A heuristic method for fast and accurate phasing and imputation of single-nucleotide polymorphism data in bi-parental plant populations

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Abstract

Key message New fast and accurate method for phasing and imputation of SNP chip genotypes within diploid bi-parental plant populations.

Abstract This paper presents a new heuristic method for phasing and imputation of genomic data in diploid plant species. Our method, called AlphaPlantImpute, explicitly leverages features of plant breeding programmes to maximise the accuracy of imputation. The features are a small number of parents, which can be inbred and usually have high-density genomic data, and few recombinations separating parents and focal individuals genotyped at low density (i.e. descendants that are the imputation targets). AlphaPlantImpute works roughly in three steps. First, it identifies informative low-density genotype markers in parents. Second, it tracks the inheritance of parental alleles and haplotypes to focal individuals at informative markers. Finally, it uses this low-density information as anchor points to impute focal individuals to high density. We tested the imputation accuracy of AlphaPlantImpute in simulated bi-parental populations across different scenarios. We also compared its accuracy to existing software called PlantImpute. In general, AlphaPlantImpute had better or equal imputation accuracy as PlantImpute. The computational time and memory requirements of AlphaPlantImpute were tiny compared to PlantImpute. For example, accuracy of imputation was 0.96 for a scenario where both parents were inbred and genotyped at 25,000 markers per chromosome and a focal F2 individual was genotyped with 50 markers per chromosome. The maximum memory requirement for this scenario was 0.08 GB and took 37 s to complete.

Abbreviations

LD Low density
HD High density
SNP Single-nucleotide polymorphism
cM Centimorgan

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Introduction

This paper presents a new heuristic method for phasing and imputation of single-nucleotide polymorphism (SNP) array data in diploid plant species. High-density SNP array data in plant breeding populations is increasingly valuable for genomic selection and for identifying regions of the genome that underlie traits of interest in genome-wide association studies. The accuracy of genomic selection and power of association studies increases with the number of individuals and with the density of SNP markers. However, the cost of genotyping many individuals at high-density is high. This high cost is a barrier to the adoption of genomic selection in plant breeding programmes where the number of selection candidates in each cycle can be very large. An effective strategy to overcome this cost barrier is to genotype a proportion of the population at high-density, phase their genotypes, and use this data for imputation of large numbers of individuals genotyped at low-density (Jacobson et al. 2014, 2015; Gorjanc et al. 2017a, b). This strategy has been widely adopted in livestock and human populations, partly because
genotype imputation tools that work well in these populations are widely available (Kong et al. 2008; Howie et al. 2009; Druet and Georges 2010; Li et al. 2010; Sargolzaei et al. 2011; Hickey et al. 2011; Cleveland and Hickey 2013; Hickey and Krisan 2013; Van Raden et al. 2015; O’Connell et al. 2016; Loh et al. 2016; Antolín et al. 2017).

Bi-parental populations that are widely used in plant breeding have four features that make them ideal for imputation. First, they are derived from only two parents. High-density genotyping of the two parents and low-density genotyping of focal individuals (i.e. descendants that are the imputation targets) is an effective low-cost strategy in these populations. Second, the number of meiosis separating parents and focal individuals is small. This means that parental haplotypes remain largely intact in focal individuals, which simplifies imputation. Third, they have well-known crossing structures that could be informative for imputation, although the process of selfing or the creation of doubled haploids can add complications that are not present in human and livestock settings. However, these “complications” can in certain situations empower imputation. Finally, parents that contribute to a bi-parental population are usually inbred. This means that they are homozygous at many loci and the majority of their genome is phased de facto.

A recent simulation study demonstrated that achieving high imputation accuracies could empower genomic selection in bi-parental populations (Gorjanc et al. 2017a, b). The high imputation accuracies with SNP array data were achieved using the PlantImpute software (Nettelblad et al. 2009; Hickey et al. 2015). The main drawback of PlantImpute is that it has large computational requirements in terms of time and memory. This makes it impractical for routine use in breeding programmes. Existing software for imputation in livestock or human populations do not have large computational requirements. However, software for imputation in livestock or human populations are not designed to leverage features of plant breeding programmes, and in some cases, cannot work where selfing and bi-sexuality is common. To our knowledge, existing imputation software for plant breeding programmes (e.g. Swarts et al. 2014) are not explicitly designed for imputation of SNP array genotypes in bi-parental populations.

