

Seeing red: the origin of grain pigmentation in US weedy rice

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Abstract

Weedy forms of crop species infest agricultural fields worldwide and are a leading cause of crop losses, yet little is known about how these weeds evolve. Red rice (*Oryza sativa*), a major weed of cultivated rice fields in the US, is recognized by the dark-pigmented grain that gives it its common name. Studies using neutral molecular markers have indicated a close relationship between US red rice and domesticated rice, suggesting that the weed may have originated through reversion of domesticated rice to a feral form. We have tested this reversion hypothesis by examining molecular variation at *Rc*, the regulatory gene responsible for grain pigmentation differences between domesticated and wild rice. Loss-of-function mutations at *Rc* account for the absence of proanthocyanidin pigments in cultivated rice grains, and the major *rc* domestication allele has been shown to be capable of spontaneous reversion to a functional form through additional mutations at the *Rc* locus. Using a diverse sample of 156 weedy, domesticated and wild *Oryzas*, we analysed DNA sequence variation at *Rc* and its surrounding 4 Mb genomic region. We find that reversion of domestication alleles does not account for the pigmented grains of weed accessions; moreover, we find that haplotypes characterizing the weed are either absent or very rare in cultivated rice. Sequences from genomic regions flanking *Rc* are consistent with a genomic footprint of the *rc* selective sweep in cultivated rice, and they are compatible with a close relationship of red rice to Asian *Oryzas* that have never been cultivated in the US.

Keywords: crop–weed evolution, de-domestication, grain colour, *Oryza sativa*, proanthocyanidin, red rice

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Introduction

Red rice is a conspecific weedy relative of cultivated Asian rice (*Oryza sativa* L.) that occurs in rice fields worldwide. In the southern US, red rice infestations can reduce crop harvests by up to 80% (Estorninos *et al.* 2005) and are estimated to create crop losses exceeding \$45 million annually (Gealy *et al.* 2002). Control of red rice has proved difficult because of the weed's morphological and agroecological similarity to cultivated rice. Moreover, the potential for gene flow from domesti-

cated rice into red rice populations threatens the long-term effectiveness of weed control strategies based on herbicide-resistant crop varieties (Rajguru *et al.* 2005; Shivrain *et al.* 2007).

A defining feature of red rice is the dark-pigmented pericarp (bran) that gives the weed its common name. Like some other weed-associated traits, such as seed shattering and seed dormancy, pericarp pigmentation is a characteristic of wild *Oryza* species that was selected against during rice domestication (Sweeney *et al.* 2006, 2007). The vast majority of modern domesticated rice varieties lack proanthocyanidin pigments in the pericarp; this domestication trait arose through human selection for loss-of-function mutations at the *Rc* locus,

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which encodes a bHLH regulatory protein in the proanthocyanidin synthesis pathway (Sweeney *et al.* 2006). Selection for non-pigmented grains in the crop may potentially reflect human aesthetic preferences, selection against seed dormancy, selection for improved taste and cooking qualities and/or selection for other domestication traits linked to *Rc* (Gu *et al.* 2004; Sweeney *et al.* 2007).

The major *rc* domestication allele, present in more than 97% of non-pigmented cultivars (Sweeney *et al.* 2007), is characterized by a 14-bp frameshift deletion in exon 7 (initially annotated as exon 6; see Sweeney *et al.* 2006; Furukawa *et al.* 2007). This mutation generates a truncated, non-functional gene product and the non-pigmented ('white') pericarp of domesticated rice (Sweeney *et al.* 2006; Furukawa *et al.* 2007). An independently evolved domestication allele, *Rc-s*, is characterized by a C → A substitution in exon 7 that creates a premature stop codon (Sweeney *et al.* 2006); this allele occurs at a frequency of less than 3% in rice varieties globally (Sweeney *et al.* 2007). Analyses of DNA sequence variation in the *Rc* genomic region by Sweeney *et al.* (2007) have revealed that the major *rc* domestication allele arose initially in *japonica* rice and was subsequently introgressed into other varieties through selective breeding during domestication. The size of this introgressed genomic sequence is <1 Mb in most varieties surveyed to date (Sweeney *et al.* 2007). In contrast, *Rc-s* most likely originated in *aus* rice, a variety-group cultivated in a limited area of the northern Indian subcontinent; dissemination of the *Rc-s* allele beyond this geographical region has been minimal (Sweeney *et al.* 2007).

Recently, two studies have documented instances of mutational reversion of the major *rc* domestication allele to a functional form. Brooks *et al.* (2008) demonstrated that a spontaneous red-pericarp variant of the US cultivar 'Wells' discovered in 2005 arose through a 1-bp deletion located 20-bp upstream of the original 14-bp deletion; this new mutation restores reading frame, protein function and the proanthocyanidin-pigmented pericarp. Similarly, Lee *et al.* (2009) have documented a 1-bp deletion located 44 bp upstream of the loss-of-function deletion in a red-pericarp revertant of the Italian cultivar 'Perla'. Given that two independent instances of *rc* spontaneous reversion have been confirmed in the four years since *Rc* was first molecularly characterized (Sweeney *et al.* 2006, 2007; Furukawa *et al.* 2007), it seems plausible that *rc* reversions could occur repeatedly during the history of rice cultivation and that this process could account for the emergence of a red-pericarp phenotype in cultivated rice fields.

To date, studies examining the evolutionary origin of red rice have focused on neutral genetic markers, and there is growing evidence from these studies that US

weed strains are closely related to Asian domesticated rice. In particular, the two major phenotypic variants within US red rice, 'blackhull awned' (BHA) strains and 'strawhull awnless' (SH) strains, appear to be more closely related to domesticated rice of the *aus* and *indica* variety groups, respectively, than to any other groups within *Oryza* (Londo & Schaal 2007; Gealy *et al.* 2010; Reagon *et al.* 2010) (see Fig. 1A). Interestingly, neither *indica* nor *aus* varieties have ever been grown commercially in the US (Mackill & McKenzie 2003; Moldenhauer *et al.* 2004; Lu *et al.* 2005), and as there are no wild *Oryza* species native to North America, these patterns suggest that US red rice most likely arose through accidental introductions of *indica*-like and *aus*-like germplasm from Asia. Cultivated rice in the southern US is exclusively of the *tropical japonica* variety group (Mackill & McKenzie 2003), which is genetically distinct from *indica* and *aus* rice (Garris *et al.* 2005; Caicedo *et al.* 2007).

While neutral markers indicate a close relationship between US red rice and Asian cultivated varieties, these data do not provide information on whether the US weed strains are derived from Asian domesticates directly or whether they are instead descended from undomesticated (wild and/or weedy) *Oryza* populations that are simply related to *aus* and *indica* rice. DNA sequences from the pericarp pigmentation gene, *Rc*, can provide a novel source of information for addressing this question. This gene underlies a phenotype that is nearly fixed in domesticated rice and absent in both wild and weedy *Oryzas*. Thus, if red rice were descended from white-pericarp crop varieties, the *Rc* alleles present in the weed would be reversions of domestication alleles back to a functional form, as has been documented in US and European cultivars (Brooks *et al.* 2008; Lee *et al.* 2009). Alternatively, if red rice were instead descended from undomesticated material or from the rare landraces that still have pigmented pericarps, the weed's *Rc* alleles would be most closely related to alleles that never underwent loss-of-function mutations. These two possibilities need not be mutually exclusive, as there could be more than one evolutionary origin for the red pericarp phenotype present in weed populations (Fig. 1B).

