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Identifying a large number of high-yield genes in rice by pedigree analysis, whole genome sequencing and CRISPR-Cas9 gene knockout

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Repeated artificial selection of a complex trait facilitates the identification of genes underlying the trait, especially if multiple selected descendant lines are available. Here we developed a pedigree-based approach to identify genes underlying the Green Revolution (GR) phenotype. From a pedigree analysis, we selected 30 cultivars including the "Miracle rice" IR8, a GR landmark, its ancestors and descendants, and also other related cultivars for identifying high-yield genes. Through sequencing of these genomes, we identified 28 ancestral chromosomal blocks that were maintained in all of the high-yield cultivars under study. In these blocks, we identified 6 genes of known function, including the GR gene sd1, and 123 loci with genes of unknown function. We randomly selected 57 genes from the 123 loci to do knockout or knockdown studies and found that a high proportion of these genes are essential or have phenotypic effects related to rice production. Notably, knockout lines have significant changes in plant height (p<0.003), a key GR trait, compared to wild-type lines. Some gene knockouts or knockdowns were especially interesting. For example, knockout of Os10g0555100, a putative glucosyltransferase gene, showed both reduced growth and altered panicle architecture. In addition, we found that in some retained chromosome blocks, several GR related genes were clustered, although they have unrelated sequences, suggesting clustering of genes with similar functions. In conclusion, we have identified many high-yield genes in rice. Our method provides a powerful means to identify genes associated with a specific trait.

high yield gene | pedigree analysis | green revolution | gene knockout

Complex traits, which might be related to survival in natural environments or crop productivity (1), are genetically difficult to dissect. This is, in part, because the effect of a single gene on a phenotype is usually small (2). To determine the genetic architecture of a complex trait (and the underlying gene networks), the most commonly employed methods are quantitative trait loci (QTL) mapping and genome-wide association studies (GWAS). QTL mapping is suitable for relatively simple quantitative traits (3), while GWAS provides valuable insights into trait architecture or candidate loci (4). Both methods have limitations, however. In OTL, the effects detected may be sensitive to external environments (5) and the span of chromosomal regions detected is often too long (owing to limited recombination events (6)) to pinpoint the causative gene(s). Similarly, in GWAS, the effects detected are sensitive to population structure, leading to both false positives and false negatives (7, 8).

Recently, a pedigree from crosses between different founding genotypes was used to fine-map QTLs in *Arabidopsis* (1, 9). The pedigree-based analysis combines linkage and association study (6). A pedigree with a founding genotype (e.g., derived from a single cross of two ancestors) and with recombination events over many generations could overcome the disadvantages inherent in QTL and GWAS. To reduce the sensitivity to environmental effects, however, it is necessary to have a clear phenotypic difference between the two ancestors. Identification of chromosomal blocks preserved in all members of the pedigree under selection for a given trait will facilitate identification of candidate genes. The question then is whether these candidates are indeed associated with the trait. The CRISPR-cas9 system (10) can in principle be used to knock out each candidate gene to get an insight into its function. Below we describe an application of this pedigree/knockout approach to the identification of high-yield genes in rice.

Our study takes advantage of the diploid rice pedigree in the Green Revolution. The Green Revolution has dramatically increased agriculture production worldwide since the 1960s, saving millions of lives from food shortage (11). The novel technologies allowed agronomists to breed high-yield varieties of maize, wheat, and rice. The yields were more than doubled in developing countries from 1961–1985 (12). Perhaps the most significant milestone of the Green Revolution was the introduction of semi-dwarfing genes into selected rice cultivars by hybridization.

The first semi-dwarf and high-yield modern rice variety (HYV) of the Green Revolution, known as the "Miracle rice" IR8, was created by crossing the Indonesian variety "Peta" with the Chinese variety "Dee-geo-woo-gen" (DGWG). It represented the first generation of the "high-yielding plant type", which provided a significantly higher yield potential for irrigated rice (13). In addition to the significant reduction in stem length, the high-yield rice cultivars have other important traits such as an early flowering time, improvement in photosynthetic allocation, and insensitivity to day length, directly or indirectly influencing the grain yield and yield stability(14, 15). These high-yield traits could

Significance

Finding the genes that control a complex trait is difficult because each gene may have only minor phenotypic effects. Quantitative trait loci mapping and genome-wide association study techniques have been developed for this purpose but are laborious and time-consuming. Here we developed a new method combining pedigree analysis, whole genome sequencing and CRISPR-Cas9 technology. By sequencing the parents and descendants of IR8, the "Miracle rice" in Green Evolution, we determined many genes that had been retained in the pedigree by selection for high yield. Knockout and knockdown studies showed that a large proportion of the identified genes are essential or have phenotypic effects related to production. Our approach provides a powerful means for identifying genes involved in a complex trait.