This paper presents a new heuristic method, called AlphaPlantImpute, for phasing and imputation of SNP array data in diploid plant species. AlphaPlantImpute works roughly in three steps. First, it identifies markers fully or partially informative for parent-of-origin. Second, it tracks the inheritance of parental alleles and haplotypes to focal individuals at informative markers. Finally, it uses this low-density information as anchor points to impute focal individuals to high-density.

We tested the accuracy of AlphaPlantImpute in simulated bi-parental populations across different scenarios. These scenarios varied in the levels of inbreeding in the parents, the number of selfing events separating parents and focal individuals, the chromosome size (i.e. recombination rate) and the number of markers on the low-density array. We calculated the accuracy of imputation within each scenario as the correlation between the true and imputed genotypes. In general, AlphaPlantImpute gave excellent accuracy of imputation and typically outperformed or performed equally as well as PlantImpute for the accuracy of imputation. The computational time and memory requirements of AlphaPlantImpute were always tiny compared to that of PlantImpute.

Materials and methods

Definitions

A focal individual is an individual that is to be imputed. A fully informative marker is one where the two parents have opposing homozygous genotypes, i.e. genotypes 0 and 2 (note that the method is agnostic of which allele is the reference allele). A partially informative marker is where one parent is homozygous and the other is heterozygous. Markers where parents are fixed for the same allele or where both parents are heterozygous are uninformative. AlphaPlantImpute considers only bi-allelic loci. Genotypes are coded as 0, 1, and 2, and alleles are coded as 0 and 1. A diplotype refers to the genotypes of a pair of SNPs and a haplotype refers to the alleles of a pair of SNPs. The high-density (HD) array is the array at which parents have genotypes and is the target array for imputation. In our test datasets, the HD array consisted of 25,000 SNP markers. The low-density (LD) array is the array at which focal individuals have genotypes. We tested eight LD arrays (see below), all of which were nested subsets of the HD array.

Description of the method

We present a new heuristic method, called AlphaPlantImpute, for phasing and imputation of SNP array data in diploid plant species. In detail, our method has five steps: (1) identify markers that are informative for parent-of-origin of alleles in focal individuals; (2) infer the most likely linked alleles at two markers; (3) phase and assign parent-of-origin for focal individual’s alleles; (4) impute focal individual to high-density using low-density anchors captured in step 3; and (5) impute markers in recombined regions. Impute markers adjacent to recombination locations. Step 1 is the only step applied to groups of focal individuals together. Steps 2, 3, 4 and 5 are applied for each focal individual separately. A description of the definitions used and of each step is given below, and a detailed schematic is given in Fig. 1.
Method steps

Step 1: Identify informative low-density markers in parents

In the first step, we determine which low-density markers are fully or partially informative in parents, which is used in the following steps to infer parent-of-origin of phased alleles in focal individuals. For example, in Fig. 1 eight of the ten markers on the HD array genotyped in the parents are fully informative and two (markers 2 and 9) are uninformative. Of the ten HD markers, five (markers 1, 3, 5, 7, 9) are also on the LD array, which was used to genotype focal individuals. Of these five LD markers, four are informative and one (marker 9) is uninformative.

Step 2: Infer the most likely linked alleles at two markers

In the second step, we infer the most likely linked alleles for all possible marker pairs for all informative markers. This information on linkage relationships between markers is used in later steps to phase and assign parent-of-origin to heterozygous markers of focal individuals. If parent haplotypes are inherited directly without recombination, the most likely linked alleles at two markers recover the parent haplotypes. When this is not the case, the most likely linked alleles at two markers indicate a potential recombination hotspot or marker map error for the population. For each pair of informative markers, we perform three steps.

(2a) First, identify focal individuals that are homozygous at the first and the second marker.

(2b) Second, count the number of times focal individuals have genotype:

- 0 for the first and 0 for the second marker (diplotype 0-0),
- 0 for the first and 2 for the second marker (diplotype 0-2),
- 2 for the first and 0 for the second marker (diplotype 2-0),
- 2 for the first and 2 for the second marker (diplotype 2-2).