In this study, we evaluate patterns of DNA sequence variation at the *Rc* locus and surrounding genomic regions to determine the origin of the red pericarp characterizing US red rice. We have compared *Rc* haplotypes in the US weed to variation present in wild and domesticated rice. Our sampling of cultivated varieties includes all major variety groups that form genetically distinct subpopulations within the crop: *indica* and *aus* varieties (together composing the *indica* subspecies); and *tropical japonica*, *temperate japonica* and *aromatic* varieties (together composing the *japonica* subspecies) (Garris *et al.* 2005; Londo *et al.* 2006; Caicedo *et al.* 2007). The cultivated

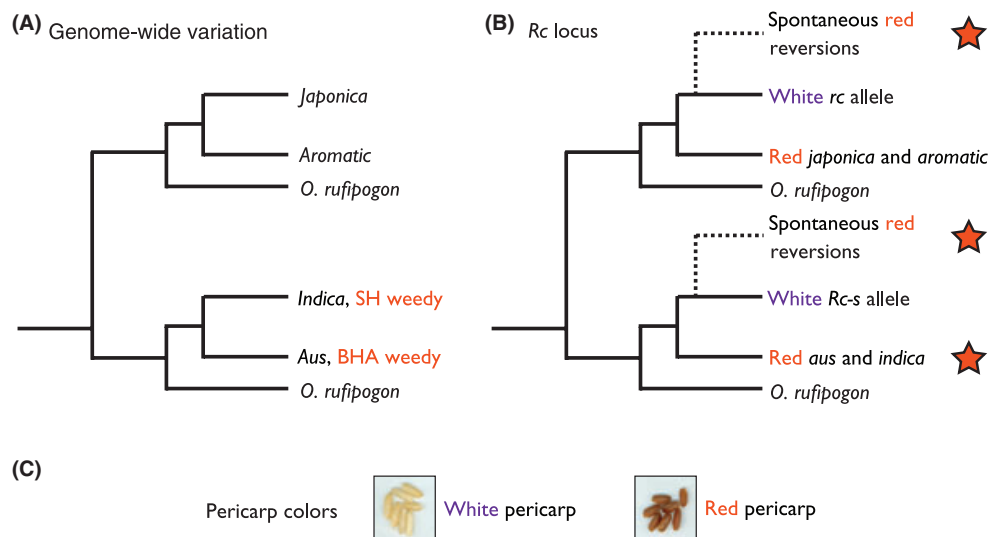


Fig. 1 Relationships among *Oryza* species and varieties based on genome-wide variation and the *Rc* locus. Schematic representations of the relationships between different *Oryza* species and variety groups based on (A) genome-wide surveys of genetic variation (Garrius *et al.* 2005; Caicedo *et al.* 2007) and (B) genetic variation at the *Rc* locus. In B, potential relationships of the US weed haplotypes to cultivated varieties are indicated by red stars. Dotted lines indicate potential origins of functional *Rc* alleles through reversion from domestication alleles. Non-pigmented (white) and pigmented (red) pericarps are shown in (C).

O. sativa sampling includes landraces with pigmented pericarps as well as the more typical white-pericarp cultivars. In addition to the *Rc* locus, we have examined DNA sequence variation at five loci that flank this gene, spanning nearly 2 Mb upstream and 2 Mb downstream of *Rc* on chromosome 7. This genomic region is expected to extend beyond the region of introgressed *japonica* sequence carrying the common *rc* domestication allele in most crop varieties (Sweeney *et al.* 2007). Taken together, this combination of sampling and loci allows us to evaluate relationships at the *Rc* locus, where most crop varieties carry the common *rc* allele, as well as relationships outside of this region of introgression, where patterns of sequence diversity reflect the phylogenetic relationships among weeds and crop varieties. Our data reveal that reversion of domestication alleles has played a minimal role, if any, in the evolution of the US red rice phenotype; that there is minimal sharing of *Rc* haplotypes between weedy rice and red-pericarp crop landraces; and that *Rc* haplotypes in the weed are genealogically distinct from those that gave rise to the domestication alleles. Together these findings strongly suggest that US red rice has not originated through reversions of white-pericarp cultivars to a feral form.

Materials and methods

Samples

Samples included both newly generated DNA sequence data and previously sequenced accessions that are pub-

licly available. For the newly generated DNA sequences, the initial sample set consisted of 138 *Oryza* accessions from around the world (Table S1, Supporting information); seven accessions were eliminated from the dataset due to poor sequence quality, yielding a final set of 131 accessions. Analyses of the *Rc* locus also included 25 previously published sequences from cultivated *O. sativa* that represented accessions not already present in our sample set (Table S1). US red rice was represented by 58 samples from throughout the range of the rice cultivation in the Southern US, including Missouri, Arkansas, Mississippi, Louisiana and Texas. These accessions were classified into four categories based on morphology (D. Gealy pers. comm.) and a genome-wide survey of sequence variation (Reagon *et al.* 2010). Classifications are as follows: 24 blackhull awned (BHA), 24 strawhull (SH), five accessions that were a rare intermediate brownhull awned phenotype (BR) and five accessions identified as crop-weed admixed individuals (MIX) (Table S1, Supporting information). Weed phenotypic descriptions are based on the morphology of the majority of accessions present in each genetically distinct group, as identified from genome-wide sequence data (Reagon *et al.* 2010). All but one of the weedy rice strains have pigmented (red) pericarps; the exception (1D10; Table S1, Supporting information) has been identified as a probable crop-weed hybrid (MIX) based on morphological characterization and genome-wide sequence data (Reagon *et al.* 2010). All weedy accessions were generously provided by Dr. David Gealy from collections maintained by the

USDA-ARS Dale Bumpers National Rice Research Center in Stuttgart, Arkansas; these accessions were all propagated through self-fertilization for two or more generations prior to use in the study (D. Gealy, pers. comm.).

Domesticated rice was represented by 13 US cultivars and 55 Asian landraces representing the diversity of varieties cultivated outside the southern US (Table 1). The US cultivars were obtained from the USDA-ARS and included eight modern cultivars released after 1930 as well as five historical crop varieties released before 1930, from which most modern US cultivars are derived. US cultivars, both modern and historical, are of the *tropical japonica* variety group, with extremely limited genetic contributions from *temperate japonica* and *indica* varieties at loci controlling traits such as semi-dwarfism and rice blast resistance (Mackill & McKenzie 2003). Sampled crop accessions from outside the US included 18 *indica*, nine *aus*, 16 *tropical japonica*, seven *temperate japonica* and five *aromatic* landraces (Table 1). *Oryza rufipogon* (including samples labelled as the annual form *O. nivara*) was represented by 32 accessions from throughout the native range of the species (Table S1, Supporting information). In addition, samples of two other closely related *Oryza* species were included as potential progenitors of the weedy form and for use as outgroups. These included the Latin American species *O. glum-aepatula* (two accessions) and the Australasian species *O. meridionalis* (two accessions). A portion of the crop accessions from outside the US and a portion of the *O. rufipogon* samples were kindly provided by

Dr Susan McCouch of Cornell University; other accessions of the crop and wild progenitor were obtained from collections maintained by the International Rice Research Institute in the Philippines. Samples of all congeners used as outgroups were obtained from the International Rice Research Institute.