Reserved for Publication Footnotes



Fig. 1. Pedigree and flowchart for the identification of geneloci under selection. (a) An abridged pedigree of the major rice cultivars used in this study. The green-highlighted cultivars were re-sequenced, while the gray-highlighted were not. OP means "the other parent" and was not sequenced. The percentage in a box shows the expected probability of a given locus inherited from DGWG (D) or Peta (P) in that generation. The bottom box indicates the expected probabilities of a locus shared by all of the 8 MH63 descendants, which are extremely low (see *SI Appendix*, Table S4). A solid arrow denotes a direct parent (i.e. IR20) and a dotted arrow indicates an indirect ancestor (i.e. IR24). (b) Flowchart of the approach used to identify candidate blocks and gene loci derived from DGWG or Peta. Numbers of blocks (B) and gene loci (G) within the high confidence blocks are shown in each step of filtering. The reported 6 genes (3 from DGWG and 3 from Peta) arethe gene loci that have clear functions reported in literature. Most of the 129 gene loci each contain only one gene, except that 28 of them have two or more overlapped genes within a locus (see Methods).

be traced from the pedigree of "Miracle rice" IR8 that consists of its parents and high-yield progenies.

We assume that the genes related to high-yield were under strong artificial selection because yield was the major target trait of rice breeding since the 1960s. In this scenario, we note: 1) if the multiple lineages descended from an original cross have all been placed under the same selection, the alleles responsible for the trait in question should be found in all of the descendants, but not in all control populations; 2) in principle these alleles can be traced back to their origination and any variants inherited in all generations can be identified; 3) a gene under strong artificial selection should be present more commonly in progeny than genes not under selection; and 4) when knocking out a high-yield gene, a changed plant phenotype (e.g., an observable change in morphology or physiological response such as sterility) should be observed. All these expectations can be tested by sequencing the cultivars at important nodes in the pedigree and then by a knockout study using the CRISPR-Cas9 system.

Using the strategy above, we studied the extended pedigree of the ancestors and descendants of IR8 and other related lines (Fig. 1A) to determine a set of genes that played a critical role in the rice Green Revolution. By resequencing 30 cultivars from the pedigree (Fig. 1), we identified 28 chromosomal blocks, including 129 candidate gene loci, that have been preserved by artificial selection (Fig. 2). Fifty-seven gene loci with unknown function were selected to do knockout using the CRISPR-Cas9 technique. If the knockout failed, then a knockdown experiment was conducted. We found that 79% (15/19) knocked out loci and 62%(10/16) knocked down loci have phenotypic changes. These studies revealed a striking enrichment in yield/morphology-associated



Fig. 2.Blocks inherited from DGWG and Peta in IR8, IR24, IR30, MH63 and the 8 descendants of MH63. Blue and red bars represent blocks derived from401DGWG and Peta, respectively. "Shared"denotes the shared regions in all of the eight MH63 descendants. The purple arrows represent the 6 genes reported402with functions related to plant type or high-yield, while the asterisks represent the 123 gene loci with unknown functions; the 6 genes are shared by all 8403MH63 descendants and 5 collateral series. Chromosomes 3, 4 and 6, which contain no regions shared by all 8 MH63 descendants, are not shown here. The
second last block on chromosome 1 was shortened using breaks.405

genes among the candidate genes. Thus, our pedigree-guided

approach provides a simple, robust and fast means to identify candidate genes under directional selection.

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Total

Table 1. Num	bers of blocks deriv	ed from	DGWG	and Pet	ta in dif	ferent d	escend	ants
Ancestor	Descendant	Chroi	nosome					
		1	2	3	4	5	6	7

DGWG	IR8	4	15	17	12	7	11	4	3	3	13	6	13	108
	IR24	4	12	9	8	6	11	4	3	3	13	3	9	85
	IR30	4	11	7	7	6	7	4	2	3	12	2	8	73
	MH63	2	5	3	2	4	2	2	1	3	7	2	3	36
	Shared ^a	2	2	0	0	3	0	0	1	0	3	1	1	13
	Genes ^b	442	75	0	0	136	0	0	64	0	28	6	34	785
Peta	IR8	4	15	17	12	6	11	5	2	2	12	5	12	103
	IR24	4	14	11	6	4	6	5	1	2	10	3	10	76
	IR30	4	12	10	6	4	2	5	0	2	10	3	8	66
	MH63	1	5	7	1	3	0	3	0	1	6	1	5	33
	Shared ^a	1	2	0	0	3	0	2	0	1	5	1	0	15
	Genes ^b	101	34	0	0	115	0	265	0	42	308	95	0	960