(2c) Third, compare the count of diplotype 0-0 to diplotype 0-2 and of diplotype 2-2 to diplotype 2-0. If the count of 0-0 is higher than 0-2 and 2-2 is higher than 2-0, then the 0 allele at the first marker is more frequently linked to the 0 allele at the second marker (haplotype 0-0) and the 1 allele at the first marker is more frequently linked to the 1 allele at the second marker (haplotype 1-1). If the count of diplotype 0-2 is higher than diplotype 0-0 and diplotype 2-0 is higher than diplotype 2-2, then the 0 allele at the first marker is more frequently linked to the 1 allele at the second marker (haplotype 0-1) and the 1 allele at the first marker is more frequently linked to the 0 allele at the second marker (haplotype 1-0). For example, in Fig. 1, 2-2 and 0-0 are the two...
most frequent diplotypes at markers 1 and 3, which suggests the most likely haplotypes are 1-1 and 0-0.

**Step 3: Phase and assign parent-of-origin for focal individual’s alleles**

In the third step, we phase alleles in focal individuals and assign their parent-of-origin. We perform this first for the homozygous markers and then for the heterozygous markers.

**3a) Phase homozygous markers**

We phase alleles at homozygous markers as the 0 allele for both haplotypes when the genotype is 0 and as the 1 allele when the genotype is 2. For example, in Fig. 1, the focal individual ID_Y has genotype 2 for marker 7 and we phase it as the 1 allele for both haplotypes.

**3b) Assign parent-of-origin to alleles at homozygous markers**

We assign parent-of-origin for phased alleles in the step 3a based on the informative markers in the step 1. For example, in Fig. 1, marker 7 is informative. At this marker, the Parent_A has the 0 allele, while the Parent_B has the 1 allele. Focal individual ID_Y has genotype 2 for marker 7 and we phase it as the 1 allele for both haplotypes.

**3c) Phase heterozygous marker**

We phase alleles at heterozygous markers iteratively based on the most likely linked alleles in the step 2. Specifically, we perform four steps. We start at the first heterozygous marker. For example, in Fig. 1, the first marker for which the focal individual ID_Y is heterozygous is marker 1. (3c1) First, phase the first heterozygous marker randomly as the 1 allele for the first haplotype and the 0 allele for the second haplotype.

(3c2) Second, phase the second heterozygous marker based on the most likely linked alleles in the step 2. For example, in Fig. 1, the second heterozygous marker is marker 3. Information from the most likely linked alleles suggest that the 0 (1) allele at marker 1 is linked to the 0 (1) allele at marker 3. Using this information, we phase marker 3 alleles of ID_Y as the 1 allele for the first haplotype and the 0 allele for the second haplotype. We continue moving from left-to-right until the last heterozygous marker is phased.

(3c3) Third, we repeat steps 3c1 and 3c2, but this time starting from the last heterozygous marker and progressing to the first heterozygous marker.

(3c4) Finally, we derive a consensus between the haplotypes derived from moving left-to-right and right-to-left along the chromosome. If they disagree, set the consensus haplotypes to missing. If only one is filled, set the consensus haplotype to the filled information.

**3d) Assign parent-of-origin to alleles at heterozygous marker**

We assign parent-of-origin for phased alleles in the step 3c based on the informative markers in the step 1. For example, in Fig. 1, focal individual ID_Y is heterozygous at marker 1. At this marker, the 1 allele on ID_Y’s first haplotype is inherited from Parent_A and the 0 allele on ID_Y’s second haplotype is inherited from Parent_B. If the marker is partially informative, we assign both the parent-of-origin and the haplotype-of-origin (i.e. first or second haplotype of the parent that is heterozygous for that marker).

**Step 4: Impute focal individual to high-density using anchors from the step 3**

**4a) Fill uninformative homozygous markers**

For uninformative homozygous markers at HD that are not genotyped in the focal individual at LD, we phase and impute the focal individual with the parental information. For example, in Fig. 1, both parents have genotype 0 for marker 2, so focal individual ID_Y is imputed as genotype 0.