DNA extraction, primer design and sequencing

Plants were grown from seed in greenhouses at Washington University in St. Louis and the University of Massachusetts, Amherst. Fresh tissue was harvested and frozen in liquid nitrogen; DNA was extracted via a modified CTAB procedure (Gross *et al.* 2009). Primers were designed to PCR-amplify and sequence the coding region of *Rc* (6.4 kb) as well as the regions approximately 1.6 kb upstream and 0.8 kb downstream of the start and stop codons. Nineteen overlapping PCR fragments (~400–600 bp) were used to sequence the region. Primers were designed using Primer3 (Rozen & Skaletsky 2000) and compared against the *indica* (93-11) and *temperate japonica* (Nipponbare) reference genome sequences in GenBank to ensure conservation and single-copy status. Primers were also designed to amplify and sequence ~500 bp portions of five genes in genomic regions upstream and downstream of *Rc*; targeted loci were approximately 2.1 Mb, 1.1 Mb and 0.3 Mb upstream, and 0.8 Mb and 2.0 Mb downstream of *Rc*. Previous research has shown that the length of the selective sweep surrounding the *Rc* locus is generally <1 Mb (Sweeney *et al.* 2007), so these flanking fragments include locations that were likely to be both within and outside of the swept region. Flanking fragment primers were located in exons, with the sequenced regions containing intronic regions of the targeted loci. Targeted loci, their genomic locations and primers are listed in Tables S2 and S3 (Supporting information).

Polymerase chain reaction (PCR) fragment amplification and Big Dye Terminator sequencing were performed according to conventional methods and separated on ABI capillary sequences by Cogenics™ in Connecticut and Texas. Missing fragments or poor quality data were finished using an ABI 3130 capillary sequencer at the Washington University Biology Departmental core facility. Sequencing techniques were standard and are available on request. One of the flanking fragments (rc3_005) was missing sequences for one *aus* and two US cultivars due to poor sequence quality; these three accessions were excluded from calculations for this locus and for calculations using concatenated flanking loci (Table S1, Supporting information). GenBank accession numbers for sequenced regions are GU260891–GU261673.

Table 1 *Oryza sativa* samples used in *Rc* analyses. Sample sizes are indicated in parentheses. For white-pericarp accessions, samples are listed by domestication allele type. Details on accessions are provided in Table S1

<i>O. sativa</i> subpopulation	Pericarp colour	
	Red	White (<i>rc</i> , <i>Rc-s</i>)
Asian domesticated variety groups (68)		
<i>indica</i> (18)	3	15, 0
<i>aus</i> (9)	4	2, 3
<i>tropical japonica</i>		
— Asian cultivars (16)	3	13, 0
— US cultivars (13)	0	13, 0
<i>temperate japonica</i> (7)	1	6, 0
<i>aromatic</i> (5)	0	4, 1
US weedy rice (57)		
Blackhull awned (BHA)	24	–
Strawhull awnless (SH)	24	–
Brownhull (BR)	5	–
Putative crop-weed hybrids (MIX)	3	1, 0

Data analysis

Sequence editing and alignment were performed using the PHRED, PHRAP and Polyphred programs (Deborah Nickerson, University of Washington) and BioLign v. 4.0.6 (Tom Hall, North Carolina State University, Raleigh, NC). As *O. sativa* is predominately self-fertilizing, the majority of crop and weed samples were homozygous at sequenced loci. However, *O. rufipogon* outcrosses more frequently, and there were some heterozygous *O. rufipogon* individuals at each locus. Because sequences were not cloned, allelic phase was not known unambiguously, and analytical phasing was not successful for the *Rc* locus due to recombination within the sequenced region. Thus, all sequences were treated as genotypic and whenever possible, unphased ambiguity codes were used for data analysis. It was necessary to generate arbitrarily phased 'pseudohaplotypes' for input into programs that could not accept ambiguity codes, but in these cases analyses were only conducted if they were not affected by SNP phase. The four weed accessions identified as putative crop-weed hybrids (MIX) (Reagon *et al.* 2010) were excluded from calculations of diversity and divergence to avoid artificially inflating values for the weed, but they were included in the phylogenetic analysis.

Nei's (1982) γ_{ST} genetic distance between groups and net sequence divergence (D_a ; Nei 1987) were calculated as measures of genetic differentiation for the *Rc* locus and for the flanking regions, both concatenated and separately. Net sequence divergence controls for the amount of within-group variation by subtracting the average within-group diversity from the total divergence between populations. Sequence diversity within groups and values of Tajima's (1989) D were also calculated for each sequenced region separately. Nucleotide diversity and divergence calculations were completed in DnaSP 5.0 (Librado & Rozas 2009).

Maximum likelihood (ML) analyses were performed for the *Rc* locus, with the best-fit model of nucleotide substitution selected in jModelTest 0.1.1 (Guindon & Gascuel 2003; Posada 2008) based on likelihood scores for 88 different models and the Akaike information criterion. ML trees were generated in RAxML (Stamatakis 2006b; Stamatakis *et al.* 2008) via the CIPRES web portal using the GTR model of molecular evolution with rate variation among sites. Because the lengths of *Rc* sequences from published GenBank accessions were shorter than the 9576 bp region sequenced for the present study, the aligned *Rc* dataset for tree construction was 6676 bp. Rate heterogeneity among sites was estimated using the GTR + CAT approximation, which is a fast computational substitution for the GTR + Γ model that can be applied to large datasets (more than 50 taxa)

(Stamatakis 2006a). Bootstrap values were calculated via 1000 replicates of the dataset, and a consensus tree was generated using Consense from Phylip 3.68 (Felsenstein 2005). Neighbour joining (NJ) analysis based on the same dataset was performed using Phylip 3.68 (Felsenstein 2005). Distances for the neighbour joining analysis were calculated using the F84 model, and bootstrap values were calculated via 1000 replicates of the data.

Results

Distribution of *Rc* alleles and pericarp pigmentation

Analyses of the *Rc* locus were based on 156 *Oryza* accessions, including 125 *O. sativa* samples (68 cultivated rice accessions and 57 weed strains), 27 accessions of the wild progenitor *O. rufipogon* and four accessions from related wild *Oryza* species used as outgroups (two accessions apiece of *O. meridionalis* and *O. glumaepatula*). A summary of *O. sativa* samples is shown in Table 1 (see also Table S1, Supporting information). Cultivated rice accessions include all five genetically distinct variety groups that are recognized in the crop (Garris *et al.* 2005; Caicedo *et al.* 2007), and the *tropical japonica* variety group is represented by both US and Asian cultivars (Table 1). Red rice accessions include 24 BHA strains, 24 SH strains, five samples that show a rare intermediate 'brownhull awned' (BR) phenotype and four samples derived from individuals that have been identified as probable crop-weed hybrids (MIX) in morphological characterizations (D. Gealy, USDA, Stuttgart, AR, pers. comm.) and in an analysis of DNA sequences at 48 sequence tagged site (STS) loci (Reagon *et al.* 2010). Together these *Oryza* samples comprise 131 newly sequenced accessions and 25 previously sequenced crop varieties (Sweeney *et al.* 2007; Table S1, Supporting information). The aligned length of the newly generated *Rc* sequences is 9576 bp, which includes ~1.6 kb of sequence immediately upstream of the *Rc* start codon, the transcribed portion of the gene, and ~0.8 kb downstream of the stop codon.

