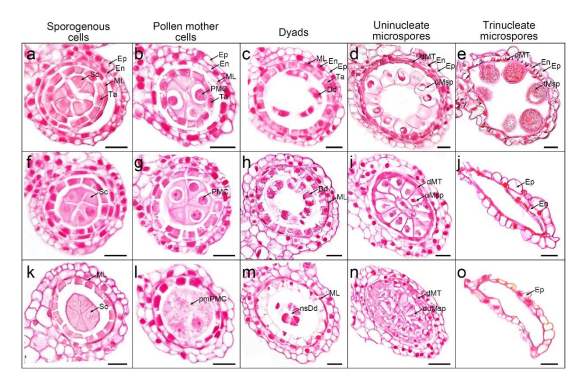
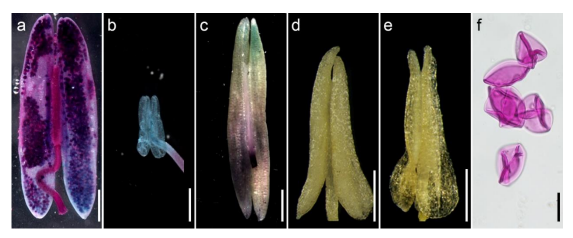
Supplementary Figures

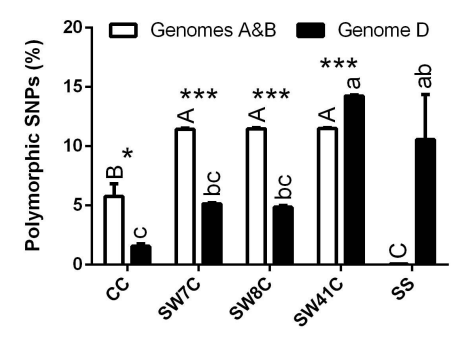


Supplementary Figure 1. Disruption of microsporogenesis in Taigu lines.

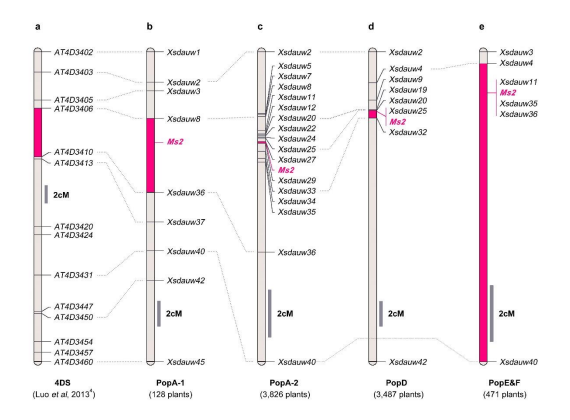
Transverse sections of anthers were prepared from wild-type male-fertile (MF) wheat (**a-e**) and male-sterile (MS) Taigu lines (**f-o**) at the five stages (marked on top of the columns). The plants are MF lines of XY6 (a,d,e) and CS (b,c), and MS lines of CS_{RMs2} (f-h,l,m,o) and $XY6_{Ms2}$ (i-k,n). The Taigu lines have various cellular disorders: 1) premeiotic degeneration of the middle layer in k, 2) plasmodial mass of pollen mother cells in l, 3) non-separating dyads in m, 4) a common degeneration of uninucleate microspores in n, and 5) a 'pollen-less type' sac¹ in j and o. Abbreviations: Ep, epidermis; En, endothecium; ML, middle layer; Ta, tapetum; PMC, pollen mother cell; Sc, sporogenous cell; Dd, dyads; nsDd, non-separating dyads; uMsp, uninucleate microspore, tMsp, trinucleate microspore; dMT, degenerated middle layer and tapetum; duMsp, degenerated uninucleate microspore; pmPMC, plasmodial mass of pollen mother cells. Scale bars = $50\mu m$.



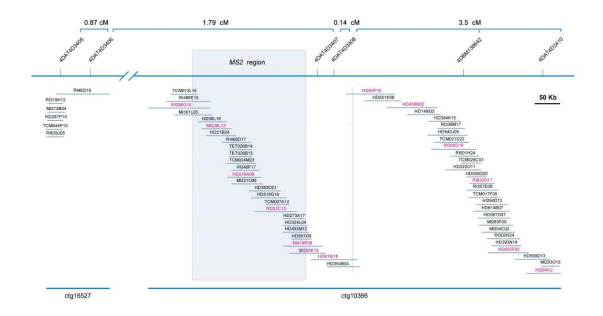
Supplementary Figure 2. Spectrum of morphology of Taigu anthers. (a) An anther from a wild-type male-fertile (MF) wheat plant stained by Alexander's method. (b, c) Anthers from male-sterile (MS) plants from popA-2 (XY6_{Ms2}/2*SW41) stained by Alexander's method with a diminutive anther (b) and a more normally-sized anther but without viable pollen (c). (d, e) Late MS florets from a MS plant in the *RMs2*-based recurrent selection population had more normally-sized anthers but the pollen (f), stained by Alexander's method, was non-viable. The anthers of LM15_{RMs2} and other Taigu lines did not typically bear any pollen. Late flowers in Taigu lines occasionally bore more normally-sized anthers²; however, the Taigu anthers were typically thin and rigid; any pollen that was produced was non-viable. Scale bars = 500 μ m (a-e) and 25 μ m (f).



