Journal of Zhejiang University-SCIENCE B (Biomedicine & Biotechnology) ISSN 1673-1581 (Print); ISSN 1862-1783 (Online) www.jzus.zju.edu.cn; www.springerlink.com E-mail: jzus@zju.edu.cn

## **Review:**



## Breeding for low cadmium accumulation cereals<sup>\*</sup>

Qin CHEN, Fei-bo WU<sup>†‡</sup>

Department of Agronomy, College of Agriculture and Biotechnology, Zhejiang University, Hangzhou 310058, China <sup>†</sup>E-mail: wufeibo@zju.edu.cn

Received Sept. 30, 2019; Revision accepted Jan. 15, 2020; Crosschecked May 25, 2020

**Abstract:** Cadmium (Cd) is an element that is nonessential and extremely toxic to both plants and human beings. Soil contaminated with Cd has adverse impacts on crop yields and threatens human health via the food chain. Cultivation of low-Cd cultivars has been of particular interest and is one of the most cost-effective and promising approaches to minimize human dietary intake of Cd. Low-Cd crop cultivars should meet particular criteria, including acceptable yield and quality, and their edible parts should have Cd concentrations below maximum permissible concentrations for safe consumption, even when grown in Cd-contaminated soil. Several low-Cd cereal cultivars and genotypes have been developed worldwide through cultivar screening and conventional breeding. Molecular markers are powerful in facilitating the selection of low-Cd cereal cultivars. Modern molecular breeding technologies may have great potential in breeding programs for the development of low-Cd cultivars, especially when coupled with conventional breeding. In this review, we provide a synthesis of low-Cd cereal breeding.

Key words: Cereals; Low Cd accumulation; Gene/quantitative trait locus (QTL) mapping; Breeding https://doi.org/10.1631/jzus.B1900576 CLC number: S184

#### 1 Introduction

Soil cadmium (Cd) contamination poses a serious problem for safe food production and has become a potential agricultural and environmental hazard worldwide. Cd contamination may be caused by industrial emissions or applications of Cd-containing sewage sludge, phosphate fertilizers, or municipal waste (Čásová et al., 2009; Roberts, 2014; Liu et al., 2015; Yousaf et al., 2016). Particularly, soils derived from marine shales or contaminated by mine waste are likely to have a high Cd content (Arunakumara et al., 2013). Cd has high soil-to-plant mobility and can easily accumulate in plant tissues (Song et al., 2015), causing reduced crop yields and threatening human health via the food chain (Clemens et al., 2013; Sun et al., 2013). In humans, Cd accumulates mainly in the kidneys and has a biological half-life of about 20 years (Clemens et al., 2013; Aziz et al., 2015). This very slow elimination of accumulated Cd can lead to a variety of serious health issues including anemia, cancer, cardiovascular disease, and renal tubular damage (Satarug et al., 2003, 2009). Extreme cases of chronic Cd toxicity can result in osteomalacia and bone fractures, such as the "Itai-Itai" disease that occurred in Japan during the 1950s and 1960s (Huang et al., 2009). As a result of these and other illnesses, Cd contamination in soils has been of increasing concern for decades in many industrialized and developing countries (Arthur et al., 2000; Wu et al., 2004; Cao et al., 2014a).

Soil Cd contamination poses considerable risks to human health via the consumption of foods containing high Cd concentrations, especially staple foods such as cereals (Stolt et al., 2003). For example, rice

442

<sup>&</sup>lt;sup>‡</sup> Corresponding author

<sup>\*</sup> Project supported by the Key Research Foundation of Science and Technology Department of Zhejiang Province of China (No. 2016C02050-9-7)

ORCID: Fei-bo WU, https://orcid.org/0000-0001-8797-3398; Qin CHEN, https://orcid.org/0000-0002-3964-321X

<sup>©</sup> Zhejiang University and Springer-Verlag GmbH Germany, part of Springer Nature 2020

has been a major dietary source of Cd for the Japanese population (Tsukahara et al., 2003), contributing about 30% of the total dietary Cd intake in Japan during the 1990s (Ikeda et al., 1999). Dabeka et al. (1987) found that the Cd content of cereals was the highest among ten classified food groups in Canada. Grain Cd concentrations as high as 1-2 mg/kg dry weight (DW) have been recorded in rice grown in some areas of central-southern and southwestern China (Du et al., 2013; Chen et al., 2018). Those levels are much higher than the maximum permissible concentration (MPC) for cereal grains as defined by the Chinese National Standard (NHFPC, 2017) and the Codex Alimentarius Commission (CAC, 2019). According to national average estimates of exposure, cereals contribute about 32% of the total Cd intake by the Chinese population (WHO, 2011). Therefore, it is imperative to reduce Cd accumulation in cereal grains so as to control Cd intake by human beings.

The development and planting of low-Cd cultivars is a cost-effective and environmentally friendly approach to minimize soil-plant transfer of Cd and produce safe food for slightly or moderately contaminated soils. For over a decade, breeding programs have been initiated to select for low-Cd crop cultivars (Oliver et al., 1995). Many studies have been conducted to screen and develop low-Cd cultivars in a variety of crop species, including rice, barley, maize, and wheat (Oliver et al., 1995; Wu and Zhang, 2002; Yu et al., 2006; Chen et al., 2007a, 2007b; Wu et al., 2007; Grant et al., 2008; Zeng et al., 2008). In this review, we focus on recent advances in selecting low-Cd cereal crops, including: (1) genes related to Cd accumulation; (2) quantitative trait loci (QTLs) associated with Cd accumulation; and (3) breeding low-Cd cultivars.

## 2 Genes associated with Cd accumulation in cereal crops

For developing low-Cd crop cultivars, it is important to identify genes related to Cd accumulation and to understand the mechanisms of Cd uptake and accumulation in plants. In recent years, a number of genes related to Cd transport in cereals have been identified (Table 1), and progress has been made in understanding the mechanisms of Cd uptake and transport (Chen JG et al., 2019). Cd is not essential for plant growth, but its transport in plants is mediated by the transporters for essential elements such as Zn, Mn, Ca, and Fe (Uraguchi and Fujiwara, 2013; Clemens and Ma, 2016). Some genes related to chelation have also been found to play roles in plant Cd homeostasis.

#### 2.1 Transporter-related genes

In plants, natural resistance-associated macrophage proteins (NRAMPs) are a family of metal transporters that are integral components of membranes and play important roles in metal uptake, translocation, and intracellular transport (Nevo and Nelson, 2006; Sasaki et al., 2012). OsNramp5, a Mn transporter located at the plasma membrane of root cells, was found to be a major transporter of Cd uptake in rice roots exposed to Cd-containing soil medium. Knockout or knockdown of OsNramp5 resulted in more than a 90% reduction in Cd uptake (Sasaki et al., 2012; Yang M et al., 2014), thereby drastically reducing the Cd content of shoots and grains (Ishimaru et al., 2012; Tang et al., 2017). Transfer DNA (T-DNA) insertion mutants of OsNramp5 led to large yield reductions of about 11% compared with wild-type plants (Sasaki et al., 2012), while OsNramp5 knockout lines developed by ion-beam irradiation (Ishikawa et al., 2012) or by the clustered regularly interspaced short palindromic repeat (CRISPR)/CRISPR-associated protein 9 (Cas9) system (Tang et al., 2017) did not affect grain yield. Orthologues of Nramp5 in other cereal crops have a similar function. TpNramp5 in dwarf Polish wheat functions as a metal transporter for Cd, Mn, and Co. When this gene was expressed in Arabidopsis, it significantly increased Cd, Co, and Mn concentrations in roots, shoots, or the whole plant (Peng et al., 2018b). Similarly, knockdown of the orthologue in barley, HvNramp5, which is preferentially expressed in the outer root cell layers and root tips, resulted in significant reductions of Mn and Cd concentrations in both roots and shoots (Wu et al., 2016), but its relative contribution appears to be smaller than that of OsNramp5 in rice. Comparison of the expression levels and transport activities of three Nramp5 orthologues from rice, wheat, and maize, showed that OsNramp5 in rice was expressed at higher levels, resulting in higher Cd transport activity than Nramp5 in wheat or maize (Sui et al., 2018). Other NRAMP members in cereal crops also show Cd

Gene	Gene name	Probable function	Reference
Oryza sativa <sup>1</sup>			
OsNramp1	Natural resistance-associated	Cd and Fe transporters	Takahashi et al., 2011a, 2011b
OsNramp5	macrophage protein	Cd, Mn, and Fe transporters	Ishikawa et al., 2012; Sasaki et al., 2012; Yang M et al., 2014; Tang et al., 2017
OsIRT1	Zinc- and iron-regulated transporter	Cd and Fe transporters	Nakanishi et al., 2006; Lee and An, 2009
OsIRT2			Nakanishi et al., 2006
OsZIP1		Cd and Zn transporters	Bashir et al., 2012; Ramegowda et al., 2013; Liu XS et al., 2019
OsZIP7		Cd and Zn accumulation	Ricachenevsky et al., 2018; Tan et al., 2019
OsHMA2	P-type heavy metal ATPase	Cd and Zn translocation	Satoh-Nagasawa et al., 2012; Takahashi et al., 2012; Yamaji et al., 2013
OsHMA3		Sequestration of Cd in root	Ueno et al., 2010; Miyadate et al., 2011; Sasaki et al., 2014; Yan et al., 2016; Shao et al., 2018; Lu et al., 2019
OsHMA9		Cd efflux	Lee et al., 2007
OsLCD	Low cadmium	Cd tolerance and accumulation	Shimo et al., 2011
OsCd1	Major facilitator superfamily	Cd uptake	Yan et al., 2019
OsLCT1	Low affinity cation transporter	Cd transporter in phloem	Uraguchi et al., 2011, 2014
OsCCX2	Cation/Ca <sup>2+</sup> exchanger 2	Cd tolerance and translocation	Hao et al., 2018
OsMTP1	Metal tolerance protein gene	Cd translocation	Yuan et al., 2012
OsPCS2	Plant chelatase synthase 2	Cd tolerance and accumulation	Das et al., 2017
OsPCS1	Plant chelatase synthase 1	Cd tolerance and accumulation	Uraguchi et al., 2017
OsMTI-1b	Metallothionein-like protein 1b	Cd tolerance and accumulation	Ansarypour and Shahpiri, 2017
CAL1	Defensin-like protein	Cd accumulation in leaf	Luo et al., 2018
Triticum aestiv	ит		
TaHMA2	P-type heavy metal ATPase	Cd and Zn transporters	Tan et al., 2013
HMA3-B1		Cd accumulation in grains	Wiebe et al., 2010
Triticum polon	icum		
TpNRAMP3	Natural resistance-associated	Cd, Mn, and Co transporters	Peng et al., 2018a
TpNRAMP5	macrophage protein	Cd, Mn, and Co transporters	Peng et al., 2018b
Zea mays			
GRMZM2G 175576	Homologous to rice OsHMA2 and OsHMA3	Cd accumulation	Zhao XW et al., 2018
Hordeum vulga	are		
HvIRT1	Zinc-and iron-regulated	Fe, Mn, Zn, and Cd transporters	Pedas et al., 2008
HvZIP3	transporter	Mn, Zn, and Cd accumulation	Sun et al., 2015
HvZIP8		Mn, Zn, and Cd accumulation	Sun et al., 2015
HvHMA2	P-type heavy metal ATPase	Zn, Fe, and Cd transporters	Barabasz et al., 2013
HvHMA3		Cd accumulation	Wu et al., 2015
HvNramp5	Natural resistance-associated macrophage protein	Mn and Cd accumulation in roots and shoots	Wu et al., 2016

Table 1 List of reported genes of cereal crops related to Cd homeostasis

<sup>1</sup> List of reported genes in rice (Oryza sativa) related to Cd homeostasis was reviewed by Chen JG et al. (2019)

uptake and transfer activities. The plasma membranelocalized iron transporter, OsNRAMP1, may assist in loading Cd into the xylem. Thus, over-expression of *OsNRAMP1* in rice increased Cd accumulation in leaves while reducing Cd levels in roots (Takahashi et al., 2011a). *TpNramp3* from dwarf Polish wheat has a similar function to *TpNramp5*, and is also a transporter for Cd, Mn, and Co. Heterologous expression of *TpNramp3* in *Arabidopsis* resulted in increased Cd, Co, and Mn concentrations in roots, shoots, and whole plants, but did not affect their translocation from roots to shoots (Peng et al., 2018a).