For the cultivated rice accessions, all white-pericarp varieties carry one of the two previously reported domestication alleles, with the rare *Rc-s* allele restricted to one *aromatic* and three *aus* accessions [Table 1 and Table S1 (Supporting information)]. Among the weed strains, a single MIX individual has a white pericarp (accession 1D10, Table S1, Supporting information); this individual carries the common *rc* domestication allele, consistent with an origin of the phenotype via crop-weed hybridization (followed by one or more generations of selfing; see 'Methods'). The other MIX accessions carry haplotypes that are either identical or nearly identical to the other red rice strains (described

below). In contrast to red-pericarp phenotypes that have been reported in US and European crop cultivars (Brooks *et al.* 2008; Lee *et al.* 2009), no red-pericarp weed accessions carry haplotypes with the 14-bp deletion characterizing the *rc* domestication allele, nor are there any other nucleotide substitutions indicating mutational reversion from a non-functional gene copy. Thus, the pericarp pigmentation that characterizes US red rice is not derived through a process of reversion from a domestication allele.

A ML tree of *Rc* haplotypes is shown in Fig. 2. Haplotype relationships are consistent between this tree and a tree generated by NJ analysis (NJ results not shown). For crop accessions, inferred haplotype relationships are concordant with previous analyses (Sweeney *et al.* 2007); the common *rc* domestication allele is grouped with haplotypes of red-pericarp *japonica* landraces (84% bootstrap support), and the *Rc-s* allele is grouped with

red-pericarp *aus* and *indica* landraces (99% bootstrap support). Red-pericarp weed accessions are characterized by eight closely related *Rc* haplotypes, which are grouped into a cluster with *O. rufipogon* and red-pericarp *aus* accessions (81% bootstrap support) (Fig. 2). Out of these eight weed haplotypes, seven are unique to red rice. A single haplotype, observed in 12 BHA strains and one MIX accession, is also present in one red-pericarp *aus* landrace and a single *O. rufipogon* accession (Fig. 2). The overall lack of haplotype sharing between weed strains and red-pericarp landraces indicates that the weeds are unlikely to be directly derived from the landraces represented by our sampling. Moreover, the genealogical distance between weed haplotypes and haplotype clusters containing the *rc* and *Rc-s* domestication alleles (Fig. 2) indicates that the weed alleles are not closely related to those characterizing common cultivated rice varieties. Similarly, the absence

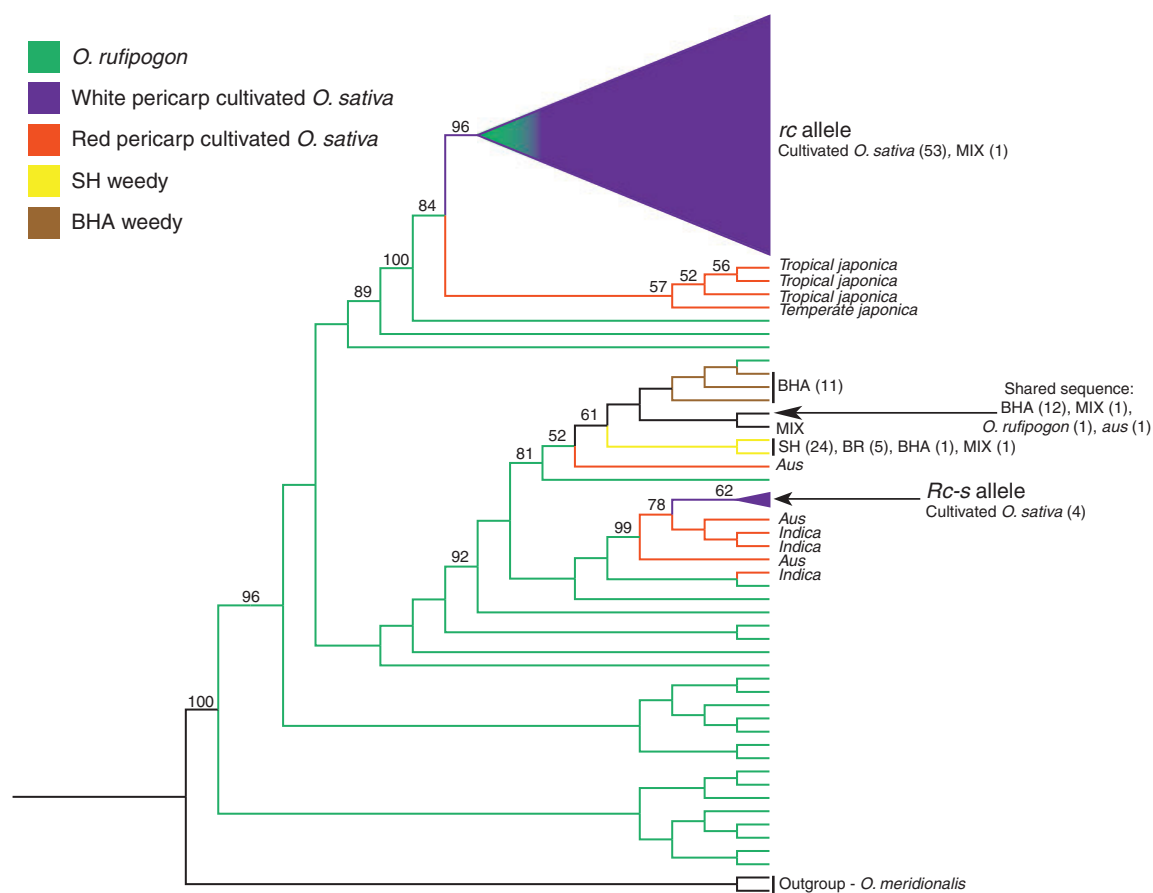


Fig. 2 Maximum likelihood (ML) tree of *Rc* haplotypes. Haplotypes carrying domestication alleles (*rc*, *Rc-s*) are grouped as triangles, with the size of the triangle roughly proportional to the number of accessions carrying the allele. Colours indicate the identities of accessions carrying a haplotype, as indicated in the key. The BR and putative crop-weed hybrid (MIX) red rice accessions are not represented by separate colours, but are labelled where they occur on the tree. Numbers following labels indicate the number of individuals sharing a haplotype. Haplotypes represented by only one individual are labelled but not numbered, except for haplotypes unique to *O. rufipogon*, which are neither labelled nor numbered. Numbers at nodes represent percent bootstrap support; bootstrap values over 50% are shown. For simplicity, only *O. meridionalis* is shown as the outgroup.

of extensive haplotype sharing with *O. rufipogon* accessions suggests that the weed strains are not directly descended from the populations of this wild species represented by our sampling.

Rc nucleotide diversity and population differentiation

Silent site (synonymous and noncoding) nucleotide diversity at *Rc* across all cultivated *O. sativa* accessions is $\pi = 1.55$ per kb (Table 2). For the subset of crop varieties carrying the common *rc* domestication allele, this value drops to $\pi = 0.03$ per kb, with a Tajima's *D* value indicating a statistically significant deviation from neutrality ($D = -1.9473$, $P < 0.05$; Table 2); these patterns are consistent with the selective sweep previously reported for the domestication allele (Sweeney *et al.* 2007). This signature of selection is also evident as negative Tajima's (1989) *D* values for individual variety groups possessing the *rc* allele (Table 2), although the pattern is statistically significant only for *tropical japonicas* ($D = -1.9778$, $P < 0.05$), which have the largest sample size ($n = 29$). For red-pericarp crop landraces, which have not been subject to domestication selection at *Rc*, the silent site nucleotide diversity is more than 100 times greater than the *rc* haplotype class ($\pi = 3.14$ per kb; Table 2), and is comparable to the genome-wide average for cultivated *O. sativa* based on a survey of 111 STS loci ($\pi = 3.20$ per kb; Caicedo *et al.* 2007). Silent site nucleotide diversity for the wild progenitor, *O. rufipogon*, is $\pi = 5.19$ per kb (Table 2); this value exactly matches the average level of polymorphism found in a survey of *O. rufipogon* accessions across 111 STS loci ($\pi = 5.19$ per kb; Caicedo *et al.* 2007).