Table 2. Phenotype when a specific gene was knocked out

Sampled ancestral block	Loci	Observed phenotypes	
DGWG chr01:37602014-39226171	Os01q0884200	Dwarf, sterile	
	Os01q0884400ª	Late heading, sterile	
	Os01q0884450	5.	
	Os01q0885000	Small, growth retarded, fewer tillers	
	Os01q0886000	Late heading, fewer tillers, sterile	
Peta chr01:40248759-40971796	Os01q0925600 ^a Os01q0925700	Rolling leaves, shorter panicle, dwarf	
	Os01g0930800	Late heading, sterile	
	Os01g0930900	No phenotypic change	
Peta chr10:21769689-21922126	Os10g0555600 ª	Dwarf	
	Os10g0555651		
	Os10g0555900ª Os10g0556000	Dwarf, late heading	
	Os10g0556200	Dwarf	
	Os10g0556900	No phenotypic change	
	Os10g0555100	Dwarf, spike shape change ,	
	Os10g0555200	Dwarf, sterile	
	Os10g0555300	Dwarf, sterile	
	Os10g0555700	Sterile	
	Os10g0556100	Small, growth retarded, leaf rolling	
Peta chr10:21992900-22072751	Os10g0558850	Rolling leaves, dwarf, weak	
	Os10g0559800ª Os10g0559833	No phenotypic change	
Peta chr11:6540176-7824094	Os11g0242400	No phenotypic change	

The 123 gene loci that passed our filtration came from 16 blocks, which ranged in size from 43kb to 1624kb. In total, 19 gene loci from 5 blocks of different sizes (80kb-1624kb) were successfully knocked out. For each gene, about 15 independently transgenic plants were obtained and on average in 79.5% of the cases the gene was knocked out in both homologous chromosomes. The phenotypic change was based on the observation of the homozygous knockout plants. No phenotypic change means no significant change in phenotype; e.g., the knockout of the locus Os01g0930900 showed shorter plants and shorter awns, but the changes were not statistically significant. In total, 15 out of the 19 knockouts exhibited phenotypes different from the wild type, suggesting that a large portion of these unknown-function gene loci are involved in flowering, fertility, leaf morphology, etc. The genotype and phenotype of each gene studied are described in Table S15. All the knockout plants in this table were in the Kasalath background ^aIn five pairs, the two genes in a pair are partly or completely overlapped. For example, Os01g0884450 is completely contained in Os01g0884400.

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Table 3. Phenotypic changes in knock-down mutants.

Locus	Abnormal phenotypes
Os01g0883900	Curled leaves, retarded growth. Died before
	matured.
Os01g0931600	Retarded growth, multiple tillers.
Os05g0170200	Retarded growth, curled leaves.
Os10g0556500	Brown and curled leaves. Died before
	matured.
Os10g0556700	Normal.
Os10g0559866	Normal.
Os02g0258900	Retarded growth, brown and curled leaves.
Os10g0391100	Normal.
Os10g0391200	Normal.
Os10g0392400	Curled leaves. Died before matured.
Os10g0554900	Normal.
Os12g0103000	Brown leaves. Died before matured.
Os12g0104250	Normal.
Os12g0104400	Brown leaves. Died before matured.
Os12g0104700	Retarded growth, curled leaves. Died before
-	matured.
Os12g0104733 ^a	Only grew roots. No seedling.
Os12g0104766 ª	Curled leaves.
^a These genes are i	ncluded in the same locus
inese genes are i	neiddeu in the same locus.

Results

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Rice cultivar selection and SNP identification

The famous "Miracle rice" IR8 is the key cultivar in our pedigree analysis (Fig. 1A). Its descendants and derivatives have been extensively used in the field, and its parents have been widely utilized to breed desired plant types (16). Another key cultivar is Minghui63 (MH63), which is a fourth generation descendant of IR8 and was the restorer line for a number of rice hybrids. MH63 accounted for >20% of the total production area in China during the 1980s and 1990s (17). Because of its wide planting areas with a stable high-yield, environmental or epigenetic effects could be excluded. IR8 and MH63 form the basis of our pedigree analysis. The pedigree further expands upward to the parents of IR8 (i.e., DGWG and Peta) and MH63 (IR30 and Gui630) and downward to the descendants of IR8 (i.e., IR24) and MH63. IR20, which has the same parents as IR8, and eight extensively used descendants of MH63 are also included in the analysis (Fig. 1A). All descendants of IR8 possessed the common feature of high-yield. To enhance the resolution in identifying genes under selection, we also sequenced four IR8 collateral series, eight tall landraces and a wild rice as the controls (SI Appendix, Fig. S1B and Table S1). The alleles present in the control groups were considered unlikely to contribute to high yield.

The 30 diploid rice accessions selected above were resequenced with a reasonable coverage depth (> $20\times$) in our study (SI Appendix, Table S1 and Fig. S2). Because pedigree information and independent re-sequencing of descendants from the same ancestor offer the unique advantage of discriminating against false markers, each inherited block of interest can be double-checked not only between successive generations but also between nodes separately by more than one generation and between lineages. Based on the linked markers in the majority of the successive generations, this approach can exclude false markers, infer correct single nucleotide polymorphisms (SNPs), and improve the accuracy of SNP identification (SI Appendix, Fig. S3). In the two most important parent-offspring trios, DGWG-Peta-IR8 and IR30-Gui630-MH63, a total of 592,603 and 481,385 high-quality SNPs were called, respectively, to detect the inherited chromosomal blocks from IR8 and its parents (see Methods and SI Appendix, Fig. S4).

