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## PERSPECTIVE

# Fertility versus disease resistance, a hard choice

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Who among us has not heard the phrase, all good things come at a cost. Luckily, the cost/benefit scenarios that we face day to day are generally minor (extra piece of cake or a slim figure) compared with having to choose between the opportunity to have children and disease immunity. The work by Wang and colleagues (Chu et al. 2006) brings into light this exact conundrum in the plant world. The authors demonstrate a trade-off between fertility and pathogen defense, depending on levels of expression of the *xa13* gene.

## Use of natural variation to clone disease resistance genes

Plants have acquired a myriad of subtle tricks to deal with their surroundings. Whether it is capturing sunlight, searching for water, or avoiding being eaten, they have evolved numerous signaling mechanisms and specialized cell types to deal with their world. Most striking are the ways they have learned to deal with pests and disease. Some plants use volatiles to signal the enemy of the herbivore that is on the attack (Kessler and Baldwin 2001). Others are resilient due to spines, toxic compounds, or an impervious cuticle. Still others play a cat and mouse game of evolution, learning to recognize the disease, then sealing it off in sacrificial death of the infected cells. The rapid evolution of these disease agents (notably bacteria, fungi, and viruses) means that the plant is often quickly outwitted and loses resistance.

*Xanthomonas oryzae* pv. *oryzae* (Xoo), the causal agent of bacterial leaf blight, is the most devastating bacterial disease in Asia and Africa and is now known to occur in Australia, the United States, and South America (Jones 1988). Fortunately for the rice plant and those of us that eat rice, plants have evolved resistance genes to combat this devastating disease. Cloning of five of these genes has been previously described (Song et al. 1995; Yoshimura et al. 1998; Iyer and McCouch 2004; Sun et al. 2004; Gu et al. 2005). In this issue, Wang and colleagues (Chu et al. 2006) describe their recent success at isolating a resistance gene to Xoo that has a surprise role in fertility.

Wang and her group used a mapping population constructed between near-isogenic lines IR24 carrying *Xa13*, and IRBB13 carrying *xa13*, to narrow the position of *Xa13* to a 9.2-kb region. Within this interval was a single predicted gene. They confirmed that it was the sought-after gene by transforming the dominant allele, *Xa13*, into IRBB13, the resistant host. The resulting transgenic lines carrying the *Xa13* allele were susceptible. Comparison of the IRBB13 and IR24 versions of *XA13* revealed one amino acid difference. To determine whether this difference was the key between susceptibility and resistance, they sequenced *xa13* and *Xa13* from 10 additional varieties. Some of the *xa13* and *Xa13* alleles had identical amino acid sequences, suggesting that the difference between susceptibility and resistance lay outside the coding sequence.

The group (Chu et al. 2006) then analyzed the promoter region of *Xa13/xa13* from seven susceptible and 11 resistant rice lines. Various differences were found that were not allele specific, but all 11 *xa13* lines had insertions, deletions, or substitutions within an 18-base-pair (bp) region, 70 bp upstream from the start of transcription. They also showed that transcript levels of *Xa13* are dramatically increased upon infection, going up sixfold at 8 h and 47-fold at 72 h. Such an induction does not occur in an *xa13* background. Other susceptible or resistant lines showed a similar trend. Thus, this 18-bp region is likely to be responsible for induction of gene expression of *Xa13* but not *xa13* alleles. Silencing of either *xa13* or *Xa13* by RNA interference (RNAi)-enhanced the resistance, supporting the hypothesis that high levels of expression of this gene are detrimental to the plant.

The *XA13* sequence does not resemble the product of any previously identified resistance gene, but does show 50% amino acid identity to the product of a nodulin *MtN3* gene that is induced by *Rhizobium* in legume nodule development (Gamas et al. 1996). Wang and colleagues suggest that *XA13* encodes a modulator of bacterial invasion. Indeed, the bacteria grow nearly 2000-fold better in plants expressing *Xa13* than in the *xa13* plants by 14 d after inoculation. Another disease-resistance allele that is inherited recessively is *mlo* in barley. The dominant *Mlo* allele encodes a membrane protein that is thought to function as an entry portal for successful pathogenesis of fungal pathogens as it accumulates at sites of fungal penetration (Bhat et al. 2005). Although

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Bart et al.