For red-pericarp weed accessions, the silent site nucleotide diversity at *Rc* is nearly an order of magnitude lower than that of red-pericarp crop landraces ($\pi = 0.41$ per kb; Table 2). However, this observed nucleotide diversity is comparable to levels observed for the same set of weed accessions across 48 randomly selected STS loci [π per kb = 1.85 ± 3.31 (mean \pm SD); Reagon *et al.* 2010]. This finding suggests that genetic variation at *Rc* is consistent with a genome-wide genetic bottleneck in weed populations, most likely associated with their introduction into North America (Reagon *et al.* 2010).

Tajima's *D* for red rice is significantly positive ($D = 2.6015$, $P < 0.05$; Table 2). This pattern is created by *Rc* haplotype structure within the weed that closely corresponds to the major phenotypic variants: for haplotypes defined by SNPs only (i.e. excluding indels), weed accessions fall into two haplogroups (differing at six of the seven polymorphic sites), with one group consisting of all BHA strains but one (accession 1E08; Table S1, Supporting information) and the other con-

sisting of all SH and BR strains plus the remaining BHA strain. This pattern is similar to the genome-wide differentiation detected between BHA and SH strains at neutral loci (Londo & Schaal 2007; Gealy *et al.* 2010; Reagon *et al.* 2010); it is therefore likely to be reflecting historical population structure between the two weed subpopulations, with rare admixture events. The *Rc* haplotype structure in red rice also accounts for an absence of SNP polymorphism observed within the SH strains (Table 2), as well as a significantly negative Tajima's (1989) *D* value for BHA strains (caused by the inclusion of the single divergent 1E08 sequence; Table 2).

In quantifications of population differentiation at the *Rc* locus, weed strains show the greatest differentiation from *japonicas* and the other cultivated varieties where the *japonica*-derived *rc* allele predominates (*aromatics*, *indicas*) and the least differentiation from the *aus* variety group and *O. rufipogon* (Table 3). These patterns are consistent between the two differentiation measures employed (γ_{ST} and D_a), and they are similar for both BHA and SH strains. They are also consistent with inferred *Rc* haplotype relationships, which indicate that the weeds are more closely related to *O. rufipogon* and *aus* accessions at this locus than to any other samples in the dataset (Fig. 2). These patterns of differentiation at *Rc* stand in contrast to genome-wide patterns, which have indicated that SH strains are least differentiated from *indica* varieties (with very low differentiation between them) and that BHA strains are least differentiated from *aus* varieties and *O. rufipogon* (with somewhat greater differentiation than is observed between SH weeds and *indicas*) (Londo & Schaal 2007; Reagon *et al.* 2010). The *Rc*-specific pattern is directly attributable to selective introgression of the *rc* allele across cultivated variety groups, as all white-pericarp *indicas* carry the *japonica*-derived domestication allele at this locus (Table 1; Fig. 2; see also Sweeney *et al.* 2007).

Flanking regions

For the set of 131 accessions newly sequenced at *Rc*, sequences were also generated for portions of five loci distributed up to 2 Mb upstream and 2 Mb downstream of the *Rc* locus [Fig. 3, Table S2 (Supporting information)]. Aligned lengths of DNA sequences range from 419 bp to 505 bp for the five regions. Silent site nucleotide diversity varies widely among these loci (Table S4 and Fig. S1, Supporting information); one downstream fragment, *rc3_011* (encoding LOC_OS07G14150), shows exceptionally high diversity due to the presence of two divergent haplogroups that transcend species and variety groups in the sample set. Because the diversity at *rc3_011* is an order of magnitude higher for some

Table 2 Values of π , θ_w , and Tajima's D for total sites and silent sites per 1000 bases at the Rc locus and average values across flanking fragments

		Cultivated <i>Oryza sativa</i>											
		<i>Oryza rufipogon</i> ($n = 27$)	All cultivated ($n = 68$)	white-pericarp (rc) ($n = 53$)	red-pericarp cultivated ($n = 11$)	<i>indica</i> ($n = 18$)	<i>aus</i> ($n = 9$)	<i>temperate japonica</i> ($n = 7$)	<i>tropical japonica</i> (All) ($n = 29$)	<i>tropical japonica</i> (US only) ($n = 13$)	All US weedy rice* ($n = 56$)	BHA ($n = 24$)	SH ($n = 24$)
<i>Rc</i> locus													
π per kb	Total sites	4.62	1.27	0.04	2.65	1.32	2.29	0.04	0.07	0.02	1.80	0.06	0.00
π per kb	Silent sites	5.19	1.55	0.03	3.14	1.60	2.71	0.05	0.05	0.02	2.18	0.07	0.00
θ_w per kb	Total sites	6.58	1.68	0.19	2.33	1.73	2.01	0.06	0.22	0.04	2.16	0.18	0.00
θ_w per kb	Silent sites	7.52	1.90	0.16	2.64	2.09	2.32	0.07	0.14	0.04	2.62	0.22	0.00
Tajima's D	Total sites	-1.0653	-0.8099	-1.9473	0.6593	-0.9714	0.7122	-1.0062	-1.9778	-1.1492	-1.2441	-2.0832	N/A [†]
Flanking fragment average (flanking fragment average without rc3_011)‡													
π per kb	Total sites	6.13	4.09	3.76	0.82	0.67	4.55	0.00	1.39	0.00	7.75	0.17	0.43
		(3.25)	(1.95)	(1.71)	(1.03)	(0.83)	(2.48)	(0.00)	(0.64)	(0.00)	(1.68)	(0.21)	(0.53)
π per kb	Silent sites	9.31	6.21	5.76	0.97	1.09	6.73	0.00	2.09	0.00	12.32	0.12	0.76
		(4.31)	(2.53)	(2.25)	(1.22)	(1.36)	(3.11)	(0.00)	(0.81)	(0.00)	(2.13)	(0.15)	(0.95)
θ_w per kb	Total sites	6.32	3.55	3.65	0.67	0.61	3.77	0.00	2.36	0.00	7.75	0.44	0.22
		(4.60)	(1.65)	(1.70)	(0.84)	(0.76)	(2.09)	(0.00)	(1.28)	(0.00)	(1.68)	(0.44)	(0.28)
θ_w per kb	Silent sites	9.11	5.40	5.56	0.80	1.01	5.58	0.00	3.50	0.00	12.32	0.39	0.39
		(5.93)	(2.14)	(2.20)	(1.00)	(1.26)	(2.62)	(0.00)	(1.60)	(0.00)	(2.13)	(0.49)	(0.49)
Tajima's D [§]	Total sites	-0.4729	0.2903	-0.0958	2.0119	0.5058	0.7963	N/A	-1.3118	N/A	0.00	0.2648	N/A
		(-0.8667)	(0.2521)	(-0.1556)	(2.0119)	(0.5058)	(0.6257)	N/A	(-1.3475)	N/A	(0.00)	(-1.3268)	(N/A)

Statistically significant values at $P < 0.05$ are indicated in bold font.