**4b) Assign parent-of-origin to HD marker alleles**

For markers on the HD array, assign parent-of-origin to marker alleles based on the parent-of-origin assignment of the two nearest marker alleles on the LD array. For example, in Fig. 1, marker 6 is not genotyped on the LD array, but the two neighbouring markers 5 and 7 are genotyped on the LD array. We have assigned the second haplotype of focal individual ID_Y to Parent_B for both markers 5 and 7. We therefore also assign marker 6 to Parent_B for the second haplotype. We have assigned the first haplotype of focal individual ID_Y to Parent_A for marker 5 and to Parent_B for marker 7. We conclude that there was a potential recombination around marker 6 at the first haplotype and we do not assign parent-of-origin for this allele.

**4c) Phase and impute HD markers using parent-of-origin assignment from step 4b**

For HD markers with assigned parent-of-origin in step 4b, we phase the allele inherited from that parent for the haplotype of the focal individual. If we have phased both alleles at a marker, we impute the genotype as the sum of the
two alleles on the two haplotypes of the focal individual. If parent-of-origin has not been assigned for one or both alleles of the focal individual, we leave the genotype as missing.

**Step 5: Impute markers in recombined regions**

We phase and impute missing HD markers in potentially recombined regions in one of two ways. We either (1) impute expected genotype dosage as the average of the alleles of the two parents; or (2) phase and impute using information from a genetic or physical map. For (2), we first identify the two closest neighbouring markers that were informative and phased, second use the distance between these two markers as a weight to phase the missing alleles as the weighted average of the alleles of the two parent haplotypes, and third impute expected genotype dosage as in (1).

**Implementation**

We have implemented the method in a programme called AlphaPlantImpute, which is controlled by a specification file that contains some user specified thresholds and the addresses of input files. The required input data are membership of individuals to the bi-parental populations, HD genotypes for parents, and LD genotypes of focal individuals. The output data are imputed genotypes, phased haplotypes, inferred parent-of-origin for focal individual haplotypes, and information on whether a marker is informative. AlphaPlantImpute implements some data editing checks, which are described in the user manual.

**Examples of implementation: description of datasets**

To test the imputation accuracy of AlphaPlantImpute, testing datasets of a subset of the scenarios described in Hickey et al. (2015) were simulated. This enabled the comparison of AlphaPlantImpute with PlantImpute without re-running PlantImpute with its large computational cost. Although the simulation design is largely a replication of that in Hickey et al. (2015), a brief description of the general structure and simulation method of the different scenarios tested is given below for completeness.

**Simulation of genomic data**

Sequence data for 100 base haplotypes for a single chromosome were simulated using the Markovian Coalescent Simulator (Chen et al. 2009) and AlphaSim (Faux et al. 2016). The base haplotypes were $10^8$ base pairs in length, with a per site mutation rate of $1.0 \times 10^{-8}$ and a per site recombination rate that varied across scenarios. The different recombination rates simulated were $0.25 \times 10^{-8}$, $0.5 \times 10^{-8}$, $1.0 \times 10^{-8}$, $1.5 \times 10^{-8}$, $2.0 \times 10^{-8}$, $3.0 \times 10^{-8}$, and $4.0 \times 10^{-8}$, resulting in chromosome sizes of 25, 50, 100, 150, 200, 300, and 400 cM, respectively. The effective population size ($N_e$) was set at specific points during the simulation to mimic changes in $N_e$ in a crop such as maize (*Zea mays* L.). These set points were: 100 in the base generation, 1000 at 100 generations ago, and 10,000 at 2000 generations ago, with linear changes in between. The resulting whole-chromosome haplotypes had approximately 80,000 segregating sites in total.

**Simulation of a pedigree**

A pedigree of 11,266 individuals was constructed. The pedigree was initiated from six outbred founders (A, B, C, D, E, F). These six founders were crossed to generate the founder bi-parental populations (A×B, C×D, E×F). These founder bi-parental populations were selfed to $F_1$, $F_2$, $F_3$, $F_{10}$, or $F_{20}$, resulting in different levels of inbreeding in the parents. To properly propagate the residual heterozygosity in these parents, they were crossed to generate 100 pairs of $F_1$ individuals. $F_1$ individuals were selfed to generate 100 $F_2$ individuals. $F_2$ individuals were selfed to generate 100 $F_3$ individuals, and selfing continued through to $F_{10}$. The focal individuals (i.e. descendants that were the imputation targets) were $F_2$, $F_4$, $F_6$, or $F_{10}$ descendants.