In addition to the NRAMP family, several members of the zinc/iron-regulated transporter-like proteins (ZIPs) family, the primary zinc and iron transporters in plants, are also involved in Cd uptake. Some Fe transporters have been found to participate in plant Cd uptake. Nakanishi et al. (2006) found that OsIRT1 and OsIRT2 play important roles in Cd uptake. Over-expression of OsIRT1 resulted in increased Cd accumulation (Lee and An, 2009). In Arabidopsis, over-expression of AtIRT1 and AtIRT2 also caused an increase in Cd accumulation (Connolly et al., 2002; Vert et al., 2009). Similarly, HvIRT1, a plasma membrane-localized transporter for Fe, Mn, and Zn, also exhibits Cd transport activity when expressed in yeast (Pedas et al., 2008). Moreover, OsZIP1 has long been considered a metal uptake transporter for Zn and Cd in rice (Bashir et al., 2012; Ramegowda et al., 2013). Recent studies showed that over-expression of OsZIP1 reduced concentrations of Zn, Cu, and Cd in rice and improved growth under metal stress (Liu XS et al., 2019). The function of OsZIP1 is similar to that of the copper efflux transporter, OsHMA9, in which it has a Cd efflux function, causing an excretion of Cd from root cells and a reduction of Cd accumulation in rice (Lee et al., 2007). OsZIP7 encodes a plasma membrane-localized protein with influx transport activity for both Zn and Cd, and plays an integral role in xylem loading in roots and inter-vascular transfer in nodes to preferentially deliver Zn and Cd to developing tissues and grains. Thus, knockout of OsZIP7 resulted in the retention of Zn and Cd in roots and basal nodes (Ricachenevsky et al., 2018; Tan et al., 2019). Suppression of the zinc transporter genes HvZIP3 and HvZIP8 by RNA interference (RNAi) silencing showed increased Cd accumulation and reduced Zn and Mn concentrations in barley grains (Sun et al., 2015). However, considering that the metal concentrations used in some studies were much higher than those found in natural soils or rhizosphere environments, these conclusions should be viewed with caution. For example, the RIT1 gene cloned from pea (Pisum sativum L.), a homolog of the Arabidopsis IRT1 Fe transporter gene, was previously thought to be associated with Fe uptake, and induced a high rate of  $Cd^{2+}$  and  $Zn^{2+}$  influx into the roots of pea seedlings (Cohen et al., 1998). However, further research showed that the enhancing effect of RIT1 on Zn and Cd existed only when these elements were present at elevated concentrations, but not at their physiologically relevant soil levels (Cohen et al., 2004).

P<sub>1B</sub>-ATPases, also called heavy metal ATPases (HMAs), play direct roles in plant heavy metal transmembrane transport. OsHMA2, a transporter for Zn and Cd, is localized in the plasma membrane and mediates Zn/Cd xylem loading and intervascular transport to grains. However, studies showed that both OsHMA2 knockout and overexpression plants had considerably reduced concentrations of Zn and Cd in leaves and grains compared with wild-type rice (Satoh-Nagasawa et al., 2012; Takahashi et al., 2012; Yamaji et al., 2013). HMA2 genes have also been identified in other crops and have a highly conserved function among cereals. For example, barley HvHMA2 has been identified predominantly in the plasma membrane and has a role in pumping Zn and Cd in yeast (Mills et al., 2012). Ectopic expression of HvHMA2 in tobacco resulted in modified Zn, Fe, and Cd homeostasis (Barabasz et al., 2013). TaHMA2 from bread wheat can transport  $Zn^{2+}$  and  $Cd^{2+}$  across membranes, and overexpression of TaHMA2 in Arabidopsis increased Zn/Cd root-to-shoot translocation, and enhanced root length and fresh weight (Tan et al., 2013). In contrast, OsHMA3 located in the tonoplast, plays a role in pumping Cd into vacuoles and has a decisive role in reducing the root-to-shoot translocation of Cd (Ueno et al., 2010; Miyadate et al., 2011). HvHMA3 has been identified as a candidate gene for Cd accumulation in barley shoots and grains (Wu et al., 2015). In durum wheat, the HMA3-B1 gene has been found to match perfectly with a major locus *Cdu1*, which can explain 80% of the variation in grain Cd accumulation (Knox et al., 2009; Zimmerl et al., 2014). Sequence analysis of this *HMA3* gene in highand low-Cd durum wheat cultivars showed that a premature stop caused by a 17-bp duplication occurred only in high-Cd cultivars. The close relationship between a severely truncated HMA3 transporter and high Cd phenotype further confirmed that *HMA3* was the best candidate gene for the *Cdu1* locus (Wiebe, 2012; Aprile et al., 2018).

OsHMA3 has high specificity for Cd rather than broad substrate-specificity. Consequently, loss-offunction or weak alleles of this gene result in increasing Cd root-to-shoot translocation (Ueno et al., 2010; Yan et al., 2016). In field trials, overexpression of OsHMA3 decreased Cd concentrations in brown rice by more than 90%, but had no significant effect on grain yield or concentrations of other essential micronutrients, including Zn, Fe, Cu, and Mn (Sasaki et al., 2014; Lu et al., 2019). Given the fact that OsHMA3 and OsHMA2 have different tissue localizations and promoter activities, driving the expression of OsHMA3 by OsHMA2 promoter could specifically reduce Cd accumulation in grains, and mitigate the adverse effect on rice yield induced by OsHMA2 knockout as well (Shao et al., 2018). These mutants might be used directly in breeding programs.

Recently, nodes in graminaceous plants such as rice and barley were found to play crucial roles in the allocation of multiple mineral nutrients, including essential and nonessential metal elements, by mediating intervascular transfers (Yamaji and Ma, 2014). Several node-expressed transporters, mainly in rice, have been found to transport Cd, and among them OsHMA2 is expressed in the phloem of nodes (Satoh-Nagasawa et al., 2012; Takahashi et al., 2012; Yamaji et al., 2013). In addition, the plasma membranelocalized gene, low-affinity cation transporter 1 of rice (OsLCT1), is highly expressed in the uppermost node, and has been shown to mediate the export of phloem Cd and certain other elements into the sieve tube and translocate them from leaf blades and nodes to the grains of rice. Thus, down-regulation of OsLCT1 resulted in an about 50% reduction of Cd in grains compared with that of control plants, whereas the contents of other metals and plant growth were little affected (Uraguchi et al., 2011, 2014). In addition, OsCCX2, a putative cation/Ca<sup>2+</sup> exchanger (CCX),

may function in loading Cd into xylem vessels, thus mediating direct root-derived Cd transport to grains. Knockout of *OsCCX2* caused a significant reduction of Cd content in rice grains (Hao et al., 2018). Shimo et al. (2011) found a novel gene, *OsLCD* (low Cd), which positively regulates Cd tolerance and accumulation in rice. A knockout via a T-DNA insertion reduced Cd accumulation by 43%–55% in grains, but had no effect on Cd accumulation in leaf blades. Yuan et al. (2012) reported that down-regulation of metal tolerance protein 1 of rice (*OsMTP1*, a bivalent cation transporter of heavy metals including Zn and Cd) markedly reduced heavy metal tolerance and accumulation in various rice tissues.

#### 2.2 Chelation-related genes

In addition to transporter proteins, metal ligands and chelators including metallothioneins (MTs), phytochelatins (PCs), organic acids, and other cysteine (Cys)-rich peptides have a role in metal transport and homeostasis (Ovečka and Takáč, 2014). MTs belong to a superfamily of intracellular metal-binding proteins, and bind metals through their Cys residues. The isoform, OsMTI-1b, a rice MT type 1 gene, when expressed in yeast cells, results in increased tolerance to CdCl<sub>2</sub> treatment and increases accumulation of Cd<sup>2+</sup> ions (Ansarypour and Shahpiri, 2017). PCs are predominantly associated with detoxification of nonessential toxic metals and metalloids such as Cd and arsenic (As), and thus significantly affect Cd accumulation in cereal plants (Das et al., 2017). Uraguchi et al. (2017) identified two independent OsPCS1 mutant rice lines (a T-DNA and a Tos17 insertion line) and found that they exhibited increased sensitivity to Cd, showing the importance of OsPCS1-dependent PC synthesis for Cd tolerance in rice. Elemental analyses of rice plants showed decreased Cd accumulation in the grains of both mutants. Recently, a defensin-like protein, CAL1, was found to be involved in Cd accumulation in rice leaves. It promotes Cd secretion into the extracellular space via chelation, then loading into the xylem and long-distance transport. thereby resulting in most Cd being deposited in leaves rather than in grains. This provides a novel and ideal mechanism for developing dual-function rice cultivars that can produce low-Cd grains and be used as Cd hyper-accumulators for remediating contaminated soils (Luo et al., 2018).

#### 3 Identification of molecular markers for low Cd accumulation in cereals

OTL mapping is acknowledged as an efficient approach for dissecting complicated agronomic traits controlled by multiple genes, and has been successfully applied to detect loci controlling Cd accumulation in some crops. A series of QTLs that control Cd concentrations have been reported in various plants including wheat, maize, barley, and rice (Table 2). Cd uptake, translocation, and accumulation in grains are complex processes that are controlled by many genes and are affected by environmental factors. Only a small number of genes related to low Cd accumulation have so far been identified, and the underlying knowledge regarding the molecular mechanisms for plant Cd uptake, translocation, and accumulation is still fragmental. Furthermore, the application of these genes to low-Cd plant breeding is still lacking and has little effect on cultivars released by breeders. The challenge now facing breeders is to identify more novel, effective low-Cd genes, and to successfully apply those already identified to produce low-Cd crop cultivars.

