignified cell walls (Suzuki et al. 2009). This QTL was found to be located between simple sequence repeat markers, Sat_093 and Sat_366, and the shattering resistance allele at *qPDH1* was proved to be valuable in different genetic backgrounds at multiple locations (Funatsuki et al. 2008). However, pdh1 for the shattering-resistant genotype is defective because it has a premature stop codon (Funatsuki et al. 2014).

Three QTLs, two on LG J and one on LG D1b were found, while QTLs on LG J were mapped quite far away from qPDH1 (SAXE et al. 1996). A QTL, identical to *qPDH1* was also detected on LG J in another study, however, its effect was not big enough (LIU et al. 2007). No major QTL but only a minor QTL only was detected either on LG J, or on other LGs in the segregating population of soybean derived from a shattering susceptible cultivar, and a shattering resistant cultivar (KANG et al. 2009). These findings suggest that (1) other identified QTLs are likely to differ from *qPDH*; (2) multiple alleles at *qPDH1* or other QTLs near *qPDH1*existed for pod dehiscence; and (3) a shattering resistant cultivar might be developed by pyramiding shattering resistance alleles at minor QTL. Fine mapping and identification of QTLs with large effects on pod dehiscence specific to soybean or legumes deserve further investigation.

Concluding remarks

Cell wall degrading enzymes, particularly β -glucanases and endopolygalacturonases play an important

role in the process of pod dehiscence. How and to what extent a specific hormone regulates DZ differentiation and the dehiscence process is not clear. Interactions among phytohormones might be more important than a single hormone in DZ differentiation and/or as triggers of pod shattering. Shattering resistance is highly heritable and is not controlled by one gene. Several QTLs associated with dehiscence have been identified in legume crops, but the identification of the underlying genetic functions lags far behind. The understanding of how pods 'unzip' at molecular level is essentially important, which could provide a further avenue for regulating pod dehiscence. Further physiological analyses of pod tissues, particularly those of the pod wall will assist in the fully understanding pod dehiscence. Plant biologists and breeders are still confronted with a great challenge in manipulating or tackling the forefront of the pod dehiscence issue.

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