In the base generation, the six founders had their chromosomes sampled from the 100 base haplotypes. In subsequent generations, the chromosomes of each individual were sampled from parental chromosomes with recombination. The recombination rate varied depending on the scenario resulting in chromosome sizes of 25, 50, 100, 150, 200, 300, and 400 cM. Recombinations occurred with a 1% probability per cM and were uniformly distributed along the chromosome.

**Simulated SNP marker arrays**

A single HD array of 25,000 SNP markers for the single chromosome was simulated. To test the effect of the number of markers on the LD array, eight LD arrays of 3, 5, 10, 20, 50, 100, 200, and 400 markers for the single chromosome were simulated. Arrays were constructed by aiming to select a set of markers that segregated in the parents and that were evenly distributed across the chromosome. All LD arrays were nested within each other and within the HD array.

**Scenarios**

The imputation accuracy of AlphaPlantImpute and PlantImpute were compared in four different scenarios (scenario 1, 2, 3, and 4). Scenarios 1, 2, and 3 were the same as scenarios 2, 4, and 5 in Hickey et al. (2015). A description of all four scenarios is provided below. In all scenarios, focal individuals genotyped at LD were imputed to the single HD array of
25,000 SNP markers. Ten replications of each scenario were performed, and the average of each replication is reported in the results.

Scenario 1: The effect of the number of selfing events separating parents and focal individuals. Parents were almost fully inbred ($F_{20}$) and chromosomes were 100 cM in length. The accuracy of imputation was assessed for $F_2$, $F_4$, $F_6$, and $F_{10}$ focal individuals.

Scenario 2: The effect of the level of inbreeding in parents. Parents were $F_1$, $F_2$, $F_4$, $F_{10}$, or $F_{20}$ and chromosomes were 100 cM in length. The accuracy of imputation was assessed for $F_2$ focal individuals.

Scenario 3: The effect of chromosome size. Parents were fully inbred ($F_{20}$) and the accuracy of imputation was assessed for $F_2$ focal individuals. Chromosomes were 25, 50, 100, 150, 200, 300, or 400 cM in size.

Scenario 4: The effect of number of focal individuals in the bi-parental population. Parents were fully inbred ($F_{20}$) and the accuracy of imputation was assessed for $F_2$ focal individuals. Subsets of focal individuals were randomly selected from the 100 focal individuals to generate bi-parental population sizes of 1, 5, 10, 25, and 50 focal individuals.

Analysis

Imputation was performed within each bi-parental population. Parents were assumed genotyped at HD and focal individuals were assumed genotyped at LD. The imputation accuracy was calculated for each focal individual as the correlation between the true and imputed genotypes. The precision in imputation accuracy was calculated as the log of the inverse of the variance in imputation accuracy within each bi-parental population.

Results

For each scenario, we first present the imputation accuracy of AlphaPlantImpute and then compare it to PlantImpute (Nettelblad et al. 2009; Hickey et al. 2015).

Effect of the number of markers on the low-density array

Increasing the number of LD markers increases the imputation accuracy of AlphaPlantImpute. Figure 2 plots the number of LD markers against the genotype imputation accuracy using AlphaPlantImpute for $F_2$ focal individuals of a bi-parental cross where the parents are $F_{20}$ inbred individuals.

Scenario 1: Effect of the number of selfing events separating parents and focal individuals

Increasing the number of selfing events separating parents and focal individuals slightly decreases the imputation accuracy of AlphaPlantImpute. Figure 3a plots the accuracy of imputation in $F_2$, $F_4$, $F_6$, and $F_{10}$ focal individuals of a bi-parental population where the parents were $F_{20}$. Figure 3a shows that with 3 LD markers, the average imputation accuracy decreased from 0.85 for $F_2$ focal individuals to 0.77 for $F_{10}$ focal individuals. Increasing the number of LD markers beyond 10 markers mitigates the decrease in the average imputation accuracy between $F_2$ focal individuals and $F_{10}$ focal individuals. Figure 3a shows that with 20 LD markers, the average imputation accuracy decreased from 0.96 for $F_2$ focal individuals to 0.95 for $F_{10}$ focal individuals.

