1 Limited haplotype diversity underlies polygenic trait architecture

2 across 70 years of wheat breeding

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23 Abstract

24 Background

25 Breeding has helped improve bread wheat yield significantly over the last century.

26 Understanding the potential for future crop improvement depends on relating segregating

- 27 genetic variation to agronomic traits.
- 28 *Results*

29 We bred NIAB Diverse MAGIC population, comprising over 500 recombinant inbred lines, descended from sixteen bread wheat varieties released between 1935-2004. We sequenced 30 31 the founders' exomes and promotors by capture. Despite being highly representative of 32 North-West European wheat and capturing 73% of global polymorphism, we found 89% of 33 genes contained no more than three haplotypes. We sequenced each line with 0.3x 34 coverage whole-genome sequencing, and imputed 1.1M high-quality SNPs that were over 35 99% concordant with array genotypes. Imputation accuracy remained high at coverage as 36 low as 0.076x, with or without the use of founder genomes as reference panels. We created 37 a genotype-phenotype map for 47 traits over two years. We found 136 genome-wide 38 significant associations, concentrated at 42 genetic loci with large and often pleiotropic 39 effects. Outside of these loci most traits are polygenic, as revealed by multi-locus shrinkage 40 modelling.

41 *Conclusions*

Historically, wheat breeding has reshuffled a limited palette of haplotypes; continued
improvement will require selection at dozens of loci of diminishing effect, as most of the
major loci we mapped are known. Breeding to optimise one trait generates correlated trait
changes, exemplified by the negative trade-off between yield and protein content, unless

selection and recombination can break critical unfavourable trait-trait associations. Finally,
low coverage whole genome sequencing of bread wheat populations is an economical and
accurate genotyping strategy.

49

50 Introduction

Bread wheat (*Triticum aestivum* L.) production is a critical component of worldwide food security. Demand for wheat is predicted to increase by 60% between 2014 and 2050[1], by which time the human population will have reached 9 billion. Breeding will be a key component of meeting this demand sustainably[2]. Over the past century, genetic gains have been responsible for between one third and two thirds of yield improvements in European wheats, amounting to a 12-120kg increase in yield (~1%) per hectare per year[3–6].

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58 Genomic data is expected to accelerate the rate of genetic gain in wheat[7,8]. Surveys of 59 global standing genetic variation include, for example, whole genome resequencing (WGS) of 60 93 accessions[9], exome capture for 870 accessions[10], genotyping by sequencing (~16k 61 markers) for ~17k breeding programme lines[11], and genotyping array data for collections 62 of 804[12] and 4,500[13] accessions (~15k and ~113k markers, respectively). Bread wheat's large genome size (17Gb) inflates the cost of collecting sequencing data and its hexaploidy 63 64 reduces the accuracy and cross-population consistency of genotyping array data[12]. The potential for genotyping by low-coverage WGS in polyploid wheat has yet to be established. 65 66

To aid genetic gain through breeding, it is crucial to link genetic data with phenotypic
information and thereby reveal genotype-phenotype associations[11,14]. Previous

69 genotypic/phenotypic datasets include five traits measured in two years for 870 global 70 accessions with exome capture data[10], and 12 traits measured in two years, six locations, 71 and three cropping intensities for 191 German varieties with genotyping array data (~9k 72 markers)[15]. Genotype-trait and trait-trait associations may be confounded by population 73 structure or hidden by low allele frequencies in studies of existing varieties or breeding lines. 74 These problems can be controlled in experimental populations produced by crossing. 75 However, mapping resolution and overall genetic diversity are typically low in experimental 76 populations. Multiparent Advanced Generation Intercross (MAGIC) populations are designed 77 to address these issues by accumulating recombination events through generations of intercrossing and capturing diversity across multiple founders[16–18]. 78

79

80 In this study we undertook a systematic approach to these challenges. We bred a new multi-81 parental population, the 'NIAB DIVERSE MAGIC' population (hereafter 'NDM') through 82 hundreds of structured inter-crosses between sixteen diverse founders. Our multi-funnel 83 crossing design creates a greater number and more uniform genome-wide distribution of recombinant haplotypes than alternative multiparent populations[19] and the relatively large 84 85 number of diverse founders samples more genetic variation. We sampled founders released 86 between 1935-2004, aiming to determine the genetic basis for historical changes in 87 agronomic traits and the potential for future improvement from within the existing pool of 88 variants. We used a cost-effective genotyping strategy by low-coverage WGS, accurately 89 imputing over 1M SNPs in over 500 recombinant inbred lines. We measured 47 phenotypes 90 in the population, of which 25 were assessed across two growing seasons. The power of NDM comes from the combination of carefully designed germplasm and dense genotypic and 91 92 phenotypic information, all of which we make publicly available.

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94 We address the following questions. First, what genetic variation exists among the exomes of 95 the NDM founders, how does it reflect global wheat diversity, and specifically how many 96 distinct haplotypes typically segregate at each locus? Second, how does this variation underlie 97 agronomic traits, as revealed through genetic mapping and genomic prediction? And third, 98 what do these models imply about the future potential for phenotypic change and to what 99 extent should we expect selection to cause correlated trait changes due to the sharing of 100 causal genetic variants between traits.

101

102 **Results**

103 NIAB DIVERSE MAGIC Founders

104 The 16 founders were selected from a panel of 94 historical varieties released in the UK over 105 a ~70 year period (and originating from the UK, France, Denmark, Sweden and the Netherlands, Supplementary Table 1) using 546 Diversity Array Technology (DArT) and 61 106 107 Single Sequence Repeat (SSR) markers[20]. We sequenced 15 founders after enrichment for 108 (a) genic regions and (b) putative promoters using a capture probe-set[21] at average 109 coverage of 22.94x of the targets (Supplementary Table 1). The remaining founder, Holdfast, 110 was sequenced by WGS, but to ensure consistency across founders, we restricted our 111 attention to the capture targets, for which coverage in Holdfast was 15.8x. We sequenced 112 using Illumina 150bp paired-end reads whose combined length often included sequence 113 differences between homeologous loci on the A, B and D subgenomes of hexaploid wheat, 114 thereby resolving otherwise ambiguous alignments. Furthermore, we only used high quality 115 alignments (mapQ>30) for coverage calculations and variant calling, and subsequently

116 excluded variant sites with missing or heterozygous calls in any founder (e.g. from 117 homeologous variation and misalignment). After quality control, we called 1.13M high-quality 118 single nucleotide polymorphisms (SNPs) across the 110,790 promoter-gene pairs targeted by 119 the capture probes (557Mb in total), summarised in Supplementary Figure 1. Only 97,727 120 SNPs (8.7%) were on the D subgenome and almost half (17,289/35,021, 49.4%) of the 121 promoter-gene pairs on the D subgenome had no SNPs passing quality control, compared to 122 26.6% (9,656/36,302) and 21.7% (8,012/36,738) on the A and B subgenomes, respectively. A 123 comparative lack of diversity is expected on the D subgenome as it was acquired in the most 124 recent allo-polyploidisation event[22].

125

126 We placed the 16 founders in the context of global wheat diversity by analysing 113,457 127 genotyping array sites that vary among 4,506 diverse global wheat accessions[13], of which 128 50,335 sites were callable across all founders. We classified global wheats into nested subsets 129 representing the UK only (n=154), North-West (NW) Europe (n=1,343), Europe (n=2,331), and 130 Global (n=4,506), to understand how allele frequencies across subsets relate to our founders 131 (Figure 1). Most Global common variants are polymorphic in the founders whereas rare alleles 132 are more likely to be fixed in the founders, particularly those scarce in NW Europe and the 133 UK. For example, 79.7% of those SNPs polymorphic within the UK subset (which includes 134 landraces) also segregate among the founders, falling to 73.4% Global sites across all 4,506 135 accessions. We next asked whether we could have selected 16 founders that more 136 comprehensively sampled the variation space. We simulated selections from the same nested 137 subsets and compared the distribution of the fraction of segregating sites with that in the 138 actual NDM founders, and found the latter capture more diversity than an average selection 139 of UK wheats, about average diversity for NW European wheats, but less than average for

wider European and Global sets (Figure 1). As the Global dataset is highly diverse, with
modern varieties (released 1960-2009, n=2,294), landraces (1800-1959, n=965), and
uncategorised/landrace germplasm (n=1,247), we conclude that NDM is representative of

143 NW European wheat germplasm.

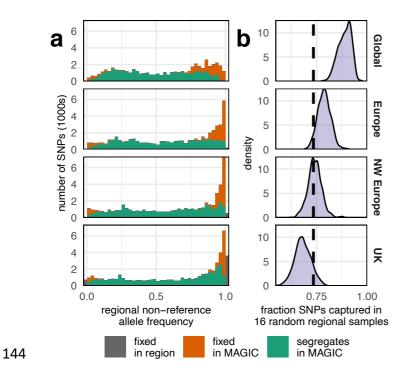


Figure 1 The NDM population is representative of NW European wheat. (a) SNPs segregating (green) or fixed (orange) in NDM at 50,335 sites in 4,506 global wheats, grouped into 'Global', 'European', 'North-West European' and 'UK' nested subpopulations and binned by the allele frequency in these subpopulations. (b) The fraction of sites that are polymorphic in 16 randomly chosen wheats from each subpopulation based on 1000 random replications. The dashed vertical black line at x=0.734 is the fraction of SNPs segregating among NDM founders.

