RESEARCH ARTICLE SUMMARY

PLANT SCIENCE

Enhanced sustainable green revolution yield via nitrogen-responsive chromatin modulation in rice

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INTRODUCTION: The green revolution of the 1960s boosted cereal crop yields in part through widespread adoption of semi-dwarf plant varieties. The beneficial semi-dwarfism is respectively conferred in wheat and rice green revolution varieties by mutant Reduced height-1 (Rht-1) and semi-dwarf1 (sd1) alleles. These alleles cause accumulation of growth-repressing DELLA proteins, the normal forms of which are characterized by the presence of an Asp-Glu-Leu-Leu-Ala amino acid motif. Resultant semi-dwarf plants resisted lodging but required high nitrogen fertilizer inputs to maximize yield. Normally, gibberellin promotes growth by stimulating DELLA degradation as regulated by the gibberellin receptor GID1 (GIBBERELLIN INSENSITIVE DWARF1), the F-box protein GID2 (GIBBERELLIN INSENSITIVE DWARF2), and the SCF (Skp, Cullin, F-box-containing) ubiquitin ligase complex. Nitrogen fertilization-induced increase in grain yield is determined by the integration of three components (tiller number. grain number, and grain weight), but exogenous application of gibberellin reduces tiller number in rice. Here, we asked how nitrogen fertilization affects the gibberellin signaling that regulates rice tillering. Nitrogen fertilization promotes crop yield, but overuse in agriculture degrades the environment. A future of sustainable agriculture demands improved nitrogen use efficiency.

RATIONALE: Increased tillering, nitrogen fertilization, and high-density planting all contribute to the high yield typical of green revolution rice varieties. Increases in tiller number despite reduced nitrogen fertilization could help to sustain vield while reducing the environmental impact of agriculture. To investigate the effect of gibberellin on nitrogen-promoted rice tillering, we used genetic screening to identify a mutation in the ngr5 (nitrogen-mediated tiller growth response 5) gene. Plants carrying the ngr5 mutant displayed fewer tillers; tiller number was insensitive to nitrogen supply. Further genetic and biochemical studies defined the mechanisms underlying the interaction between nitrogen- and gibberellinmediated effects on tiller number.

RESULTS: We found that increased nitrogen supply enhanced transcription and abundance of the rice APETALA2-domain transcription factor encoded by an *NGR5 (NITROGEN-MEDIATED TILLER GROWTH RESPONSE 5)* allele. NGR5 interacts with a component of the polycomb repressive complex 2 (PRC2) and alters the genome-wide histone H3 lysine 27 trimethylation (H3K27me3) pattern response to





changes in nitrogen availability. The extent of this alteration was reduced in *ngr5* plants or by gibberellin treatment. RNA sequencing and chromatin immunoprecipitation (ChIP)–polymerase chain reaction analysis showed that an increase in nitrogen supply reduced the abundance of mRNAs specified by strigolactone signaling and

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Read the full article at http://dx.doi. org/10.1126/ science.aaz2046 other branching-inhibitory genes [such as *Dwarf14* (*D14*) and *squamosa promoter binding proteinlike-14* (*OsSPL14*)] in a dosage-dependent manner. Lack of *D14* or *OsSPL14*

function was epistatic to *ngr5* in regulating rice tillering. We next found that the DELLA-mediated enhancement of nitrogen-induced tiller number, typical of green revolution rice varieties, was abolished in plants with the *ngr5* mutation.

These observations suggest that NGR5-driven recruitment of PRC2 promotes repressive H3K27me3 modification of target branchinginhibitory genes, thus causing an increase in tiller number. On the other hand, a nitrogen-induced NGR5-dependent increase in tiller number is enhanced in green revolution rice varieties, and this effect is inhibited by gibberellin treatment. Although NGR5 abundance is negatively associated with gibberellin amount, gibberellinpromoted destabilization of NGR5 is neither dependent on nor downstream of gibberellininduced DELLA destruction. Moreover, NGR5 interacts with the gibberellin receptor GID1 and DELLA proteins; this suggests that gibberellinpromoted proteasomal destruction of NGR5 is not due to gibberellin-promoted destruction of DELLAs, but is due to a gibberellin-potentiated interaction of NGR5-GID1, leading to polyubiquitination of NGR5 and subsequent destruction in the proteasome. Accumulation of DELLA proteins competitively inhibited the GID1-NGR5 interaction, thus stabilizing NGR5 by reducing its gibberellin-GID1-mediated destruction.

CONCLUSION: We conclude that nitrogen fertilization alters genome-wide reprogramming of H3K27me3 methylation via NGR5-dependent recruitment of PRC2. In rice, methylation represses genes that inhibit tillering and consequently promotes an increase in tiller number. NGR5 is a target of gibberellin-GID1-promoted proteasomal destruction. Modulation of competitive interactions among NGR5, DELLA proteins, and GID1 enables enhanced grain yield in elite rice varieties despite reduced nitrogen fertilizer inputs. Such shifts in yield and input use could promote agricultural sustainability and food security.

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RESEARCH ARTICLE

PLANT SCIENCE

Enhanced sustainable green revolution yield via nitrogen-responsive chromatin modulation in rice

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Because environmentally degrading inorganic fertilizer use underlies current worldwide cereal yields, future agricultural sustainability demands enhanced nitrogen use efficiency. We found that genome-wide promotion of histone H3 lysine 27 trimethylation (H3K27me3) enables nitrogen-induced stimulation of rice tillering: APETALA2-domain transcription factor NGR5 (NITROGEN-MEDIATED TILLER GROWTH RESPONSE 5) facilitates nitrogen-dependent recruitment of polycomb repressive complex 2 to repress branching-inhibitory genes via H3K27me3 modification. NGR5 is a target of gibberellin receptor GIBBERELLIN INSENSITIVE DWARF1 (GID1)–promoted proteasomal destruction. DELLA proteins (characterized by the presence of a conserved aspartate-glutamate-leucine-leucine-alanine motif) competitively inhibit the GID1-NGR5 interaction and explain increased tillering of green revolution varieties. Increased NGR5 activity consequently uncouples tillering from nitrogen regulation, boosting rice yield at low nitrogen fertilization levels. NGR5 thus enables enhanced nitrogen use efficiency for improved future agricultural sustainability and food security.

he agricultural green revolution of the 1960s enhanced cereal crop yields, fed a growing world population, and was in part due to increased cultivation of semidwarf green revolution varieties (1-4). The beneficial semi-dwarfism is conferred by mutant alleles at the wheat Reduced height-1 (Rht-1) (5, 6) and rice Semi-dwarf1 (SDI) (7, 8) loci that enhance the activity of growth-repressing DELLA proteins (DELLAs). Normally, the phytohormone gibberellin stimulates the destruction of DELLAs (9, 10), thus promoting plant growth. However, the mutant wheat DELLA protein Rht-1 likely resists gibberellin-stimulated destruction (5), whereas the rice sd1 allele reduces gibberellin abundance (11, 12) and increases accumulation of the rice DELLA protein SLR1 (SLENDER RICE1) (13). The result is plants that are shorter than normal, which, because they are shorter, are more resistant to lodging (the flattening of plants by wind and rain) (4). However, green revolution rice varieties require a high-nitrogen fertilizer supply to achieve maximum vield potential, and the drive toward