Expected and observed proportions of inherited blocks

With the pedigree information, the probability of a block or a gene being passed on to the next generation can be computed using classical genetic theory. One can then compare the computed probability with the observed proportion (see SI Appendix, SI Materials and Methods). In the absence of selection, the probabilities of a gene locus in MH63 from DGWG and Peta are expected to be 3.9% and 13.4%, respectively (Fig. 1A and SI Appendix, Table S2). The probability of one or more DGWG or Peta blocks being present in all eight descendants of MH63 is extremely low (4.71×10⁻⁸ or 1.62×10⁻⁷) (Fig. 1A, Table 1 and *SI Appendix*, Table \$3, Table S4). Therefore, every block retained in all of the MH63 progenies is likely to have been targeted by artificial selection for the high-yield phenotype.

Theoretically, the heterozygosity of the F₁ hybrid will be reduced to half in its F₂ progeny through selfing and will eventually be reduced to almost zero in an inbred line (e.g., IR8 or MH63). Therefore, the crossover events can be detected in both IR8 and MH63 to determine the origin of each block (SI Appendix, Table S5 and Table S6). The block information in MH63 enabled us to exclude the genetic blocks from Gui630 and identify those from DGWG or Peta based on the pedigree in Fig. 1A. In MH63, we found 57 and 59 blocks that were derived from 36 DGWG and 33 Peta blocks in IR8, respectively (Fig. 2). Thus, many of the original inherited blocks from DGWG and Peta had been fragmented into smaller ones in MH63 by recombination. The average length is 483 kb for the 57 DGWG-derived blocks and 398 kb for the 59 Peta-derived blocks, which are 5.45- and 3.20fold shorter than the average lengths in the original blocks in IR8, respectively (SI Appendix, Table S7 and Table S8). Among those original blocks, only a total of 6.26 Mb DGWG and a total of 8.76 Mb Peta segments are inherited in all of the 8 MH63 descendants. They were 2.39- and 1.55-fold shorter than the inherited blocks observed in MH63, respectively. The sequences shared by all 8 MH63 descendants contained 785 DGWG- and 960 Peta-specific genes (Fig. 1B and Fig. 2).

Identification of candidate genes for the high-yield phenotype When only a limited number of genes in a block are under selection, the ancestral block will become shorter and shorter over generations because of recombination events. Fig. 2 includes an example in which a block on Peta chromosome 5 became shorter and shorter by crossover events from IR8 to MH63. Interestingly, a candidate gene, GW5, which is responsible for rice grain width, shape, quality and yield, is located near recombination hotspots (18) but has been retained. The pattern displays efficient selection on this block.

659 In a block with many genes, some alleles that are not sub-660 jected to selection may be inherited due to linkage (i.e., hitchhik-661 ing). Several strategies were employed to exclude the hitchhiked 662 genes and identify the genes that were most likely the target of 663 selection, including those with un-annotated functions (Fig. 1B). 664 π (polymorphic sites/informative sites) was calculated for each 665 10-kb window to compare the diversity values within and between 666 different groups. First, we assumed that targeted alleles should 667 have been retained also in the IR8 collateral series because those 668 cultivars are also of high-yield plant types. With this assumption, 669 we selected four cultivars of the IR8 collateral series (SIAppendix, 670 Fig. S1 and Table S1) and calculated the nucleotide diversity 671 of these candidate genes between MH63 and each of the four 672 collateral cultivars together with IR26, a progeny of IR24, and 673 a sister line of UPR221 (a parent of IR30 in Fig. 1A). Only the 674 genes that had an average diversity < 0.0001 and were identical 675 in the majority of collateral series (≥ 3) for the compared pairs 676 were kept. Second, we assumed that a gene with an extremely 677 low diversity among wild rice lines and cultivars should be ex-678 cluded because it is more likely to be essential for fundamental 679 biological processes rather than being responsible for the high-680