151 We next estimated the haplotypic diversity in the founders at the 1.13M sites. First, we 152 clustered the founders by their haplotypic similarity at the 73,982/110,790 (66.7%) promoter-153 gene loci with at least two haplotypes. Assuming that founders carry the same haplotype 154 when their genotypic similarity exceeds 95%, 38,535 loci (52% of loci with variants) had only 155 two haplotypes, 61,438 loci (83%) had at most three haplotypes, and 70,602 loci (95%) had 156 four haplotypes at most (Figure 2b). Second, we estimated haplotype diversity by a dynamic 157 programming algorithm that adjusted locus/block boundaries (Figure 2c, Supplementary 158 Figure 2) to minimise the number of distinct haplotypes within a locus, while balancing transitions between calling identical versus non-identical haplotypes. Over a wide range of parameters, the average number of haplotypes present at any locus rarely exceeded two (Supplementary Figure 2: 81.2% of 1.13M sites inferred to have just two haplotypes). This analysis found slightly fewer haplotypes than the gene-based analysis because it can infer one haplotype (4.1% of sites) when nearby variation is inconsistent, and split genes with high haplotypic diversity into multiple blocks.

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166 For comparison, the 19 natural accessions that founded the Arabidopsis thaliana MAGIC[23] 167 display much greater haplotypic polymorphism[24]. In A. thaliana, genic haplotypes were determined at the level of protein sequence similarity (>95% similarity within 168 169 haplotypes)[24]. On average there were 4.8 protein haplotypes per gene and 7,263/13,919 170 (52.2%) of genes with two (n=4,825) or three (n=2,438) haplotypes (excluding genes with no 171 variation). Our estimates for the NDM founders are 2.7 haplotypes per gene and 83% of 172 variable genes having at most three haplotypes. Protein-level differences are lower than DNA 173 level differences making this comparison conservative, and thus the true difference is even 174 greater.

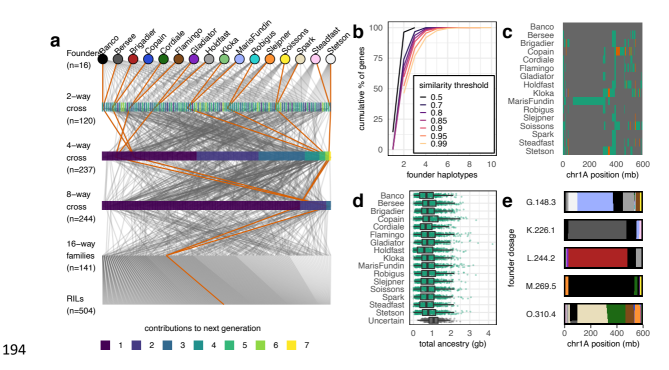
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176 The NIAB DIVERSE MAGIC Population

We developed a total of 596 Recombinant Inbred Lines (RILs), each descended from all 16 founders via a crossing funnel (Figure 2a). After 6 generations of inbreeding, all 596 RILs were initially genotyped using the Axiom 35k wheat breeders' SNP genotyping array[12]. We called SNPs at 20,688 sites, of which 5,747 overlapped with the 1.13M SNP calls made in the founders. These overlapping sites suggested that only 59.8% of genotyping array probes could have been unambiguously placed using BLASTn[25], underlining the difficulty of using short probes in polyploids (Supplementary Table 2). We used the overlapping sites as a truth genotype set to find sample misidentifications and estimate the accuracy of sequence-based genotyping in the RILs.

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We excluded 46 RILs excessively similar (>92%) to other RILs, indicating possible errors during population development. We sequenced the remaining 550 RILs after 7 generations of inbreeding by low coverage WGS (mean 0.304X) and called variants at the 1.13M founder SNP sites using sequence alignments. A further 46 RILs were excluded as their genotypic concordance with the initial 35k array data was below 95%, leaving 504 RILs in 141 families (RILs in the same 'family' are derived from the same 16-way cross), from which we based our main analyses.



195 Figure 2 NDM population design and haplotypic diversity. (a) Pedigree showing the construction of 504 196 Recombinant Inbred Lines (RILs). One exemplar pedigree is highlighted to show how all 16 founders are 197 intercrossed into each RIL. (b) Founder haplotype groups at 73,982 promoter-gene loci with SNP variation, where 198 founders with the same haplotype have genotypic similarity fractions that exceed the corresponding threshold. 199 (c) Pairwise similarity/dissimilarity between founders on chromosome 1A, determined using a dynamic 200 programming algorithm to infer founder similarity and breakpoint position. Founders that are inferred to have 201 similar haplotypes for each region are the same colour. (d) The total length of genomic blocks in NDM lines 202 inferred to come from each founder; uncertain ancestry blocks have a maximum founder dosage of <90%. (e)

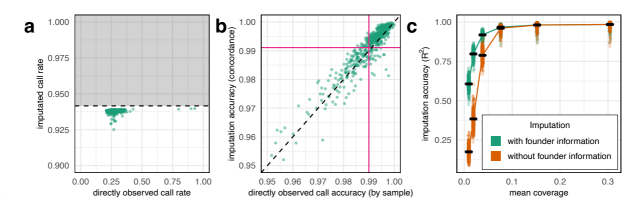
203 Inferred founder dosage and ancestry mosaics across chromosome 1A for five example RILs, with founders 204 coloured as in (a).

205 We imputed RIL genotypes using STITCH[26] by inferring the founder haplotype carried by 206 each line at each location. Figure 2c shows the haplotypic similarity among founders on 207 chromosome 1A, indicating that a small number of haplotypes have been heavily recombined during the 69 years of breeding history that separates the founders. Most recombination is 208 209 located towards the distal ends of the chromosomes, as expected[27]. Only limited further 210 recombination occurs during MAGIC population construction and the haplotype blocks 211 inherited from each founder are relatively long (Supplementary Figure 2) and therefore 212 distinguishable from one another. Thus, it was necessary to assume 16 unique haplotypes 213 were segregating to obtain the highest imputation accuracy (Supplementary Figure 2). 214 Founder haplotypes could be confidently assigned (i.e. with >90% dosage from a single 215 founder) at over 92.2% of sites (Figure 2d). These haplotype assignments implied that an 216 average of 4.8-13.7 recombination events occurred per RIL per chromosome (mean 8.7 sd 2), 217 giving an average of 183 (sd 36.3) recombination events per RIL in total. Consistent with 218 estimated genetic map lengths of 35-37.4M[12,28], 4.9-5.2 recombination events were 219 observed per Morgan, in line with the predicted ~5-fold increase in 16-parent MAGIC 220 populations compared to two-way crosses[29]. Example founder haplotype mosaics across 221 chromosome 1A are shown in Figure 2e.

222

The fraction of sites called directly (i.e. without imputation) for 501 RILs varied between 20.9-42.7% (mean 27.8% sd 3.4%), as expected for 0.3x-coverage sequence data. A further three RILs were sequenced to higher depth (2.7x, 4.0x, and 4.3x) and had call rates of 79.9%, 90.0%, and 93.0%, respectively (Figure 3a). After imputation, 94.2% of the 1.13M SNPs (i.e. 1.07M) were called across all 504 RILs and the effective call rate of imputed sites was 99.6%, with

228 5.8% of the SNP sites inaccessible or removed by quality control: 0.93% of sites are on the 229 "Un" chromosome in the wheat reference (excluded from imputation), 1.36% were removed 230 by imputation QC (info score <0.4) and 3.52% had imputed minor allele frequencies below 231 2.5% and/or missingness above 90%. Figure 3b shows that the concordance between array 232 and imputed genotypes (AI) and between array and directly called genotypes (AD) are 233 strongly correlated, suggesting that instances of poorer concordance are unlikely to be caused by imputation. Overall, imputation marginally improved accuracy versus direct calls (mean AI 234 235 99.1% versus mean AD 99.0%) but increased the call rate three-fold. Downsampling read 236 coverage showed the founder haplotype space and recombination mosaics could be 237 accurately inferred from coverage as low as 0.076x per sample (Figure 3c); above this level imputation accuracy was independent of whether founder haplotypes were included as a 238 239 reference panel (mean AI 98.7%) or ignored (mean AI 98.5%).