Regardless of the number of selfing events separating parents and focal individuals, the accuracy of imputation for AlphaPlantImpute was higher than for PlantImpute when the number of LD markers was low. Figure 3b plots the average imputation accuracy of AlphaPlantImpute on the y-axis and for PlantImpute on the x-axis. The colours represent the different number of LD markers and the shapes represent the number of selfing events separating the parents and the focal individuals. The red diagonal line indicates when the imputation accuracy of the two methods is equal. Points above the line indicate when the accuracy of imputation was higher for AlphaPlantImpute than for PlantImpute and vice versa. Figure 3b shows that with 3 LD markers, the average accuracy of imputation was 0.85 for AlphaPlantImpute and 0.76 for PlantImpute for $F_2$ focal individuals and was 0.77.

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for AlphaPlantImpute and 0.70 for PlantImpute for $F_{10}$ focal individuals.

For all numbers of selfing events separating parents and focal individuals, increasing the number of LD markers reduced and in some cases reversed the advantage of AlphaPlantImpute over PlantImpute. This was most obvious for $F_{10}$ focal individuals for medium number of LD markers where the imputation accuracy with PlantImpute was slightly higher than with AlphaPlantImpute. Figure 3b shows that with 10 LD markers, the average imputation accuracy was 0.93 for AlphaPlantImpute and 0.94 for PlantImpute for $F_2$ focal individuals and was 0.90 for AlphaPlantImpute and 0.92 for PlantImpute for $F_{10}$ focal individuals. Increasing the number of LD markers beyond 100 markers meant that the average accuracy of imputation for AlphaPlantImpute equalled that for PlantImpute. Figure 3b shows that with 100 LD markers, the average imputation accuracy was 0.99 for both AlphaPlantImpute and PlantImpute for $F_2$ focal individuals and for $F_{10}$ focal individuals.

For all numbers of selfing events separating parents and focal individuals, the precision of imputation accuracy (i.e. consistency across focal individuals) for AlphaPlantImpute was higher than for PlantImpute when the number of LD markers was low. Figure 3c is similar to Fig. 3b and plots the log of the precision of imputation accuracy for AlphaPlantImpute on the y-axis and PlantImpute on the x-axis. Points above the line indicate better precision (i.e. less variance) for AlphaPlantImpute than for PlantImpute, and vice versa. Figure 3c shows that with 3 LD markers, the precision of imputation was 1.62 for AlphaPlantImpute and 1.08 for PlantImpute for $F_2$ focal individuals and was 1.32 for AlphaPlantImpute and 1.11 for PlantImpute for $F_{10}$ focal individuals.

Figure 3c also shows that for medium number of LD markers, the precision of imputation accuracy for AlphaPlantImpute was higher than for PlantImpute for $F_2$ focal individuals but was lower when the number of selfing events was higher. With 20 LD markers, the precision of imputation accuracy was 2.48 for AlphaPlantImpute and 2.00 for PlantImpute for $F_2$ focal individuals and was 2.57 for AlphaPlantImpute and 2.80 for PlantImpute for $F_{10}$ focal individuals. With the highest number of LD markers (400), the precision of imputation accuracy was 3.84 for AlphaPlantImpute and 4.00 for PlantImpute for $F_2$ focal individuals and was 5.40 for both AlphaPlantImpute and PlantImpute for $F_{10}$ focal individuals.