*Xa13* and *MLO* proteins show no similarity, they may carry out similar tasks. *Xoo* invades rice plants through wounds and lives in the vascular system (Huang and De Cleene 1989) thus enhanced expression in membranes of these cells could lead to infection. In fact, *Xa13* protein localizes to the plasma membrane in callus bombardment experiments and GUS staining shows it is expressed in parenchyma cells surrounding the vascular cells of leaves. It is therefore in a perfect position to enhance bacterial invasion.

### *Xa13* is essential during pollen development

A question that immediately comes to mind is: Why is *Xa13* maintained if it makes the plant more susceptible? Interestingly, in the absence of pathogen infection, both alleles are expressed at low levels in leaves but at high levels in panicles (flower-bearing structures) and anthers, the male reproductive structures. Examination of the RNAi plants showed a loss of fertility that was correlated with loss of *Xa13* expression (Fig. 1). The finding that *Xa13* is required for fertility explains why expression of the dominant allele has been maintained. It also highlights the selective advantage of the recessive *xa13* alleles, in which high levels of expression exists in panicles, but no pathogen induction occurs.

An understanding of the fertility defect in *Xa13* RNAi plants requires some understanding of plant sex. Anthers produce pollen, which contain sperm cells, and carpels produce embryo sacs, which contain the egg cell. Meiosis occurs in both anthers and carpels and is followed by mitotic divisions for both the microspore (male spore) and megaspore (female spore). The unicellular microspore divides asymmetrically to produce a generative cell within the cytoplasm of the vegetative cell. The gen-

erative cell undergoes a second round of division to produce a tricellular cell containing two sperm cells and the vegetative cell (Fig. 1). One sperm cell will fertilize the egg and the other will fertilize the central cell, thus producing the endosperm. The vegetative cell produces the pollen tube for fertilization (McCormick 2004; Ma et al. 2005).

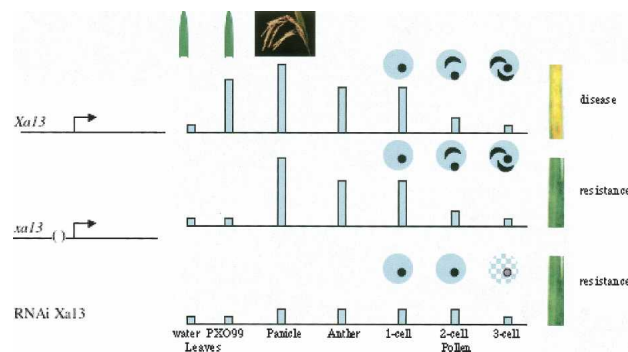
Possible clues for a role for *Xa13* in fertility come from the timing of the defect in *Xa13* RNAi plants and in situ expression patterns. Pollen from RNAi plants looks normal at the unicellular stage but often fails to progress beyond this stage. At a time when tricellular pollen is visible in controls, RNAi lines show unicellular microspores or degenerating pollen grains. Thus, it appears that *Xa13* is needed for the mitotic divisions of the pollen. Other male sterile mutants have been described that show normal meiosis but are blocked during microspore mitotic divisions (Ma et al. 2005). Similar phenotypes have also been found in plants that carry chromosomal defects (Kindiger et al. 1991). It will be important to determine whether *Xa13* is also expressed at high levels in carpels and whether there is any lack of fertility in female transmission.