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	Locus	Mutant height (cm)	WT height (cm)	Height change: (mutant height-WT height)/WT heigh
4 positive controlsª	Os01g0883800	62.8	132.4	-52.6%
	Os01g0884300	65.5	129.3	-49.3%
	Os05g0170000	47.8	130.3	-63.4%
	Os02g0260200	98.3	123.3	-20.3%
18 target gene loci ^b	Os01g0884200	110.7	129.3	-14.40%
	Os01g0884400	125.3	127.3	-1.6%
	Os01g0884450			
	Os01g0886000	127.7	129.6	-1.5%
	Os01g0925600	93.3	125.6	-25.7%
	Os01g0925700	ICCI		
	Os01g0930800	125.3	131.2	-4.5%
	Os01g0930900	130.2	129.1	0.9%
	Os10g0555100	99.6	130.1	-23.4%
	Os10g0555200	99.7	130.3	-23.5%
	Os10g0555300	105.2	129.2	-18.6%
	Os10g0555600	95.2	130.2	-26.9%
	Os10g0555651			
	Os10g0555700	127.1	130.6	-2.7%
	Os10g0555900	64.6	132.4	-51.2%
	Os10g0556000			
	Os10g0556100	65.2	127.3	-48.8%
	Os10g0556200	73.7	122.1	-39.6%
	Os10g0556900	131.2	129.1	1.6%
	Os10g0558850	117.2	130.2	-10.0%
	Os10g0559800	130.1	129.3	0.6%
	Os10g0559833			
	Os11g0242400	129.4	128.6	0.6%
10 random controls ^c	Os01g0936100	130.3	131.5	-0.9%
	Os05g0375600	134.0	132.4	1.2%
	Us05g0571700	126.5	129.2	-2.1%
	Us05g05/3600	132.0	130.1	1.5%
	Us10g0341750	134.0	130.1	3.0%
	Us10g0342300	132.0	129.2	2.2%
	Os10g0341700	133.0	130.2	2.2%
	Us05g05/1300	134.3	132.4	1.5%
	Os10g0558400	128.5	132.4	-2.9%
	Us10g0342650	131.3	131.5	-0.1%

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On average, positive controls showed 46.4% reduction in plant height (p=0.017, two-tail t-test, 95% confidence interval: -99.6 -20.84), while 18 target gene loci showed 16.4% reduction in plant height (p=0.0013, two-tail t-test, 95% confidence interval: -31.98 -9.22). 10 random controls only showed a slight difference (knockout effect) (average 0.5%, p=0.42, two-tail t-test, 95% confidence interval: -1.16 - 2.54).

^a Four of the 6 positive controls were knocked out in Kasalath. GW5 was knocked out in Wuyungeng and rl14 was not successfully knocked out.

^b The knockout plant (Os01g0885000) died before the tilling stage, and the plant height could not be compared with the others. Therefore, only 18 target mutants were measured.

^c10 genes adjacent to the target blocks were randomly chosen as controls.



Fig. 3. Photos of knockout mutants with changed phenotypes.These photos show 6 examples as shorter plants, rolling leaves, a later heading date, changed panicles and empty seeds compared with the wild type. The other 9 knockout mutants with observable phenotypic changes and the controls are shown in *SI Appendix*, Fig. S7. Supporting informationThe following materials are available in the online version of this article.**SI Materials and Methods**

yield phenotype. Therefore, we further filtered out the bottom

50% of genes in terms of the diversity between MH63 and the

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953 11 wild rice varieties. Third, we filtered further by comparison 954 to tall cultivars as follows. All the re-sequenced cultivars in this 955 study were grown in the field and their heights were measured. 956 Because the semi-dwarfism trait was specifically selected for the 957 Green Revolution, we expect that the alleles related to the Green Evolution would be divergent with tall cultivars and only be kept 958 959 in the genes showing a diversity higher than the median between 960 MH63 and each of the 8 tall cultivars (SI Appendix, Table S1 and 961 Table S9).

962 The above filtering procedure identified 129 gene loci, which 963 can be divided into 101 single loci and 28 loci with overlapping 964 genes (where two or more genes overlap completely or partly 965 within the same locus). As an example of overlapping genes, 966 the coding sequence of Os01g0883850 is completely contained 967 in the reported gene sd1 (Os01g0883800). These two genes are 968 thus considered as a single entity in our analysis. Each locus is 969 named by one gene it contains. Of the 129 gene loci, 44 are 970 from DGWG- and 85 from Peta-specific blocks (Fig. 1B and SI 971 Appendix, Table S10). These 129 gene loci are located on 17 blocks 972 which are inherited in all 8 descendants of MH63. Six of the 129 973 gene loci contain genes with known functions, including the semi-974 dwarf gene, sd-1, known as the "green revolution gene". This gene 975 encodes gibberellin 20-oxidase, the key enzyme in the gibberellin 976 biosynthesis pathway. Another gene, larger panicle (lp), which 977 controls the panicle architecture (19), has recently been found 978 to be a target of selection in Indica cultivars by a GWAS study 979 of 1479 rice accessions (20). The others are GW5, bc10, rl14 and 980 OsNAC6, responsible for grain width, brittle culm, leaf rolling 981 and stress tolerance, respectively (18, 21-24). Interestingly, half 982 of these six genes were identified from natural mutants in contrast 983 to the fact that most functional genes were identified commonly 984 from T-DNA insertion and mutagen induced mutants (roughly 985 accounting for 90% of genes reported with known function). This 986 suggests that the identified genes from a pedigree analysis could 987 better reflect the real targets of selection in plant breeding than 988 the genes identified from artificial mutants. 989

Knockout phenotypes of candidate gene loci

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To determine whether a gene locus with unknown function has a phenotypic effect when knocked out, 57 of the 123 loci with unknown function were randomly sampled to do knockout by the CRISPR-cas9 system. Of these, 19 had knockout mutants, which were confirmed by PCR and Sanger sequencing. However, in the other 38 gene loci, no knockout mutants were obtained even after at least two independent transformations. We suspected that many of these genes are essential in callus development, so that no transformant survived. This possibility is supported by the observation that most (91.2%) of these genes had medium or high expression levels in callus (SI Appendix, Table S11).