#### 3.1 QTLs associated with Cd accumulation in rice

Ishikawa et al. (2005) detected three putative QTLs controlling Cd concentrations in brown rice located on chromosomes 3, 6, and 8. Kashiwagi et al. (2009) found that the interaction of two putative QTLs on chromosome 4 increased Cd concentrations in brown rice. Xue et al. (2009) identified six QTLs associated with Cd tolerance and three QTLs for root and shoot Cd concentrations in rice seedlings. Ueno et al. (2009b) identified a major QTL on chromosome 11 controlling the translocation of Cd from roots to shoots in seedlings that explained 16.1% of the detected variation in Cd accumulation. Ueno et al. (2009a) isolated another novel major QTL with a large effect on Cd distribution in roots and shoots of rice using an F<sub>2</sub> population derived from Anjana Dhan×Nipponbare. This QTL, detected on the short arm of chromosome 7, explained as much as 85.6% of the phenotypic variation. High-resolution mapping and complementation analysis identified OsHMA3 as being the causal gene for this QTL locus (Ueno et al., 2010). Ishikawa et al. (2010) localized a major-effect QTL (named qGCd7) to the short arm of chromosome

7, which explained 35.5% of the observed variation, and enhanced grain Cd accumulation in rice grown in a Cd-polluted paddy field. Abe et al. (2011) detected a major OTL, *qCdp7* (accounting for 31%–54% of the phenotypic variance), on chromosome 7, that significantly increased Cd accumulation in both the grain and straw of rice. Zhang et al. (2011) identified three QTLs for Cd accumulation in brown rice. Sato et al. (2011) reported two QTLs for increasing Cd accumulation in brown rice: qLCdG11 explained 9.4%-12.9% of the phenotypic variation, while qLCdG3 accounted for 8.3%-13.9%. Yan et al. (2013) constructed a recombinant inbred (F7) line (RIL) population to identify QTLs controlling Cd accumulation and distribution. Five significant QTLs were detected: scc10 was responsible for Cd accumulation in shoots; gcc3, gcc9, and gcc11 were closely associated with Cd accumulation in grains; and sgr5 controlled the translocation of Cd from roots to shoots. Among them, sgr5 had the greatest effect on the distribution of Cd in grains. Using 46 chromosome segment substitution lines (CSSLs), Abe et al. (2013) detected a major QTL, *qlGCd3*, on the long arm of chromosome 3, which was responsible for reduced Cd concentration in rice grains. Using 204 RILs, Hu et al. (2018) identified 22 QTLs affecting the accumulation of Cd, Cu, Fe, Mn, and Zn in brown rice, among which qCd5 was a novel QTL that regulated low Cd accumulation. Luo et al. (2018) identified a QTL named CAL1 (Cd accumulation in leaf 1) on chromosome 2 using a doubled haploid (DH) population derived from TN1 and CJ06. Cd levels in grains and leaves of the over-accumulating cultivar (TN1) were 3-4-fold higher than those in the under-accumulating cultivar (CJ06). The authors localized CAL1 and cloned the causal gene using map-based cloning. Liu WQ et al. (2019) identified two QTLs (qCd-2 and qCd-7) for Cd accumulation in rice grains. McCouch et al. (2016) established an open-access resource for genome-wide association studies (GWAS) in rice. Using this resource, Zhao JL et al. (2018) identified 14 QTLs for low Cd accumulation in rice grains evenly distributed in indica and japonica subpopulations consisting of 312 diverse rice accessions. These QTLs colocalized with functional genes, including OsHMA3, OsNRAMP1, OsNRAMP5, and OsLCD. Also, a novel QTL, qCd3-2, may encode a member of the rice NRAMP gene family (OsNRAMP2). Liu WQ et al.

Crop	Mapping population	Maker	Traits	Chromosome	QTL	PVE (%)	Reference
Rice	39 CSSLs, Kasalath/	129 RFLPs	Cd accumulation in	3, 6, 8	Putative QTLs		Ishikawa
D.	Koshihikari	DELD 1	grains		for grain Cd		et al., 2005
Rice	98 BILs, Kasalath/	RFLP and	Cd concentration in	4, 11	<i>qcd4-1</i> , <i>qcd4-2</i> ,		Kashiwagi
Diag	Nipponbare	SSK DELD and	leaves and culms	1 2 5 6 7	qcall 4 OTL a		et al., 2009
Rice	127 DHS, JAT //24 Q8	SSR	roots and shoots or their ratio	1, 5, 5, 6, 7, 8, 10	4 QILS		2009
Rice	184 F <sub>2</sub> , Badari Dhan/ Shwe War	141 SSRs	Cd concentration in shoots	2, 5, 11	A major QTL	16.1	Ueno et al., 2009b
Rice	177 F <sub>2</sub> , Nipponbare/ Anjana Dhan	SSR	Root-to-shoot Cd translocation	7	A major QTL	85.6	Ueno et al., 2009a
Rice	965 F <sub>2</sub> , Nipponbare/ Anjana Dhan	SSR	Cd concentration in shoots	7	OsHMA3		Ueno et al., 2010
Rice	85 BILs, Sasanishiki/ Habataki	SSR	Cd concentration in shoots and grains	2, 7, 12	qGCd2, qGCd7, qSCd12	7.2–35.5	Ishikawa et al., 2010
Rice	144 F <sub>2</sub> , Cho-Ko-Koku 9 Akita 63	SSR	Root-to-shoot Cd translocation	7	qCdT7	88.0	Tezuka et al., 2010
Rice	F <sub>2</sub> , Cho-Ko-Koku 9 Akita 63	CAPS	Root-to-shoot Cd translocation	7	qCdT7		Miyadate et al., 2011
Rice	103 BILs, Koshihikari/ Jarjan	169 SSRs	Cd concentration in grains	7	qCdp7	31.0-54.0	Abe et al., 2011
Rice	127 DHs, JX17/ZYQ8	RFLP and SSR	Cd concentration in grains	3, 4, 6	qCdc3, qCdc4, qCdc6	10.8-41.7	Zhang et al., 2011
Rice	126 RILs, Fukuhibiki/ LAC23	SSR and CAPS	Cd concentration in grains	3, 11	gLCdG3, gLCdG11	8.3–13.9	Sato et al., 2011
Rice	91 RILs, SNU-SG1/ Suwon490	124 SSRs	Cd concentration in shoots and grains	3, 5, 9, 10, 11	gcc3, srg5, gcc9, gcc11, scc10	16.1–24.9	Yan et al., 2013
Rice	46 CSSLs, Koshihikari/ LAC23	345 SNPs	Cd concentration in shoots	3	glGCd3		Abe et al., 2013
Rice	204 RILs, ZS97B/MY46		Cd concentration in grains	5	qCd5		Huang et al., 2018
Rice	DH, YK17/D50	170 SSRs	Cd concentration in brown or milled rice	2, 3, 4, 5, 7, 9	32 QTLs		Hu et al., 2018
Rice	119 DHs, 3651 BC3F3, Tainan1/Chunjiang06	RFLP	Cd concentration in leaves	2	CAL1	13.1	Luo et al., 2018
Rice	115 RILs, Xiang 743/ Katy	SSR	Cd concentration in grains	2,7	qCd-2, qCd-7		Liu WQ et al., 2019
Wheat	155 DHs, W9262-260D3/ Kofa	SSR	Cd concentration in grains	5B	Cdu1		Knox et al., 2009
Wheat	DH, W9262-260D3/ Kofa	STS	Cd concentration in grains	5B	<i>Cdu1</i> and a minor QTL	80.0	Wiebe et al., 2010
Wheat	103 RILs, Ch/Sh		Cd concentration in roots	4A, 5D	2 QTLs	10.0–17.5	Ci et al., 2012

Table 2 List of reported quantitative trait loci (QTLs) attributing to Cd accumulation in cereal crops

List of reported genes in rice (*Oryza sativa*) related to Cd homeostasis was reviewed by Chen JG et al. (2019). PVE: phenotypic variation explained; CSSL: chromosome segment substitution line; RFLP: restriction fragment length polymorphism; SSR: simple sequence repeat; BIL: backcross inbred line; RIL: recombinant inbred line; DH: doubled haploid; CAPS: cleaved amplified polymorphic sequence; SNP: single nucleotide polymorphism; STS: sequence tagged site

(2019) used 276 accessions with 416000 single nucleotide polymorphisms (SNPs) and performed GWAS of grain Cd concentrations in rice grown in heavily multi-contaminated farmland. Seventeen QTLs were found to be responsible for the grain Cd concentration. Yan et al. (2019) performed GWAS with 127 rice cultivars and identified a QTL on chromosome 3 that explained about 20.7% of the phenotypic variation. Using gene ontology (GO) and yeast spot analyses, they found a member of the major facilitator

superfamily, *OsCd1*, that takes part in rice Cd uptake and grain Cd accumulation and was the causal gene for QTL3. These results provide a good reference for cloning potentially useful candidate genes and molecular breeding of low-Cd cereal crops.

# 3.2 QTLs associated with Cd accumulation in wheat

In contrast to multiple-gene control, durum wheat grain Cd accumulation seems to be controlled

by a single dominant gene, Cdu1, which was first found on a linkage group using the random amplified polymorphic DNA marker (OPC20) (Penner et al., 1995). Later studies reported that its expression was highly heritable (0.84-0.88) and exhibited no significant effect on agronomic performance or the uptake of other micronutrients (Clarke et al., 1997, 2002). Cdu1 is located on the long arm of chromosome 5B (Knox et al., 2009). The same major QTL, which is in the vicinity of Cdu1, has been repeatedly identified with different mapping populations and markers, explaining most of the Cd variation in durum wheat grain (Wiebe et al., 2010; AbuHammad et al., 2016; Salsman et al., 2018). With regard to the QTLs associated with Cd accumulation in vegetative organs in wheat, Ci et al. (2012) identified two QTLs for root Cd accumulation at the germination and seedling stages using 103 RILs derived from a cross of Ch×Sh. AbuHammad et al. (2016) also identified a major QTL contributing a low-Cd uptake allele and explaining 54.3% of phenotypic variation. Oladzad-Abbasabadi et al. (2018) identified a novel low-Cd uptake locus on chromosome 5BL, and the candidate causal gene was homologous to an aluminum-induced protein-encoded gene rather than to the closely linked Cdu1-B gene.

# 3.3 QTLs associated with Cd accumulation in maize and barley

In maize, Soric et al. (2009) detected a QTL associated with Cd uptake on chromosome 2, which could explain 49.8% of the phenotypic variation. Zhao XW et al. (2018) found 63 loci associated with leaf Cd accumulation through GWAS, and successfully identified a major QTL on chromosome 2 that contained 40 SNPs highly associated with leaf Cd concentration, which could explain over 38% of the phenotypic variation. In barley, Wu et al. (2015) detected nine QTLs for root Cd accumulation, 21 for shoot Cd accumulation, and 15 for grain Cd accumulation, via GWAS mapping of 100 accessions from the International Barley Core Selected Collection (BCS). Wang et al. (2019) identified a single QTL, *qShCd7H*, localized on chromosome 7H, responsible for shoot Cd accumulation, using a DH population derived from a cross between Suyinmai 2 (Cdsensitive) and Weisuobuzhi (Cd-tolerant). Furthermore, a novel gene, HvPAA1, related to shoot Cd concentration, was identified from *qShCd7H*. Functional identification showed that *HvPAA1* plays a role in the detoxification of Cd in barley. The results provide a molecular basis for understanding Cd accumulation in barley and will contribute to the development of molecular markers that can be used in breeding for low-Cd-accumulating cultivars.

### 4 Breeding cereal cultivars with low Cd accumulation in grains

Although the use of agronomic practices and chemical regulators can reduce plant Cd uptake, it has been generally accepted that the development of crop cultivars with low Cd accumulation in edible parts is the most efficient approach to deal with medium or slightly Cd polluted farmland. Low-Cd-accumulating cultivars should accumulate Cd in edible parts only to a level below maximum permissible concentrations (MPC) for safe consumption, even when grown in Cd-contaminated soil. The MPCs are set by either the relevant national authorities or regional and international organizations. For example, the Cd standard limit for polished rice set by the Chinese National Standard (NHFPC, 2017) is 0.2 mg/kg DW, and the limit set by the Codex Alimentarius Commission (CAC, 2019) is 0.4 mg/kg DW. In addition to MPCs, further criteria have also been proposed for the selection of low-Cd cultivars. For instance, the yield and eating quality of selected low-Cd cultivars should not be reduced significantly when grown in Cdpolluted soils.