*Excluding the white-pericarp MIX accession.

†N/A, cannot be calculated due to lack of polymorphisms.

‡Average values without locus rc3_011 are indicated in parentheses.

§Average across all fragments for which a Tajima's D value could be calculated.

Table 3 Percent net sequence divergence (D_n) between groups (above diagonal) and γ_{ST} genetic distance between groups (below diagonal), for (A) the *Rc* locus and (B) the flanking fragments (concatenated sequences). The US weedy group is not compared to BHA or SH weeds, as it comprises those groups

	<i>O. rufipogon</i>	<i>aus</i>	<i>indica</i>	<i>japonica</i>	<i>aromatic</i>	US cultivars	All US weedy	BHA	SH
(A) <i>Rc</i> gene									
<i>O. rufipogon</i>		0.087	0.153	0.178	0.191	0.190	0.161	0.168	0.180
<i>aus</i>	0.070		0.220	0.261	0.280	0.280	0.074	0.074	0.108
<i>indica</i>	0.126	0.436		0.008	0.018	0.017	0.354	0.352	0.387
<i>japonica</i>	0.170	0.528	0.096		0.002	0.001	0.408	0.412	0.446
<i>aromatic</i>	0.068	0.419	0.113	0.035		0.001	0.431	0.437	0.470
US cultivars	0.191	0.594	0.219	0.049	0.152		0.431	0.437	0.470
US weedy	0.296	0.237	0.690	0.785	0.568	0.827		N/A	N/A
BHA	0.256	0.366	0.856	0.929	0.931	0.976	N/A		0.062
SH	0.279	0.441	0.891	0.952	0.994	0.997	N/A	0.907	
(B) Flanking fragments									
<i>O. rufipogon</i>		0.179	0.128	0.474	0.269	0.654	0.155	0.195	0.143
<i>aus</i>	0.091		0.104	0.156	0.169	0.278	0.085	0.096	0.097
<i>indica</i>	0.102	0.251		0.459	0.392	0.661	0.030	0.078	0.007
<i>japonica</i>	0.313	0.235	0.628		0.074	0.013	0.521	0.575	0.490
<i>aromatic</i>	0.094	0.213	0.503	0.140		0.126	0.461	0.518	0.427
US cultivars	0.402	0.522	0.923	0.081	0.368		0.727	0.784	0.693
US weedy	0.241	0.171	0.127	0.669	0.406	0.825		N/A	N/A
BHA	0.245	0.299	0.527	0.777	0.627	0.968	N/A		0.052
SH	0.187	0.253	0.075	0.707	0.512	0.912	N/A	0.475	

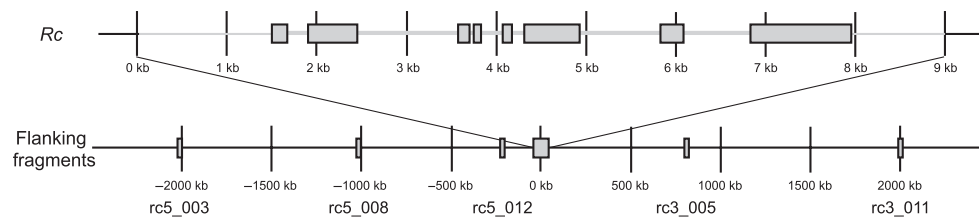


Fig. 3 Schematic of sequenced regions. Locations of the *Rc* locus with up- and downstream sequenced regions and flanking fragments on chromosome 7. At the *Rc* locus (top), exons are indicated as rectangles and the length of the sequenced region is shown in light grey. Distance of flanking fragments from the sequenced region of the *Rc* locus and names of the fragments are indicated below. Distances are approximately to scale and are based on the Nipponbare reference sequence.

groups compared to the other flanking fragments, average nucleotide diversity measures for flanking fragments were considered both with and without this locus for comparison purposes (Table 2; see also Table S4 and Fig. S1, Supporting information).

For cultivated *O. sativa*, average silent site nucleotide diversity across the flanking regions ($\pi = 2.53$ per kb; rc3_011 excluded) is similar to the previously reported genome-wide average of 3.20 (Caicedo *et al.* 2007) (Table 2). This pattern stands in contrast to the low diversity for cultivars at *Rc* (see above), an indication that the domestication-associated *rc* selective sweep does not dominate throughout the broader 4 Mb genomic region. Average values of silent site nucleotide diversity for *O. rufipogon* and US red rice ($\pi = 4.31$ and 0.63 per kb, respectively, with rc3_011 excluded; Table 2) are similar to values both at the *Rc* gene and

from genome-wide assessments (see above) (Caicedo *et al.* 2007; Reagon *et al.* 2010). When rc3_011 is included, values of π are elevated for populations containing both haplogroups (*aromatic*, *aus*, *tropical japonica* cultivated varieties; *O. rufipogon*), but are roughly unchanged for all other groups (Table 2).

For weed strains, genetic differentiation averaged across the flanking loci matches patterns revealed by genome-wide neutral markers (Londo & Schaal 2007; Reagon *et al.* 2010), with very close genetic similarity between SH strains and *indica* cultivars and moderately close similarity between BHA strains and *aus* varieties plus *O. rufipogon*. For a dataset consisting of concatenated sequences of the five flanking loci, γ_{ST} measures (Nei 1982) indicate that BHA is least differentiated from *O. rufipogon* followed by *aus* and that SH is least differentiated from *indica* followed by *O. rufipogon* (Table 3).

For measures of D_{av} , BHA and SH are collectively least differentiated from *indica* followed by *aus* varieties (Table 3). *Oryza rufipogon* shows low differentiation from both weedy and cultivated *O. sativa* in these measures; this is as expected, given that both groups are ultimately derived from *O. rufipogon* and share many polymorphisms with the wild species.

Of the five flanking loci, the nearest upstream locus, rc5_012 (encoding LOC_OS07G10600), is located ~310 kb away from *Rc* (Fig. 3; Table S2, Supporting information); it is therefore predicted to lie within the genomic region of *rc* allele introgression for some, but not all, introgressed *rc* haplotypes (Sweeney *et al.* 2007). The remaining four flanking loci are predicted to occur outside the ~1 Mb boundaries of introgression for most *rc* haplotypes (Sweeney *et al.* 2007). To assess the extent to which the *rc* selective sweep affects patterns of weed-crop differentiation across the 4 Mb genomic region, we examined SH-*indica* and BHA-*aus* population differentiation at *Rc* and at each of the flanking loci, and we compared these patterns to the strength of the *rc* selective sweep at each locus (measured as the proportion of nucleotide diversity in the *rc* allele class compared to the diversity in all cultivated accessions at that locus) (Fig. 4). For *indica* varieties, where virtually

every white-pericarp cultivar carries the introgressed *rc* allele (Sweeney *et al.* 2007), the degree of population differentiation from SH weed strains closely matches the genomic footprint of the *rc* selective sweep: differentiation is highest at *Rc*, lower at the nearest flanking locus (rc5_012), and lowest at the loci falling outside the region of *rc* introgression. In contrast, for BHA-*aus* differentiation, correspondence to the footprint of the *rc* selective sweep is much less evident (Fig. 4). This difference is consistent with the fact that, unlike *indic*as, most *aus* varieties in the sample set do not carry an introgressed *rc* allele (Table 1), so that the genomic impact of the *rc* selective sweep is lessened in this variety group.