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increased agricultural sustainability necessitates reduced nitrogen fertilizer use (13). Grain yield is the sum of the multiplicative integration of three major components [tiller numbers per plant, grain numbers per panicle, and 1000-grain weight (14)], and an increased tillering ability in high-density planting conditions contributes to the high-yield properties of green revolution rice varieties (1, 15). Further increase in tiller (lateral branch) numbers at low nitrogen supply is therefore important for future agricultural sustainability and is a key cereal breeding goal. Here, we first define the mechanisms underlying the promotive effects of nitrogen on tiller bud outgrowth. We then show how genetic modulation of these mechanisms can enable increased grain yield of green revolution varieties despite reduced nitrogen input, thus advancing agricultural sustainability.

Nitrogen promotes rice tillering via NGR5

We found that the tiller number per plant of *indica* rice variety Nanjing6 (NJ6) increased with increasing nitrogen supply (Fig. 1A). Additional effects of increased nitrogen on NJ6 included increases in grain number (per panicle) and yield (per plant) (13) (fig. S1, A to C). NJ6-sd1 (a NJ6 isogenic line containing the sd1 allele) also displayed nitrogen-dependent tiller number increases: increased tiller numbers per plant under different nitrogen fertilization levels, with tiller numbers being consistently higher in NJ6-sd1 than in NJ6 (Fig. 1A). The *Rht-B1b* (formerly termed *Rht-1*) allele conferred similar properties on wheat





(versus the Rht-Bla control allele; Fig. 1B and fig. S1, D to F). In contrast, either exogenous gibberellin treatment or overexpression of the rice gibberellin receptor GID1 (16) inhibited nitrogen-promoted tillering (fig. S2). Thus, the enhanced DELLA function typical of green revolution varieties increases nitrogen-induced promotion of tiller number. Further analysis showed that nitrogeninduced increases in tiller number in the elite sd1-containing indica rice variety 9311 (fig. S3, A and B) were not due to nitrogen-responsive increases in numbers of lateral buds, but rather to increased numbers of buds initiating outgrowth and tiller branch extension (17) (fig. S3. C to E).

We next screened an ethyl methane sulfonate (EMS)-mutagenized 9311 population for mutants displaying an altered tiller number nitrogen response. Among such mutants, ngr5 (nitrogen-mediated tiller growth response 5) displayed a reduced tiller number that was insensitive to changes in nitrogen supply (Fig. 1, C and D). Map-based cloning (fig. S4, A and B) and genetic complementation (Fig. 1, C and D, and fig. S4, C and D) revealed the NGR5 allele to encode an APETALA2 (AP2)-domain transcription factor [NGR5, previously known as SMOS1 (SMALL ORGAN SIZE1) and RLA1 (REDUCED LEAF ANGLE1)] (18-20), thus identifying an unknown function for NGR5 in nitrogen-responsive tillering regulation. The *ngr5* allele carries a $G \rightarrow A$ nucleotide substitution conferring a $Gly \rightarrow Arg$ mutant protein (fig. S4B), which fails to complement ngr5 phenotypes (fig. S4, C to F). In addition to its effect on tiller number (Fig. 1, C and D, and fig. S4D), NGR5 is required for nitrogeninduced promotion of panicle branching and grain number (fig. S4, E and F). Accordingly, whereas 9311 grain yield per plot increased progressively with increasing nitrogen supply (13), this effect was abolished in ngr5 plants (fig. S4G). Further analysis showed that lack of NGR5 (in ngr5 plants) had no effect on the formation of tiller buds (lateral bud initials; fig. S3C) but reduced the number of buds initiating lateral outgrowth and tiller branch extension (fig. S3, D and E), thus confirming that nitrogen-responsive regulation of tillering is dependent on NGR5.

We next found that an increasing nitrogen supply increased NGR5 abundance at both mRNA and protein levels (Fig. 1, E and F). First, increasing nitrogen supply increased NGR5 mRNA abundance, and this effect was abolished in *ngr5* plants (Fig. 1E). Second, although nitrogen supply had no effect on NGR5-HA (hemagglutinin-tagged fusion gene) mRNA abundance in plants transgenically expressing *p35S::NGR5-HA* (fig. S5), accumulation of NGR5-HA fusion protein increased with increasing nitrogen supply (Fig. 1F). Furthermore, NGR5 positively regulated tillering over a wide expression range, because the *p35S::NGR5* transgene increased 9311 tiller number (thus mimicking the effect of increasing nitrogen supply on tillering capacity; Fig. IG). We conclude that nitrogen promotes increased NGR5 abundance, which in turn promotes tiller bud outgrowth. In addition, because *ngr5* suppressed the *sd1*-conferred tillering phenotype of 9311 (Fig. 1D), NGR5 is necessary for the DELLA-promoted increase in tiller number characteristic of green revolution varieties.

NGR5 represses branching-inhibitory genes

RNA sequencing (RNA-seq) analysis next revealed that lack of NGR5 causes genome-wide change in mRNA abundance, with multiple differentially expressed genes displaying an increase in mRNA abundance in ngr5 (fig. S6A). Further gene set enrichment analysis revealed a correlation between genes up-regulated in ngr5 and the set of H3K27me3 (histone H3 lysine 27 trimethylation)-marked genes already known to be normally repressed by histone modification (fig. S6B), with H3K27me3 marks occurring at both TSS (transcription start site) and gene body regions of ngr5-upregulated genes (fig. S6C). These results suggest that NGR5 may be involved in PRC2 (polycomb repressive complex 2)-mediated epigenetic repression. Among genes up-regulated in ngr5 (table S1), we identified D14 [Dwarf14, encoding the receptor for the phytohormone strigolactone (SL)] (21), D3 [Dwarf3, encoding the F-box component of the Skp, Cullin, F-box-containing (SCF) ubiquitin ligase that targets the DWARF53 repressor of SL signaling for proteasomal destruction] (22-24), OsTB1 (TEOSINTE BRANCHEDI, encoding a TCP domain transcription factor) (25), and OsSPL14 (squamosa promoter binding protein-like-14, encoding an SBP-domain transcription factor) (26, 27) genes, all of which are already known to inhibit lateral branching and tiller number. Quantitative real-time polymerase chain reaction (qRT-PCR) analysis showed that high nitrogen supply reduced the abundances of mRNAs specified by these shoot branchinginhibitory genes, and this effect was abolished by lack of NGR5 function (in ngr5; fig. S6D). In addition, we found that lack of D14 or OsSPL14 function (conferred by d14 or osspl14 alleles) (28, 29) is epistatic to ngr5 in regulating lateral branching (fig. S7). Thus, D14 and OsSPL14 function downstream of NGR5, and NGR5 mediates nitrogen-promoted increase in tiller number by repressing the inhibitory functions of D14 and OsSPL14 (and likely of other) branching-regulatory genes.