Supplementary Figure 3. DNA polymorphisms amongst parental lines that were used to map Ms2. DNA polymorphisms were detected using the wheat 90K iSelect SNP assay³. Comparisons were made within common wheat (CC), and the synthetic hexaploid wheat (SS), and between the synthetic hexaploid wheat and common wheat (e.g. SW7C). Taigu lines are in the common wheat lineage. For each X-axis group, a mean polymorphism rate was obtained by averaging the within group comparisons (n=3). The error bar is one standard error of the mean (s.e.m.). Significant differences between genotypes in the percentage of SNPs were determined by a Tukey-Kramer HSD (α =0.05, capital letters for the AB genomes and small letters for the D genomes). Analysis of significant differences in the percentage of SNPs for each genotype in the AB versus D genomes was done with paired t-tests (asterisks: * P<0.05, *** P<0.001).

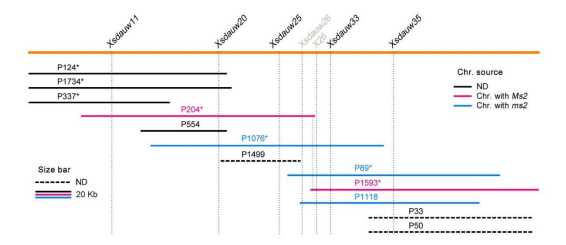


Supplementary Figure 4. Mapping of *Ms2* **in Taigu lines.** (a) The 4DS linkage map of *Ae. tauschii*⁴. (b) A preliminary map from popA-1 (XY_{Ms2}/2*SW41) using 43 plants for the linkage map and 85 plants for validating the *Xsdauw8-Xsdauw36* interval. (c) Fine map of the *Ms2* region based on popA-2 (XY_{Ms2}/2*SW41, 3,826 plants). (d) Fine map of the *Ms2* region based on popD (XY_{Ms2}/2*SW7, 3,487 plants). (e) An integrated map from popE (LM_{RMs2}/2*SW41, 235 plants) and popF (CS_{RMs2}/2*SW41, 236 plants). *Xsdauw32* and *Xsdauw33* are derived from the same region, where the *Xsdauw33* range is nested within *Xsdauw32*. The pink region represents the *Ms2* interval. The *Xsdauw4/Xsdauw5-Xsdauw40* intervals are genetically comparable, about 10.5 to 10.8 cM, in both *Ms2*- and *RMs2*-based populations (popA-2, popE, and popF). The *Xsdauw11-Xsdauw36* interval is about 5.1 cM and spans *Ms2* in popA-2, but both markers were completely linked to *Ms2* in the *RMs2*-based populations popE and popF.

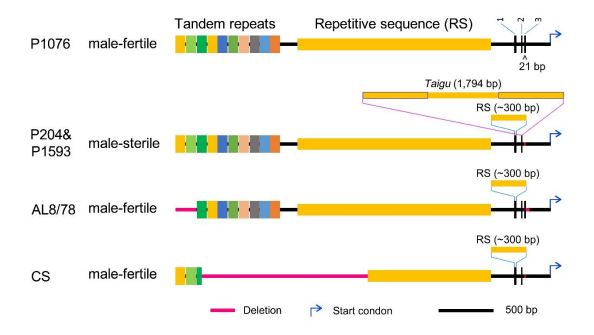


Supplementary Figure 5. Physical contigs of the ms2 region in Ae. tauschii.

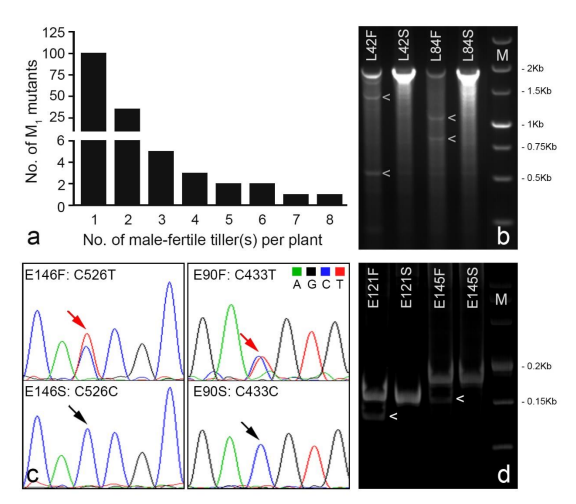
Genetic distances and physical contigs are from a previous report⁴. Overlapping BACs in pink or purple were sequenced to span the *Ms2* interval. BACs in black were not sequenced. The *MS2* region is highlighted by gray shading. A previous study revealed a complete linkage between 4DBM138642 and 4DAT4D3410⁴.



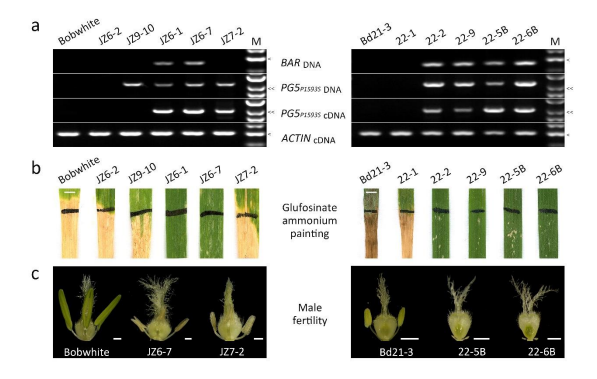
Supplementary Figure 6. Physical map of the *Ms2* **region in LM15**_{RMs2}. Five markers in black were used to screen the LM15_{RMs2} BAC library; two markers in gray (*Xsdauw26* and *X26*) assisted in differentiating between the *ms2* and *Ms2* associated chromosomes. P204 and P1593 shared 1,212 bases and formed a complete physical map of the *Xsdauw11-Xsdauw35* interval in the *Ms2*-associated chromosome (BACs in pink). P89 and P1076 formed a complete physical map of the *Xsdauw20-Xsdauw35* interval in the *ms2*-associated chromosome (BACs in blue). The chromosome source was not determined (ND) for seven clones (BACs in black).