240

Figure 3 Call rate and accuracy of genotypes after imputation and after downsampling. (a) Imputed call rate (y-axis) vs direct call rate (x-axis. Only 28.1% of the 1,131,251 SNP sites can be genotyped directly from the low coverage sequence data, whereas 93.8% of sites had genotypes after imputation. 5.8% of sites (grey region and horizontal dashed line in a) were removed by quality control filters after imputation or on the unimputed 'Un' chromosome (0.93%). (b,c) Accuracy as evaluated at 5,747 sites that overlap with the Axiom 35k array. (c) Imputation before/after downsampling was performed with (green) and without (orange) using the genotypes of the founders as a reference panel.

249 Introgressions and Segregation Bias

250 Several recent studies have used genomic data (e.g., SNP density[9]) to study the 251 introgression of genetic material into hexaploid bread wheat from the secondary and tertiary 252 gene pool[9,10,30]. We examined evidence for introgressions in previously reported 253 locations[9,31–33] using founder coverage and non-reference allele frequency. Because we 254 developed RILs, we were also able to examine segregation bias, which often accompanies 255 wheat introgressions[34,35]. We found evidence for at least six introgressions covering 256 ~1.1Gb segregating in the population, five of which showed segregation bias (Supplementary 257 Table 3).

258

259 Phenotypic Characterisation and QTL Mapping

260 We measured 47 phenotypes in replicated field trials over two years (Table 1, Supplementary 261 Tables 4, 5, and 6), including the 10 time points at which Green Leaf Area (GLA) was measured. 262 Of these, 25 phenotypes were collected in both years and two were also measured in smaller 263 1x1m nursery plots (Yellow Rust infection, YR, and Juvenile Growth Habit, JGH) to give a total 264 of 73 phenotypic measurements. Phenotype distributions are shown in Supplementary Figure 265 3, showing that some RILs have more extreme phenotypes than any founder (transgressive segregation) for almost all phenotypes (RIL maximum \geq founder maximum for 61/73 266 267 phenotypes and RIL minimum \leq founder minimum for 68/73 phenotypes). All phenotypes 268 have significant (p<0.05, Pearson's correlation test) correlations with at least one other phenotype (Supplementary Figure 4). 269

270 Table 1 Phenotypes collected

ABBREVIATION	TRAIT	ABBREVIATION	TRAIT
BIS	Basal infertile spikelets	GS55	Ear emergence date
EL	Ear length	GS65	Anthesis date
ETA	Ear taper	GW	Grain width
ETS	Ear tip sterility	GY	Yield

EW	Ear weight	HEB	Height to ear base
FLA	Flag leaf angle	HET	Height to ear tip
FLED	Flag leaf to ear distance	HFLB	Height to flag leaf base
FLF	Flag leaf floppiness	JGH	Juvenile growth habit
FLL	Flag leaf length	LOD	Lodging
FLS	Flag leaf senescence	PHS	Pre-harvest sprouting
FLW	Flag leaf width	PIG	General pigmentation
GA	Grain area	SH	Spring habit
GL	Grain length	SPIG	Stem pigmentation
GLA#	Green leaf area (10 time points, Nov–Mar)	SW	Specific weight
GLAU	Glaucousity	TGW	Thousand grain weight
GPC	Grain protein content	TIS	Tip infertile spikelets
GR	Germination rate	TS	Total spikelets
GS39	Flag leaf emergence date	YR	Yellow rust infection

271

From the 1.07M imputed SNPs, we selected a subset of 55,067 pruned by linkage 272 273 disequilibrium (LD). Using genome-wide association scans (GWAS) on both SNP and founder 274 haplotype data, we mapped 136 Quantitative Trait Loci (QTLs) across the 73 phenotype/year 275 combinations that were genome-wide significant at the 5% level. Many QTLs overlapped for 276 different phenotypes, clustering into 42 distinct genome locations. For 25 phenotypes that 277 were measured in both years, we found 48 QTLs in year 1 and 49 QTLs in year 2, of which 28 278 were mapped to the same location and were genome-wide significant in both years. For 279 example, in replicated trials lacking fungicide treatment we mapped yellow rust (Puccinia 280 striiformis) susceptibility to four QTLs in year 2 (on chromosomes 2A[31,36], 2B[37], 3B, and 6A), of which three were also mapped in year 1 (2A, 3B, and 6A); only one (6A) was also 281 282 mapped in trials treated with fungicide. 126/136 QTLs at 40/42 genomic locations were 283 mapped using SNP-based associations, whereas 87/136 QTLs at 30/42 genomic locations 284 were mapped using haplotype-based association tests. That is, 10 QTLs and two genomic 285 locations were only identified from haplotype-based association whereas 49 QTLs and 12 286 genomic locations were only identified from SNP-based association. This is consistent with 287 the limited gene-level haplotypic diversity observed among the founders.

288

289 Figure 4b summarises the 40 loci with genome-wide significant SNP-based associations. We 290 were able to assign 21 of these, including most of those with the strongest effects, to 291 previously reported QTLs. In 11 high confidence cases, candidate genes have been reported 292 and/or validated experimentally. In other cases, QTLs contained homeologs or paralogs of 293 these high confidence candidates, or previous studies had reported associations to a genetic 294 map using marker data, but not firmly anchored these locations on the reference genome 295 assembly (low confidence co-localisation, n=10). We checked six high confidence candidate 296 loci with annotated reference genome locations (RHT-1[38], RHT-2[39], WAPO-A1[40], ALI-297 1[41], TaMyb10-B1[42], Yr7/Yr5/YrSP[37], PPD-D1[43]), all of which were within our mapping 298 intervals. We created a genotype-phenotype map for community use by placing all QTLs on 299 the physical map (Supplementary Table 7) to a median interval of 9.2Mb.

300

Most loci with strong effect co-localise with previously reported QTLs. Some large effects are commonly associated with adaptation of the founders to the geographic and temporal range they sample. For example, the early flowering allele at the photoperiod locus *PPD-D1* carried by the founder Soissons is favoured in southern Europe to avoid the summer drought[44]. The modern semi-dwarfing alleles at *RHT-B1* or *RHT-D1* that have been favoured globally since the Green Revolution[45] are absent from founders Banco, Bersee, Copain, Flamingo, Holdfast, Kloka, Spark, Steadfast and Stetson.

308

To examine the pleiotropic effects of the relatively few genome-wide significant QTLs, we took the most strongly associated SNP at each locus and then tested for associations with all other phenotypes, requiring a lower threshold for evidence of association (p<0.05) than was initially used to establish genome-wide significance. The results are visualised in Figure 4d,

which shows that loci significant for one phenotype are also common to other phenotypes,
consistent with extensive pleiotropy and the shared genetic control of correlated phenotypes.

316 *Gene Deletions*

317 Our analysis of SNP variation ignored sites that could not be called reliably in all 16 founders, 318 possibly due to whole-gene deletions relative to the reference genome. We obtained no 319 coverage from at least one founder at 8,019 (7.2%) of genic regions and 1,095 (1.1%) of 320 promoter-gene pairs, suggesting possible structural variations (Supplementary Figure 1). 321 Based on the deviation in gene coverage from that expected given the mean coverage for the 322 founder, we computed a quantitative gene deletion score (GDS) for each gene and founder 323 and imputed the scores into the RILs using the founder ancestry mosaics. We tested the association between each GDS and each phenotype in order to identify potential causal 324 325 deletions. Across 27/73 phenotypes we found 30 GDS associations with p-values $<10^{-6}$ 326 (Supplementary Table 8). Significant associations almost always occurred within QTLs 327 previously mapped by SNP association, so this analysis only identified candidate genes with 328 deletion status consistent with the pattern of action across the founders of a QTL. Of these, 329 at 10 loci the peak GDS logP association was at least 90% of the peak SNP logP. Thus most 330 QTLs are not likely to be caused by gene deletions. However, the GDS is based on empirical 331 read coverage, and so is likely to be affected by stochastic experimental variations hence it is possible that the association at a true causal GDS might appear weaker than that of a tagging 332 333 SNP. A further caveat is that deletions are always inferred relative to the reference genome 334 of Chinese Spring, such that insertions or functional genes missing from the reference genome 335 annotation will not be captured.