Chromatin immunoprecipitation–PCR (ChIP-PCR) experiments revealed binding of NGR5-HA to gene body and promoter regions of *D14* and *OsSPL14* [fig. S6E; confirmed in EMSA (electrophoretic mobility shift assays), fig. S6F].

Furthermore, the extent and effect of binding on NGR5-target gene repression correlated with increasing nitrogen supply (fig. S6, G and H). D14 mRNA abundance decreased with increasing nitrogen supply in NGR5 (but not in ngr5; fig. S6G), and the extent of NGR5 binding and the level of H3K27me3 modification at D14 were correspondingly increased in a nitrogen-dependent manner (but not in ngr5; fig. S6G). Similar effects were observed for OsSPL14 (fig. S6H), which suggests that NGR5 promotes tillering in response to increasing nitrogen supply by binding to target branching-inhibitory genes, thus causing their repression through regulation of H3K27me3 modification.

NGR5 recruits PRC2 for H3K27me3 deposition

To determine how NGR5 regulates nitrogenpromoted H3K27me3 modification, we first performed a yeast two-hybrid screen for NGR5 interactors, identifying LC2 (leaf inclination2, a component of the PRC2 complex) (30) among many others (table S2). NGR5-LC2 interactions were confirmed in bimolecular fluorescence complementation (BiFC; Fig. 2A) and coimmunoprecipitation (Co-IP; Fig. 2B) experiments. Furthermore, a CRISPR/Cas9-generated LC2 reduced-function allele (lc2; fig. S8A) was shown, like ngr5, to abolish nitrogen-promoted increase in tiller number (Fig. 2, C and D). lc2 also suppressed the increased tiller number conferred by p35S::NGR5 (Fig. 1G and Fig. 2, C and D), whereas lack of D14 or OsSPL14 function (conferred by d14 or osspl14) was epistatic to lc2 (fig. S9). Taken together, these results suggest that NGR5-dependent nitrogen-promoted increase in tiller number depends on LC2 (PRC2 complex) function. Because PRC2 regulates genome-wide patterns of H3K27me3 methylation, we next conducted genome-wide surveys of H3K27me3 methylation in response to varying nitrogen supply. Although increasing nitrogen supply altered the genome-wide H3K27me3 methylation pattern, the extent of this alteration was reduced in ngr5 (Fig. 2E), which suggests that nitrogen-mediated genomewide reprogramming of H3K27me3 methylation is NGR5-dependent.

We next performed ChIP-sequencing experiments and identified a total of 453 binding sites shared in common by NGR5 and LC2 (Fig. 2F and tables S3 and S4). Further analysis identified potential target-site recognition motifs shared by NGR5 and LC2 (Fig. 2, G and H), with a predominant shared GCCGCC motif being common in the gene body regions of both *ngr5*-up-regulated and nitrogen-induced genes (Fig. 2E). Accordingly, increasing nitrogen supply progressively reduced *D14* and *OsSPL14* mRNA abundance in wild-type plants, and these effects were abolished in *lc2* plants (Fig. 2, I and J, top), just as they were in *ngr5* plants (fig. S6, G and H). Furthermore, increasing nitrogen





supply progressively increased H3K27me3 modification of both *D14* and *OsSPL14* in wild-type plants, and these effects were also abolished in *lc2* plants (Fig. 2, I and J, bottom). Taken together, these observations suggest that NGR5-driven recruitment of the PRC2 complex (of which LC2 is a component) to *D14* and *OsSPL14* results in repressive H3K27me3 modification of these genes in response to increased nitrogen supply, thereby promoting bud outgrowth and increasing tiller number.

NGR5 is a target of gibberellin receptor GID1

As shown above, nitrogen-induced increase in tiller number was enhanced in green revolu-

tion varieties (Fig. 1, A and B), and this effect was inhibited by exogenous gibberellin treatment (fig. S2A). Analysis of both RNA-seq and ChIP sequencing revealed multiple common gene targets to be co-regulated by NGR5 and gibberellin treatment (tables S5 and S6). Furthermore, gibberellin treatment altered the change in genome-wide H3K27me3 modification pattern due to increasing nitrogen supply in a manner similar to the alteration conferred by *ngr5*, whereas a partially restored H3K27me3 modification pattern was induced by treatment with paclobutrazol (PAC, an inhibitor of gibberellin biosynthesis; Fig. 2E). In addition, gibberellic acid (GA₃), like *ngr5* and *lc2*, inhibited the nitrogen-dependent increase in H3K27me3 modification and consequent repression of expression of shoot branching inhibitor genes such as *D14* (Fig. 2K) and *OsSPL14* (Fig. 2L). These observations suggest the existence of a mechanistic link between nitrogen- and gibberellin-mediated effects on tiller number.

In canonical gibberellin signaling, gibberellin binds its receptor GID1, thus recruiting DELLAs for polyubiquitination by the F-box protein GIBBERELLIN INSENSITIVE DWARF2 (GID2) and the Skp, Cullin, F-box-containing (SCF) ubiquitin ligase complex (SCF^{GID2}) and subsequent destruction in the 26S proteasome, thus promoting plant growth (9, 10, 16, 31-34). We next found that reduced GID1 function in a NJ6-gid1-10 mutant (gid1 loss-of-function mutant; fig. S8B) led to an increased tiller number above that of NJ6 controls in both high and low nitrogen supply (similar to NJ6-sd1; Fig. 3, A and B). Conversely, transgenic NJ6-sd1 plants overexpressing GID1 under the control of the cauliflower mosaic virus (CaMV) 35S promoter exhibited nitrogen-insensitive responses, with lower tiller number than in nontransgenic controls (Fig. 3B). Although gibberellin repressed tiller number in both NJ6 and NJ6-sd1 plants, it had no effect on tiller number in NJ6-gid1-10 or NJ6-sd1-ngr5 plants, nor in NJ6-sd1 plants overexpressing GID1 (Fig. 3B). Furthermore, either gibberellin-induced inhibition or GID1mediated repression of tillering mimicked the effect of ngr5 (Fig. 3B). These results suggest that gibberellin-GID1-mediated repression of tiller number is dependent on the nitrogenregulated function of NGR5.