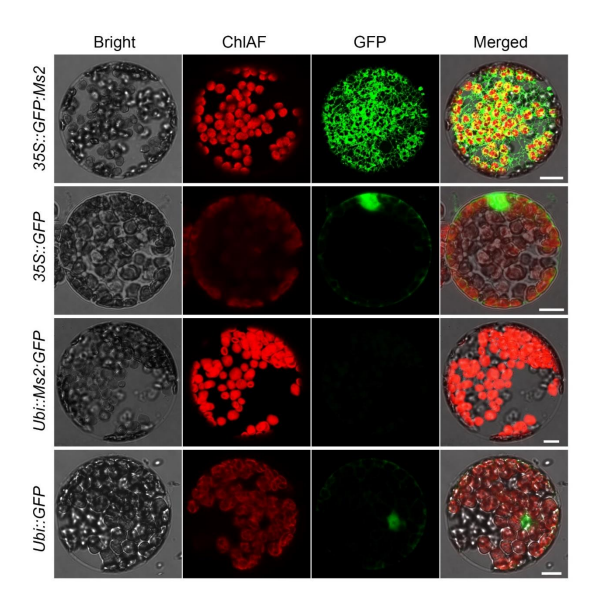
Supplementary Figure 7. Major structural variations in the PG5 promoter region. A 3,502-bp region was analyzed. The first 2,894-bp contains three to ten tandem repeats and then a large repetitive sequence (RS). The remaining 608bp is not repetitive overall, and has three structural variations: 1) a RS insertion in all entries except for MF P1076, 2) a Taigu retroelement insertion in MS P204 and P1593, and 3) a small fragment deletion in all entries except for P1076. Here, P204 and P1593 have the only male-sterility allele. All other lines (AL8/78, CS, and LM15) are male-fertile. The base numbers are based on the $PG5_{P1076F}$ (= LM15).



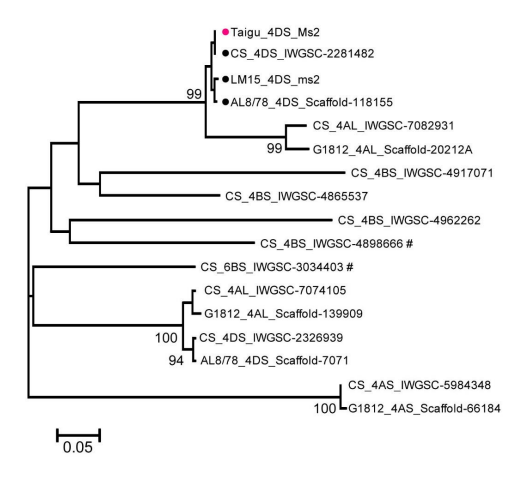
Supplementary Figure 8. EMS-induced mutagenesis of the $PG5_{P1593S}$ gene. (a) Distribution of male-fertile (MF) tillers in the CS_{RMs2} subset. (b) TILLING screening (P133/P134, 1,937 bp) of the segregating tillers of the same M_1 plants (S: sterile tiller; F: fertile tiller). (c) Sequence chromatograms of selected mutations. (d) Marker-assisted validation of the identified mutations (T1 and T2 markers for the C158T and G437A mutations in E121 and E145, respectively; Supplementary Data 4 and 5). Arrows and arrow heads highlight the *Cel*I-digested band (b), the base change (c), and the Hpy188III-digested band (d). M, the DS5000 DNA Ladder (Dongsheng, Guangzhou, China; in b) or the DM500 DNA Ladder (CWBIO, Beijing, China; in d).



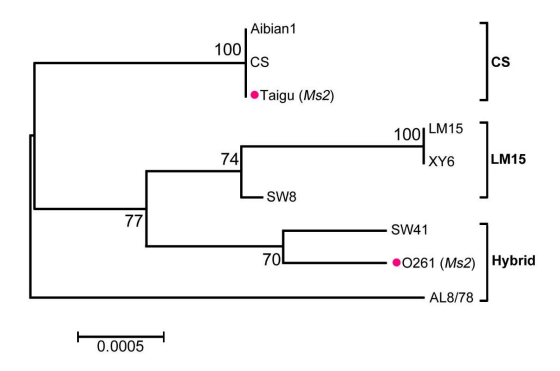
Supplementary Figure 9. Transformation of wheat (left panel) and *Brachypodium* (right pane) with the $PG5_{P1593S}$ gene. (a, b) Co-occurrence of $PG5_{P1593S}$ DNA, mRNA and herbicide tolerance in transgenic male-sterile (MS) lines and negative phenotypes in two male-fertile (MF) non-transformed controls: 'Bobwhite' wheat and 'Bd21-3' Brachypodium. (a) PCR confirmation of transgene integration (BAR and $PG5_{P1593S}$ DNA) and transgene transcription ($PG5_{P1593S}$ and ACTIN cDNA) in T₀ plants. A 36-cycle PCR was performed to detect the intergration of BAR (P121/P122, 412 bp) and $PG5_{P1593S}$ (P133/P141, 1,086 bp, or the HT5 marker), and to detect transcripts of $PG5_{P1593S}$ (P129/P130, 997 bp) and ACTIN (P142/P143, 503 bp). (b) BAR-based bioassay for herbicide tolerance; the herbicide glufosinate ammonium was applied below the black line. (c) Male fertility in selected transgenic T₀ plants. Other MS transgenics are shown in Fig. 3. PCR primers are described in Methods and Supplementary Data 2. M, the 250bp DNA Ladder ('<' = 0.5kb, '<<' = 1kb). Scale bars = 2.5mm (b) and 1mm (c).



