Lysine is considered an ‘essential amino acid’ required at sufficient levels to prevent malnutrition and serious diseases, particularly in developing countries. It is mostly obtained from various plant foods. In the current issue (pages 4285–4296) Yang and colleagues report on successful stimulation of lysine biosynthesis and suppression of its catabolism in transgenic rice plants without changing the plant phenotype. This approach led to the production of high-lysine rice plants, rendering them as nutritionally favourable crops.

Lysine is considered the first limiting essential amino acid in cereal and legume crops – i.e. it is present in the smallest quantity (Galili et al., 1994). This restricted content significantly reduces the nutritional values of these crops to 50–70% compared with those containing more balanced levels (Galili and Amir, 2013). As an ‘essential amino acid’, not produced in the bodies of humans and farm animals, lysine must be obtained from other sources. It is quite extensively present in livestock-derived foods, such as meat, eggs and cheese. However, where diets rely on plant-derived foods – which is the case for huge populations living in poverty in developing countries – people suffer from insufficient lysine levels, leading to vulnerability to disease, decreased levels of blood proteins and retarded mental and physical development in young children (Galili and Amir, 2013). To prevent this, enriching the content of lysine in those crop plants which serve as the major sources of human foods and livestock feed in these countries is essential. Among the cereal crops, rice (Oryza sativa) is a stable source of calories and protein intake for approximately one-third of the world’s population (Kusano et al., 2015).

Lysine metabolism in plants has been studied for over 50 years, since the discovery of the maize high-lysine mutant opaque-2 (o2), which contains low levels of lysine-poor seed storage proteins (zeins) and consequently an increased lysine and tryptophan content compared with the wild type (Mertz et al., 1964). However, enrichment of lysine levels in crops using classical genetics and breeding approaches is difficult because: (i) lysine synthesis is highly negatively regulated by a feedback inhibition loop in which lysine feedback inhibits the activity of dihydrodipicolinate synthase (DHPS), the first enzyme of the lysine biosynthesis pathway, slowing down its synthesis (Box 1); and (ii) lysine is efficiently degraded by its catabolism into the tricarboxylic (TCA) cycle, a pathway initiated by the bi-functional enzyme LKR/SDH (Box 1), which exhibits both lysine-ketoglutarate reductase (LKR) and saccharopine dehydrogenase (SDH) activities (Karchi et al., 1995). Nevertheless, it is by careful engineering of both of the above enzymes (DHPS and LKR/SDH) that Yang et al. (2016) have now developed two pyramid transgenic lines with free lysine content elevated to 25-fold compared to wild type without changing the plant phenotype. This is an important achievement.

Enhancing lysine levels

A recent study reported the lysine biofortification of seeds of rice by overexpressing genes encoding two endogenous rice histone proteins that are naturally enriched in lysine levels. Expression was achieved using a modified rice glutelin 1 promoter, which is specifically expressed at a relatively low level in developing seeds where the glutelin proteins are synthesized. The choice of a weak seed-specific promoter was intended to prevent physiological problems, such as germination efficiency and vegetative growth, enabling an enhancement of lysine level by a meaningful (up to 35%) amount with no negative impact on the levels of other amino acids. This increase could also meet the nutritional standards of the World Health Organization, offering improved solutions for consumption of human foods and livestock feeds (Wong et al., 2015).

A different approach to that taken by Wong et al. (2015) is through metabolic engineering. A number of the earliest of these studies aiming to improve lysine content in plants concentrated on expressing lysine feedback-insensitive DHPS enzymes of lysine biosynthesis under the control of constitutive or seed-specific promoters, so important for cereals where the intention is fortification for human consumption (Karchi et al., 1994; Falco et al., 1995). Expression in soybean and canola seeds led to a notable increase of seed free lysine (Falco et al., 1995). Among these early studies, Keeler et al. (1997) also indicated that de novo expression of alpha-helical coiled-coil proteins might also increase the accumulation of lysine in mature tobacco seeds.
Elevating lysine content by enhancing its synthesis is one way of improving the nutritional quality of crops, but another useful approach is to prevent degradation (catabolism) into the TCA cycle. A third way forward is to combine these in the same crop plant. For example, Zhu and Galili (2004) further expressed the bacterial DHPS (with embryo-specific promoter) in an Arabidopsis knockout mutant lacking LKR/SDH and therefore getting round the problem of lysine catabolism. This approach caused a significant, 64-fold increase in lysine level in the seeds, but was also associated with a prominent reduction of germination rate, indicating that more moderate enrichments of lysine levels would be likely to be more successful. Hence, the balance of synthesis and degradation of lysine is a key regulatory point not only for controlling lysine level, but also improvement of the nutritional quality of crops, and similar findings were also previously reported in maize (Frizzi et al., 2008).