We next found that NGR5 abundance is negatively associated with gibberellin level: NGR5-HA accumulation was increased in relatively gibberellin-deficient NJ6-sd1 plants (versus NJ6), whereas it was reduced by exogenous gibberellin treatment (Fig. 3C). Conversely, a gibberellin-mediated decrease in NGR5-HA abundance was inhibited by treatment with the proteasome inhibitor MG132 (carbobenzoxy-Leu-Leu-leucinal), such that NGR5-HA accumulation was increased above that of NJ6-sd1 plants (Fig. 3C). Accordingly, Western blot analysis detected the accumulation of polyubiquitinated NGR5-HA in the presence of MG132 (Fig. 3D), which suggests that gibberellin promotes polyubiquitination and subsequent proteolysis of NGR5 in the 26S proteasome. In addition, gibberellin-induced degradation of NGR5-HA was inhibited in the NJ6-gid1-10 mutant (Fig. 3E), indicating that gibberellininduced promotion of NGR5 polyubiquitination and proteasome destruction is dependent on the GID1 function.

Gibberellin responses are conventionally considered to be activated by GID1-mediated destruction of DELLAS (*9*, *10*). However, we found that gibberellin-mediated degradation of NGR5-HA occurs either in the absence of



Fig. 3. Gibberellin-GID1-SCF^{GID2} targets NGR5 for destruction. (A) Mature plants grown in low (90 kg/ha; LN) versus high (180 kg/ha; HN) nitrogen supply. Scale bar, 20 cm. (B) Tiller number. Data are means ± SE (n = 20). (C) Immunodetection of NGR5-HA. (D) Immunodetection of poly-ubiquitinated NGR5-HA. (E) Accumulation of NGR5-HA. (F) Effects of gibberellin on NGR5-HA and SLR1 accumulation. In (C) to (F), total protein was extracted from tiller buds of 3-week-old plants treated for 4 hours with 1µM GA₃ and/or 100µM MG132; HSP90 serves as loading control. (G and J) SFLC assays. nLUC-tagged GID1 (G) or nLUC-tagged GID2 (J) was co-transformed into tobacco leaves along with cLUC-targeted NGR5. (H and K) Co-IP assays. Flag-tagged GID1 (H) or Flag-tagged GID2 (K) was co-transformed into rice protoplasts along with HA-targeted NGR5.
(I) Pull-down assays. (L) In vitro ubiquitination assay. The immunoprecipitated GID2-Flag proteins transiently expressed in rice protoplasts were used in an in vitro ubiquitination reaction in the presence of E1, E2 (UbcH5B), His-Ub, and GST-NGR5. (M) Gibberellin-GID1-SCF^{GID2} destabilizes GST-NGR5. Lysates from NJ6, NJ6-*gid1-c1*, and NJ6-*gid2-c1* plants were co-incubated with GST-NGR5 in the presence of 100µM GA₃ and 100µM MG132. The lysates were harvested at various incubation times and immunoblotted to assess the accumulation of NGR5 and HSP90.

DELLAs in a loss-of-function *slr1* mutant (Fig. 3E) (11) or in the presence of the high-level DELLA accumulation conferred by the *Slr1-d6* gain-of-function mutation (Fig. 3F) (35). Although the mutant SLR1 DELLA encoded by *Slr1-d6* was relatively resistant to gibberellin-

mediated destruction, NGR5-HA was still destabilized by exogenous gibberellin treatment (Fig. 3F). Thus, gibberellin-promoted destabilization of NGR5, although dependent on GID1, is neither dependent on nor downstream of gibberellin-induced destruction of DELLAs. We therefore explored the possibility of an alternative, previously unknown, DELLA-independent mechanism whereby the SCF^{GID2} E3 ubiquitin ligase directly mediates gibberellin-promoted destruction of NGR5, finding that GID1 interacts directly with NGR5 [as assayed by both split firefly luciferase complementation (SFLC) and Co-IP; Fig. 3, G and H] and that the strength of this interaction is potentiated by increasing concentrations of gibberellin (Fig. 3, H and I). Thus, as with the DELLAs, gibberellin enhances the interaction between NGR5 and GID1, thereby identifying NGR5 as a potential alternative substrate for GID1promoted polyubiquitination.

Although NGR5 lacks the specific DELLA motif that enables the GID1-DELLA interaction (33, 34), we found a motif within the AP2-R2 (repeated units 2) (18) domain of NGR5 to enable the GID1-NGR5 interaction (fig. S10). Furthermore, NGR5 also interacted with the GID2 F-box component of the SCF^{GID2} E3 ubiquitin ligase that normally targets SLR1 for destruction in the 26S proteasome (Fig. 3, J and K). Accordingly, an in vitro ubiquitination assay showed that GST (glutathione S-transferase)-NGR5 fusion protein is polyubiquitinated by GID2-Flag (Asp-Tyr-Lys-Asp-Asp-Asp-Lys peptide) fusion protein in the presence of ubiquitin-activating enzyme E1, ubiquitinconjugating enzyme E2, and ubiquitin, but not in the absence of GID2-Flag (Fig. 3L), which suggests that NGR5 is a substrate of the $\mathrm{SCF}^{\mathrm{GID2}}$ E3 ubiquitin ligase. Additional time-course experiments showed that gibberellin promotes the progressive degradation of GST-NGR5 but that this degradation is inhibited both by MG132 and in gid1-c1, a CRISPR/Cas9-generated GID1 loss-of-function mutant (Fig. 3M and fig. S8B). Finally, lack of GID2 function (in a gid2-c1 mutant; fig. S8C) also inhibits gibberellinmediated degradation of GST-NGR5 (Fig. 3M). We conclude that the gibberellin-mediated regulation of NGR5 is not due to gibberellinpromoted destruction of DELLAs, but is due to a previously unknown direct and gibberellinpotentiated interaction of NGR5-GID1, leading to polyubiquitination of NGR5 by the SCF^{GID2} E3 ubiquitin ligase and subsequent destruction in the proteasome.

DELLA-NGR5 modulation of tiller N response

We next found that NGR5 interacts directly with SLR1 [in yeast two-hybrid screens (table S2) and in BiFC and Co-IP assays (fig. S11, A and B)]. Nonetheless, we additionally found that the LC2-NGR5 interaction is not inhibited by the presence of SLR1 (fig. S12), which suggests that the SLR1-NGR5 interaction does not directly interfere with the LC2-NGR5 interaction that determines NGR5 function. Further experiments showed that the LHR1 (leucine heptad repeat 1) motif of the DELLA protein is necessary for the NGR5-SLR1 interaction

(fig. S11C). Thus, in addition to both being substrates of gibberellin-GID1-SCFGID2, SLR1 and NGR5 interact directly with one another. With the LHR1 motif being conserved in all the GRAS [derived from three initially identified members, GAI (gibberellin-insensitive), RGA (repressor of ga1-3), and SCARECROW] proteins, we next found that NGR5 interacts with two additional GRAS proteins [DWARF AND LOW-TILLERING (DLT) and MONOCULM1 (MOC1)] previously shown to regulate tiller number (17, 19, 36) (fig. S13A). The competitive nature of the SLR1-NGR5 relationship with respect to GID1 (and hence with respect to GID1-promoted destruction) (fig. S13B) caused accumulation of NGR5-HA (conferred by *p35S::NGR5-HA*) to further increase the accumulation of SLR1 in 9311 (fig. S13C).