Discussion

Rc and the evolutionary origin of red pericarps in US weedy rice

No visual trait so distinguishes US red rice from domesticated rice as the proanthocyanidin-pigmented pericarp that gives the weed its common name. Recent advances in our understanding of the genetic basis of pericarp colour in wild and domesticated rice (Sweeney *et al.* 2006, 2007; Furukawa *et al.* 2007; Brooks *et al.* 2008; Lee *et al.* 2009) have now made it possible to examine the genetic basis and origin of this weed-associated trait. Pericarp colour is largely controlled by the *Rc* locus: a functional *Rc* allele is required for the production of pericarp proanthocyanidins present in wild and weedy *Oryzas*, and the widespread nonfunctional *rc* allele results in the white pericarp that characterizes nearly all domesticated rice (Sweeney *et al.* 2006, 2007). If red rice were directly descended from de-domesticated (feral) white-pericarp varieties, the alleles present in the weed would likely be reversions of *rc* null alleles (Fig. 1B). If red rice were instead derived from undomesticated material or from the rare landraces that still have red pericarps, the *Rc* alleles in the weed would be related to ancestral, functional alleles that did not undergo loss-of-function mutations (Fig. 1B). *A priori*, at least two lines of evidence would seem to favour a de-domestication scenario. First, studies of genome-wide variation in US weeds have consistently shown that both BHA and SH weed forms are closely related to domesticated varieties of *O. sativa* (*aus* and *indica* varieties, respectively; Fig. 1A) (Londo & Schaal 2007; Gealy *et al.* 2010; Reagon *et al.* 2010); this would suggest that the weeds might well be directly descended from the crops with which they co-occur—the vast majority of which have white pericarps. Second, two independent instances of spontaneous *rc* allele reversion have already been documented in the four years since *Rc*

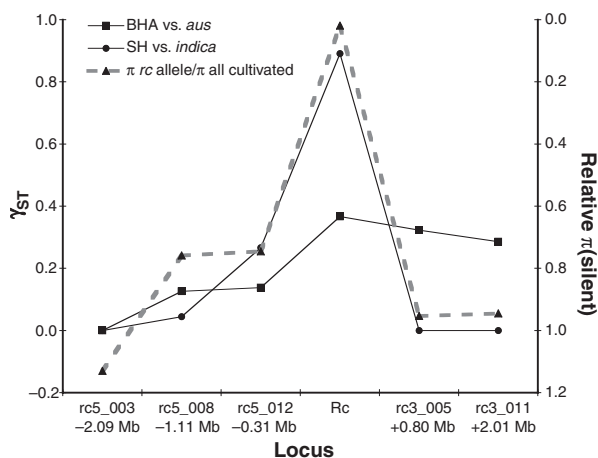


Fig. 4 Selection and weed-crop differentiation across the *Rc* genomic region. Solid black lines indicate genetic differentiation (γ_{ST} ; left-hand Y axis) between red rice phenotypic variants (BHA and SH) and their closest crop relatives (*aus* and *indica*, respectively). The dashed gray line indicates an index of the *rc* selective sweep in cultivated rice (right-hand y-axis); the index is calculated as silent-site π for crop accessions carrying the *rc* domestication allele as a proportion of π for all crop accessions at that locus. Numbers below flanking fragment names indicate their approximate location in relation to the *Rc* locus (see Fig. 3). Zero γ_{ST} values for rc5_003 indicate that all red rice, *aus* and *indica* accessions share an identical sequence at this locus.

was characterized at the molecular level (Brooks *et al.* 2008; Lee *et al.* 2009), and one of these reversion mutations likely occurred no earlier than 2005 (Brooks *et al.* 2008). Thus, mutational reversions from the common domestication allele have apparently occurred recurrently in cultivated rice fields.

Nonetheless, our examination of *Rc* alleles in US red rice and its possible progenitors does not support a crop reversion scenario, for either BHA or SH weeds. With the exception of a single white-pericarp weed accession that is a probable crop-weed hybrid (Reagon *et al.* 2010), no US weeds carry the widespread 14-bp deletion in exon 7, nor do any accessions show polymorphisms consistent with reversion from non-functional alleles. Thus, mutational reversion from *rc* domestication alleles does not account for the weed-associated phenotype in US red rice. It is important to note that this conclusion is for North American weed populations and may not necessarily hold true for all weedy rice populations worldwide. Red rice infests crop fields in virtually all world regions where rice is grown, and while currently limited, available genetic diversity data suggest that the genetic composition of the weed may vary widely by geographical region [e.g. Cao *et al.* (2006); reviewed by Gross & Olsen (2009)]. Given the apparent ease with which the *rc* domestication allele can revert to a functional form (Brooks *et al.* 2008; Lee *et al.* 2009), it seems likely that this mechanism could account for the red pericarp in at least some weed strains on a global scale. Expanded sampling of weed populations from diverse geographical regions will be useful for testing this hypothesis.

Given that the functional *Rc* alleles in the US weeds are not derived from white-pericarp crop varieties, what can be determined about their origin? One possibility is that they are derived from crop landraces that still possess red-pigmented pericarps. We included red-pericarp *indica*, *aus*, *temperate japonica* and *tropical japonica* landraces in our sampling to evaluate this possibility. With the exception of one weed haplotype that is shared with a single *aus* accession, most of the *Rc* haplotype diversity in the weeds is distinct from that characterizing red-pericarp landraces (Fig. 2). In the one instance of crop-weed haplotype sharing, the haplotype is also present in the wild species *O. rufipogon* (Fig. 2), suggesting that this may be a case of shared ancestral polymorphism or possibly recent gene flow between the *Oryza* populations. Thus, to the extent that our sampling of crop accessions can be considered representative of cultivated rice, our data suggest a limited role, if any, for red-pericarp landraces in the origin of the US weed alleles.

It should be noted that this finding does not rule out altogether a potential role for domesticated varieties in

the origin of the weed *Rc* alleles. It is possible that *Rc* haplotype variation in the weeds represents variation that was once present in the crop (or in incipiently domesticated red-pericarp populations) but which was lost as the white-pericarp domestication alleles were selectively favoured. It is also possible that more intensive sampling of extant red-pericarp landraces—specifically, *indica* and *aus* varieties—might reveal *Rc* haplotype similarity to weed strains that is not detected in the present analysis. In either of these cases, the weed populations would be descended primarily from domesticated germplasm, which is consistent with the close SH-*indica* and BHA-*aus* relationships observed in analyses of genome-wide neutral loci (Londo & Schaal 2007; Gealy *et al.* 2010; Reagon *et al.* 2010).

Other than domesticated rice, the other potential origin for the red rice *Rc* alleles is undomesticated *Oryza* populations. In our dataset, just as most *Rc* haplotype variation in US red rice is not shared with domesticated rice, it is also largely absent from the *O. rufipogon* samples (Fig. 2). However, whereas our sampling of domesticated rice is arguably representative, the same cannot necessarily be said for *O. rufipogon*. As a geographically widespread, outcrossing wild species, *O. rufipogon* shows much higher genetic diversity than the crop (Table 2; Caicedo *et al.* 2007) and the full genetic diversity of this species remains poorly characterized. Expanded collecting of *O. rufipogon* populations might be useful not only for further characterizing *Rc* haplotype diversity in comparison to red rice, but also for neutral marker assessments to clarify the roles of wild vs. domesticated populations in the weed's evolutionary origins.