#### 4.1 Methodologies for breeding for low-Cd cultivars

Breeding for low grain Cd cultivars can be achieved by both conventional and modern breeding approaches. Considerable genetic variation in Cdaccumulating ability has been widely reported in a range of cereal crops (Grant et al., 2008; Zhao JL et al., 2018; Zhao XW et al., 2018). Thus, in conventional breeding, low-Cd cultivars can be selected based on the determination of Cd content, coupled with measurements of morphological and physiochemical parameters. To handle segregating accessions, such breeding strategies as mass, pure line, and recurrent selection methods can be effectively adopted in developing low-Cd cultivars. In addition, breeders adopt random cross combinations of genotypes to create genetic variation for screening low Cd accumulation genotypes. However, a more efficient strategy is to screen for low-Cd cultivars by measuring the grain Cd content of cultivars that have been or will be in commercial application (Chen et al., 2007a, 2007b; Cao et al., 2014b). Cultivars selected as low-Cd types from among existing commercial cultivars with excellent yield and agronomic traits can be directly used in commercial production. Conventional breeding methods have been partially successful in developing low-Cd cultivars. For example, a Cd-pollution-safe spring durum wheat cultivar named "Strongfield" was commercially released in Canada in 2004 (Clarke et al., 2006). However, this breeding method typically requires a large amount of effort, and a time-consuming selection process.

Recently, molecular breeding approaches have accelerated progress in breeding low-Cd cultivars by exploiting technologies such as marker-assisted selection, allele discovery, allele pyramiding, genome mutation, genome selection, genome-wide association mapping, and gene editing. The adoption of novel technologies, such as genetic modification, markerassisted selection, and gene editing, has received increasing interest (Randhawa et al., 2013; Oladzad-Abbasabadi et al., 2018). Furthermore, the successful integration of molecular breeding with conventional breeding methods will accelerate the development of low-Cd cultivars.

## 4.2 Genetic differences and screening for low-Cd cereal cultivars—conventional breeding approaches

It is well documented that there are significant differences in Cd accumulation among genotypes. Hence, screening crop cultivars for low Cd content is a practical strategy for the selection of low-Cd cultivars and genotypes. This conventional breeding method has found its place in the development of low-Cd cereals. In rice, Arao and Ae (2003) investigated genotypic variation in grain Cd content, and identified three cultivars with the lowest Cd concentrations. Since then, many researchers have investigated the genotypic and environmental variation in Cd accumulation and distribution in rice and attempted to select low-Cd genotypes (Liu et al., 2005, 2007; He et al., 2006; Yu et al., 2016; Cao et al., 2014b; Pinson et al., 2015; Li et al., 2017;

Zhou et al., 2017). In general, durum wheat tends to accumulate more Cd in grains than bread wheat (Stolt et al., 2003; Greger and Löfstedt, 2004). Wiebe et al. (2010) even suggested that the risk of high grain Cd threatening human health was limited to durum wheat. Thus, more attention and effort have been paid to breeding low-Cd durum wheat than bread wheat. Several dozen low-Cd durum wheat cultivars have been identified worldwide (Eriksson, 1990; Grant et al., 2008; Arduini et al., 2014; Zimmerl et al., 2014; Kubo et al., 2016; Perrier et al., 2016; Hirzel et al., 2017, 2018; Vergine et al., 2017; Oladzad-Abbasabadi et al., 2018; Yue et al., 2018).

Due to the similar chemical properties and electron structures of Zn and Cd (Järup and Åkesson, 2009; Nordberg, 2009), they often present in soils and move into plants via similar uptake and transport pathways (Waters and Sankaran, 2011; Khan et al., 2014). Thus, down-regulation of the pathways to reduce Cd accumulation in cereal grains may at the same time inadvertently decrease Zn transport (Palmgren et al., 2008; Sebastian and Prasad, 2014). Similarly, efforts to remove grain Cd during food processing by leaching and physical separation have reduced the content of some beneficial elements. For example, milling to produce white flour not only reduces Cd concentration by 50.0%, but also leads to reductions of 78.3% in Fe and 69.0% in Zn concentrations (Guttieri et al., 2015a). Fortunately, recent research on grain Cd and Zn contents in hard winter wheat germplasm indicated that their accumulation seems to be independently regulated by different regions of the genome, indicating that it may be possible to breed for low-Cd hard winter wheat genotypes without reducing Zn content (Guttieri et al., 2015b). In barley, our previous research successfully identified low grain-Cd barley genotypes such as Beitalys and Shang 98-128 (Chang et al., 1982; Wu and Zhang, 2002; Chen et al., 2007a, 2007b; Wu et al., 2007). There is also wide genetic variation in Cd content among maize cultivars. Hinesly et al. (1978) first reported large genetic variation in leaf and grain Cd concentrations among maize inbred lines, and found that hybrid maize line B73×R805 had the lowest Cd accumulation in grains when grown on sludge-amended soil (Hinesly et al., 1982). In a study of 19 inbred maize lines, a large genetic variation observed in shoot Cd concentration was attributed to

root-to-shoot translocation rather than to Cd uptake (Florijn and van Beusichem, 1993a, 1993b). From evaluation of Cd concentrations (Zhang et al., 2008; Yang YM et al., 2014; Fahad et al., 2015; Retamal-Salgado et al., 2017), Chunyou30 was identified as a low-Cd maize genotype (Zhang et al., 2008). In short, conventional breeding methods have had partial success in the development of low-Cd cereals.

As they were selected from among commercial cultivars, these low-Cd genotypes can be used either directly or for hybridization to transfer the low Cd accumulation trait (Yu et al., 2006). However, with these breeding methods, breeders often prioritize the selection of low-Cd cultivars at the expense of yield and quality. Hence, some selected low-Cd cereal cultivars often exhibit reduced grain yield and lower nutritional quality (Chen et al., 2007a, 2007b). White and Broadley (2005) and Murgia et al. (2013) noticed a decrease in essential minerals such as Fe, Zn, Mn, and Ca in low-Cd cultivars. Fortunately, there are low-Cd cultivars with normal levels of essential minerals. Luo et al. (2018) evaluated 212 rice accessions, and identified CJ06 as a low-Cd rice cultivar. Further elemental analysis revealed that CJ06 contains levels of Fe, Mn, Zn, and Cu in grains similar to those in high-Cd-accumulating cultivars (i.e., TN1). However, it may be difficult to breed low-Cd cultivars when undesirable characters are linked to low-Cd-related genes, but problems of this type can now be overcome using novel technologies, including gene editing.

### 4.3 Molecular marker-assisted selection for low-Cd cereal cultivars

Molecular marker-assisted selection (MAS) should be a practical and acceptable alternative option. A typical example is the markers for a single major locus, *Cdu1*, related to Cd accumulation in durum wheat. The markers include a random amplified polymorphic DNA (RAPD) marker (OPC-20), an SNP marker which was able to identify 89% of the low-Cd lines in the Strongfield×Alkabo population, and two user-friendly competitive allele-specific polymerase chain reaction (KASP) markers (Cad-5B and Ex\_ c1343\_2570756) for grain Cd accumulation, with an average prediction accuracy of 84%–88%. These markers have assisted the successful selection of low-Cd durum wheat (Knox et al., 2009; Wiebe et al., 2010; AbuHammad et al., 2016; Salsman et al., 2018). Likewise, qShCd7H, a major QTL controlling shoot Cd accumulation in barley, may facilitate the selection of low-Cd barley cultivars in the future (Wang et al., 2019). However, an insufficient number of validated markers have been the primary obstacle hindering the application of MAS to the selection of low-Cd progenies of staple food crops. Fortunately, the abundant germplasm resources of the main cereal crops has made GWAS possible, and with advanced linkage mapping technology, QTL identification and verification has become more powerful (Ueda et al., 2015; Nicod et al., 2016; Zhang et al., 2016). In addition, with the rapid development of next-generation sequencing and RNA sequencing (RNA-seq) profiling, the construction of high-density molecular markers and the quantification of expression profiles under Cd exposure have become possible (Wang et al., 2017; Aprile et al., 2018; Cao et al., 2019). Next generation sequencing is expected to enable the identification of further novel QTLs and molecular markers controlling Cd uptake, root-to-shoot translocation, and grain Cd accumulation.

## 4.4 Application of gene editing to breeding low-Cd cultivars

Progress has been made in the identification and cloning of low-Cd-related genes in the past two decades. For example, AtIRT1 and AtIRT2 (Connolly et al., 2002), OsIRT1 and OsIRT2 (Nakanishi et al., 2006), and OsNRAMP1 (Curie et al., 2000) were identified and found to play roles in Cd uptake. Over-expression of OsNRAMP1 in rice increased Cd accumulation in the leaves, indicating that OsNRAMP1 was an important protein associated with high Cd accumulation in rice (Takahashi et al., 2011a). The knockout of OsNRAMP5 or overexpressing OsHMA3 significantly reduced Cd concentration in roots, shoots, and grains when compared with that of their wild-type relatives (Ueno et al., 2010; Sasaki et al., 2012, 2014; Yang M et al., 2014; Yan et al., 2016; Lu et al., 2019). The down-regulation of OsLCT1 resulted in an about 50% reduction of Cd in grains compared with that of control plants (Uraguchi et al., 2011, 2014). Manipulating the expression of these transporter genes might be an efficient approach to reduce Cd content in grain crops. However, processes involved in the uptake and transport of Cd are closely associated with those

involved in the uptake and transport of essential micronutrients such as Zn, Fe, and Mn (Nakanishi et al., 2006; Sasaki et al., 2012). Thus, there is still a great challenge, as reducing the Cd concentration in grains may simultaneously interfere with the balance of some essential mineral elements. One typical example is knockout mutations of the major transporter responsible for Mn and Cd uptake in rice-OsNramp5. These resulted in a large reduction of Cd in rice grains compared with that of wild-type plants (Ishikawa et al., 2012; Sasaki et al., 2012), but simultaneously, a decline in Mn concentration was detected (Ishikawa et al., 2012). Because it is an essential micronutrient for plant growth and development, Mn deficiency could inhibit growth and reduce yield. Similarly, the T-DNA insertion lines and RNAi lines of OsNRAMP5 within a japonica cultivar Zhonghua 11, exhibited severe growth inhibition and a significant decrease in grain yield (grain yield of the T-DNA insertion mutant was only 11% of that of wild-type plants) (Sasaki et al., 2012). Supplying Mn fertilizer might alleviate Mn deficiency, but this problem may also be solved by inducing different mutation sites or using various genetic backgrounds. A good example is the lcd-kmt mutants of OsNRAMP5 (produced by carbon ionbeam irradiation of a popular Japanese temperate japonica rice cv. Koshihikari), which show few adverse effects on growth with less than 100 mg/kg Mn in their straw (Ishikawa et al., 2012). Another example is a Zn and Cd transporter OsHMA2, which plays a role in Zn and Cd loading into the xylem and participates in root-to-shoot translocation of these metals in rice. Overexpression of OsHMA2 driven by a 35S promoter can significantly reduce both Zn and Cd contents in rice grains. However, unexpectedly, when OsHMA2 was expressed under the control of OsSUT1, Cd concentration in the grains of SUT1-rice was only about half of that in the wild type, while the concentrations of Zn and other elements remained almost the same (Takahashi et al., 2012). The OsSUT1 promoter is thought to enhance expression of the Zn transporter, improving its efficiency in accumulating Zn in rice grains (Scofield et al., 2007). In addition, OsZIP7 seems to cooperate with OsHMA2 to load Cd and Zn into sieve tubes in the phloem region of nodes, and thus OsSUT1-OsHMA2 is thought to be beneficial for reducing the Cd content of rice grains (Tan et al.,

2019). However, further research is needed to determine whether OsSUT1-OsZIP7 constructs can reduce Cd concentrations in rice grains.