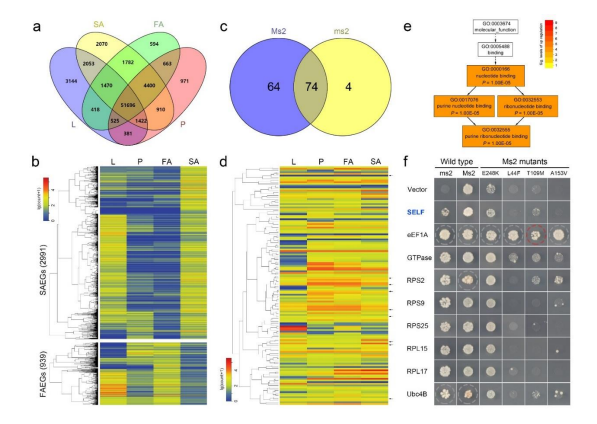
Supplementary Figure 10. Subcellular location of the GFP:Ms2 protein in mesophyll protoplasts of tobacco. The GFP signal is shown in green and the chlorophyll autofluorescence (ChlAF) is shown in red. pMDC43 (35S::GFP) serves as a control for the 35S::GFP:Ms2 construct (PC983). PC134 (Ubi::GFP) serves as a control for the Ubi::Ms2:GFP construct (PC972). In the upper row, the localization of GFP:Ms2 (from a gene with a 35S promoter) appears to be consistent with the rough endoplasmic reticulum. Scale bars = $10\mu m$.



Supplementary Figure 11. Phylogenetic analysis of the MS2-like fragments in **Triticeae species.** We used sequence data of accessions of common wheat 'Chinese Spring' (CS) and LM15, and wheat progenitor species: Aegilops tauschii accession 'AL8/78' and Triticum urartu accession 'G1812'. Phylogenetic analysis was performed on fragments that align with the first five exons of Ms2. The program MEGA 6.06⁵ was used to align, remove indels and construct a neighbor-joining tree with 1,000 bootstrap iterations. The distance scale bar indicates nucleotide differences per unit length. Bootstrap numbers (>80) are shown at the respective nodes. On the figure, accessions are named as one of the four types (either CS, LM15, Al8/78, or G1812)_the chromosome arm_ and the gene name or accession number from the wheat genome sequence (IWGSC1.0+popseq.28, http://plants.ensembl.org). The dominant Ms2 (pink circle) and recessive ms2 (black circle) alleles are shown. Predicted exons, reported in wheat expressed sequence tags, are marked by a pound sign (#). In IWGSC-3034403, the fourth and fifth pseudo-exons are switched; they were switched back for better alignment. In wheat homoeologous group 4, the chromosome arm 4AL is homologous to 4BS and 4DS⁶.



Supplementary Figure 12. Phylogenetic analysis of the MS2 full-length genes in wheat 4DS chromosomes. We selected members of the common wheat 'Chinese Spring' (CS) and LM15 haplogroups, and wheat progenitor species Aegilops tauschii accession 'AL8/78'. Phylogenetic analysis was performed using the full-length genomic DNAs (aligned with -805..+507 from the CS genomic sequence, Supplementary Data 3). The program MEGA 6.06^5 was used to align and construct a neighbor-joining tree with 1,000 bootstrap iterations. The distance scale bar indicates nucleotide differences per unit length. Bootstrap numbers (\geq 70) are shown in the respective nodes. In this figure, Taigu and O261 are the only male-sterile accessions, each with the dominant Ms2 (pink circle); the rest of the lines are male-fertile. The MS2 genes within the CS group are nearly identical; the only difference is in the promoter region of MS2 between the sterile and fertile lines (Supplementary Fig. 7). The MS2 gene haplogroups (CS, LM15 and Hybrid on right) are defined in Table 1 and Supplementary Data 8.

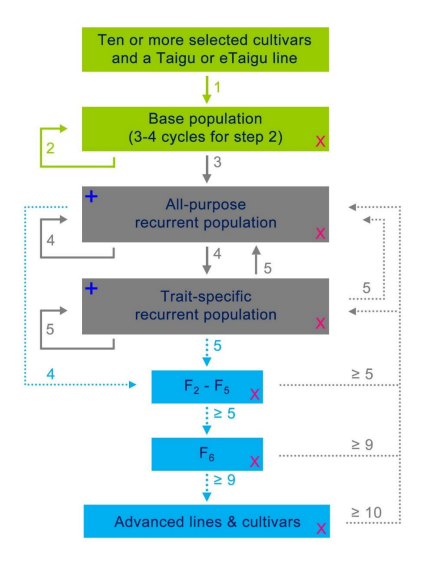


Supplementary Figure 13. *MS2* **networks.** (a) Venn diagram of 72,499 expressed genes in leaf (L), pistil (P), sterile anther (SA), and fertile anther (FA). (b) Heat maps of anther-enriched genes. (c) Venn diagram of 142 MS2-interacting proteins. (d) Heat map of the 142 MS2-interacting proteins. (e) GO analysis of the 142 MS2-interacting proteins. The color of a node matches the significance (sig.) bar and a *P*-value was included for the significant node. (f) Protein interactions between MS2 (bait, on top) and representative full-length proteins (prey, on left). Yeast colonies (50-fold dilution) were grown on the triple dropout medium. Dashed circles indicate yeast growth on the quadruple dropout medium, and a dashed red circle indicates weak growth. All tested Ms2 variants confers male-fertility. Vector, the prey vector; SELF, the same protein used as the bait; RPSs and RPLs, ribosomal proteins; Ubc4B, ubiquitin-conjugating enzyme.