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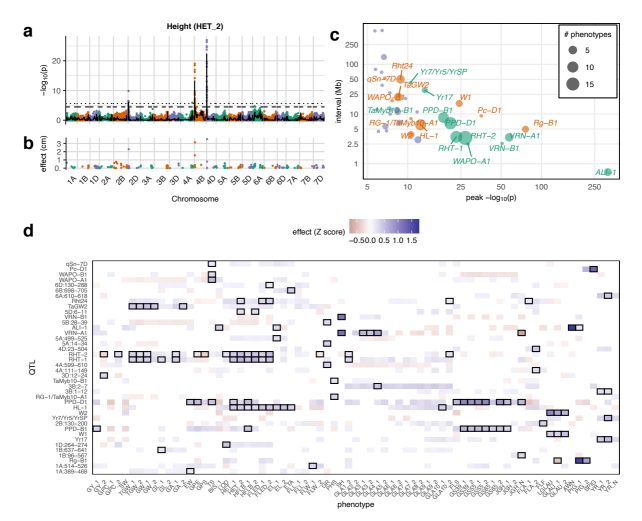
337 Genomic Prediction

338 We next performed phenotypic prediction all 55,067 tagging SNPs, to predict the potential 339 for genetic improvement within the NDM. We trained genomic prediction models using three 340 shrinkage methods: ridge regression (RR), least absolute shrinkage and selection operator 341 (LASSO) and Elastic Nets (EN), using 50-fold cross-validation with randomly-selected training 342 sets comprising 90% of RILs and test sets of the remaining 10%. LASSO and EN had almost 343 identical prediction accuracies but EN included on average 26% more SNPs than LASSO 344 (Supplementary Figure 5). Accordingly, we only report the LASSO results. LASSO prediction 345 accuracies for all traits are shown in Figure 5b, alongside the proportion of heritable variation 346 explained by QTLs (Figure 5a). Across traits, LASSO had higher average prediction accuracy 347 than RR (Figure 5c), particularly for phenotypes where a larger fraction of variation can be 348 explained by genome-wide significant QTLs (Figure 5d), as expected for a model selection 349 method. LASSO prediction accuracies (correlation coefficients) varied from 0.13-1 (mean 350 0.43) across phenotypes, using models with 1-465 SNPs (mean 155 SNPs). The number of 351 SNPs in the LASSO model is higher for phenotypes where the overall heritability estimate 352 greatly exceeds the fraction of variation that can be explained by genome-wide significant 353 QTLs (Figure 5e).

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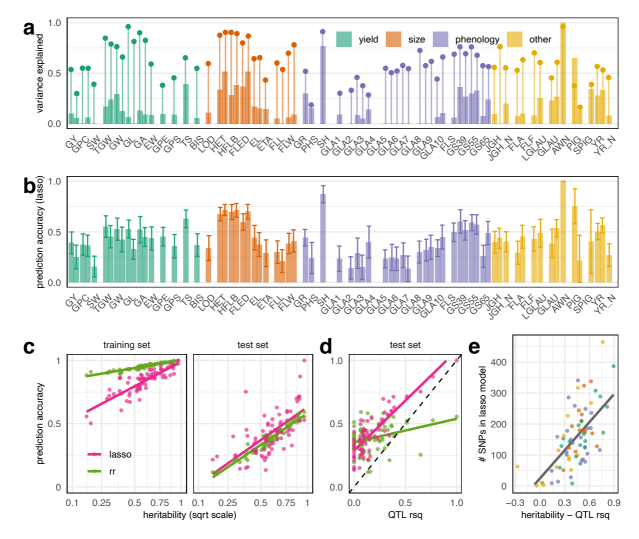
Out-of-sample test set prediction confirms that polygenic LASSO SNPs have predictive power and are therefore likely to be tagging genetic variants affecting phenotypic variation. Most phenotypes are polygenic; their prediction models exhibited a mixture of a few large effect and many smaller effect loci. A typical example (for height) of the 193 non-zero LASSO SNP effects is shown in Figure 4c. In contrast, the Mendelian AWN phenotype is fully explained and predicted using a single genome-wide significant QTL.

362	The reduced accuracy of RR compared to the LASSO is expected in the absence of significant
363	population structure. There will be reduced variation in kinship among RILs compared to the
364	wider germplasm from which the founders are usually selected. Much of the prediction
365	accuracy of RR results from exploiting kinship rather than from tagging causative variants[46]
366	so there is less opportunity for high prediction accuracy in MAGIC populations. In these
367	circumstances, a feature selection method such as the LASSO can more accurately identify
368	and tag haplotypes contributing to trait variation and give greater prediction accuracies. The
369	LASSO also accurately predicts traits determined predominantly by a few QTL of large effects,
370	in which circumstances RR performs poorly (Figure 5d). The LASSO is therefore better for
371	genomic prediction in MAGIC.



372

373 Figure 4 Genotype-phenotype associations. (a) Exemplar Manhattan plot of the genome-wide -log₁₀ p values of 374 association (logP) between the height to ear tip phenotype from year 2 (HET 2) and 55,067 LD-pruned SNP 375 dosages (dots) or founder haplotype dosages (line). The horizontal lines show the 5% genome-wide significance 376 thresholds for SNPs (dotted) and haplotypes (dashed). (b) The 193 non-zero estimated LASSO SNP effects for 377 HET 2. (c) The 40 genomic locations where genome-wide significant SNP associations were found for at least 378 one phenotype, classified by effect size (logP; x-axis) and genomic interval width (Mb; y-axis). Each circle 379 represents one locus, and its size shows the number overlapping QTLs; the smallest interval width and p value is 380 shown where there are multiple overlapping phenotype associations. Labels indicate QTLs that colocalise with 381 previously described QTLs or candidate genes; green indicates high-confidence colocalization (n=11) and purple 382 low-confidence colocalization (n=10). (d) Pleiotropy across 40 loci: those loci without names are labelled by 383 chromosome and position in Mb) and 73 phenotypes. Shades indicates the significant (p<0.05) locus phenotypic 384 effects expressed as the number of standard deviations (Z-score). Genome-wide significant QTLs are highlighted 385 with boxes.



386

387 Figure 5 Genetic architectures of 73 trait/year combinations (47 distinct traits) as revealed by QTL mapping and 388 genomic prediction. (a) Phenotypic variation explained by all genome-wide significant QTLs (thick bars) and by 389 the full SNP-based genetic relationship matrix (heritability, thin bars and dots). Phenotypes measured in year 1 390 and year 2 are paired, shifted to the left and right, respectively. (b) LASSO prediction accuracy (correlation 391 coefficients) across 50-fold cross validation; error bars show sds. (c) Prediction accuracy correlations (y-axis) and 392 sqrt(heritability) (x-axis) and in the test and training sets under ridge regression (rr) and LASSO genomic 393 prediction models. Prediction into the test set is generally higher with LASSO, especially for traits where more 394 variation is explained by genome-wide significant QTLs (d). (e) LASSO models usually include more SNPs when 395 more heritable variation is unaccounted by genome-wide significant QTLs (x-axis is difference between 396 heritability and QTL R²).

397 We used these genomic prediction models to explore the potential for selection in a much

398 larger simulated population of 20,160 MAGIC RILs, 40 times larger than the real population.

399 These were created by permuting the founder identities in the founder genome mosaics

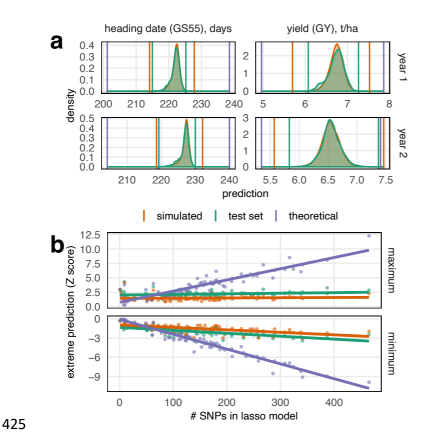
400 inferred in the real RILs, preserving linkage through the genetic map. Phenotypes were

- 401 predicted for the test set of real RILs (10% of all lines) and in the simulated RILs for all 50
- 402 prediction models (training/test set jackknife resamples). Figure 6a shows, for two example

403 phenotypes, that the distribution of predicted phenotypes is almost identical in the real (test 404 set) and simulated RILs. As expected, the most extreme predicted values (maximum and 405 minimum) in the simulated RILs exceed than those in the real dataset because novel allelic 406 combinations are generated in the larger simulated population. However, the average 407 improvement in extrema between the test set and simulated phenotype predictions is only -408 0.5 (for the minimum) and +0.68 standard deviations (for the maximum). This is in line with 409 extreme-value distribution theory and shows that blind-breeding a very large population in 410 the hope of generating novel combinations of beneficial alleles is inefficient.