With advances in genomics and gene editing technology, molecular breeding for low Cd accumulation with high yield and quality may become a reality (Shan et al., 2014; Li et al., 2018). Tang et al. (2017) and Liu SM et al. (2019) demonstrated that it is feasible to generate yield-competitive and low-Cd rice lines through CRISPR/Cas9-mediated mutagenesis of *OsNramp5*, using the rice cultivars Huazhan and Xidao 1 as the parental wild-type lines. However, genetic modification does not always meet with favorable public perceptions (Chen JG et al., 2019; Chen KL et al., 2019).

Conventional and molecular breeding strategies, used alone and in combination, have been designed to develop low-Cd cereal cultivars to minimize Cd uptake and toxicity. Conventional breeding is still an attractive and potential tool for screening low-Cd crop cultivars. Molecular markers are a powerful technology for the selection of low-Cd cereal cultivars. In addition, modern molecular breeding technologies have great potential in breeding programs for the development of low-Cd cultivars, especially when coupled with conventional breeding. Within the next decade, the emphasis on low-Cd cereal breeding is likely to create more genetic variation for effective selection of low-Cd cultivars and further identification of unknown Cd accumulation-related genes and novel major QTLs controlling grain Cd accumulation. This will lead to a balanced use of phenotypic and genotypic selection and conventional and molecular breeding procedures. Both enhanced evaluation systems and regulations are necessary to determine whether low-Cd cultivars can be introduced into commercial production.

#### Contributors

Qin CHEN shaped the tables, wrote and edited the manuscript. Fei-bo WU wrote and edited the manuscript. Both authors have read and approved the final manuscript and, therefore, have full access to all the data in the study and take responsibility for the integrity and security of the data.

#### Acknowledgments

The authors would like to thank Prof. Guo-ping ZHANG (Zhejiang University, Hangzhou, China) for instructing the writing of this manuscript.

#### **Compliance with ethics guidelines**

Qin CHEN and Fei-bo WU declare that they have no conflict of interest.

This article does not contain any studies with human or animal subjects performed by either of the authors.

#### References

Abe T, Taguchi-Shiobara F, Kojima Y, et al., 2011. Detection of a QTL for accumulating Cd in rice that enables efficient Cd phytoextraction from soil. *Breed Sci*, 61(1): 43-51.

https://doi.org/10.1270/jsbbs.61.43

Abe T, Nonoue Y, Ono N, et al., 2013. Detection of QTLs to reduce cadmium content in rice grains using LAC23/ Koshihikari chromosome segment substitution lines. *Breed Sci*, 63(3):284-291.

https://doi.org/10.1270/jsbbs.63.284

- AbuHammad WA, Mamidi S, Kumar A, et al., 2016. Identification and validation of a major cadmium accumulation locus and closely associated SNP markers in North Dakota durum wheat cultivars. *Mol Breed*, 36(8):112. https://doi.org/10.1007/s11032-016-0536-1
- Ansarypour Z, Shahpiri A, 2017. Heterologous expression of a rice metallothionein isoform (*OsMTI-1b*) in *Saccharomyces cerevisiae* enhances cadmium, hydrogen peroxide and ethanol tolerance. *Braz J Microbiol*, 48(3):537-543. https://doi.org/10.1016/j.bjm.2016.10.024
- Aprile A, Sabella E, Vergine M, et al., 2018. Activation of a gene network in durum wheat roots exposed to cadmium. *BMC Plant Biol*, 18:238.

https://doi.org/10.1186/s12870-018-1473-4

- Arao T, Ae N, 2003. Genotypic variations in cadmium levels of rice grain. Soil Sci Plant Nutr, 49(4):473-479. https://doi.org/10.1080/00380768.2003.10410035
- Arduini I, Masoni A, Mariotti M, et al., 2014. Cadmium uptake and translocation in durum wheat varieties differing in grain-Cd accumulation. *Plant Soil Environ*, 60(1):43-49. https://doi.org/10.17221/416/2013-PSE
- Arthur E, Crews H, Morgan C, 2000. Optimizing plant genetic strategies for minimizing environmental contamination in the food chain. *Int J Phytoremediation*, 2(1):1-21. https://doi.org/10.1080/15226510008500027

Arunakumara KKIU, Walpola BC, Yoon MH, 2013. Current status of heavy metal contamination in Asia's rice lands. *Rev Environ Sci Bio/Technol*, 12(4):355-377. https://doi.org/10.1007/s11157-013-9323-1

Aziz R, Rafiq MT, Li TQ, et al., 2015. Uptake of cadmium by rice grown on contaminated soils and its bioavailability/ toxicity in human cell lines (Caco-2/HL-7702). J Agric Food Chem, 63(13):3599-3608. https://doi.org/10.1021/if5055572

https://doi.org/10.1021/jf505557g

Barabasz A, Wilkowska A, Tracz K, et al., 2013. Expression of *HvHMA2* in tobacco modifies Zn-Fe-Cd homeostasis. J *Plant Physiol*, 170(13):1176-1186. https://doi.org/10.1016/j.jplph.2013.03.018

Bashir K, Ishimaru Y, Nishizawa NK, 2012. Molecular mecha-

nisms of zinc uptake and translocation in rice. *Plant Soil*, 361(1-2):189-201.

https://doi.org/10.1007/s11104-012-1240-5

- CAC (Codex Alimentarius Commission), 2019. General Standard for Contaminants and Toxins in Food and Feed, Codex Standard 193-1995.
- Cao FB, Chen F, Sun HY, et al., 2014a. Genome-wide transcriptome and functional analysis of two contrasting genotypes reveals key genes for cadmium tolerance in barley. *BMC Genomics*, 15:611. https://doi.org/10.1186/1471-2164-15-611
- Cao FB, Wang RF, Cheng WD, et al., 2014b. Genotypic and environmental variation in cadmium, chromium, lead and copper in rice and approaches for reducing the accumulation. *Sci Total Environ*, 496:275-281. https://doi.org/10.1016/j.scitotenv.2014.07.064
- Cao ZZ, Lin XY, Yang YJ, et al., 2019. Gene identification and transcriptome analysis of low cadmium accumulation rice mutant (*lcd1*) in response to cadmium stress using MutMap and RNA-seq. *BMC Plant Biol*, 19:250. https://doi.org/10.1186/s12870-019-1867-y
- Čásová K, Černý J, Száková J, et al., 2009. Cadmium balance in soils under different fertilization managements including sewage sludge application. *Plant Soil Environ*, 55(8):353-361.

https://doi.org/10.17221/142/2009-PSE

Chang AC, Page AL, Foster KW, et al., 1982. A comparison of cadmium and zinc accumulation by four cultivars of barley grown in sludge-amended soils. *J Environ Qual*, 11(2):409-412.

https://doi.org/10.2134/jeq1982.00472425001100030018x

Chen F, Wu FB, Dong J, et al., 2007a. Cadmium translocation and accumulation in developing barley grains. *Planta*, 227(1):223-232.

https://doi.org/10.1007/s00425-007-0610-3

Chen F, Dong J, Wang F, et al., 2007b. Identification of barley genotypes with low grain Cd accumulation and its interaction with four microelements. *Chemosphere*, 67(10): 2082-2088.

https://doi.org/10.1016/j.chemosphere.2006.10.014

Chen HP, Yang XP, Wang P, et al., 2018. Dietary cadmium intake from rice and vegetables and potential health risk: a case study in Xiangtan, southern China. *Sci Total Environ*, 639:271-277.

https://doi.org/10.1016/j.scitotenv.2018.05.050

- Chen JG, Zou WL, Meng LJ, et al., 2019. Advances in the uptake and transport mechanisms and QTLs mapping of cadmium in rice. *Int J Mol Sci*, 20(14):3417. https://doi.org/10.3390/ijms20143417
- Chen KL, Wang YP, Zhang R, et al., 2019. CRISPR/Cas genome editing and precision plant breeding in agriculture. *Annu Rev Plant Biol*, 70:667-697. https://doi.org/10.1146/annurev-arplant-050718-100049
- Ci DW, Jiang D, Li SS, et al., 2012. Identification of quantitative trait loci for cadmium tolerance and accumulation in wheat. *Acta Physiol Plant*, 34:191-202.

https://doi.org/10.1007/s11738-011-0818-5

Clarke JM, Leisle D, Kopytko GL, 1997. Inheritance of cadmium concentration in five durum wheat crosses. *Crop Sci*, 37(6):1722-1726.

https://doi.org/10.2135/cropsci1997.0011183X003700060008x

- Clarke JM, Norvell WA, Clarke FR, et al., 2002. Concentration of cadmium and other elements in the grain of nearisogenic durum lines. *Can J Plant Sci*, 82(1):27-33. https://doi.org/10.4141/P01-083
- Clarke JM, McCaig TN, DePauw RM, et al., 2006. Registration of 'Strongfield' durum wheat. Crop Sci, 46(5):2306-2307.

https://doi.org/10.2135/cropsci2005.12.0454

Clemens S, Ma JF, 2016. Toxic heavy metal and metalloid accumulation in crop plants and foods. *Annu Rev Plant Biol*, 67:489-512.

https://doi.org/10.1146/annurev-arplant-043015-112301

Clemens S, Aarts MGM, Thomine S, et al., 2013. Plant science: the key to preventing slow cadmium poisoning. *Trends Plant Sci*, 18(2):92-99.

https://doi.org/10.1016/j.tplants.2012.08.003 Cohen CK, Fox TC, Garvin DF, et al., 1998. The role of irondeficiency stress responses in stimulating heavy-metal

transport in plants. *Plant Physiol*, 116(3):1063-1072. https://doi.org/10.1104/pp.116.3.1063

Cohen CK, Garvin DF, Kochian LV, 2004. Kinetic properties of a micronutrient transporter from *Pisum sativum* indicate a primary function in Fe uptake from the soil. *Planta*, 218(5):784-792.

https://doi.org/10.1007/s00425-003-1156-7

- Connolly EL, Fett JP, Guerinot ML, 2002. Expression of the IRT1 metal transporter is controlled by metals at the levels of transcript and protein accumulation. *Plant Cell*, 14(6):1347-1357.
  - https://doi.org/10.1105/tpc.001263
- Curie C, Alonso JM, le Jean M, et al., 2000. Involvement of NRAMP1 from *Arabidopsis thaliana* in iron transport. *Biochem J*, 347:749-755.

https://doi.org/10.1042/bj3470749

Dabeka RW, McKenzie AD, Lacroix GMA, 1987. Dietary intakes of lead, cadmium, arsenic and fluoride by Canadian adults: a 24-hour duplicate diet study. *Food Add Contam*, 4(1):89-101.

https://doi.org/10.1080/02652038709373618

Das N, Bhattacharya S, Bhattacharyya S, et al., 2017. Identification of alternatively spliced transcripts of rice phytochelatin synthase 2 gene OsPCS2 involved in mitigation of cadmium and arsenic stresses. *Plant Mol Biol*, 94(1-2):167-183.

https://doi.org/10.1007/s11103-017-0600-1

Du Y, Hu XF, Wu XH, et al., 2013. Affects of mining activities on Cd pollution to the paddy soils and rice grain in Hunan province, Central South China. *Environ Monit Assess*, 185(12):9843-9856.

https://doi.org/10.1007/s10661-013-3296-y

Eriksson JE, 1990. A field study on factors influencing Cd

levels in soils and in grain of oats and winter wheat. *Water Air Soil Poll*, 53(1-2):69-81.

https://doi.org/10.1007/BF00154992

Fahad S, Hussain S, Saud S, et al., 2015. Grain cadmium and zinc concentrations in maize influenced by genotypic variations and zinc fertilization. *CLEAN-Soil Air Water*, 43(10):1433-1440.