		DDO(LT)		T	TDO(LTH)		QDO(LTHA)		A)	
PREY (AD)	BAIT (BD)	×	50X	500X	×	50X	500X	×	50X	500X
Ms2	Vector	•	•	•	•			0		
Ms2	Ms2	•	•	•	•	•		•		
Ms2	ms2	•	•,	•	•	0		•	of .	
Vector	Ms2	•	•	•	•	•		•		
ms2	Ms2	•	•	•	•	•	•	0	•	4
ms2	Vector	•	•	65	•			0		
ms2	ms2	•	•	•	•	•	0	•		
Vector	ms2	•	•	•	•			•		
Positive	control	•	•	*	•	•	*	•	•	4
Negative	e control	•	•	8	•			•	198	

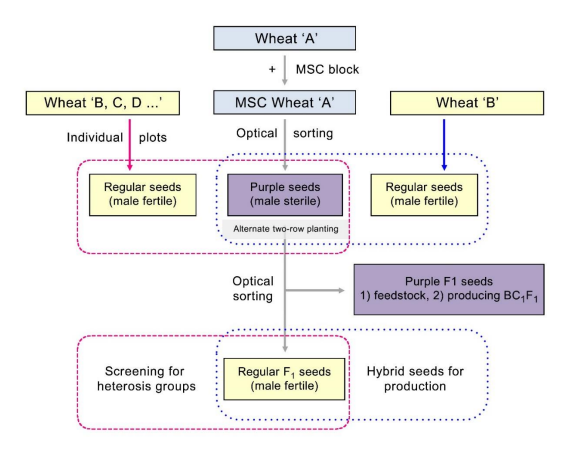
Supplementary Figure 14. Yeast two-hybrid analysis of the MS2 to ms2 proteins.

Ms2 is from PG5_{P1593S}, and ms2 is from PG5_{P1076F}. Yeast two-hybrid analysis was performed on a Gal4 system including the AD vector (prey) and BD vector (bait). Successful co-transformation of AD and BD plasmids was verified by yeast growth in the DDO medium, whereas autoactivation or interaction was tested in TDO and QDO media. The yeast colonies were progressively diluted from the original suspension (1x), 1:50-fold dilution (50x), and 1:500-fold dilution (500x). The Y2H assay included the positive control (BD-wXb12 and AD-wXb12IP2) and negative control (BD-wXb12 and AD-wRAR1)⁷. Ms2 and ms2 had no autoactivity on the TDO and QDO media except for Ms2 as a bait on TDO. Ms2 (bait) also interacted with ms2 (prey) on QDO; ms2 interacted with itself on TDO. Potentially, MS2 proteins may form a dimer or multimer.

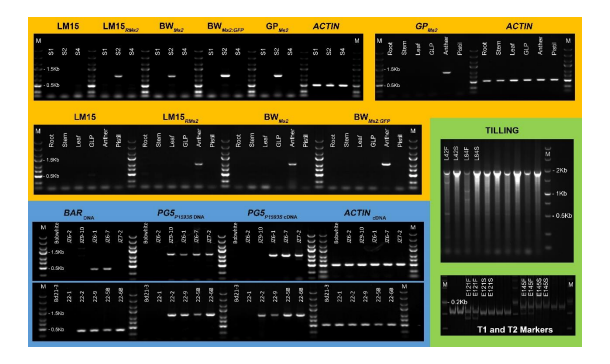


Supplementary Figure 15. Ms2-based recurrent selection in wheat. This

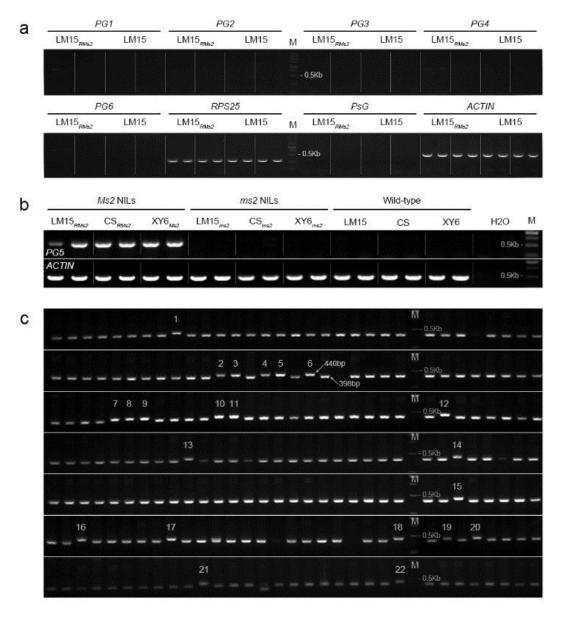
procedure, adapted from previous studies⁸⁻¹¹, includes the use of either wild-type Taigu or 'engineered Taigu' (eTaigu) lines that have a *Ms2* gene. The procedure includes the development of an initial "base" population (green), the development and maintenance of recurrent populations (gray), and the release of advanced lines and cultivars (light blue). The number of steps in the simplest path is indicated in the center. The number of steps in more complicated paths is marked by a 'greater than or equal to' sign (≥). A solid step line indicates bulk seeds from selected male sterile (MS) plants and a dashed step line indicates seeds from selected male fertile (MF) plants. Seeds of MF plants can be used to maintain the recurrent populations (dashed lines in gray) and to develop advanced lines and cultivars (dashed lines in light blue). To enrich the genetic variability in recurrent populations, new genotypes/lines can be added into the recurrent populations, as marked by a plus sign (+). During the course of recurrent selection and progeny tests, undesirable plants/lines will be removed, as labeled by an x-shaped cross.