411

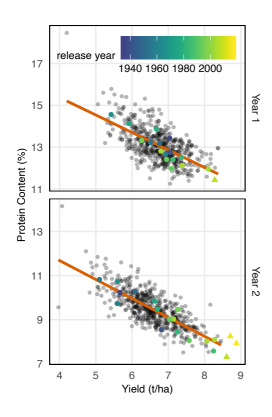
412 Next, we predicted the theoretical extreme phenotypic values that it is possible to create 413 from segregating variation if unlimited recombination were possible. That is, we computed 414 the phenotypic prediction in an imaginary line that carries all the alleles predicted to 415 increase/decrease each phenotype. For this exercise, we trained the prediction models on 416 the full set of 504 RILs so they differ slightly from those used to predict phenotypes in the test 417 set. In the test set and simulated RILs, the predicted phenotypic extremes generally reflect 418 the population size, which determines the probability that a single line happens to sample 419 many alleles with positive/negative effects. However, Figure 6b shows that the theoretical 420 maximum/minimum phenotypic prediction is linearly related to the complexity of the LASSO 421 model (i.e. the number of non-zero SNP coefficients in the model). This suggests that 422 hundreds of loci would need to be selected over multiple generations to generate any large 423 phenotypic shifts, in line with the decades of breeding that has been required to produce 424 genetic gain.



426 Figure 6 Predicted potential for phenotypic change. (a) We predicted the phenotypes of real MAGIC recombinant 427 inbred lines, RILs (green distribution), and a large population of 20,160 simulated MAGIC RILs (orange 428 distribution). These distributions largely overlap but more combinations are made in the simulated dataset such 429 the the extreme values are more extreme. Nevertheless, the highest/lowest phenotypic prediction in the 430 simulated population of 20,000 is generally only ~0.5 standard deviations higher/lower than the trait predictions 431 in the real dataset of 504 lines. Upper graphs: predictions based on year 1 phenotype, lower graphs predictions 432 based on year 2 phenotype (b). We also estimated the externes of the phenotype predictions that are possible 433 given the full lasso genomic prediction models (purple line in a). Large deviations from the current population 434 mean are predicted to be possible but only through the fixation of a large number of loci, with less potential for 435 change predicted at less-highly polygenic traits.

436 For essentially all crops where yield and yield quality are high priority traits, a trade-off is 437 evident between these two phenotypes and this is recognised as a longstanding problem in 438 wheat. Thus, identifying opportunities to break this trade-off is important[47,48]. We estimate that yield has increased by 0.021 t ha⁻¹ year⁻¹ based on a regression of average yield 439 440 on founder release year (p=0.006, n=16, R²=0.43). The highest yields measured in founders 441 and RILs exceeds the maximum predicted yield from the genomic prediction models (Figure 6) due to shrinkage in estimating SNP effects. However, high grain yield (GY) is correlated with 442 low grain protein content (GPC) among the founders (Pearson's correlation coefficient -0.94, 443

444 p<0.001, n=16), Figure 7. Founder genetic material is reshuffled without selection in the RILs, 445 but the GY-GPC relationship continues (correlation -0.77, p<0.001, n=504), suggesting some 446 pleiotropy in the underlying genetic effects. To investigate the segregating genetic variation 447 that may be available to break this trade-off, we analysed the deviation from the trend (PYD: 448 distance from symmetrical Thiel-Sen regression between GPC and GY, after Z-score 449 normalisation). The heritability for PYD was 0.41 in year one and 0.25 in year two and could 450 be predicted with accuracy 0.26 (sd 0.11) in year one and 0.13 (sd 0.11) in year two. These estimates are lower than those for GY and GPC analysed separately (GY heritability 0.54 and 451 452 0.30, prediction accuracy 0.39 and 0.25; GPC heritability 0.55 and 0.55, prediction accuracy 453 0.375 and 0.36, Figure 5. PYD of the founders did not correlate with release date, but these 454 results suggest modest potential to break the yield-protein trade off, requiring strong and 455 targeted breeding effort[47,48].



458 Figure 7 Negative trade-off across two years between grain yield (GY; x-axis) and grain protein content (GPC; y 459 axis) in 504 NIAB DIVERSE MAGIC RILs, 16 founders, and 3 more recently developed varieties (triangles, only
 460 one measured in year 1).

461 Discussion

457

462 We report five main findings. First, imputation from low coverage WGS is a cost-effective and 463 straightforward genotyping strategy for crops, at least in multiparental populations. Despite 464 its large, repetitive and hexaploid genome, wheat genotypes can be reliably imputed from 465 WGS with average per-sample coverage in the range of 0.075x-0.3x and without the use of 466 reference panels[26]. Thus there is no absolute requirement to even know the identities of, 467 let alone sequence, the population founders, although this may be desirable for other 468 purposes such as pan-genome assembly and re-annotation[18,49]. In this study, we were 469 able to impute genotypes and founder haplotypes at >1M SNP sites in >500 NDM RILs, which 470 proved ample for genetic mapping and genomic prediction.

472 Second, based on SNPs called from exome capture, no more than three haplotypes segregate 473 at most genes in commercial NW European bread wheats released since 1935. There appears 474 to be little or no variation at about a quarter of genes on the A and B subgenomes, and at 475 about half on the D subgenome. Complete re-assembly and re-annotation of the 16 founders 476 of the NDM would yield more complete insights into the extent and impact of coding 477 variation. Limits on haplotypic variation are probably the result of historical selection and 478 population bottlenecks that reduced the effective population size before the onset of 479 intensive breeding programmes[12,50], as well as the close relatedness among breeding 480 materials in more recent wheat pedigrees[51]. However, it appears that the low overall level of genetic diversity has not been further reduced during the 20th Century[52,53]. 481

482

483 Third, as a consequence, most QTLs are accounted for by bi-allelic SNPs rather than haplotype 484 differences. For comparison, about 40% of QTLs identified in a multi-founder population of 485 rats were attributed to multi-allelic/haplotypic effects[54]. Furthermore, most genome-wide 486 significant QTLs had pleiotropic effects. Extensive pleiotropy suggests that naïve selection on 487 one phenotype is likely to induce correlated responses in other phenotypes. In particular, we 488 found improved yields in recent varieties has come at the cost of a decline in protein content 489 (Figure 7; increasing yield by one t/ha reduces protein content by about 1%). Despite 490 reshuffling haplotypes without selection, this trade-off continues in the NDM, which indicates 491 directed selection would be required to break the yield-quality trade-off, potentially creating 492 varieties with improved nitrogen use efficiency[55,56].

493

Fourth, across 47 phenotypes, we found a wide range of underlying genetic architectures. For
traits such as awns, pigmentation, spring habit and yellow rust resistance, almost all of the

496 heritable phenotypic variance could be explained by one to four genome-wide significant 497 QTLs (Figure 5). In other cases, a few loci with large phenotypic effects were accompanied by 498 dozens of loci with smaller effects on traits such as flowering time and height (Figure 4a). The 499 loci with very large effects have mostly been reported before (Figure 4b) because we 500 recapitulate key historical steps such as the introduction of photoperiod sensitive and semi-497 dwarfing alleles from Japan[44,45]. Traits such as yield were polygenic with the majority of 502 heritable variation coming from many loci of smaller effect (Figure 5).

503

504 Fifth, our genomic prediction models suggest that hundreds of loci will need to be selected 505 and fixed to achieve large phenotypic changes in polygenic traits in the future (Figure 6B). We 506 achieved reasonable prediction accuracy with modest numbers of SNPs; the mean out-of-507 sample prediction accuracy was 0.43, using on average only 155 SNPs per phenotype, out-508 performing ridge-regression which considers all markers simultaneously. Other crop and 509 livestock studies have also found very sparse markers can be sufficient for useful genomic 510 prediction[11,57,58]. Here, rather than using low marker densities, we trained models that 511 select a few hundreds of markers from ~55k tagging SNPs. In part, this sparsity is a 512 consequence of the design and construction of MAGIC populations, eliminating rare alleles 513 and creating blocks of markers that can be easily tagged in prediction models[15]. These 514 factors may be responsible for the use of far fewer markers than used to generate polygenic 515 prediction scores in humans[59], where there is a long tail of rare variation and less linkage 516 disequilibrium.