https://doi.org/10.1002/clen.201400376

- Florijn PJ, van Beusichem ML, 1993a. Uptake and distribution of cadmium in maize inbred lines. *Plant Soil*, 150(1):25-32. https://doi.org/10.1007/BF00779172
- Florijn PJ, van Beusichem ML, 1993b. Cadmium distribution in maize inbred lines: effects of pH and level of Cd supply. *Plant Soil*, 153(1):79-84. https://doi.org/10.1007/BF00010546
- Grant CA, Clarke JM, Duguid S, et al., 2008. Selection and breeding of plant cultivars to minimize cadmium accumulation. *Sci Total Environ*, 390(2-3):301-310. https://doi.org/10.1016/j.scitotenv.2007.10.038
- Greger M, Löfstedt M, 2004. Comparison of uptake and distribution of cadmium in different cultivars of bread and durum wheat. *Crop Sci*, 44(2):501-507. https://doi.org/10.2135/cropsci2004.5010
- Guttieri MJ, Seabourn BW, Liu CX, et al., 2015a. Distribution of cadmium, iron, and zinc in millstreams of hard winter wheat (*Triticum aestivum* L.). *J Agric Food Chem*, 63(49): 10681-10688.

https://doi.org/10.1021/acs.jafc.5b04337

- Guttieri MJ, Baenziger PS, Frels K, et al., 2015b. Prospects for selecting wheat with increased zinc and decreased cadmium concentration in grain. *Crop Sci*, 55(4):1712-1728. https://doi.org/10.2135/cropsci2014.08.0559
- Hao XH, Zeng M, Wang J, et al., 2018. A node-expressed transporter OsCCX2 is involved in grain cadmium accumulation of rice. *Front Plant Sci*, 9:476. https://doi.org/10.3389/fpls.2018.00476
- He JY, Zhu C, Ren YF, et al., 2006. Genotypic variation in grain cadmium concentration of lowland rice. J Plant Nutr Soil Sci, 169(5):711-716. https://doi.org/10.1002/jpln.200525101
- Hinesly TD, Alexander DE, Ziegler EL, et al., 1978. Zinc and Cd accumulation by corn inbreds grown on sludge amended soil. *Agron J*, 70(3):425-428.

https://doi.org/10.2134/agronj1978.00021962007000030015x

Hinesly TD, Alexander DE, Redborg KE, et al., 1982. Differential accumulations of cadmium and zinc by corn hybrids grown on soil amended with sewage sludge. *Agron J*, 74(3):469-474.

https://doi.org/10.2134/agronj1982.00021962007400030017x

- Hirzel J, Retamal-Salgado J, Walter I, et al., 2017. Cadmium accumulation and distribution in plants of three durum wheat cultivars under different agricultural environments in Chile. J Soil Water Conserv, 72(1):77-88. https://doi.org/10.2489/jswc.72.1.77
- Hirzel J, Retamal-Salgado J, Walter I, et al., 2018. Effect of soil cadmium concentration on three Chilean durum wheat

cultivars in four environments. *Arch Agron Soil Sci*, 64(2): 162-172.

https://doi.org/10.1080/03650340.2017.1337892

- Hu DW, Sheng ZH, Li QL, et al., 2018. Identification of QTLs associated with cadmium concentration in rice grains. J Integr Agric, 17(7):1563-1573. https://doi.org/10.1016/S2095-3119(17)61847-1
- Huang FL, Wei XJ, He JW, et al., 2018. Mapping of quantitative trait loci associated with concentrations of five trace metal elements in rice (*Oryza sativa*). *Int J Agric Biol*, 20(3):554-560.

https://doi.org/10.17957/IJAB/15.0524

Huang YZ, Hu Y, Liu YX, 2009. Heavy metal accumulation in iron plaque and growth of rice plants upon exposure to single and combined contamination by copper, cadmium and lead. Acta Ecol Sin, 29(6):320-326. https://doi.org/10.1016/j.chnaes.2009.09.011

Ikeda M, Zhang ZW, Higashikawa K, et al., 1999. Background exposure of general women populations in Japan to cadmium in the environment and possible health effects.

- *Toxicol Lett*, 108(2-3):161-166. https://doi.org/10.1016/S0378-4274(99)00084-3
- Ishikawa S, Ae N, Yano M, 2005. Chromosomal regions with quantitative trait loci controlling cadmium concentration in brown rice (*Oryza sativa*). *New Phytol*, 168(2):345-350. https://doi.org/10.1111/j.1469-8137.2005.01516.x
- Ishikawa S, Abe T, Kuramata M, et al., 2010. A major quantitative trait locus for increasing cadmium-specific concentration in rice grain is located on the short arm of chromosome 7. *J Exp Bot*, 61(3):923-934. https://doi.org/10.1093/jxb/erp360
- Ishikawa S, Ishimaru Y, Igura M, et al., 2012. Ion-beam irradiation, gene identification, and marker-assisted breeding in the development of low-cadmium rice. *Proc Natl Acad Sci USA*, 109(47):19166-19171. https://doi.org/10.1073/pnas.1211132109
- Ishimaru Y, Takahashi R, Bashir K, et al., 2012. Characterizing the role of rice NRAMP5 in manganese, iron and cadmium transport. *Sci Rep*, 2:286. https://doi.org/10.1038/srep00286
- Järup L, Åkesson A, 2009. Current status of cadmium as an environmental health problem. *Toxicol Appl Pharmacol*, 238(3):201-208.

https://doi.org/10.1016/j.taap.2009.04.020

Kashiwagi T, Shindoh K, Hirotsu N, et al., 2009. Evidence for separate translocation pathways in determining cadmium accumulation in grain and aerial plant parts in rice. *BMC Plant Biol*, 9:8.

https://doi.org/10.1186/1471-2229-9-8

Khan MA, Castro-Guerrero N, Mendoza-Cozatl DG, 2014. Moving toward a precise nutrition: preferential loading of seeds with essential nutrients over non-essential toxic elements. *Front Plant Sci*, 5:51.

https://doi.org/10.3389/fpls.2014.00051

Knox RE, Pozniak CJ, Clarke FR, et al., 2009. Chromosomal location of the cadmium uptake gene (*Cdu1*) in durum

wheat. *Genome*, 52(9):741-747. https://doi.org/10.1139/G09-042

Kubo K, Kobayashi H, Fujita M, et al., 2016. Varietal differences in the absorption and partitioning of cadmium in common wheat (*Triticum aestivum* L.). *Environ Exp Bot*, 124:79-88.

https://doi.org/10.1016/j.envexpbot.2015.12.007

- Lee S, An G, 2009. Over-expression of OsIRT1 leads to increased iron and zinc accumulations in rice. Plant Cell Environ, 32(4):408-416. https://doi.org/10.1111/j.1365-3040.2009.01935.x
- Lee S, Kim YY, Lee Y, et al., 2007. Rice P<sub>1B</sub>-type heavy-metal ATPase, OsHMA9, is a metal efflux protein. *Plant Physiol*, 145(3):831-842. https://doi.org/10.1104/pp.107.102236
- Li K, Yu HY, Li TX, et al., 2017. Cadmium accumulation characteristics of low-cadmium rice (*Oryza sativa* L.) line and F<sub>1</sub> hybrids grown in cadmium-contaminated soils. *Environ Sci Pollut Res*, 24(21):17566-17576. https://doi.org/10.1007/s11356-017-9350-5
- Li S, Liu SM, Fu HW, et al., 2018. High-resolution meltingbased TILLING of  $\gamma$  ray-induced mutations in rice. J Zhejiang Univ-Sci B (Biomed & Biotechnol), 19(8):620-629.

https://doi.org/10.1631/jzus.B1700414

- Liu JG, Zhu QS, Zhang ZJ, et al., 2005. Variations in cadmium accumulation among rice cultivars and types and the selection of cultivars for reducing cadmium in the diet. J Sci Food Agric, 85(1):147-153. https://doi.org/10.1002/jsfa.1973
- Liu JG, Qian M, Cai GL, et al., 2007. Uptake and translocation of Cd in different rice cultivars and the relation with Cd accumulation in rice grain. *J Hazard Mater*, 143(1-2): 443-447.

https://doi.org/10.1016/j.jhazmat.2006.09.057

- Liu SM, Jiang J, Liu Y, et al., 2019. Characterization and evaluation of OsLCT1 and OsNramp5 mutants generated through CRISPR/Cas9-mediated mutagenesis for breeding low Cd rice. Rice Sci, 26(2):88-97. https://doi.org/10.1016/j.rsci.2019.01.002
- Liu WQ, Pan XW, Li YC, et al., 2019. Identification of QTLs and validation of *qCd-2* associated with grain cadmium concentrations in rice. *Rice Sci*, 26(1):42-49. https://doi.org/10.1016/j.rsci.2018.12.003
- Liu WX, Shang SH, Xue F, et al., 2015. Modulation of exogenous selenium in cadmium-induced changes in antioxidative metabolism, cadmium uptake, and photosynthetic performance in the 2 tobacco genotypes differing in cadmium tolerance. *Environ Toxicol Chem*, 2015, 34(1): 92-99.

https://doi.org/10.1002/etc.2760

- Liu XS, Feng SJ, Zhang BQ, et al., 2019. OsZIP1 functions as a metal efflux transporter limiting excess zinc, copper and cadmium accumulation in rice. *BMC Plant Biol*, 19:283. https://doi.org/10.1186/s12870-019-1899-3
- Lu CN, Zhang LX, Tang Z, et al., 2019. Producing cadmium-free

*Indica* rice by overexpressing *OsHMA3*. *Environ Int*, 126: 619-626.

https://doi.org/10.1016/j.envint.2019.03.004

Luo JS, Huang J, Zeng DL, et al., 2018. A defensin-like protein drives cadmium efflux and allocation in rice. *Nat Commun*, 9:645.

https://doi.org/10.1038/s41467-018-03088-0

- McCouch SR, Wright MH, Tung CW, et al., 2016. Open access resources for genome-wide association mapping in rice. *Nat Commun*, 7:10532. https://doi.org/10.1038/ncomms10532
- Mills RF, Peaston KA, Runions J, et al., 2012. HvHMA2, a P<sub>1B</sub>-ATPase from barley, is highly conserved among cereals and functions in Zn and Cd transport. *PLoS ONE*, 7:e42640.

https://doi.org/10.1371/journal.pone.0042640

- Miyadate H, Adachi S, Hiraizumi A, et al., 2011. OsHMA3, a P<sub>1B</sub>-type of ATPase affects root-to-shoot cadmium translocation in rice by mediating efflux into vacuoles. *New Phytol*, 189(1):190-199.
- https://doi.org/10.1111/j.1469-8137.2010.03459.x Murgia I, de Gara L, Grusak MA, 2013. Biofortification: how can we exploit plant science and biotechnology to reduce micronutrient deficiencies? *Front Plant Sci*, 4:429. https://doi.org/10.3389/fpls.2013.00429
- Nakanishi H, Ogawa I, Ishimaru Y, et al., 2006. Iron deficiency enhances cadmium uptake and translocation mediated by the Fe<sup>2+</sup> transporters OsIRT1 and OsIRT2 in rice. *Soil Sci Plant Nutr*, 52(4):464-469. https://doi.org/10.1111/j.1747-0765.2006.00055.x
- Nevo Y, Nelson N, 2006. The NRAMP family of metal-ion transporters. *Biochim Biophys Acta*, 1763(7):609-620. https://doi.org/10.1016/j.bbamcr.2006.05.007
- NHFPC (National Health and Family Planning Commission), 2017. National Safety Standard for Contaminants in Foods, GB2762-2017. National Standards of People's Republic of China (in Chinese).
- Nicod J, Davies RW, Cai N, et al., 2016. Genome-wide association of multiple complex traits in outbred mice by ultra-low-coverage sequencing. *Nat Genet*, 48(8):912-918. https://doi.org/10.1038/ng.3595
- Nordberg GF, 2009. Historical perspectives on cadmium toxicology. *Toxicol Appl Pharmacol*, 238(3):192-200. https://doi.org/10.1016/j.taap.2009.03.015
- Oladzad-Abbasabadi A, Kumar A, Pirseyedi S, et al., 2018. Identification and validation of a new source of low grain cadmium accumulation in durum wheat. *G3*, 8(3):923-932. https://doi.org/10.1534/g3.117.300370
- Oliver DP, Gartrell JW, Tiller KG, et al., 1995. Differential responses of Australian wheat cultivars to cadmium concentration in wheat grain. *Aust J Agric Res*, 46(5):873-886.