We next determined whether competitive SLR1-NGR5-GID1 relationships also condition gibberellin and nitrogen effects on tiller number. As shown in Fig. 1B, the enhanced DELLA function conferred by the Rht-B1b allele results in increased tiller number. We therefore tested the possibility that the effect of Rht-B1b on tiller number might be due to differential effects on NGR5 stability. Accordingly, using FRET (Förster resonance energy transfer) analvsis, we found that the extent of the interaction between GID1 and NGR5 is reduced by the presence of Rht-B1b (Fig. 4, A and B). We then confirmed the expectation that a reduced GID1-NGR5 interaction in the presence of Rht-B1b reduces the rate of proteasome-dependent GST-NGR5 destruction (Fig. 4C), and we found similar reductions to be conferred by accumulation of wild-type Rht-Bla or SLR1 proteins (Fig. 4, A to C). Further comparative studies showed His-NGR5 (His-tagged fusion protein) destruction to be more rapid than that of His-SLR1 (His-tagged fusion protein; Fig. 4D). Thus, although gibberellin-promoted NGR5 destruction was DELLA-independent (Fig. 3, E and F), competition between NGR5 and SLR1 for GID1 interaction reduced the extent of NGR5-GID1 interaction (fig. S13B). Moreover, the abundance of NGR5-GFP (green fluorescent protein) fusion protein (conferred by p35S::NGR5-GFP) was increased with PAC treatment but was reduced in response to combined gibberellin and PAC treatments (fig. S13, D and E). This is also consistent with the observations that ngr5 exhibits a higher ratio of gibberellin-induced leaf sheath growth, whereas transgenic plants overexpressing NGR5-HA display reduced sensitivity to PAC treatment relative to wild-type controls (fig. S14). We conclude that the enhanced DELLA function characteristic of both wheat and rice green revolution varieties competitively inhibits the GID1-NGR5 interaction, thus stabilizing NGR5 by reducing gibberellin-GID1-mediated destruction.

To determine whether DELLA promotion of rice tillering is NGR5-dependent, we generated

9311 NILs (near-isogenic lines) carrying various combinations of different *SD1*, *GID1*, and *NGR5* alleles. Although the increased SLR1 accumulations in both 9311-*NGR5-gid1-10* and 9311-*NGR5-sd1* increased the tiller numbers of plants grown in either low or high nitrogen supply (versus 9311-*NGR5-SD1*), there was almost no difference in tiller number when 9311-*ngr5-sd1*, 9311-*ngr5-SD1*, and 9311-*ngr5-gid1-10* were compared (Fig. 4E). Thus, the DELLA-mediated enhancement of nitrogen-induced tiller number increase typical of green revolution rice varieties is dependent on NGR5 function. Accordingly, comparisons of NGR5-regulated mRNA abundance and H3K27me3 modification of branching-inhibitory *D14* (Fig. 4F) and *OsSPL14* (Fig. 4G) genes in 9311 (containing *sd1*) versus 9311-*SD1* revealed mRNA abundance and modification status at 0.6N in 9311-*SD1* to be roughly equivalent to that at 0.2N in



Fig. 4. Balanced DELLA-NGR5 interactions improve nitrogen use efficiency. (**A**) FRET images. Scale bars, 200 μ m. (**B**) Mean N-FRET data for GID1-CFP and NGR5-YFP channels. Data are means ± SE (n = 5). (**C**) Time-course analysis of GST-NGR5 degradation. HSP90 serves as loading control. (**D**) Degradation rate of His-NGR5 and His-SLR1. (**E**) Tiller number. (**F** and **G**) Relative mRNA abundance and H3K27me3 modification of *D14* (F) and *OSSPL14* (G). Transcript abundance values are relative to that of 9311-*sd1* in 1N (set to 1). Data are means ± SE (n = 3). (**H**) Natural allelic variation at *NGR5*. (**I**) Tiller number and grain yield. Data are means ± SE (Hap.1, n = 305; Hap.2, n = 84; Hap.3, n = 62; Hap.4, n = 138; Hap.5, n = 97). (**J**) Relative mRNA abundance of *NGR5*. Abundance shown is relative to that of *OsActin1*. (**K**) Plant height. Data in (E) to (K) are means ± SE (n = 20). (**L** and **M**) Grain yield per plot. Data are means ± SE of six plots (each plot contained 220 plants) per line per nitrogen level. In (B), (E) to (G), and (I) to (M), different letters denote significant differences (P < 0.05, Duncan multiple range test).

9311-*sd1*. Thus, the enhanced DELLA function of *sd1* increases tiller number in response to nitrogen supply by increasing the stability of NGR5 (Fig. 3, C and E), which in turn inhibits the expression of shoot branching inhibitor genes, thereby promoting tiller number.

NGR5 improves yield and nitrogen use efficiency

We next determined whether an increase in NGR5 abundance beyond that seen in elite rice varieties (e.g., sd1-containing 9311) could further increase tiller number and yield in reduced nitrogen fertilizer inputs. First, we surveyed publicly available rice varietal genome sequence data for natural genetic variation at NGR5 (37), distinguished five distinct haplotypes (Hap.1 to Hap.5; Fig. 4H), and found that Hap.2 was associated with increased NGR5 mRNA abundances in both low and high nitrogen conditions (fig. S15), together with increases in tiller number and field-grown grain vield of 686 diverse Asian cultivated rice accessions (Fig. 4I). Further analysis showed that Hap.2-containing Guichao2 [Guichao2(Hap.2), one of the highest-yielding of indica varieties cultivated in China since the 1980s] displayed a greater NGR5 mRNA abundance than did Guichao2(Hap.1) (a NIL carrying Hap.1 in the Guichao2 genetic background) and other lines (including Hap.5-containing 9311), even at low and moderate nitrogen supply (Fig. 4J). In addition, we showed that a transgenic mimic of Hap.2 (expression from a p35S::NGR5 transgene) enhanced 9311 grain yield in a range of nitrogen supply conditions, without affecting the characteristic and beneficial semi-dwarfism of 9311 (Fig. 4, K and L); this suggests that breeding with Hap.2 is a feasible future strategy toward improving nitrogen use efficiency of the elite rice varieties. Finally, having recently shown that allelic variation at GRF4 (encoding the rice GROWTH-REGULATING FACTOR 4 transcription factor) enhances grain yield and nitrogen use efficiency through coordinating effects on carbon and nitrogen metabolic regulation (13), we investigated the genetic interaction between GRF4 and NGR5. We found that increased abundances of both GRF4 and NGR5 further enhanced 9311 yield and nitrogen use efficiency, particularly at relatively low levels of nitrogen supply (Fig. 4M).