Getting the balance right

Enhancing the synthesis or reducing the catabolism of lysine can elevate lysine content levels in plants, but lysine overproduction results in increased lysine degradation. Therefore, how to balance the synthesis and catabolism of lysine? It’s the key point of control for lysine levels.

Long et al. (2013) showed that the level of LKR/SDH was significantly enhanced in the developing seeds of rice Asp kinase (AK)/DHPS overexpression lines. This means that the effect of transgene AK/DHPS was counterbalanced by the activity of lysine catabolism in sustaining a steady level of lysine. Overexpression of AK and DHPS only increased the free lysine level by 1.1-fold compared with the wild type; however, the LKR-RNAi line showed a 10-fold increase, while combined expression of AK and DHPS and interference of LKR/SDH to achieve both metabolic effects meant a substantial increase in the free lysine content to 60-fold. Similar findings were previously reported in Arabidopsis (Zhu and Galili, 2004) and maize (Frizzi et al., 2008).

Taking this one step further, Yang et al. (2016) show a perfect example of meaningful enhancement of lysine levels in transgenic rice by using a combined enhancement of lysine synthesis and suppression of its catabolism. However, in previous studies, the high lysine transgenic plants were typically accompanied by obvious changes of seed phenotype, including oil content, germination and yield (Shaul and Galili, 1993; Tzchori et al., 1996; Zhu and Galili, 2003; Angelovici et al., 2011). In the present study, there was no difference in major agronomic traits, including yield. Furthermore, the promoter used enabled specific accumulation of lysine in the seeds, the main organs consumed in human foods and livestock feeds.

The potential is there for this to solve major problems of malnutrition worldwide, but particularly in developing countries. So far, a maize cultivar, LY038, developed by genetic engineering to have a high lysine content, has been approved for commercial use in a number of countries, including in Africa (Dizigan et al., 2007). The findings of Yang et al. (2016) are similarly favourable to commercialization. Meanwhile, there is strong public debate regarding the safety of genetically modified plants and, as with all other GM crops, the opportunities for use of GM rice with enhanced lysine will depend on public acceptance.

Key words: Aspartate kinase (AK), dihydrodipicolinate synthase (DHPS), lysine, lysine ketoglutaric acid reductase/saccharopine dehydrogenase (LKR/SDH), rice (Oryza sativa).
Endogenous auxin biosynthesis and de novo root organogenesis

Ya Lin Sang, Zhi Juan Cheng and Xian Sheng Zhang*

State Key Laboratory of Crop Biology, College of Life Sciences, College of Forestry, Shandong Agricultural University, Taian, Shandong 271018, China

* Correspondence: zhangxs@sdau.edu.cn

Induction of adventitious roots is essential for vegetative propagation of plants, and auxin has long been used as an exogenous root-inducing agent. In this issue of Journal of Experimental Botany, Chen et al. (pages 4273–4284) demonstrate that different members of the YUCCA family orchestrate the endogenous auxin biosynthesis that is required for the induction of adventitious roots.

Sun Wukong, also known as the Monkey King, is the main character in the classical Chinese novel Journey to the West. As a fabled deity, he was endowed with magical properties allowing each of his hairs to be transformed into clones of himself as needed. Plants also possess the remarkable ability of multiplication, with detached pieces of adult tissues capable of forming an entire plant body (Gordon et al., 2007; Birnbaum and Sánchez Alvarado, 2008; Sugimoto et al., 2010). This unique ability is mainly based on de novo organogenesis, in which adventitious shoots or roots are generated from isolated tissues or organs (Duclercq et al., 2011; Cheng et al., 2013; Xu and Huang, 2014).

De novo organogenesis can be induced under both natural and tissue culture conditions (Chen et al., 2014). Plant organs, such as stems and leaves, give rise to adventitious roots under natural growth conditions and this property has long been used for vegetative propagation of elite genotypes in agriculture, forestry and horticulture (De Klerk et al., 1999). Six decades ago, Skoog and Miller demonstrated that the entire plant could be regenerated by tissue culture (Skoog and Miller, 1957). They showed that culturing explants in medium containing high levels of cytokinin induced the formation of adventitious shoots, whereas medium with high levels of auxin triggered initiation of adventitious roots. This classic system laid the foundations for plant micropropagation and genetic transformation (Duclercq et al., 2011; Li et al., 2011). In both cases, a key step ensuring the success of plant regeneration is de novo root organogenesis.

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