The timing of expression fits with a requirement for *Xa13* to progress through pollen development. *Xa13* is abundant in the tapetum, the sporophytic nutritive tissue that surrounds the developing unicellular microspore, and in the unicellular pollen. By the time the pollen is tricellular, expression has disappeared except for some trace in the vascular tissue. Thus, *Xa13* transcripts appear at stages before the defect is visible. Although this study describes the first disease-resistance gene that is also required for pollen development, other examples exist of genes expressed during pollination or fertilization that are also up-regulated by stress (Lan et al. 2005).

### Rice disease-resistance genes

*xa13* belongs to a broad family of plant genes known as resistance (R) genes. R genes have been identified from a variety of plant species and provide resistance to all groups of pests and pathogens (insects, bacteria, fungi, viruses) (Kaloshian 2004). Of the ~30 *Xoo* targeting R genes identified in rice (Makino et al. 2006), two-thirds function as dominant resistance alleles and the others provide resistance when recessive, similar to *xa13*. Of the five *Xoo* R genes that were previously cloned, only resistance conferred by the dominant *Xa27* gene is controlled at the level of transcriptional regulation (Gu et al. 2005). Like *xa13*, *Xa27* and *xa27* encode identical proteins but have differences in the promoter. *Xa27* encodes a protein of 113 amino acids that does not resemble any other R gene and has no significant sequence similarity to any proteins of known function. In contrast to *Xa13*, in which the susceptible allele is induced by pathogen infection, the resistant allele of *Xa27* is expressed after challenge with the pathogen.

Recessively inherited resistance does not always follow the pattern established with *xa13* and *mlo*; i.e., resistance by avoidance of pathogen. For example, *xa5*, the first *Xoo* targeting recessive R allele to be cloned, is constitutively expressed at the same level as the dominant



**Figure 1.** *Xa13* expression and consequence. On the left is a schematic of the difference between *xa13* and *Xa13* promoters. In the middle is a representation of expression levels. *Xa13* leaves (green ovals) that are treated with PX099 pathogenic bacteria express high levels of transcript compared with water control and *xa13* leaves. The expression of *Xa13* upon infection leads to disease as shown by a chlorotic (yellow) leaf at the far right. Transcript levels are high in panicles (shown in photo) and anthers and decrease during stages of pollen development (represented as blue circles that progress from unicellular to bicellular to tricellular). RNAi plants have increased resistance but also increased sterility due to a failure to progress from the unicellular stage of pollen.

susceptible allele, *Xa5*, and neither allele is induced or suppressed in response to Xoo (Iyer and McCouch 2004; Jiang et al. 2006). *xa5* was identified as a natural mutation conserved among different varieties of rice and encodes the  $\gamma$  subunit of transcription factor IIA (TFIIA $\gamma$ ). The proteins encoded by the recessive and dominant alleles differ by a single amino acid change from valine to glutamic acid, representing a change from hydrophobic to hydrophilic at that location. Jiang et al. (2006) suggest that this change disrupts a protein-protein interaction necessary for disease.

Perhaps the most striking aspect of the *xa13* study is the unusual connection between disease resistance and pollen development. Although these topics are generally thought to be distinct, other examples exist where development and disease intersect. For example, resistance controlled by *Xa21*, which encodes a receptor-like kinase with a transmembrane domain and a leucine-rich repeat (RLK-LRR) extracellular domain (Mazzola et al. 1994; Song et al. 1995; Wang et al. 1998) is developmentally regulated. Century et al. (1999) showed that the *Xa21* transcript is expressed at a constant low level throughout development and is not induced by pathogen infection; however, disease resistance increases with age in *Xa21*-expressing plants. Interestingly, the stability of the *XA21* protein itself is developmentally regulated, and is associated with proteolytic activity during development. Similarly, Chern et al. (2005) showed that overexpression of the rice gene *NRR*, a negative regulator of resistance, affects both resistance and development. As a rice plant matures, basal levels of resistance increase (Koch and Mew 1991). In plants overexpressing *NRR*, this age-related resistance was compromised. This decrease in age-related resistance correlated with a decrease in fertility. Taken together, these studies show that development and disease resistance are often intimately connected.

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