We have shown that nitrogen determines genome-wide chromatin status (specific H3K27me3 histone modification) via NGR5dependent recruitment of the polycomb complex PRC2 to target genes, among which are tiller branch-repressing genes. In consequence, repression of tiller outgrowth is reduced in increasing nitrogen supply, causing increased tillering. We have also shown that NGR5 is a non-DELLA target of gibberellin-GID1-SCF^{GID2}-mediated proteasomal destruction and that competitive NGR5-DELLA-GID1 interactions cause the NGR5-dependent yieldenhancing tillering increases typical of green revolution rice varieties. Because NGR5 is already known to be involved in the cross-talk between auxin and brassinosteroid signaling (19, 20), our discoveries add to a growing understanding of how diverse modes of molecular and functional cross-talk between multiple phytohormonal signaling and fertilizer use responses function in the environmentally adaptive regulation of plant growth and development. Finally, we have shown that increasing NGR5 expression or activity provides a breeding strategy to reduce nitrogen fertilizer use while boosting grain yield above what is currently sustainably achievable.

Materials and methods Plant materials and growth conditions

A nitrogen-insensitive rice mutant, designated ngr5 (nitrogen-mediated tiller growth response 5), was isolated from the progeny of EMSmutagenized indica cultivar 9311. NILs carrying allelic combinations of the NGR5, SD1, and GID1 loci were obtained by backcrossing to recurrent parent 9311 (or NJ6) six times. Details of the germplasm used for the positional cloning and haplotype analysis are described in (13, 37-39). Paddy-grown rice plants, including 686 diverse Asian cultivated rice accessions (37), were planted in rows 20 cm apart and raised in standard paddy conditions at two experimental stations, one in Lingshui (Hainan Province), the other in Hefei (Anhui Province).

Hydroponic culture conditions

Hydroponic culture conditions were as described (13). Rice seeds were surface-sterilized with 20% sodium hypochlorite solution for 30 min, then rinsed and soaked in water for 3 days to allow the seeds to germinate. Surfacesterilized seeds were then germinated in moist Perlite. Seven-day-old seedlings were transplanted to PVC pots containing 40 liters of nutrient solution (1.25 mM NH₄NO₃, 0.5 mM NaH₂PO₄·2H₂O, 0.75 mM K₂SO₄, 1 mM CaCl₂, 1.667 mM MgSO4·7H2O, 40 µM Fe-EDTA (Na), 19 µM H₃BO₃, 9.1 µM MnSO₄·H₂O, 0.15 µM ZnSO₄·7H₂O, 0.16 µM CuSO₄, and 0.52 µM (NH₄)₃Mo₇O₂₄·4H₂O, pH 5.5), and growth was continued in a greenhouse. Compositions of nutrient solutions containing different levels of supplied nitrogen were as follows: 1N, 1.25 mM NH₄NO₃; 0.6N, 0.75 mM NH₄NO₃; 0.2N, 0.25 mM NH₄NO₃; 0N, 0 mM NH₄NO₃. All nutrient solutions were changed twice per week; pH was adjusted to 5.5 every day. The temperature was maintained at 30°C day and 22°C night, and the relative humidity was 70%.

Map-based cloning of NGR5

Fine-scale mapping of ngr5 was based on 600 F₂ plants and 1256 BC₁F₂ plants derived

from a cross between the *ngr5* mutant and the *japonica* rice cultivar Lansheng (recurrent parent). Genomic DNA sequences in the candidate region were compared between 9311, Nipponbare, and Lansheng. Primer sequences used for map-based cloning and genotyping assays are given in table S7.

Transgene constructs

Wild-type NGR5 and ngr5 mRNA-encoding sequences (together with intron sequences and/or promoter regions lying 3 kbp upstream of the transcription start site) were amplified from 9311 and ngr5 mutant plants, respectively. These amplified genomic DNA fragments were inserted into the p35S::HA-nos (40) and pCAMBIA1300 (CAMBIA, www.cambia.org) vectors to respectively generate pNGR5::NGR5, pNGR5::ngr5, pNGR5::NGR5-HA, and p35S::NGR5 constructs. Full-length cDNAs of NGR5, GID1, and LC2 cDNAs were amplified from 9311 plants and inserted into p35S::HA-nos (40) or p35::GFP-nos (39-41) vector to respectively generate p35S::GID1, p35S::LC2-HA, p35S::NGR5-HA, and p35S::NGR5-GFP constructs. gRNA constructs required for CRISPR/Cas9-mediated generation of GID1, GID2, and LC2 mutant alleles were made as described (13, 41). The transgenic rice plants were generated by Agrobacterium-mediated transformation as described (42). Relevant primer sequences are given in table S8.

qRT-PCR analysis

Total RNAs were extracted from tiller buds of 3-week-old rice plants using the TRIzol reagent (Invitrogen) according to the manufacturer's protocol, and treated with RNase-free DNase I (Invitrogen) to remove contaminating genomic DNAs. The full-length cDNAs were then reverse-transcribed using a cDNA synthesis kit (TRANSGEN, AE311-02). Subsequent qRT-PCR processing steps were performed according to the manufacturer's instructions (TRANSGEN, AQ101), with each qRT-PCR assay being replicated at least three times with three independent RNA preparations. Rice Actin1 gene (OsActin1, LOC Os03 g50885) transcripts were used as an internal reference. Relevant primer sequences are given in table S9.

Yeast two-hybrid assays

Yeast two-hybrid screening was performed as described (*38*). The full length *NGR5* cDNA was amplified and subcloned into *pGBKT7* (Takara Bio Inc.), then transformed into yeast strain AH109. The NGR5 protein was used as a bait to screen a cDNA library prepared from equal amounts of poly(A)-containing RNA sampled from various rice tissues/organs, including tiller buds, roots, leaves, shoot apical meristem (SAM), young panicles, etc. Experimental procedures for screening and plasmid isolation were performed according to the

manufacturer's user guide. cDNAs encoding various deleted and nondeleted versions of SLR1 were amplified and then subcloned into the *pGBKT7* vector (Takara Bio Inc.). Bait and prey vectors were co-transformed into yeast strain AH109, and experimental procedures were performed according to the manufacturer's instructions (Takara Bio Inc.). Relevant primer sequences are given in table S8.

Bimolecular fluorescence complementation (BiFC) assays

As described (13), full-length cDNAs of NGR5, LC2, SLRI, DLT, and MOC1 were amplified from 9311 and inserted into pSY-735-35S-cYFP-HA or pSY-736-35S-nYFP-EE vectors (YFP, yellow fluorescent protein) to generate fusion constructs. Cotransfection of constructs (e.g., nYFP-NGR5 and cYFP-LC2) into tobacco leaf epidermal cells by Agrobacterium-mediated infiltration enabled testing for protein-protein interaction. After 48 hours of incubation in the dark, the YFP signal was captured using a confocal microscope (Zeiss LSM710). Each BiFC assay was repeated at least three times. Relevant primer sequences are given in table S8.