Scenario 2: Effect of the level of inbreeding in parents

Increasing the level of inbreeding in the parents increases the imputation accuracy for AlphaPlantImpute. Figure 4a plots the accuracy of imputation in $F_2$ focal individuals of a
bi-parental population where the parents were \( F_1, F_2, F_4, F_{10} \) or \( F_{20} \). Figure 4a shows that with 20 LD markers, the average imputation accuracy increased from 0.81 for \( F_1 \) parents to 0.96 for \( F_{20} \) parents. Figure 4a also shows that increasing the level of inbreeding in the parents beyond \( F_4 \) did not increase the average accuracy of imputation for \( F_2 \) focal individuals. The average imputation accuracy with 20 LD markers was approximately 0.96 for \( F_2 \) focal individuals when parents were \( F_4, F_{10}, \) and \( F_{20} \).

For all levels of inbreeding in the parents and all numbers of LD markers, the average imputation accuracy with AlphaPlantImpute was almost always higher than with PlantImpute. Figure 4b is similar to Fig. 3b and plots the average imputation accuracy for AlphaPlantImpute on the y-axis and for PlantImpute on the x-axis. The shapes represent the level of inbreeding in the parents. Figure 4b shows that with 20 SNP LD markers, the average imputation accuracy was 0.81 for AlphaPlantImpute and 0.74 for PlantImpute for \( F_2 \) focal individuals when parents were \( F_1 \), 0.95 for AlphaPlantImpute and 0.91 for PlantImpute when parents were \( F_4 \), and 0.96 for AlphaPlantImpute and 0.94 for PlantImpute when parents were \( F_{10} \). In two cases, the average imputation accuracy with PlantImpute was slightly higher than with AlphaPlantImpute. This was when parents were \( F_4 \) and with 3 and 5 LD markers. The average imputation accuracy was 0.84 for AlphaPlantImpute and 0.80 for PlantImpute with 3 LD markers and was 0.87 for AlphaPlantImpute and 0.85 for PlantImpute with 5 LD markers.

For all levels of inbreeding in the parents and all numbers of LD markers, the precision of imputation accuracy with AlphaPlantImpute was almost always higher than with PlantImpute. Figure 4c is similar to Fig. 3c and plots the log of the precision of imputation accuracy for AlphaPlantImpute on the y-axis and PlantImpute on the x-axis. Figure 4c shows that with 20 LD markers, the precision of imputation accuracy was 2.16 for AlphaPlantImpute and 1.92 for PlantImpute. In a few cases, the precision of imputation accuracy for PlantImpute was slightly higher than AlphaPlantImpute. This was mainly when parents were \( F_{20} \) and with 50, 200, and 400 LD markers. The precision of imputation accuracy was 3.04 for AlphaPlantImpute and 3.40 for PlantImpute with 50 LD markers, was 3.71 for AlphaPlantImpute and 4.00 for PlantImpute with 200 LD markers, and was 3.84 for AlphaPlantImpute and 4.00 for PlantImpute with 400 LD markers.

**Scenario 3: Effect of chromosome size**

Increasing the chromosome size (in cM) decreased the imputation accuracy for AlphaPlantImpute. This was most
apparent when the number of LD markers was 10 or less. Figure 5a plots the imputation accuracy for seven chromosome sizes of 25, 50, 100, 150, 200, 300, and 400 cM for $F_2$ focal individuals of a bi-parental population where the parents were $F_{20}$. Figure 5a shows that with 3 LD markers, quadrupling the chromosome size from 25 to 100 cM decreased the average imputation accuracy from 0.95 to 0.85, and quadrupling from 100 to 400 cM decreased the average imputation accuracy from 0.85 to 0.55. The reduction in the imputation accuracy was less or non-existent when the number of LD markers was higher than 10. Figure 5a shows that the imputation accuracy was approximately 0.98 for all chromosome sizes when the number of LD markers was 50.

When the chromosome size was 300 cM or less, the average imputation accuracy was higher for AlphaPlantImpute than for PlantImpute. Figure 5b is similar to Fig. 3b and plots the average imputation accuracy for AlphaPlantImpute on the $y$-axis and for PlantImpute on the $x$-axis. The shapes represent the chromosome sizes. Figure 5b shows that with 3 LD markers, the average imputation accuracy was 0.95 for AlphaPlantImpute and 0.69 for PlantImpute when the chromosome size was 25 cM and was 0.61 for AlphaPlantImpute and 0.57 for PlantImpute when the chromosome size was 300 cM. The exception to this was when the chromosome size was 150 cM, where the average imputation accuracy was 0.70 for AlphaPlantImpute and 0.83 for PlantImpute. When the chromosome size was 400 cM, the average imputation accuracy was 0.55 for AlphaPlantImpute and 0.51 for PlantImpute when 3 LD markers were used but was 0.61 for AlphaPlantImpute and 0.68 for PlantImpute when 5 LD markers were used.