As positive controls, we also attempted to knock out the 6 1002genes with known functions. As expected, 5 knockout mutants 1003 exhibited similar or stronger phenotypic changes compared to 1004 previous studies (18, 19, 21, 23, 24) (SI Appendix, Table S12). 1005 However, one of them, RL14, had no knockout mutant (see SI 1006 Appendix, Table S12). In a previous report, rl14, which carries 1007 a single amino acid mutation, exhibited severe leaf rolling and 1008 therefore RL14 may have essential functions, so that its knockout 1009 could not survive (22). In addition, as random controls, 10 genes 1010 were randomly sampled from the 1kb-300kb regions (SI Appendix, 1011 Table S13) adjacent to the retained ancestor blocks (which were 1012 shared by all 8 descendants of MH63). The near-neighbor con-1013 trols may be considered as conservative random controls for, 1014 unlike true random controls, these controls in part allow for 1015 possibly important position effects, such as the clustering of genes 1016 with similar expression profiles (25). In all 10 cases the knockout 1017 mutant showed no phenotypic changes (SI Appendix, Table S14), 1018 in contrast to 79% (15/19) of the unknown gene loci that showed 1019 1020 observable phenotypic changes when the gene was knocked out

(Table 2 and detailed changes in phenotypes and genotypes in SI Appendix, Table S15).

1023 High yield plants are typically dwarf, as dwarfism reduces 1024 investment into stalk, thereby potentially increasing investment 1025 into seeds. Therefore, we studied the growth difference between 1026 the mutated and unmutated version. We compared plant heights in knockout and wild type lines by the paired t-test (Table 4). 1028 As expected, the random control genes showed no difference 1029 in height between mutant and non-mutant versions (P = 0.42, 1030 95% confidence interval, -1.15cm, 2.54cm), while the positive 1031 controls showed a significant shorter height in mutants than in 1032 wild type (P = 0.017, confidence interval: -99.6 cm to -20.84 cm). 1033 Importantly, for the test group we also saw a strong dwarfism 1034 phenotype (P = 0.0013; 95% CI: -31.98 to -9.22 cm). As these 1035 were random samples from the 123 unknown gene loci, it implies that a high proportion of the 123 loci have a phenotype similar 1037 to that of the well-described positive control genes identified by the same method. However, the extent of the dwarfism is reduced in the test sample compared with the positive controls (t-test on percentage difference comparing positive control and test samples, P = 0.029, 95% CI: -67.82 -5.48). These genes may have weaker effects than the previously reported ones, and this may be why they have not been identified.

A gene of particular interest is Os10g0555100, as its knockout showed a different panicle architecture and a 23% reduction in height. Note that one of the reported genes, larger panicle (lp), showed an altered panicle architecture as well. The protein product may be a glycogenin glucosyltransferase (see ic4r.org), suggesting a possible role in controlling free glucose and glucose storage. This speculation, however, requires further analysis. Among the other genes some, such as Os10g0558850, had rolled leaves (Fig. 3) but a relatively modest reduction in plant height $(\sim 10\%)$. All the 15 unknown gene loci with knockout phenotypes have various protein-level motifs with unknown function, suggesting that the plant type and the high-yield phenotype are controlled by many types of genes.

Interestingly, the physically-proximal gene loci, although showing no sequence similarity, have similar functions. For example, 3 of the 6 gene loci on chromosome 1 (from Os01g0884400 to Os01g0930900) had knockouts resulted in late heading and 6 of the 11 loci on chromosome 10 (Table 2 and SI Appendix, Table S15) had knockouts resulted in dwarf phenotypes relative to the background line. This clustering mirrors the previously observed clustering of QTL signals (26). The clustering may reflect selection for coordinated gene expression or may possibly be owing to epistatic effects. Importantly, this result also suggests a strategy for finding genes with similar functions: if you have found one, investigate its neighbors.