Split firefly luciferase complementation (SFLC) assays

Full-length cDNAs of GID1 and GID2 and cDNAs encoding various deleted and nondeleted versions of NGR5 were amplified from 9311, and then inserted into pCAMBIA1300-35S-Cluc-RBS or pCAMBIA1300-35S-HA-Nluc-RBS vectors (39) to generate fusion constructs. Two different vectors (e.g., nLUC-GID1 and cLUC-NGR5) enabling testing of protein-protein interaction, together with the p19 silencing plasmid, were cotransfected into tobacco leaf epidermal cells by Agrobacterium-mediated infiltration. After 48 hours of incubation in the dark, the injected leaves were sprayed with 1 mM luciferin (Promega, E1605) and the LUC signal was captured using a cooled CCD imaging apparatus (Berthold, LB985). Each assay was repeated at least three times. Relevant primer sequences are given in table S8.

FRET (Förster resonance energy transfer) assays

FRET assays were performed as described (13). Cauliflower mosaic virus (CaMV) 35S promoterdriven fusion constructs with C-terminal tagging CFP (cyan fluorescent protein) or YFP were created to generate the donor vectors *p35S::GID1-CFP* (or *p35S::LC2-CFP*), and the acceptor vector *p35S::NGR5-YFP*. Donor and acceptor vectors, with or without *p35S::Rht-BIa* (*p35S::Rht-BIb*, or *p35S::SLR1*) vector, were cotransformed into tobacco leaf epidermis cells by *Agrobacterium*mediated vacuum infiltration to provide the FRET measurements. Transformation with the *p35S::GID1-CFP* vector provided only the donor channel, and with the *p35S::NGR5-YFP* vector only the acceptor channel. The FRET signal was detected and photographed using a confocal microscope (Zeiss LSM710). Relevant primer sequences are given in table S8.

Western blotting and coimmunoprecipitation (Co-IP) assays

Protein extracts were electrophoretically separated by SDS-PAGE and transferred to a nitrocellulose membrane (GE Healthcare). Proteins were detected by immunoblot using the following antibodies: anti-SLR1 (ABclonal Technology), anti-HA (MBL, M180-7), anti-Ubiquitin (Abcam, ab134953), anti-GST (Santa Cruz, sc-138), anti-GFP (Abcam, ab6673), and anti-HSP90 (BGI), respectively. For Co-IP experiments, full-length NGR5, LC2, SLR1, GID1, and GID2 cDNAs were amplified, then inserted into either the pUC-35S-HA-RBS or the pUC-35S-Flag-RBS vector as described (13, 39). Rice protoplasts were transfected with 100 µg of plasmid DNA and then incubated overnight in the dark. Total protein was extracted from harvested protoplasts with a lysis buffer [50 mM HEPES (pH 7.5), 150 mM KCl, 1 mM EDTA, 0.5% Triton X-100, 1 mM DTT, proteinase inhibitor cocktail (Roche LifeScience)]. Lysates were incubated with magnetic beads conjugated with an anti-DDDDK-tag antibody (MBL, M185-11) or anti-HA-tag antibody (MBL, M180-11) at 4°C for 4 hours. The magnetic beads were then washed 5 times with TBS-T buffer [500 mM NaCl, 20 mM Tris-HCl (pH 8.0), 0.1% Tween 20] and eluted with 3×Flag peptide (Sigma-Aldrich, F4709). Immunoprecipitates were electrophoretically separated and specific proteins detected by immunoblotting with anti-HA (MBL, M180-7) or anti-DDDDK (MBL, M185-7) antibodies. Relevant primer sequences are given in table S8.

In vivo pull-down

A full-length rice GID1 cDNA was amplified and then inserted into the pGEX-4T-1 vector (GE Healthcare). The recombinant GST-GID1 protein was expressed in Escherichia coli BL21 (DE3) (Transgen, CD701-01), and then purified and immobilized on Glutathione Sepharose 4B beads (GE Healthcare, 17-0756) following the manufacturer's instructions. The beads were divided into four equal aliquots and incubated with the same amount of NGR5-HA (or NGR5-Flag) protein lysate, together with His-SLR1 or with different concentrations of GA₃ (0, 2, 5, 10 µM) and/or MG132 (50 µM) for 2 hours at 4°C. The beads were subsequently washed five times with TBS-T buffer, followed by elution with 50 ul of elution buffer (50 mM Tris-HCl, 10 mM reduced glutathione, pH 8.0). Supernatants were resolved by 12% SDS-PAGE and subjected to immunoblotting using anti-GST (Santa Cruz, sc-138) and anti-HA (MBL, M180-7) antibodies. Relevant primer sequences are given in table S8.

EMSA assays

EMSA assays were performed as described (39). A full-length NGR5 cDNA was amplified and inserted into the *pGEX-4T-1* vector (GE Healthcare). Recombinant GST-NGR5 protein was expressed in *E. coli* BL21 (DE3) strain and purified using Glutathione Sepharose 4B (GE Healthcare, 17-0756) following the manufacturer's instructions. DNA probes (D5 fragment for the *D14* gene, S3 fragment for the *OsSPL14* gene) were amplified and labeled using a biotin label kit (Biosune). DNA gel shift assays were performed using the LightShift Chemiluminescent EMSA kit (Thermo Fisher Scientific, 20148). Relevant primer sequences are given in table S10.

Cell-free protein degradation assays

Three-week-old NJ6-gid1-c1 and NJ6-gid2-c1 seedlings (together with NJ6) were harvested and ground into a fine powder in liquid nitrogen. Lysates were subsequently extracted with lysis buffer [25 mM Tris-HCl (pH 7.5), 10 mM NaCl, 10 mM MgCl₂, 4 mM PMSF, 5 mM DTT, and 10 mM ATP] as described (43); total protein extracts were adjusted to be at equal concentration in the lysis buffer for each assay. Rice cell lysates (200 µl) were incubated with 100 ng of purified GST-NGR5 (or His-NGR5) fusion protein in the presence or absence of the recombinant Rht-B1b-Flag, Rht-Bla-Flag, SLR1-Flag, or His-SLR1 proteins. Proteins were extracted from lysates that had or had not been exposed to treatments with 100 µM GA₃ and/or 100 µM MG132 for a series of incubation times and then subjected to SDS-PAGE and Western blotting using an anti-GST antibody (Santa Cruz, sc-138). HSP90 was used as a loading control.