Supplementary Figure 16. A schematic representation of a system for producing hybrid wheat seed using Ms2. The system is based on the use of Ms2 color wheat (MSC). The MSC cassette is composed of two linked genes: Ms2 for male-sterility and an aleurone-specific gene for pigmented grain. Wheat 'A' is first transformed to MSC wheat, amplified, and then planted in alternate rows. Within individual plots, pollen donors of a specific cultivar are planted in each row adjacent to the MSC wheat. All seed in the MSC rows will be a F₁ hybrid, and will be planted in the field and examined for desirable traits. F₁s of each specific cross between one cultivar and an MSC line will be evaluated in replicated plots to identify specific heterotic groups of interest (dashed pink lines). Specific MSC wheat and its heterotic partner will be used to generate a large amount of F₁ seeds (dashed blue lines), and their hybrid vigor will be determined on the population level. The most promising heterotic groups from specific cultivars will be scaled up for hybrid seed production. The system also generates MSC-type F₁ hybrids, which can be used directly as feedstocks or used to prepare BC₁F₁ populations. During the process, high throughput optical sorting is used to separate pigmented seeds from wild type seeds.



Supplementary Figure 17. Uncropped images with molecular markers. Gels with an orange background have treatments in Fig. 4a,b, those with a green background have treatments in Supplementary Fig. 8b,d, and those with a blue background have treatments shown in Supplementary Fig. 9a. M, the 250bp DNA Ladder (those with an orange or blue background), the DS5000 DNA Ladder (TILLING with a green background), and the DM500 DNA Ladder (T1 and T2 Markers with a green background).



Supplementary Figure 18. Uncropped images with molecular markers. (a) The expression of seven potential genes in the *Xsdauw24-Xsdauw32* interval. (b) The expression of *PG5* in the *Ms2* near-isogenic lines (NIL) and the wild-type lines. The top panel is for *PG5* (P115/P136, 504 bp), and the lower panel is for *ACTIN* control (P142/P143, 503bp). (c) A CAPS marker was used to differentiate the *PG5* cDNA clones (P104/P100, 440 bp). M, the DS5000 DNA Ladder (Dongsheng). H2O, negative control for RT-PCR.

ID/Accession No.	Abbreviation	Species (pedigree)	Genome
AL8/78	-	Aegilops tauschii Coss.	DD
CIae 9	-	Aegilops tauschii Coss.	DD
PI 560536	-	Aegilops tauschii Coss.	DD
Langdon / CItr 13165	LDN	Triticum turgidum L.	AABB
LDN-DIC 2B(521)	$LDN_{\mathrm{DIC521-2B}}{}^{b}$	Triticum turgidum L.	AABB
SW7	SW7 ^c	Synthetic hexaploid wheat	AABBDD
SW8	SW8 ^c	Synthetic hexaploid wheat	AABBDD
SW41	SW41 ^c	Synthetic hexaploid wheat	AABBDD
Chinese Spring+RMs2	CS_{RMs2}	Triticum aestivum L.	AABBDD
Lumai 15+RMs2	$LM15_{RMs2}$	Triticum aestivum L.	AABBDD
Xiaoyan 6+Ms2	$XY6_{Ms2}$	Triticum aestivum L.	AABBDD
BC ₁ F ₁ population A	popA	$(XY6_{Ms2}/2*SW41)$	AABBDD
BC ₁ F ₁ population D	popD	$(XY6_{Ms2}/2*SW7)$	AABBDD
BC ₁ F ₁ population E	popE	$(LM15_{RMs2}/2*SW41)$	AABBDD
BC ₁ F ₁ population F	popF	$(CS_{RMs2}/2*SW41)$	AABBDD
Chinese Spring	CS	Triticum aestivum L.	AABBDD
Lumai 15	LM15	Triticum aestivum L.	AABBDD
Xiaoyan 6	XY6	Triticum aestivum L.	AABBDD
Bobwhite	BW	Triticum aestivum L.	AABBDD
Golden Promise / PI 343079	GP	Hordeum vulgare L.	НН
Bd21 / PI 254867	BD	Brachypodium distachyon L.	-
-	-	Nicotiana benthamiana Domin	_

^aWheat cultivars/lines used for haplotype analysis and genetic diversity are listed in Supplementary Data 8. ^bThe 2B chromosome in Langdon was replaced by the one from wild emmer wheat 'PI 481521' (*Triticum turgidum* ssp. *dicoccoides*). ^cPedigrees of SW7, SW8, and SW41 are Langdon/*Ae. tauschii* CIae22, Langdon/*Ae. tauschii* CIae25, and Langdon/*Ae. tauschii* RL5570, respectively. '-' not applicable or not accessible.

Supplementary Table 2. BAC library of LM15 $_{RMs2}$

Batch	Plate	Empty	Insert	Clone	Genome
Codes	Quantity	Rate (%)	Size (kb) ^a	Quantity	Coverage
A	1112	0	118	427,008	2.97
В	330	0.81	132.2	126,720	0.98
C	255	2.59	147.6	97,920	0.83
D	142	0	102.1	54,528	0.33
Total	1839	0.5	123.32	706,176	5.1

^aInsert size was averaged among non-empty clones.