517

518 Our results suggest that dramatic genetic improvement over 70 years of breeding has largely 519 been achieved through the fine shuffling of a low number of haplotypes to recombine

polygenic alleles of small effect, combined with the introduction of alien introgressions from wide crosses. The introgression of large genomic segments from related species has most commonly been for sources of resistance to specific diseases[9,33,34]. Breeders now have a choice whether to continue with the same strategy, i.e. selecting from within existing variation and introgressing selected exotic alleles, or to ambitiously expand the pool of available haplotype diversity genomewide.

526

527 Methods

528 NDM Population Creation

The 16 NDM founders were chosen to capture the greatest genetic diversity using 529 530 PowerMarker genetic analysis software[60]. They were chosen from 94 NW European wheats released in the UK that were genotyped with 546 DArT and 61 SSR markers; the full panel also 531 532 included 96 US and 50 Australian varieties, which were excluded based on STRUCTURE 533 analysis[61]. The founder selection process was run iteratively with the varieties 'Robigus' 534 and 'Soissons' first fixed to be included to coincide with the founders of the 8-founder NIAB 535 Elite MAGIC population[62]. Then the most frequently selected additional 4, then 9, and 12 536 varieties were fixed in multiple iterative selection runs and finally the most frequently 537 selected 16 were chosen. Seed for the founding varieties was sourced from the John Innes Centre Germplasm Resource Unit (GRU http://www.jic.ac.uk/germplasm/). 538

539

540 These founders were inter-crossed in a balanced funnel crossing scheme, based on a Latin 541 square field trial design, over four generations to create 16-way crosses with all the founders 542 equally represented in their pedigree. First, all 120 possible 2-way crosses between founders

543 were made in a half diallel scheme. Two-way plants were then crossed in 60 4-way 544 combinations. Multiple plants from each family were used in crossing from 2-way onwards, 545 in order to maintain maximum founder allelic diversity within the population. 30 crossing 546 combinations were made between 4-way plants to create 8-way crosses, making between 547 five and eight replicate crosses per combination using different plants. These were intercrossed in 15 combinations to create balanced 16-way crosses, with each combination 548 replicated between six and fifteen times using different 8-way plants. This resulted in 174 16-549 550 way plants from which one to sixteen inbred lines per 16-way family were made through 551 single seed descent (SSD). 596 RILs were advanced to the F_7 stage when seed for phenotyping was multiplied in 1x1m nursery plots. Supplementary Table 9 gives details the number of 552 553 plants involved in each cross and Figure 2a shows the pedigree for the 504 RILs used in our 554 main analysis only.

555

556 *Phenotyping*

RILs from the population were phenotyped in field trials over multiple environments near 557 558 Cambridge, UK. Yield trials were conducted in the growing seasons 2016-2017 and 2017-2018, 559 hereafter year 1 and year 2 (phenotype suffix codes 1 and 2). Information on location, soil 560 type, key dates and inputs for both years are given in Supplementary Table 4. Yield plot 561 dimensions were 2m wide and 4m long and plots were sown at a density aiming to achieve 300 plants m⁻². In year 1, 596 lines were included in two replicates, the sixteen founders in 562 563 four replicates and the commercial control variety 'KWS Santiago' in 24 replicates in a 564 randomised nested block design with 16 main blocks of 80 adjacent plots which comprised 565 each row in the trial and eight sub-blocks of ten plots nested within each main block. In year 566 2 trials, 596 lines and the 16 founders were included in two and four replicates respectively 567 but three control varieties ('KWS Santiago', 'Skyfall' and 'Shabras') were all included in four 568 replicates. Plots were again randomised in a nested block design but including additional plots 569 making a larger trial, consisting of 20 main blocks of 115 adjacent plots, which comprised each 570 row, and 23 sub-blocks of five plots nested within each main block.

571 Disease observation trials (DOTs) were conducted near Cambridge, UK in the same years as 572 the yield trials to assess resistance to crop diseases. These plots consisted of two 1.2m length 573 rows, treated with no fungicide but otherwise standard inputs. Due to local conditions, DOTs 574 were considered to have natural high pressure of yellow rust (*Puccinia striiformis* f.sp. tritici). 575 In both years, DOTs included two replicates of 596 RILs, four replicates of the 16 founders and 68 additional replicates of the susceptible founder 'Robigus'. Trial designs included two main 576 577 blocks of 660 plots, with 11 sub-blocks of 60 plots nested within main blocks. All trial designs 578 for both yield and disease observation trials were made using the package 'blocksdesign' in 579 R. Phenotyping of some traits was also carried out in 1x1m seed nursery plots where lines 580 were not replicated but the founders were in three replicates and randomised across the 581 nurseries (phenotype code N).

582

A wide range of traits were phenotyped across the field trials, including traits for crop developmental morphology, phenology, plant stature and canopy architecture, yield and yield components such as spike and grain morphology, disease resistance, pigmentation, plant glaucosity, indications of stress response, lodging, grain protein content and vernalisation requirement. A summary of these traits and abbreviations are presented in Table 1 and details of phenotyping methods are listed in Supplementary Table 5.

589

590 Trials Analysis

591 Adjusted phenotype values were calculated as Best Linear Unbiased Estimates (BLUEs) for 592 each trait separately for each trial year using mixed effects models with ASRemL[63]. 593 Genotype was considered a fixed effect whilst experimental blocking structure as well as 594 other covariates such as harvesting day, where relevant, were included as random effects. 595 Spatial models including first- and second-order auto-regressive spatial models were also 596 used. Model simplification was carried out where models with all possible combinations of 597 random effect terms and spatial terms for row and column were run and the best fitting 598 model was chosen based on Akiake Index Criteria (AIC). Model residuals were visually checked 599 for normality and equal variance to fitted values distribution. Best Linear Unbiased Estimates 600 (BLUEs) for all phenotypes for the 16 founders and for the 504 RILs used in our main analysis 601 (see below) are provided in Supplementary Table 6. We used symmetrical Thiel-Sen 602 regression (implemented in the 'deming' R package) after phenotype normalisation to 603 characterise the relationship between protein content (GPC) and yield (GY). The Protein-Yield 604 Deviation (PYD) phenotype is calculated as the Euclidian distance from this regression line.

605

606 Genotyping Array Data

All DNA extraction was performed using the Qiagen DNeasy Plant Kit on leaf tissue samples taken from emerging leaves of seedlings. First, genotyping was performed at the Bristol Genomics Facility using the Axiom 35k wheat breeders' array[12]. Initially, two 384-sample plates were genotyped. Seed from the plants used as founders were genotyped on each plate (32 samples) along with extra seed from the original varietal seed stock used (28 samples) and seed from founders propagated to 2017 (16 samples). In addition, 596 RILs were genotyped after 5 generations of selfing (F₆). To account for genotyping failures and to ensure

614 the accuracy of sample labels, 150 RILs were re-genotyped in the F₇ generation along with a
615 further replicate of each founder.

616

617 Genotype calling was performed using the Affymetrix Power Tools (v1.19) and SNPolisher R 618 packages, following the recommended Axiom analysis pipeline. All samples except two-way 619 crosses were given the standard inbreeding penalty, 4, which penalises calling heterozygous 620 genotypes. Four samples failed the 'dish quality control' threshold (0.82) and a further 28 621 samples with call rates were below 97% were excluded. Marker classifications were 622 performed using "ps-classification", and ps-classification-Supplementary" functions with options --species-type polyploid --hom-ro false. All calls were adjusted using the standard 623 624 0.025 confidence threshold using the Ps CallAdjust function.

625

Samples were compared to one another using the 14,935 markers classified as 'PolyHighResolution' only. Overall, 46 RIL pairs were found to be >92% similar (mean 98.5% genotype similarity), where all other comparisons between MAGIC lines were, at most, 84% similar (mean 67.8%). These apparently duplicated genotypes could indicate genotyping, labelling, or propagation errors so only one RIL from each pair was used for sequencing (550 RILs). To ensure pedigree accuracy, we chose the RIL in each pair that was genotypically most similar to other RILs derived from the same 16-way cross (i.e. in the same family).