https://doi.org/10.1071/AR9950873

Ovečka M, Takáč T, 2014. Managing heavy metal toxicity stress in plants: biological and biotechnological tools. *Biotechnol Adv*, 32(1):73-86. https://doi.org/10.1016/j.biotechadv.2013.11.011

Palmgren MG, Clemens S, Williams LE, et al., 2008. Zinc biofortification of cereals: problems and solutions. *Trends Plant Sci*, 13(9):464-473. https://doi.org/10.1016/j.tplants.2008.06.005

Pedas P, Ytting CK, Fuglsang AT, et al., 2008. Manganese efficiency in barley: identification and characterization of the metal ion transporter HvIRT1. *Plant Physiol*, 148(1): 455-466.

https://doi.org/10.1104/pp.108.118851

Peng F, Wang C, Cheng YR, et al., 2018a. Cloning and characterization of *TpNRAMP3*, a metal transporter from Polish wheat (*Triticum polonicum* L.). *Front Plant Sci*, 9:1354.

https://doi.org/10.3389/fpls.2018.01354

Peng F, Wang C, Zhu JS, et al., 2018b. Expression of TpNRAMP5, a metal transporter from polish wheat (*Triticum polonicum* L.), enhances the accumulation of Cd, Co and Mn in transgenic Arabidopsis plants. Planta, 247(6):1395-1406.

https://doi.org/10.1007/s00425-018-2872-3

- Penner GA, Bezte LJ, Leisle D, et al., 1995. Identification of RAPD markers linked to a gene governing cadmium uptake in durum wheat. *Genome*, 38(3):543-547. https://doi.org/10.1139/g95-070
- Perrier F, Yan B, Candaudap F, et al., 2016. Variability in grain cadmium concentration among durum wheat cultivars: impact of aboveground biomass partitioning. *Plant Soil*, 404(1-2):307-320. https://doi.org/10.1007/s11104-016-2847-8
- Pinson SRM, Tarpley L, Yan WG, et al., 2015. Worldwide genetic diversity for mineral element concentrations in rice grain. *Crop Sci*, 55(1):294-311. https://doi.org/10.2135/cropsci2013.10.0656
- Ramegowda Y, Venkategowda R, Jagadish P, et al., 2013. Expression of a rice Zn transporter, *OsZIP1*, increases Zn concentration in tobacco and finger millet transgenic plants. *Plant Biotechnol Rep*, 7(3):309-319. https://doi.org/10.1007/s11816-012-0264-x
- Randhawa HS, Asif M, Pozniak C, et al., 2013. Application of molecular markers to wheat breeding in Canada. *Plant Breed*, 132(5):458-471. https://doi.org/10.1111/pbr.12057
- Retamal-Salgado J, Hirzel J, Walter I, et al., 2017. Bioabsorption and bioaccumulation of cadmium in the straw and grain of maize (*Zea mays* L.) in growing soils contaminated with cadmium in different environment. *Int J Environ Res Public Health*, 14(11):1399. https://doi.org/10.3390/ijerph14111399
- Ricachenevsky FK, Punshon T, Lee S, et al., 2018. Elemental profiling of rice FOX lines leads to characterization of a new Zn plasma membrane transporter, OsZIP7. *Front Plant Sci*, 9:865.

https://doi.org/10.3389/fpls.2018.00865

Roberts TL, 2014. Cadmium and phosphorous fertilizers: the issues and the science. *Procedia Eng*, 83:52-59.

https://doi.org/10.1016/j.proeng.2014.09.012

- Salsman E, Kumar A, AbuHammad W, et al., 2018. Development and validation of molecular markers for grain cadmium in durum wheat. *Mol Breed*, 38(3):28. https://doi.org/10.1007/s11032-018-0788-z
- Sasaki A, Yamaji N, Yokosho K, et al., 2012. Nramp5 is a major transporter responsible for manganese and cadmium uptake in rice. *Plant Cell*, 24(5):2155-2167. https://doi.org/10.1105/tpc.112.096925
- Sasaki A, Yamaji N, Ma JF, 2014. Overexpression of OsHMA3 enhances Cd tolerance and expression of Zn transporter genes in rice. J Exp Bot, 65(20):6013-6021. https://doi.org/10.1093/jxb/eru340
- Satarug S, Baker JR, Urbenjapol S, et al., 2003. A global perspective on cadmium pollution and toxicity in nonoccupationally exposed population. *Toxicol Lett*, 137(1-2): 65-83.
  - https://doi.org/10.1016/S0378-4274(02)00381-8
- Satarug S, Garrett SH, Sens MA, et al., 2009. Cadmium, environmental exposure, and health outcomes. *Environ Health Persp*, 118(2):182-190. https://doi.org/10.1289/ehp.0901234
- Sato H, Shirasawa S, Maeda H, et al., 2011. Analysis of QTL for lowering cadmium concentration in rice grains from 'LAC23'. *Breed Sci*, 61(2):196-200. https://doi.org/10.1270/jsbbs.61.196
- Satoh-Nagasawa N, Mori M, Nakazawa N, et al., 2012. Mutations in rice (*Oryza sativa*) heavy metal ATPase 2 (*OsHMA2*) restrict the translocation of zinc and cadmium. *Plant Cell Physiol*, 53(1):213-224. https://doi.org/10.1093/pcp/pcr166
- Scofield GN, Hirose T, Aoki N, et al., 2007. Involvement of the sucrose transporter, OsSUT1, in the long-distance pathway for assimilate transport in rice. *J Exp Bot*, 58(12): 3155-3169.
  - https://doi.org/10.1093/jxb/erm153
- Sebastian A, Prasad MNV, 2014. Cadmium minimization in rice. A review. Agron Sustain Dev, 34:155-173. https://doi.org/10.1007/s13593-013-0152-y
- Shan QW, Wang YP, Li J, et al., 2014. Genome editing in rice and wheat using the CRISPR/Cas system. *Nat Protoc*, 9(10):2395-2410.

https://doi.org/10.1038/nprot.2014.157

- Shao JF, Xia JX, Yamaji N, et al., 2018. Effective reduction of cadmium accumulation in rice grain by expressing OsHMA3 under the control of the OsHMA2 promoter. J Exp Bot, 69(10):2743-2752. https://doi.org/10.1093/jxb/ery107
- Shimo H, Ishimaru Y, An G, et al., 2011. Low cadmium (*LCD*), a novel gene related to cadmium tolerance and accumulation in rice. *J Exp Bot*, 62(15):5727-5734. https://doi.org/10.1093/jxb/err300
- Song WE, Chen SB, Liu JF, et al., 2015. Variation of Cd concentration in various rice cultivars and derivation of cadmium toxicity thresholds for paddy soil by speciessensitivity distribution. J Integr Agric, 14(9):1845-1854.

https://doi.org/10.1016/S2095-3119(14)60926-6

- Soric R, Loncaric Z, Kovacevic V, et al., 2009. A major gene for leaf cadmium accumulation in maize (*Zea mays* L.). The Proceedings of International Plant Nutrition Colloquium XVI, UC Davis: Department of Plant Sciences.
- Stolt JP, Sneller FEC, Bryngelsson T, et al., 2003. Phytochelatin and cadmium accumulation in wheat. *Environ Exp Bot*, 49(1):21-28. https://doi.org/10.1016/S0098-8472(02)00045-X
- Sui FQ, Chang JD, Tang Z, et al., 2018. Nramp5 expression and functionality likely explain higher cadmium uptake in rice than in wheat and maize. Plant Soil, 433(1-2):377-389.

https://doi.org/10.1007/s11104-018-3849-5

- Sun HY, Cao FB, Wang NB, et al., 2013. Differences in grain ultrastructure, phytochemical and proteomic profiles between the two contrasting grain Cd-accumulation barley genotypes. *PLoS ONE*, 2013, 8(11):e79158. https://doi.org/10.1371/journal.pone.0079158
- Sun HY, Chen ZH, Chen F, et al., 2015. DNA microarray revealed and RNAi plants confirmed key genes conferring low Cd accumulation in barley grains. *BMC Plant Biol*, 15:259.

https://doi.org/10.1186/s12870-015-0648-5

- Takahashi R, Ishimaru Y, Senoura T, et al., 2011a. The Os-NRAMP1 iron transporter is involved in Cd accumulation in rice. *J Exp Bot*, 62(14):4843-4850. https://doi.org/10.1093/jxb/err136
- Takahashi R, Ishimaru Y, Nakanishi H, et al., 2011b. Role of the iron transporter OsNRAMP1 in cadmium uptake and accumulation in rice. *Plant Signal Behav*, 6(11):1813-1816.

https://doi.org/10.4161/psb.6.11.17587

- Takahashi R, Ishimaru Y, Shimo H, et al., 2012. The OsHMA2 transporter is involved in root-to-shoot translocation of Zn and Cd in rice. *Plant Cell Environ*, 35(11):1948-1957. https://doi.org/10.1111/j.1365-3040.2012.02527.x
- Tan JJ, Wang JW, Chai TY, et al., 2013. Functional analyses of TaHMA2, a P<sub>1B</sub>-type ATPase in wheat. *Plant Biotechnol J*, 11(4):420-431.

https://doi.org/10.1111/pbi.12027

- Tan LT, Zhu YX, Fan T, et al., 2019. OsZIP7 functions in xylem loading in roots and inter-vascular transfer in nodes to deliver Zn/Cd to grain in rice. Biochem Biophys Res Commun, 512(1):112-118. https://doi.org/10.1016/j.bbrc.2019.03.024
- Tang L, Mao BG, Li YK, et al., 2017. Knockout of OsNramp5 using the CRISPR/Cas9 system produces low Cdaccumulating *indica* rice without compromising yield. Sci Rep, 7:14438.

https://doi.org/10.1038/s41598-017-14832-9

Tezuka K, Miyadate H, Katou K, et al., 2010. A single recessive gene controls cadmium translocation in the cadmium hyperaccumulating rice cultivar Cho-Ko-Koku. *Theor Appl Genet*, 120(6):1175-1182. https://doi.org/10.1007/s00122-009-1244-6