In vitro protein ubiquitination assays

Full-length GID2 cDNA was amplified and inserted into PUC-35S-flag-RBS vector (39). Rice protoplasts were transfected with 100 µg of plasmid and incubated for 24 hours. Total protein was extracted from harvested protoplasts in the lysis buffer [50 mM HEPES (pH 7.5), 150 mM KCl, 1 mM EDTA, 0.5% Triton X-100, 1 mM DTT, and proteinase inhibitor cocktail (Roche LifeScience)]. Lysates were incubated with agarose-conjugated anti-Flag antibodies (Sigma-Aldrich, A2220) at 4°C for 2 hours, rinsed 6 times in the PBS-T buffer, and then eluted with 3×Flag peptide (Sigma-Aldrich). Purified GID2-Flag proteins were used for in vitro ubiquitination assays as described (44). Crude extracts containing recombinant GST-NGR5, purified E3 (GID2-Flag), 6×His-tagged Ubiquitin (Ub), E1 and E2 (Ubiquitinylation kit; Enzo Life Science, BML-UW9920-0001) were used. Buffer at a final concentration of 50 mM Tris-HCl (pH 7.4), 5 mM MgCl₂, and 2 mM ATP was also added to the system. The reactions were incubated at 30°C for 2 hours, and terminated by adding SDS sample buffer with β -mercaptoethanol. Reaction products were separated with 12% SDS-PAGE and subjected to immunoblotting using anti-ubiquitin antibody (Abcam, ab134953) and anti-GST antibody (Santa Cruz, sc-138). Relevant primer sequences are given in table S8.

ChIP-PCR assay

ChIP assays were performed as described (13). A 2-g sample of 2-week-old rice plants was collected and immediately fixed with 1% (v/v) formaldehyde under vacuum for 15 min at 25°C, and then homogenized in liquid nitrogen. After the nuclei were isolated and lysed, the chromatin was ultrasonically fragmented on ice to an average size of 500 bp. Immunoprecipitations were performed with an anti-HA antibody (Santa Cruz, sc-7932x) and an anti-H3K27me3 antibody (Millipore, 07-449) overnight at 4°C. At the same time, an equal volume of the supernatant was prepared without any antibody as a mock sample. The bound DNA fragments were then reversely released and amplified by real-time quantitative PCR. Relevant primer sequences are listed in table S11.

RNA-seq

Total RNAs were extracted from tiller buds of 3-week-old 9311 plants treated with and without gibberellin and ngr5 mutants grown in high N (1.25 mM NH₄NO₃) supply conditions using the TRIzol reagent (Invitrogen) according to the manufacturer's instructions. Libraries were constructed and sequenced using the BGISEQ-500 sequencer. Raw sequencing reads were cleaned by removing adaptor sequences, reads containing poly-N sequences, and low-quality reads, and clean reads were then mapped to the Nipponbare reference genome as described (13).

ChIP-seq

ChIP-Seq analysis was performed as described (13). Approximately 2 g of 2-week-old transgenic plants carrying the p35S::LC2-HA and pNGR5::NGR5-HA constructs grown in high N (1N, 1.25 mM NH₄NO₃) supply conditions, and of ngr5 and wild-type plants grown in low (0.2N, 0.25 mM NH₄NO₃) or high nitrogen (1N, 1.25 mM NH₄NO₃) supply conditions with or without 100 µM GA₃ (Sigma-Aldrich, G1025) and 10 µM PAC (Sigma-Aldrich, 19847) treatments, were fixed with 1% (v/v) formaldehyde under vacuum for 15 min at 25°C, and then homogenized in liquid nitrogen. After cell lysis and nucleic acid isolation, cross-linked chromatin fibers were ultrasonically fragmented into fragments of an average size of 500 bp. Immunoprecipitations were performed with anti-HA antibodies (Santa Cruz, sc-7932) and anti-H3K27me3 antibodies (Millipore, 07-449) overnight at 4°C. The precipitated DNA was recovered by centrifugation (13,000g, 5 min, 25°C) and dissolved in sterile distilled water. Illumina sequencing libraries were constructed according to the manufacturer's instructions, and then sequenced on the BGISEQ-500 platform. Sequencing reads were mapped to the reference genome as described (*13*). The precipitated DNA samples also served as template for quantitative real-time PCR. Relevant primer sequences are given in table S11.

Processing of ChIP- and RNA-sequencing data

Sequencing reads were cleaned with Trimmomatic (version 0.36) (45) and Sickle, including elimination of bases with low-quality scores (< 25) and irregular GC contents, as well as removal of sequencing adapters and short reads. The remaining clean reads were mapped to the genome of japonica rice (MSU7.0 release) with the Burrows-Wheeler Alignerbacktrack (version 0.7.16a-r1181) (46) for ChIPsequencing data and HISAT2 2.1.0 (47) for RNA-seq data. MACS (version 1.3.7) (48) was used to identify read-enriched regions (peaks) of ChIP-sequencing databased on the following combined criteria: P value < 0.00001 and fold-change > 32. Target genes were defined as genes with a peak within or near the gene body $(\pm 2 \text{ kb})$. DESeq (49) was applied to determine the significance of the differential expression between samples with the combined criteria: fold change > 2 and adj. P < 0.05.

Gene set enrichment analysis

To determine the enrichment of H3K27me3 targets in *NGR5*-regulated genes (i.e., differentially expressed genes in ngr5), we performed gene set enrichment analysis (GSEA), which is a robust computational method that determines whether an a priori gene set shows statistically significant and concordant differences between two samples (*50*). Briefly, *NGR5* regulated genes were ranked by the quantitative expression change in *ngr5*, followed by calculation of the fraction of regulated genes that are targeted by H3K27me3. The enrichment score is normalized by the size of the gene set (NES). *P* value is estimated by permutating genes.

Statistical analysis

Data were statistically analyzed and multiple comparisons were made using Duncan's multiple range test as described (13). P values of less than 0.05 were considered to indicate statistical significance. Statistical calculations were performed using Microsoft Excel 2010.

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SUPPLEMENTARY MATERIALS

science.sciencemag.org/content/367/6478/eaaz2046/suppl/DC1 Materials and Methods Figs. S1 to S15 Tables S1 to S11 20 August 2019; accepted 18 December 2019 10.1126/science.aaz2046



Enhanced sustainable green revolution yield via nitrogen-responsive chromatin modulation in rice

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Decoupling tillering and fertilization

For rice as an agricultural crop, more tillers, or branches that carry grains, are desired, as is less demand for nitrogen fertilization. Unfortunately, for many rice varieties, the number of tillers depends on the amount of nitrogen fertilization. Wu *et al.* now show that nitrogen status affects chromatin function through modification of histones, a process in which the transcription factor NGR5 recruits polycomb repressive complex 2 to target genes. Some of these genes regulate tillering, such that with more nitrogen, the plants develop more tillers. NGR5 is regulated by proteasomal destruction and mediates hormone signaling. An increase in NGR5 levels can drive increases in rice tillering and yield without requiring increases in nitrogen-rich fertilizer.

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