633

634 Sequencing Data

For whole genome sequencing, DNA was extracted from 550 RILs at the F_7 generation. DNA for RILs that failed quality control were extracted again at the F_8 generation (n=50). Sequencing and library preparation was performed at Novogene, where libraries were

generated from 1.0µg DNA per sample using the NEBNext DNA Library Prep Kit. Sequencing
was performed on a NovaSeq 6000 instrument (Illumina) to get at least 6Gb of raw sequence
data (2x150bp paired end reads) per sample. One founder (Holdfast) was sequenced to 15.8x
coverage using the same method.

642

The other founders were sequenced after capture using two recently designed probe sets 643 targeting promoter and genic regions, respectively[21]. Capture was performed at the 644 645 Earlham Institute following the SeqCap EZ Library SR v5.1 protocol (Roche NimbleGen Inc., Madison, WI, USA) with 1µg of genomic DNA sheared to 300bp[21]. Four captures were 646 647 performed using 8 samples per set (2x promoter captures and 2x genic captures). Samples for 648 the founder Stetson were included on all four capture experiments so roughly double the 649 sequence data was obtained for this variety (Supplementary Table 1). Sequencing with 650 2x150bp reads was performed at the Earlham Institute on a NovaSeq 6000 instrument 651 (Illumina) with 16 promoter capture libraries on one lane and 16 genic capture libraries on 652 another lane.

653

654 Variant Calls and Imputation

All reads were aligned to the bread wheat reference genome from cv. Chinese Spring (RefSeq v1.0)[27] using bwa-mem (version 0.7.12)[64] and sorted using samtools (version 1.3.1)[65], which was also used to calculate coverage. For compatibility with the bam file format, we split each chromosome in the reference genome at the halfway point before alignment. We called variants from the founder sequences within the high confidence gene, promoter and 5' UTR regions targeted by the capture probes[21] using GATK (version 4.0.8.0)[66] HaplotypeCaller and GenotypeGVCFs (options --interval-padding 100 --minimum-mapping-quality 30). We used vcftools (version 0.1.15) to include only bi-allelic single nucleotide polymorphisms
(SNPs) with average coverage depth between 5 and 60 (all per sample coverages between 2
and 120) and no missing calls. We also filtered with bcftools (version 1.2)[67] using standard
quality control options --exclude 'QD<2 || FS>60.0 || MQRankSum<-12.5 ||
ReadPosRankSum<-8.0 || SOR>3.0 || MQ<40'. This left 1.78M SNPs, of which we only use
the 1.13M sites with no heterozygous calls (--genotype ^het option) for our main analyses.

668

669 We first called genotypes in the RILs at these 1.13M SNP sites directly using GATK 670 HaplotypeCaller in GENOTYPE-GIVEN-ALLELES mode, using the same options as above. We assessed the concordance between array genotypes and these direct calls (AD) at overlapping 671 sites (see below). For 10 RILs, the directly called sequencing variants most closely matched 672 673 genotyping array data for a different line than expected. These were excluded because the 674 source of the discrepancy (sequence data or array data) cannot be established. The 675 concordance between our genotyping array data and direct calls (AD) was below 95% for a 676 further 36 RILs, which were excluded (mean AD 84.7% for removed lines), leaving 504 RILs. 677 We estimated heterozygosity in these 504 RILs using only genotypes called from at least four 678 reads. Of 2.6M such genotype calls, only 0.67% were called as heterozygotes.

679

We imputed genotypes at the 1.13M SNP sites using the alignments and STITCH software (version 1.5.7)[26]. Because alignments were to a reference genome with chromosomes split in half, we first ran STITCH with the generateInputOnly option, and then joined the input files for each chromosome half before imputation. For all runs, we used the parameters nGen=3, minRate=0.001, bqFilter=30, method='diploid-inbred' and then filtered all sites with an info score below 0.4, minor allele frequency below 2.5%, or missingness above 10%. For our main

686 analysis, we used the genotype calls in the founders as a reference panel and outputted the 687 estimated ancestry dosages of each founder at each position in each RIL using the 688 outputHaplotypeProbabilities and output haplotype dosages options. When using the 689 founders as a reference panel, we removed options that estimate and update the haplotypes 690 population (shuffleHaplotypeIterations, reference shuffleHaplotypeIterations, in the 691 refillIterations). To test accuracy when reference panels aren't available, we re-ran 692 imputation without the founder haplotypes, using 40 iterations to estimate the haplotype 693 space and recombination mosaics. We also used the downsampleFraction option to randomly 694 sample a fraction of alignments with/without using the founder reference panel. Finally, we tested imputation accuracy (without a reference panel), when fewer than sixteen haplotypes 695 696 were assumed to segregate in the population by varying the K parameter (Supplementary 697 Figure 2).

698

699 Genotype Comparisons

700 For comparison against the sequencing dataset, we used all genotyping array markers. 701 Replicates of founders and MAGIC RILs (where available) were used to make a consensus call 702 where the most common genotype across replicates was taken as the consensus and only 703 retained when more than 50% of the non-missing calls were in agreement. In addition, 704 markers where one homozygous genotype was missing from all RILs were converted such that 705 all heterozygous calls were assumed to be in the missing homozygous class. The failure to 706 detect a homozygous class is likely to be a result of polyploidy, which can reduce 707 differentiation between the three genotype classes and make them hard to distinguish. 708 Finally, to get plausible physical positions for the genotyped markers, BLASTn v2.2.30[25] was 709 used to compare the 75bp probe sequences (cerealsdb.uk.net)[12] against the reference genome[27]. When matching the SNP array data with the sequenced SNPs, array sites were
excluded if there had missing or heterozygous founder calls or if the genotypes and targeted
SNP alleles did not match the founder sequence data. We found 5,877 sites that overlapped
between the genotyping array data and the sequencing data (Supplementary Table 2).

714

715 To compare against global wheat diversity, we called founder genotypes at 113,457 716 genotyping array sites that were polymorphic among 4,506 diverse global wheat 717 accessions[13]. We called genotypes from alignments with mapping quality scores of at least 718 30 using GATK HaplotypeCaller in EMIT ALL SITES mode with the -emit-ref-confidence 719 BP RESOLUTION option, providing a bed file of the 113,139 genotyping array sites[13]. We 720 only considered sites where genotypes could be called in all 16 founders (n=56,063). We used 721 genotyping array calls for cv. Chinese Spring to determine reference/non-reference alleles on 722 the genotyping array, ignoring sites called as heterozygous (n=109) or missing (n=306) in 723 Chinese Spring. Seven of the MAGIC founders were also present in the global genotype set 724 (Brigadier, Copain, Maris Fundin, Soissons, Spark, Steadfast, Stetson)⁷. The average 725 concordance of the global genotype calls and our sequencing calls for these founders was 726 94.3% (sd 0.63%). We excluded 5,491 (9.8%) sites that had mismatches across these founders, 727 many of which are likely to reflect differences in the underlying genetic variation picked up 728 by the different genotyping technologies. Two other founder variety names were in the 729 genotyping array dataset⁷ (Banco and Holdfast) but the genotyping calls did not match 730 (concordances 74.2% and 71.4%, respectively), which may reflect differences in the seed 731 stock used.

732

733 Founder Haplotype Diversity

First, we used the SNPs called within each promoter-gene pair to estimate haplotypic diversity 734 735 of the founders. We calculated absolute (Manhattan) pairwise genetic distances between 736 founders at each site and then used complete linkage clustering to define haplotypic groups using dist and hclust functions implemented in R statistical software (version 3.6.0)[68]. This 737 738 was repeated using different similarity thresholds to define haplotypes. Second, we 739 determined haplotype breakpoints using a dynamic programming algorithm. For each 740 pairwise founder combination, our algorithm calculates a mosaic of genotypic 741 similarity/dissimilarity akin to the Viterbi path from a hidden Markov model. Genotype 742 matches and mismatches are allocated a score (1 by default). To prevent excessive switching 743 between states, there is also a 'transition penalty' for inferring a change between matching and mismatching states. Based on their pairwise matching/mismatching states, we then infer 744 745 the total number of haplotypes inferred at each site. We repeat this procedure with different 746 parameter choices (Supplementary Figure 2).