- Tsukahara T, Ezaki T, Moriguchi J, et al., 2003. Rice as the most influential source of cadmium intake among general Japanese population. *Sci Total Environ*, 305(1-3):41-51. https://doi.org/10.1016/S0048-9697(02)00475-8
- Ueda Y, Frimpong F, Qi YT, et al., 2015. Genetic dissection of ozone tolerance in rice (*Oryza sativa* L.) by a genome-wide association study. *J Exp Bot*, 66(1):293-306. https://doi.org/10.1093/jxb/eru419
- Ueno D, Koyama E, Kono I, et al., 2009a. Identification of a novel major quantitative trait locus controlling distribution of Cd between roots and shoots in rice. *Plant Cell Physiol*, 50(12):2223-2233.
- https://doi.org/10.1093/pcp/pcp160 Ueno D, Kono I, Yokosho K, et al., 2009b. A major quantita-
- tive trait locus controlling cadmium translocation in rice (*Oryza sativa*). *New Phytol*, 182(3):644-653. https://doi.org/10.1111/j.1469-8137.2009.02784.x
- Ueno D, Yamaji N, Kono I, et al., 2010. Gene limiting cadmium accumulation in rice. *Proc Natl Acad Sci USA*, 107(38):16500-16505.
  - https://doi.org/10.1073/pnas.1005396107
- Uraguchi S, Fujiwara T, 2013. Rice breaks ground for cadmiumfree cereals. *Curr Opin Plant Biol*, 16(3):328-334. https://doi.org/10.1016/j.pbi.2013.03.012
- Uraguchi S, Kamiya T, Sakamoto T, et al., 2011. Low-affinity cation transporter (*OsLCT1*) regulates cadmium transport into rice grains. *Proc Natl Acad Sci USA*, 108(52):20959-20964.

https://doi.org/10.1073/pnas.1116531109

- Uraguchi S, Kamiya T, Clemens S, et al., 2014. Characterization of OsLCT1, a cadmium transporter from *indica* rice (*Oryza sativa*). *Physiol Plant*, 151(3):339-347. https://doi.org/10.1111/ppl.12189
- Uraguchi S, Tanaka N, Hofmann C, et al., 2017. Phytochelatin synthase has contrasting effects on cadmium and arsenic accumulation in rice grains. *Plant Cell Physiol*, 58(10): 1730-1742.

https://doi.org/10.1093/pcp/pcx114

- Vergine M, Aprile A, Sabella E, et al., 2017. Cadmium concentration in grains of durum wheat (*Triticum turgidum* L. subsp. durum). J Agric Food Chem, 65(30):6240-6246. https://doi.org/10.1021/acs.jafc.7b01946
- Vert G, Barberon M, Zelazny E, et al., 2009. Arabidopsis IRT2 cooperates with the high-affinity iron uptake system to maintain iron homeostasis in root epidermal cells. *Planta*, 229(6):1171-1179.
  - https://doi.org/10.1007/s00425-009-0904-8
- Wang XK, Gong X, Cao FB, et al., 2019. *HvPAA1* encodes a P-Type ATPase, a novel gene for cadmium accumulation and tolerance in barley (*Hordeum vulgare* L.). *Int J Mol Sci*, 20(7):1732.

https://doi.org/10.3390/ijms20071732

Wang Y, Wang XL, Wang C, et al., 2017. Transcriptomic profiles reveal the interactions of Cd/Zn in dwarf Polish wheat (*Triticum polonicum* L.) roots. *Front Physiol*, 8:168. https://doi.org/10.3389/fphys.2017.00168 Waters BM, Sankaran RP, 2011. Moving micronutrients from the soil to the seeds: genes and physiological processes from a biofortification perspective. *Plant Sci*, 180(4):562-574.

https://doi.org/10.1016/j.plantsci.2010.12.003

White PJ, Broadley MR, 2005. Biofortifying crops with essential mineral elements. *Trends Plant Sci*, 10(12):586-593.

https://doi.org/10.1016/j.tplants.2005.10.001

- WHO (World Health Organization), 2011. WHO Food Additives Series 64: Safety Evaluation of Certain Food Additives and Contaminants. Seventy-third Meeting of the Joint FAO/WHO Expert Committee on Food Additives (JECFA), Geneva.
- Wiebe K, 2012. Molecular characterization of *Cdu-b1*, a major locus controlling cadmium accumulation in durum wheat (*Triticum turgidum* L. var *durum*) grain. MS Thesis, University of Saskatchewan, Saskatoon, Canada.
- Wiebe K, Harris NS, Faris JD, et al., 2010. Targeted mapping of *Cdu1*, a major locus regulating grain cadmium concentration in durum wheat (*Triticum turgidum* L. var *durum*). *Theor Appl Genet*, 121(6):1047-1058. https://doi.org/10.1007/s00122-010-1370-1
- Wu DZ, Sato K, Ma JF, 2015. Genome-wide association mapping of cadmium accumulation in different organs of barley. *New Phytol*, 208(3):817-829. https://doi.org/10.1111/nph.13512
- Wu DZ, Yamaji N, Yamane M, et al., 2016. The HvNramp5 transporter mediates uptake of cadmium and manganese, but not iron. *Plant Physiol*, 172(3):1899-1910. https://doi.org/10.1104/pp.16.01189
- Wu FB, Zhang G, 2002. Genotypic differences in effect of Cd on growth and mineral concentrations in barley seedlings. *Bull Environ Contam Toxicol*, 69(2):219-227. https://doi.org/10.1007/s00128-002-0050-5
- Wu FB, Wu HX, Zhang GP, et al., 2004. Differences in growth and yield in response to cadmium toxicity in cotton genotypes. *J Plant Nutr Soil Sci*, 167(1):85-90. https://doi.org/10.1002/jpln.200320320
- Wu FB, Zhang GP, Dominy P, et al., 2007. Differences in yield components and kernel Cd accumulation in response to Cd toxicity in four barley genotypes. *Chemosphere*, 70(1): 83-92.

https://doi.org/10.1016/j.chemosphere.2007.06.051

Xue DW, Chen MC, Zhang GP, 2009. Mapping of QTLs associated with cadmium tolerance and accumulation during seedling stage in rice (*Oryza sativa* L.). *Euphytica*, 165(3):587.

https://doi.org/10.1007/s10681-008-9785-3

Yamaji N, Ma JF, 2014. The node, a hub for mineral nutrient distribution in graminaceous plants. *Trends Plant Sci*, 19(9):556-563.

https://doi.org/10.1016/j.tplants.2014.05.007

Yamaji N, Xia JX, Mitani-Ueno N, et al., 2013. Preferential delivery of zinc to developing tissues in rice is mediated by P-type heavy metal ATPase OsHMA2. *Plant Physiol*, 162(2):927-939.

https://doi.org/10.1104/pp.113.216564

- Yan HL, Xu WX, Xie JY, et al., 2019. Variation of a major facilitator superfamily gene contributes to differential cadmium accumulation between rice subspecies. *Nat Commun*, 10(1):2562. https://doi.org/10.1038/s41467-019-10544-y
- Yan JL, Wang PT, Wang P, et al., 2016. A loss-of-function allele of OsHMA3 associated with high cadmium accumulation in shoots and grain of Japonica rice cultivars. *Plant Cell Environ*, 39(9):1941-1954. https://doi.org/10.1111/pce.12747
- Yan YF, Choi DH, Kim DS, et al., 2010. Genotypic variation of cadmium accumulation and distribution in rice. *J Crop Sci Biotechnol*, 13(2):69-73. https://doi.org/10.1007/s12892-010-0036-5
- Yan YF, Lestari P, Lee KJ, et al., 2013. Identification of quantitative trait loci for cadmium accumulation and distribution in rice (*Oryza sativa*). *Genome*, 56(4):227-232. https://doi.org/10.1139/gen-2012-0106
- Yang M, Zhang YY, Zhang LJ, et al., 2014. OsNRAMP5 contributes to manganese translocation and distribution in rice shoots. *J Exp Bot*, 65(17):4849-4861. https://doi.org/10.1093/jxb/eru259
- Yang YM, Nan ZR, Zhao ZJ, 2014. Bioaccumulation and translocation of cadmium in wheat (*Triticum aestivum* L.) and maize (*Zea mays* L.) from the polluted oasis soil of Northwestern China. *Chem Spec Bioavailab*, 26(1):43-51. https://doi.org/10.3184/095422914X13888342841789
- Yousaf B, Liu GJ, Wang RW, et al., 2016. Bioavailability evaluation, uptake of heavy metals and potential health risks via dietary exposure in urban-industrial areas. *Environ Sci Pollut Res*, 23(22):22443-22453. https://doi.org/10.1007/s11356-016-7449-8
- Yu H, Wang JL, Fang W, et al., 2006. Cadmium accumulation in different rice cultivars and screening for pollution-safe cultivars of rice. *Sci Total Environ*, 370(2-3):302-309. https://doi.org/10.1016/j.scitotenv.2006.06.013
- Yuan LY, Yang SG, Liu BX, et al., 2012. Molecular characterization of a rice *metal tolerance protein*, OsMTP1. Plant Cell Rep, 31(1):67-79. https://doi.org/10.1007/s00299-011-1140-9
- Yue JY, Wei XJ, Wang HZ, 2018. Cadmium tolerant and sensitive wheat lines: their differences in pollutant accumulation, cell damage, and autophagy. *Biol Plant*, 62(2): 379-387.

https://doi.org/10.1007/s10535-018-0785-4

- Zeng FR, Mao Y, Cheng WD, et al., 2008. Genotypic and environmental variation in chromium, cadmium and lead concentrations in rice. *Environ Pollut*, 153(2):309-314. https://doi.org/10.1016/j.envpol.2007.08.022
- Zhang L, Zhang L, Song FB, 2008. Cadmium uptake and distribution by different maize genotypes in maturing

stage. Commun Soil Sci Plant Anal, 39(9-10):1517-1531. https://doi.org/10.1080/00103620802006651

- Zhang XH, Warburton ML, Setter T, et al., 2016. Genomewide association studies of drought-related metabolic changes in maize using an enlarged SNP panel. *Theor Appl Genet*, 129(8):1449-1463. https://doi.org/10.1007/s00122-016-2716-0
- Zhang XQ, Zhang GP, Guo LB, et al., 2011. Identification of quantitative trait loci for Cd and Zn concentrations of brown rice grown in Cd-polluted soils. *Euphytica*, 180(2): 173-179.

https://doi.org/10.1007/s10681-011-0346-9

Zhao JL, Yang W, Zhang SH, et al., 2018. Genome-wide association study and candidate gene analysis of rice cadmium accumulation in grain in a diverse rice collection. *Rice*, 11(1):61.

https://doi.org/10.1186/s12284-018-0254-x

Zhao XW, Luo LX, Cao YH, et al., 2018. Genome-wide association analysis and QTL mapping reveal the genetic control of cadmium accumulation in maize leaf. *BMC Genomics*, 19:91.

https://doi.org/10.1186/s12864-017-4395-x

Zhou Q, Shao GS, Zhang YX, et al., 2017. The difference of cadmium accumulation between the indica and japonica subspecies and the mechanism of it. *Plant Growth Regul*, 81(3):523-532.

https://doi.org/10.1007/s10725-016-0229-0

Zimmerl S, Lafferty J, Buerstmayr H, 2014. Assessing diversity in *Triticum durum* cultivars and breeding lines for high versus low cadmium content in seeds using the CAPS marker usw47. Plant Breed, 133(6):712-717. https://doi.org/10.1111/pbr.12218

## <u>中文概要</u>

- 题 目:禾谷类作物低镉积累育种研究进展
- 概 要: 镉(Cd)是毒性最强和农田受污染最普遍的重金属之一。土壤镉污染严重影响作物的产量和品质,并可通过食物链富集,从而危害人体健康。选育和种植食用部位镉低积累的作物品种是经济有效利用镉污染土壤,保证农产品安全生产,有效降低镉进入食物链的一条经济、有效的途径。本文主要介绍了近年来国内外在禾谷类作物镉低积累相关数量性状位点(QTL)定位、基因挖掘及其在低镉积累育种中的应用的最新成果;概述了镉低积累作物品种的定义、特征与选育方法,为低镉积累分子育种与生产提供理论指导。
- 关键词:禾谷类;低镉;数量性状位点(QTL)定位;育 种