Supplementary Table 3. Plasmid constructs used in this study

DC NO	E C 44- 8	T b	GenBank	Vector
PC NO.	Expression Cassette ^a	Target Gene ^b	Acc. d	Backbone
pMDC43	35S::GFP6	GFP6	-	pMDC43
PC134	Ubi::GFP	GFP	-	pPZP200
PC970	$Ubi::PG5_{P1593S}$	$PG5_{P1593S} = Ms2$	KX533929	pCAMBia1300
PC971	$Ubi::PG5_{P1076F}$	$PG5_{P1076F} = ms2$	KX533930	pCAMBia1300
PC972	$Ubi::PG5_{P1593S}:GFP$	$PG5_{PI593S} = Ms2$	KX533929	pCAMBia1300
PC973	<i>NP1::PG5</i> _{P1593S} : <i>GFP</i>	$PG5_{P1593S} = Ms2^{c}$	KX585234	pCAMBia1300
PC976	$NP1::PG5_{P1593S}$	$PG5_{P1593S} = Ms2^{c}$	KX585234	pCAMBia1300
PC983	35S::GFP:Ms2	Ms2	KX533929	pMDC43
PC322	ADH1::AD:wRAR1	wRAR1	EF202841	pGADT7
PC324	ADH1::AD:wXb12IP2	wXb12IP2	JX424313	pGADT7
PC332	ADH1::BD:wXb12	wXb12	JX424306	pGBKT7
PC986	ADH1::AD:Ms2	Ms2	KX533929	pGADT7
PC987	ADH1::BD:Ms2	Ms2	KX533929	pGBKT7
PC988	ADH1::AD:ms2	ms2	KX533930	pGADT7
PC989	ADH1::BD:ms2	ms2	KX533930	pGBKT7
PC1034	$ADH1::AD:Ms2^{E248K}$	$Ms2^{E248K}$	-	pGADT7
PC1035	$ADH1::BD:Ms2^{E248K}$	$Ms2^{E248K}$	-	pGBKT7
PC1036	$ADH1::AD:Ms2^{L44F}$	$Ms2^{L44F}$	-	pGADT7
PC1037	$ADH1::BD:Ms2^{L44F}$	$Ms2^{L44F}$	-	pGBKT7
PC1040	$ADH1::AD:Ms2^{T109M}$	$Ms2^{T109M}$	-	pGADT7
PC1041	$ADH1::BD:Ms2^{T109M}$	$Ms2^{T109M}$	-	pGBKT7
PC1044	$ADH1::AD:Ms2^{A153V}$	$Ms2^{A153V}$	-	pGADT7
PC1045	$ADH1::BD:Ms2^{A153V}$	$Ms2^{A153V}$	-	pGBKT7
PC991	 ADH1::AD:eEF1A	 eEF1A	KX533924	pGADT7
PC995	ADH1::AD:GTPase	GTPase	KX533926	pGADT7
PC1016	ADH1::AD:RPS2	RPS2	KX533927	pGADT7
PC1018	ADH1::AD:RPS9	RPS9	HX055302	pGADT7
PC1020	ADH1::AD:RPS25	RPS25	HX019260	pGADT7
PC1022	ADH1::AD:RPL15	RPL15	HX250311	pGADT7
PC1024	ADH1::AD:RPL17	RPL17	HX188548	pGADT7
PC1032	ADH1::AD:Ubc4B	Ubc4B	HX066634	pGADT7

^a*Ubi*, the maize ubiquitin promoter¹²; *NP1*, a 5,578-bp native promoter upstream of the start codon of the $PG5_{PI593S}$ gene; 35S, 2×35 S promoter¹³; *ADH1*, a truncated promoter of the *alcohol dehydrogenase 1* gene from *Saccharomyces cerevisiae*¹⁴.

^bMost are cDNA copies. $PG5_{PI593S} = Ms2$, $PG5_{PI076F} = ms2$, and both from LM15_{RMs2}; $Ms2^{L44F}$, $Ms2^{TI09M}$, $Ms2^{A153V}$, and $Ms2^{E248K}$, single-base mutations of the wild-type Ms2 gene; GFP, green fluorescent protein gene¹⁵; GFP6, a modified GFP gene¹⁶;

eEF1A, eukaryotic elongation factor 1A; *GTPase*, GTP-binding protein; *RPSs* and *RPLs*, ribosomal proteins; *Ubc4B*, ubiquitin-conjugating enzyme; *wRAR1*, *wXb12*, and *wXb12IP2*, control genes used for yeast-two analysis⁷. ^cgenomic DNA. ^dGenBank accessions of the target genes. '-' not applicable or not accessible.

Supplementary Table 4. Homologous DNA of the MS2 gene in Triticeae

Species/Cu ltivars	Contig or Scaffold ^a	Chr.	Homologous Exons b	Chromosome Location ^c
Ta/CS	7082931	4AL	Ex1-8	4A:171806160:171813880 ^j
Tu/G1812	20212A	4AL	Ex1-8	4AL_scaff_20212:158766:161615 ^k
Ta/CS	7074105	4AL	Ex1-8; Ex7 6 ^f	4A:170957681:170967985
<i>Tu</i> /G1812	139909	4AL	Ex1-8; Ex7 6 ^f	4AL_scaff_139909:8409:18024
Ta/CS	4898666 ^d	4BS	Ex1-6; Ex7+8 ^g	4B:1148925:1155030
Ta/CS	4962262	4BS	Ex1-5; Ex7+8 ^g	4B:1473786:1479383
Ta/CS	2281482	4DS	Ex1-8 (ms2)	4DS_scaff_2281482:24011:31879
Ae/AL8/78	118155	4DS	Ex1-8 (ms2)	4DS_scaff_118155:33768:42646
Ta/CS	2326939	4DS	Ex1-8; Ex7 6 ^f	4DS_scaff_2326939:2426:9990
<i>Ae</i> /AL8/78	7071	4DS	Ex1-8; Ex7 6 ^f	4DS_scaff_7071:68422:76053
Ta/CS	3034403 ^e	6BS	Ex1-5; Ex5 4 ^f	6B:98153352:98157312
Ta/CS	5984348	4AS	Ex1-5, Ex4 ^h ; Ex7 ⁱ	4AS_scaff_5984348:5509:12263
Tu/G1812	66184	4AS	Ex1-5, Ex4 $^{\rm h}$; Ex7 $^{\rm i}$	4AS_scaff_66184:12694:21363
Ta/CS	7148227	4AL	Ex1; Ex2 i	4A:172121151:172121574
<i>Tu</i> /G1812	20212B	4AL	Ex1; Ex2 i	4AL_scaff_20212:134552:134789
Ta/CS	4865537	4BS	Ex1-3, 5-6	4BS_scaff_4865537:5609:11586
Ta/CS	4917071	4BS	Ex1-2, 4-5, 7, 8	4B:820974:824787
Ta/CS	4871850	4BS	Ex1	4B:2203566:2203808
Ta/CS	4945153	4BS	Ex4	4B:90873:90954
Ta/CS	2998791	6BS	Ex5	6B:85762514:85762595
Ta/CS	4928922	4BS	Ex6	4B:176356757:176356887
Ta/CS	4860697	4BS	Ex3, 6	4B:412957:414442
Ta/CS	4951082	4BS	Ex3, 6	4B:1613684:1614023
Ta/CS	4882842	4BS	$Ex7+8^g$	4B:1224732:1224857
Ta/CS	4910553	4BS	Ex7+8 ^g	4BS_scaff_4910553:2510:2622