Knockdown phenotypes of gene loci with no knockout transformant

1073 To investigate the 38 loci with no knockout mutants, we ran-1074 domly selected 26 loci to knock down their expression level, using 1075 the dCas9 knockdown technique (27). Similar to the knockout 1076 results, in 10 of the 26 loci (38.5%) no knockdown mutants 1077 were obtained due to the death of the transformed callus after 1078 hydromycin selection. Most of the 26 loci also have medium or 1079 higher expression levels in callus (SI Appendix, Table S16). More-1080 over, even in the 16 loci with knockdown transgenic plants, 10 1081 knockdown plants showed distinct negative phenotypic changes 1082 and 7 of them died during plant regeneration (Table 3 and SI Ap-1083 pendix, Fig. S5). As expected, the qRT-PCR study confirmed that 1084 the expression of these target loci in knockdown transformants 1085 was indeed down-regulated (SI Appendix, Fig. S6). These results 1086 suggest that most of the 38 candidate genes are essential genes in 1087 rice. 1088

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1089 Discussion

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1090 Determining the genes that explain complex traits has never been 1091 easy. The two much used methods, QTL and GWAS, have both 1092 led to important discoveries, but such analyses are typically very 1093 labor intensive. Indeed, during the past decades, much effort 1094 has gone into dissecting the genetic basis of high-yielding traits 1095 based on molecular linkage maps, e.g., the identification of many 1096 quantitative trait loci (QTLs) (28-31), but had identified rela-1097 tively few genes. The pedigree-based method that we expanded 1098 here has, for some cases, short-cut much of the effort. It requires 1099 a good pedigree and consistent directional selection, however. 1100 Confirmation of such results would until recently also have been 1101 verv time consuming, but CRISPR-Cas9 can greatly reduce the 1102 amount of work. In this study. We have not only identified the 1103 3 well known loci for the Green Revolution (the "green revolu-1104 tion gene" sdl, grain size-related gene GW5 and domestication 1105 gene lp), but also identified over 100 candidates. Among the 57 1106 candidate genes selected for knockout and knockdown studies, 1107 we found that many of them are essential genes or showed 1108 phenotypic effects. Thus, the pedigree approach seems to be 1109 highly efficient for identifying candidate genes that were subject 1110 to strong selection .. 1111

While the knockout analysis suggested a low false positive rate, the false negative rate, by contrast, is unknown and probably quite high as our filters are quite stringent. Indeed, when we look at two genes that failed to pass the diversity cutoff, we find that one of them resulted in phenotypic change when knocked out. This suggests that slight relaxation of the stringent filtering will result in more candidates, but potentially a higher false positive rate too. More generally, we do not know how many genes are essential for the rice Green Revolution. As a consequence, the method should then be considered a technique to greatly enrich for selectively relevant genes rather than a method for an exhaustive search.

This study showed that rice is unusually well suited to this pedigree method. First, the well-documented pedigree information can be used to calculate the expected proportions of blocks (or loci) being transmitted from an ancestor to a descendant (e.g., Fig. 1A). By comparing the expected and observed proportions, the gene loci that were most likely to have been the target of artificial selection could then be identified. For example, the probability of a DGWG block appearing in all of the eight MH63 descendants was estimated to be nearly zero. Thus, if a block is observed in the re-sequencing data, it was very likely subjected to strong artificial selection. Second, from the relationships in a pedigree, SNP markers can be verified and corrected by comparing the sequences of parents and offspring between generations (demonstrated in Fig. 2 and SI Appendix, Fig. S1). In rice we are fortunate to have access to the stocks of the prior generations. Third, pedigree analysis focuses on tracing relatively longer blocks from the parents to the offspring instead of single SNPs or genes. It is therefore not difficult to identify selected targets. Finally, the CRISPR-Cas9 system provides an effective way of gene knockout to find a set of genes relevant to complex traits. In conclusion, our approach should be useful for many breeding projects.

Our choice of our model organism was not just motivated by the fact that the conditions for pedigree analysis were met, but also by the enormous impact of the Green Revolution, as indicated by the generation of high-yielding plant types through breeding. The introduction of dwarfing genes has resulted in plants that possess short and strong stalks, which are less liable to lodging. The stability of shorter plants dramatically reduces the need for photosynthetic investment in the stem. Assimilates are then redirected to grain production, resulting in a better plant type and increased yield (32). The candidate genes identified

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in this study will be useful for understanding the underlying 1157 mechanism of this physiology. 1158

1159 Importantly then, we have identified many new genes respon-1160 sible for high-yield, an economically most important trait. Most of 1161 these gene loci have not yet been functionally annotated, although 1162 a few of them belong to the β -expansin family or contain a zinc 1163 finger domain, which are known to play an important role in plant height, flower development, and light-regulated morphogenesis 1164 1165 (33-35). We highlighted Os10g0555100, the knockout of which 1166 showed a different panicle architecture and a 23% reduction in height. We also note that our results suggest that the genes 1167 1168 identified from cultivated lines in a pedigree could better reflect 1169 the real targets in plant breeding than the genes identified from artificial mutants. Our catalogue of 123 unannotated gene loci 1170 1171 provides choices for downstream analysis. Our knockout and 1172 knockdown study of about half of these loci revealed that most 1173 of the genes in these loci are essential for rice phenotypes or 1174 for normal growth. Among the 159 genes we identified, there 1175 are at least 31 yield related genes, including 15 identified by 1176 knockout, 10 by knockdown and 6 previously reported. This 1177 proportion (19.5%) is significantly higher than the expectation (2.33 in 159) = 1.5%) based on the reported yield related genes 1178 in the rice genome (p < 0.001, $\chi 2=334$, df=1, Chi-squared test with Yate's correction, see details in the SI Appendix, SI Materials 1179 1180 and Methods). However, the alleles contributing to the Green 1181 1182 Evolution are not necessarily null alleles, so our knockout and 1183 knockdown studies did not directly test the contribution of allelic 1184 changes to the Green Revolution. Gene replacements in IR8 or 1185 MH63 would directly reveal the contributions of the alleles, but 1186 IR8 and MH63 are difficult to transform and gene replacement is currently difficult in rice. 1187