Why is red rice red?

The red-pigmented pericarp is nearly ubiquitous in US weedy rice, and aside from occasional reports of white-pericarp weed strains in Asia (Suh *et al.* 1992; Ushiki *et al.* 2005), proanthocyanidin pigmentation characterizes the great majority of red rice populations worldwide. This raises the question of whether proanthocyanidin pigmentation can be considered a weed-adaptive trait. Several lines of evidence suggest that the red pericarp is selectively favoured, either directly or indirectly, in the US weed populations. First, we find no evidence for long-term persistence of non-functional *rc* alleles in US red rice populations. We observed the *rc* domestication allele in only one MIX weed accession (1D10; Table S1, Supporting information), and that accession shows genome-wide evidence of descent from a very recent crop-weed hybridization event (Reagon *et al.* 2010). Like cultivated rice, red rice is predominantly self-fertilizing, and so the recessive

white-pericarp phenotype would be expressed in weed populations in generations following a crop-to-weed introgression event. Given observations from other studies suggesting outcrossing rates from cultivated to weedy rice of up to 1% (Messeguer *et al.* 2001; Zhang *et al.* 2003; Chen *et al.* 2004; Shivrain *et al.* 2007), the near absence of the crop *rc* allele in weed populations is potentially consistent with selection against the white-pericarp phenotype. Indeed, at the molecular level we observe haplotype structure consistent with long-term selective maintenance of functional *Rc* alleles in the divergent SH and BHA weed populations (Table 2).

Additional evidence for the potential adaptive role of the red pericarp is provided by studies indicating that pericarp pigmentation may be important for seed dormancy. Strong seed dormancy is considered a classic weed-adaptive trait, allowing for the persistence of viable seeds over successive seasons (Baker 1965); the dormancy of weedy rice in particular makes it difficult to prevent re-infestation of a field once the weed has been introduced (Cohn & Hughes 1981). Work involving weedy rice, *Arabidopsis* and wheat has all indicated that testa/pericarp pigmentation contributes to seed dormancy (Gfeller & Svejda 1960; Debeaujon *et al.* 2000; Groos *et al.* 2002). Moreover, genetic mapping in Asian weedy rice has localized a QTL for seed dormancy to the genomic region containing the *Rc* locus (Gu *et al.* 2004, 2005, 2006). Thus, a loss of pericarp pigmentation may be selectively deleterious in weedy rice populations. Alternatively, the possibility remains that any patterns seen at the *Rc* locus in weedy rice are actually driven by selection at a closely linked locus; this can be evaluated through future genetic and functional studies.

Patterns of variation in the *Rc* genomic region

Examination of *Rc* in the context of its surrounding 4 Mb genomic region reveals contrasting evolutionary histories, not only for *Rc* in relation to flanking loci, but also for the weed populations in relation to cultivated *O. sativa*. At *Rc*, human selection for the white pericarp in domesticated rice has dramatically reshaped nucleotide diversity and haplotype distributions among variety groups. Consistent with a previous analysis by Sweeney *et al.* (2007), we observe patterns corresponding to a strong selective sweep associated with the *japonica*-derived *rc* domestication allele (Table 2) and a redistribution of haplotypes among variety groups resulting from selective introgression of this allele into all white-pericarp *indicas* as well as some white-pericarp *aromatic* and *aus* varieties (Table 1, Fig. 2). The footprint of this *rc* selective sweep can be traced across the 4 Mb genomic region, with a declining impact on weed-crop genetic differentiation at increasing distances

from *Rc* (Fig. 4). This genomic signature closely matches expectations based on the work of Sweeney *et al.* (2007), who found that most selectively introgressed *rc* haplotypes extend <1 Mb around the *Rc* locus.

Between the *indica* and *aus* variety groups, human selection for the *rc* domestication allele has disproportionately affected *indica* varieties: all white-pericarp *indicas* examined to date carry the *japonica*-derived *rc* allele, whereas white-pericarp *aus* varieties may carry either the *rc* allele or the independently-evolved *Rc-s* allele (Table 1; Sweeney *et al.* 2007). This difference between variety groups in the impact of *rc* selection has shaped the profile of weed-crop differentiation across the broader *Rc* genomic region. For the heavily-impacted *indica* varieties, the degree of differentiation from SH weeds closely matches the genomic footprint of the *rc* selective sweep: differentiation is highest at the *Rc* locus (where all white-pericarp *indicas* carry the introgressed *japonica* haplotype), and it declines at flanking loci in a pattern closely correlated with the declining impact of the *rc* sweep (Fig. 4). In contrast, for *aus* varieties, a much weaker corresponding pattern is observed with respect to differentiation from BHA strains (Fig. 4), consistent with the smaller impact of the *rc* sweep on this variety group.

The origin of weed characteristics

With the exception of herbicide resistance (Rajguru *et al.* 2005; Tan *et al.* 2007; Whaley *et al.* 2007), very little is known about the molecular basis and evolution of the traits that characterize conspecific crop weeds. This study represents a step in the direction of understanding the mechanisms by which these traits can arise in weed populations. For the case of US red rice pericarp colour, our analyses of the *Rc* genomic region have revealed that this weed-associated trait reflects the presence of ancestral functional alleles, rather than novel mutations that arose from *rc* domestication alleles. Interestingly, this suggests a previously unrecognized constraint on the emergence of weed strains in the US: in marked contrast to North American crop weeds such as feral rye (Burger *et al.* 2006) and weedy *Brassica* (Gulden *et al.* 2008), red rice appears extremely unlikely to evolve *in situ* from US cultivated varieties—all of which carry the *rc* domestication allele. It will be interesting to see what future research reveals about the mechanisms of weedy trait evolution for other traits and other weed species.

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Supporting information

Additional supporting information may be found in the online version of this article.

Table S1 Accessions included in the study. Accession numbers, species names, variety and cultivar designations, country of origin, and pericarp colour for samples included in the study. Unless otherwise noted, white-pericarp accessions carry the common *rc* domestication allele. Shaded samples were not sequenced completely and were excluded from data analysis

Table S2 Genomic context of sequenced loci. Names of sequenced loci, name of gene where sequenced locus is located, description of gene function, location and length of locus based on Nipponbare reference sequence and approximate distance from *Rc*

Table S3 Primer sequences used to amplify and sequence the *Rc* locus, including upstream and downstream regions and flanking fragments. Primers are named starting at the 5' end of the sequence (*rc_001*) and ending at the 3' end of the sequence (*rc_019*). Locations based on the Nipponbare reference genome are listed for the original forward and reverse sequence for each fragment and redesigned/optimized primers listed below original primers are located in roughly the same regions

Table S4 Values of π , θ_W , and Tajima's *D* for total sites and silent sites at flanking fragments

Fig. S1 Nucleotide diversity across the *Rc* genomic region. Values of π for silent sites at *Rc* and sequenced flanking loci for *O. rufipogon*, cultivated *O. sativa* (all accessions), cultivated *O. sativa* (*rc* allele class) and US weedy rice. The white-pericarp hybrid weed accession is excluded from all calculations.

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