^aA representative contig/scaffold is listed. ^bThe order of exons was based on the dominant *Ms2* gene (GenBanck: KX585234). ^cChromosome of 'Chinese Spring' (IWGSC1.0+popseq.28, http://plants.ensembl.org). ^dShares five exons with wheat EST AK331503. ^eShares two exons with wheat EST BE500370. ^fPredicted exons inverted in order. ^gPredicted exons mapped to the same region. ^hExon duplicated. ⁱPredicted exon partially retained. ^jAlternative locations (4A:171550005:171557187 and 4A:172222864:172224346). ^kAlternative scaffolds (scaffold101798:1:2597 and scaffold139909:61872:70085).

Note: Ae/AL8/78, Aegilops tauschii accession 'AL8/78'; Ta/CS, Triticum aestivum cultivar Chinese Spring; Tu/G1812, Triticum urartu accession 'G1812'. Fonts in gray indicate a predicted chromosome arm and its location in a sequence scaffold.

Supplementary Table 5. Counts of paired-end (PE) reads

Samples	Trials Clean Pairs		Clean Bases (bp)	Read Length (bp)	GC Content (%)
	1	37,863,983	9,465,995,750	125	54.40%
Fertile	2	44,837,668	11,209,417,000	125	54.80%
anther (FA)	3	38,790,150	9,697,537,500	125	54.00%
C4	1	63,136,384	15,784,096,000	125	55.20%
Sterile	2	38,953,021	9,738,255,250	125	56.30%
anther (SA)	3	41,563,567	10,390,891,750	125	54.90%
	1	35,551,128	8,887,782,000	125	54.30%
Leaf (L)	2	39,012,451	9,753,112,750	125	53.90%
	3	36,765,025	9,191,256,250	125	54.90%
	1	33,563,458	8,390,864,500	125	54.70%
Pistil (P)	2	38,232,211	9,558,052,750	125	55.10%
	3	35,980,271	8,995,067,750	125	54.30%
Total	12	484,249,317	121,062,329,250	125	54.76%

Supplementary Table 6. Counts of mapped paired-end (PE) reads

	Trials	Unique Mapped Reads		Multiple M	apped Reads	Unmapped Reads		
Samples		Clean Pairs	Percentage	Clean Pairs	Percentage	Clean Pairs	Percentage	
Fertile	1	29,162,254	77.02%	3,692,938	9.80%	5,008,791	13.23%	
anther (FA)	2	34,806,413	77.63%	4,171,441	9.30%	5,859,814	13.07%	
anulei (I'A)	3	29,829,956	76.90%	3,787,733	9.80%	5,172,461	13.33%	
Ctarila	1	48,501,465	76.82%	5,567,749	8.80%	9,067,170	14.36%	
Sterile	2	28,733,927	73.77%	3,444,548	8.80%	6,774,546	17.39%	
anther (SA)	3	31,655,229	76.16%	3,604,604	8.70%	6,303,734	15.17%	
	1	21,875,846	61.53%	3,589,986	10.10%	10,085,296	28.37%	
Leaf (L)	2	23,131,090	59.29%	4,057,438	10.40%	11,823,923	30.31%	
	3	24,059,675	65.44%	3,593,468	9.80%	9,111,882	24.78%	
	1	25,488,877	75.94%	3,461,341	10.30%	4,613,240	13.74%	
Pistil (P)	2	29,098,662	76.11%	4,075,446	10.70%	5,058,103	13.23%	
	3	27,348,856	76.01%	3,744,926	10.40%	4,886,489	13.58%	
Total	12	353,692,250	73.04%	46,791,618	9.66%	83,765,449	17.30%	

	Expressed	Gene Ex	xpression	Expressed Genes			
Samples	Genes (%) ab	Mean (SE d)	Median	5th	10th	50th	(%) bc
Fertile anther (FA)	69,459 (62%)	275 (5)	47	0.28	0.80	47	61,548 (55%)
Sterile anther (SA)	73,300 (65%)	309 (7)	44	0.43	1.04	44	65,803 (58%)
Leaf (L)	65,591 (58%)	469 (19)	58	0.64	1.55	58	61,109 (54%)
Pistil (P)	68,522 (61%)	247 (3)	49	0.28	0.83	49	60,968 (54%)
All ^a	79,153 (70%)	323 (5)	49	0.52	1.06 e	49	na
All ^c	na	358 (6)	67	1.87	3.29	67	72,499 (64%)

^aResults based on the normalized counts before removing false positives. ^bAt least one nonzero count value (count > 0) in each of three trials. ^cResults based on the corrected counts after removing false positives. ^dStandard error (SE) of the mean. ^eA count cutoff value was determined as the 10th percentile of expression levels¹⁷. Percentages were calculated based on 112,495 genes. na, not applicable.

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