Our results also highlight clustering of unrelated genes with similar yield-associated phenotypes in the genome. This observation is of relevance to those hunting for complex trait genes and for those interested in genome evolution. For the former, it suggests that looking at neighbors of functionally relevant genes would be an effective way to look for functionally related genes. The clustering may reflect epistasis between genes or selection for co-expression. Previous QTL analysis also suggested that genes of similar phenotypic effects tend to cluster together (26), but this could also reflect allelic versions of control elements for a single gene. The fact that the knockouts of the clustered genes tend to have similar phenotypes suggests it is not the case. 1188

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Methods

Detailed materials and methods are outlined in SI Appendix, SI Materials and Methods

Plant materials and sequencing

The seeds of all rice accessions were obtained from International Rice Research Institute (IRRI) and China National Rice Research Institute (CN-RRI) (Dataset S1). Pedigree information was obtained from the germplasm databases of IRRI and CNRRI. All rice varieties were grown in the paddyfield. DNA samples were prepared from fresh leaves of a single plant using the Cetyltrimethyl Ammonium Bromide (CTAB) method and were sequenced at BGI-Shenzhen. Briefly, paired-end sequencing libraries with an insert size of ~500 bp were constructed for each plant, following the BGI-Shenzhen's instructions, and 2×100bp paired-end reads were generated on an Illumina HiSEq 2000. The sequencing reads of the 30 rice accessions have been deposited to the NCBI Short Read Archive under the Accession Numbers PRINA271253 and SR1060330. Indica cultivar 9311 callus RNA-seq data was downloaded from NCBI, BioProject PRJNA117345, SRR037711~SRR037724.

Construction of CRISPR Genome-Editing Vectors and target gene loci knockout

1216 For each target locus, gRNAs were designed to target specific sites at the 1217 beginning of exons to cause a frame shift mutation. For each target, a pair 1218 of DNA oligonucleotides with appropriate cloning linkers were synthesized (BGI, Inc). Each pair of oligonucleotides were phosphorylated, annealed, and 1219 then ligated into Bsal-digested pRGEB31 vectors (addgene No.7722) (36). 1220 After transformation into Escherichia coli DH5-alpha, the resulting constructs 1221 were purified with Plasmid Mini kit (Genebase. Inc) for subsequent use in 1222 rice callus transformation. We selected Kasalath and Wuyungeng24 to be 1223 the background because they have a high transformation success rate while IR8 and MH63 are difficult to transform. Besides, Kasalath has a rather high 1224 stature and it is easy to observe when it becomes dwarf. Each construct was
transformed into calli of Kasalath (an Indica) or Wuyugeng24 (a Japonica)
by the method reported in a previous study (37). About ten transformed
individuals were produced in two recipients for each vector (details in *SI Appendix*, Table S13, Table S14 and Table S15).

Genotype confirmation and phenotype observation

The transgenic plants were examined under natural field conditions in the Experimental station of Nanjing University, Nanjing, China. For each plant, genomic DNA was extracted from fresh leaves by the CTAB method. In order to get double knockout mutants, we amplified the target region by PCR and confirmed the genotypes by Sanger sequencing. Primers were designed to make PCR products of ~1kb that contain the target sites. The results showed that 82.1% of transgenic plants had a knockout allele and 79.5% had double knockout mutants. Phenotypes of the mutants were observed at different stages. Plant phenotypes were observed every three days to determine the changes in comparison with wild type rice plants. Plant height was measured after the stage of heading. Fertility and spike shape were observed when seeds were mature.

Footnotes

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Author contributions: D.T, L.D.H. and W-H.L. designed the experiments and analyses. J.H. and J.L. organized all aspects of the project, J.H., J.L., L.W. and S.Y. analyzed the sequence data. J.H., J.L. and J.Z. prepared plant materials and performed experimental confirmations. J.H., D.T., L.D.H. and W-H.L. wrote the paper.

The authors declare no conflict of interest.

Data deposition: Sequences have been deposited in the Sequence Read Archive, www.ncbi.nlm.nih.gov/sra (accession no. SRP051581).

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