747

748 Genetic Mapping and Heritability

For mapping, we used the full set of 1,065,185 high-quality SNP sites called in 504 RILs after imputation and quality control filters. We also selected a subset of 55,067 SNPs such that every other SNP was tagged at R²>0.99 by a member of the subset using PLINK (version 1.90) with option --indep-pairwise 500 10 0.99. These tagging SNPs were used to calculate the genetic relationship matrix K = GG'/p. The phenotypic variance-covariance matrix for a given vector y of standardised phenotype values was modelled as $V = K\sigma_q^2 + I\sigma_e^2$ where 755 σ_g^2, σ_e^2 are the additive genetic and environmental variance components, estimated by 756 maximum-likelihood[69]. The heritability of a trait was defined as $h^2 = \sigma_g^2/(\sigma_g^2 + \sigma_e^2)$. The 757 matrix square root of the variance matrix was calculated by eigendecomposition of V as $A^2 =$ 758 V, and the mixed model transformation of the data performed i.e. $y \to A^{-1}y, G \to$ 759 $A^{-1}G, V \to I$ to remove the inflationary effects of unequal relatedness on genetic 760 associations before association mapping.

761

762 We performed association tests at the level of both SNPs and founder haplotypes using R 763 statistical software (version 3.6.0)[68]. Initially, we tested the null hypothesis of no 764 association at each SNP site in the tagging set (~55k sites). We then determined genome-wide 765 thresholds for statistical significance using 1,000 permutations on the transformed phenotypes. If any association exceeded the 0.05 threshold (smaller p value than found across 766 767 at least 950 phenotypic permutations), then we repeated the association test at all of the 768 ~1.1M SNPs on the chromosome with the strongest association signal (lowest p value). 769 Mapping intervals were defined to include SNPs surrounding the peak SNP, with log₁₀(p) 770 values within d units of x using $d = \max\{2, 0.1x\}$ where x is the peak $\log_{10}(p)$ value. The 771 interval for haplotype-based tests includes the range of sites that have log₁₀(p) values within 772 d units of x. SNP-based intervals were calculated using the same measure but then extended by the minimum of 5Mb or the distance to the next SNP in either direction that the same 773 'strain distribution pattern'[54] as any highly-associated SNPs (SNPs with log₁₀(p) values 774 within d units of x). The 'strain distribution pattern' is the pattern of major/minor alleles 775 776 across founders. This procedure is designed to capture the uncertainty in the positioning of 777 relevant recombination events either side of the QTL peak. We fitted QTLs in a stepwise 778 manor by fitting the phenotype against the most strongly associated SNP (or haplotype

dosage) whenever genomewide significant QTLs were detected. The above association test procedure was then repeated using the phenotype residuals after fitting all previously identified QTLs. This allows closely-linked QTLs to be detected when they have different patterns of causal variants among RILs. Where QTL associations were found for different genotypes, they were judged to be at the same locus if they had overlapping mapping intervals and at least one matching strain distribution pattern at highly-associated SNP sites.

785

786 Genomic Prediction

787 To evaluate the accuracy of trait prediction within our magic population and estimate the 788 extent of polygenic variation beyond genomewide significant QTLs, we conducted genomic 789 prediction across all phenotypes using three shrinkage-based methods: ridge regression (RR), 790 Elastic Nets (EN) and least absolute shrinkage and selection operator (LASSO). We note that with appropriate choice of ridge parameter $\lambda = \sigma_e^2 / \sigma_a^2$, RR is equivalent to a mixed model in 791 792 the sense that the RR estimated SNP effects are identical to the mixed-model Best Linear 793 Unbiased Predictors (BLUPs)[70,71]. This explains the linear correlation between estimates of 794 heritability and RR prediction accuracy (Figure 5c). For each method, we conducted 50 rounds of cross validation by randomly sampling 90% of the RILs (n=454) as a training set in each 795 796 round to train the model, which was then used to predict the remaining 10% of RILs (n=50) -797 the test set. For the three methods, the model equation can be written generally as $y = \mu + \mu$ 798 $\beta G + \varepsilon$, where y is the estimated trait value, μ is the model intercept, β is the vector of SNP 799 effects, G is the genotype dosage matrix, and ε is the residual error.

800

801 The genomic prediction models were trained using the R package glmnet[72], which 802 estimates an optimal lambda shrinkage value for all three genomic prediction methods based 803 on the training set. We then predicted phenotypes in the test set by multiplying all SNP 804 coefficients estimates by their corresponding genotypes in the test set (and adding the 805 intercept term). We report the training and test set prediction accuracy as the mean Pearson 806 correlation coefficient of the predicted trait values and the actual phenotype values over 50 807 rounds of cross validation.

808

We used these genomic prediction models to simulate the potential for phenotypic change. 809 810 First, we permuted the population founder haplotypes identities 40 times across 504 RILs and 811 then projected the permuted founder genotypes onto the new lines. This creates new genetic combinations while retaining the genetic map and linkage found in the real population. We 812 813 then used the three models trained as described above to predict phenotypes for the 814 simulated MAGIC RILs. We further calculated the theoretical maximum and minimum 815 phenotype values that are possible given the genomic prediction models and the variants 816 segregating in the population. To estimate the maximum and minimum potentially achievable 817 phenotype values, we trained new genomic prediction models using the full data set of 504 818 RILs for all phenotypes. We then calculated the maximum/minimum predicted phenotypes 819 by summing the estimated effects for all positive/negative SNP coefficients.

820

821 Gene Deletion Analysis

We examined the power of gene-level coverage variation among founders to explain phenotypic variation. In each founder f and at each gene feature g, we computed a deletion index D_{gf} based on the number of reads aligning to the associated capture sequences, normalised by the overall coverage for that founder. The gene deletion score (GDS) for each MAGIC RIL i and feature j was computed as $S_{ij} = \sum_{f} H_{ijf} D_{if}$, where H_{ijf} is the haplotype dosage for founder *f* in RIL *i* at gene *j*, as computed by STITCH. For each phenotype a mixedmodel GWAS was performed, using the GDS in place of SNP dosages and with a genetic relationship matrix computed from the GDS (Supplementary Table 8). We also repeated the genomic prediction analysis described above by replacing the SNP genotype dosage matrix with the GDS matrix (Supplementary Figure 5).

832

833 Introgressions

834 The presence of introgressions were determined using summary statistics (coverage, non-835 reference allele frequency in founders and RILs) calculated in 10Mb windows moved in 5Mb 836 steps. Within introgressions, carriers have a high proportion of non-reference alleles due to the alignment of inter-specific genetic material to the bread wheat reference genome. The 837 838 introgression extent was determined as the extent of 10Mb windows where all introgression 839 carriers had a higher proportion of non-reference alleles than all non-carriers. Within these regions, we then checked the relative coverage of carriers and the extent to which the alleles 840 841 of carriers are over- or under-represented among the RILs. This evidence is summarised in 842 Supplementary Table 3. For example, the founder Maris Fundin carries a large introgression (640Mb) from *Triticum timopheevi* on chromosome 2B that inflates the total number of SNPs 843 844 called on chromosome 2B, relative to the other chromosomes (Supplementary Figure 1), this 845 introgression is substantially over-represented among RILs, as expected[34].

846

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858 Author Contributions

Project conception and funding: IM, PH (BB/E007201/1) and RM, JC (BB/M011666/1,
BB/M011585/1). Project management: JC, RM, with input from MS, NF, PH, KG and AB.
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phenotyping, trials analysis and DNA extraction: NF. Small plant phenotyping: FC. Promotergene capture pull downs and library preparation: TB. Genetic analyses: MS with support from
RM. Genomic prediction: OL. Manuscript writing: MS, OL, NF, RM, with inputs from KG, IM
and JC. All authors edited and approved the manuscript.

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867 Availability of data and materials

- 868 The sequence datasets supporting the conclusions of this article are available in the ENA
- 869 repository, under project number PRJEB39021 (temporarily embargoed),
- 870 <u>https://ebi.ac.uk/ena</u>.
- 871

- 872 The Genotyping array genotypes for founders and MAGIC RILs are available from
- 873 <u>http://mtweb.cs.ucl.ac.uk/mus/www/MAGICdiverse/index.html</u> in text tabular format.
- 874
- 875 The Imputed SNP genotypes and founder haplotype dosages are available from
- 876 mtweb.cs.ucl.ac.uk/mus/www/MAGICdiverse/MAGIC diverse FILES/MAGIC PLINK.tar.gz,
- 877 and
- 878 mtweb.cs.ucl.ac.uk/mus/www/MAGICdiverse/MAGIC diverse FILES/MAGIC HAPLOHAPLO.
- 879 <u>tar.gz</u> (temporary links).
- 880
- 881 The remaining datasets supporting the conclusions of this article are included within the
- 882 article and its additional files.
- 883
- 884 Custom analysis scripts (mixed model and haplotype dynamic programming algorithm) are
- available from github.com/michaelfscott/DIVERSE_MAGIC_WHEAT .

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