For all chromosome sizes and numbers of LD markers, the precision of imputation accuracy for AlphaPlantImpute was generally higher than for PlantImpute. Figure 5c is similar to Fig. 3c and plots the precision of imputation accuracy for AlphaPlantImpute on the $y$-axis and for PlantImpute on the $x$-axis. Figure 5c shows that with 3 LD markers, the precision of imputation accuracy was 0.71 for AlphaPlantImpute and 1.78 for PlantImpute when the chromosome size was 25 cM, was 1.08 for AlphaPlantImpute and 1.62 for PlantImpute when the chromosome size was 100 cM and was 1.59 for AlphaPlantImpute and 1.20 for PlantImpute when the chromosome size was 400 cM. The exception to this was when the chromosome size was 150 cM, where the precision of imputation accuracy was 1.17 for AlphaPlantImpute and 1.46 for PlantImpute.

**Scenario 4: Effect of the number of focal individuals in the bi-parental population**

Increasing the number of focal individuals in the bi-parental population slightly increased the imputation accuracy for

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**Fig. 5** Effect of chromosome size. **a** The genotype imputation accuracy using AlphaPlantImpute in $F_2$ focal individuals from a bi-parental cross of $F_{20}$ parents against seven chromosome sizes of 25, 50, 100, 150, 200, 300, and 400 cM. **b** Comparison of the average genotype imputation accuracy using AlphaPlantImpute ($y$-axis) versus using PlantImpute ($x$-axis). The colours represent the different LD arrays. The shapes represent the chromosome size. The red diagonal line indicates when the accuracy of PlantImpute equals AlphaPlantImpute. Points above the line are higher with AlphaPlantImpute and points below the line are when imputation accuracy is higher with PlantImpute. **c** Comparison of the precision in imputation accuracy using AlphaPlantImpute ($y$-axis) versus using PlantImpute ($x$-axis). The colours represent the different LD arrays. The shapes represent the chromosome size. The red diagonal line indicates when precision of PlantImpute equals AlphaPlantImpute. Points above the line indicate when the precision in accuracies is higher in AlphaPlantImpute and points below the line are when the precision in accuracies is higher in PlantImpute.
AlphaPlantImpute. This was most apparent when the number of LD markers was low. Figure 6 plots the accuracy of imputation for $F_2$ focal individuals of a bi-parental cross with 1, 5, 10, 25, 50 or 100 focal individuals. Figure 6 shows that increasing the number of focal individuals from 5 to 100 increased the average imputation accuracy from 0.83 to 0.85 when 3 LD markers were used. Figure 6 also shows that when the 10 or more LD markers were used, increasing the number of focal individuals had no effect on the imputation accuracy. When the number of LD markers was 400, the average imputation accuracy was 0.96 with 5 or 100 focal individuals in the bi-parental population.

Figure 6 also shows that when we only imputed one focal individual, the imputation accuracy fluctuated according to the focal individual that was sampled. As a result, increasing the number of LD markers did not always increase the imputation accuracy. For example, the average imputation accuracy was 0.95, 0.91, or 0.94 when 3, 5, or 10 LD markers were used. When 400 LD markers were used, the average accuracy of imputation was 0.997.

### Computational requirements of AlphaPlantImpute

Table 1 summarises the computational requirements of AlphaPlantImpute for twelve datasets across the three scenarios. Datasets were chosen to reflect the extremes in the number of selfing events separating parents and focal individuals ($F_2$ vs. $F_{10}$), the level of inbreeding in the parents ($F_1$ vs. $F_{20}$) and the number of LD markers (3, 50, or 400). Table 1 shows that the average run time for AlphaPlantImpute was 22.13 s with a maximum of 49.33 s. The average memory requirement for AlphaPlantImpute was 0.08 GB with a maximum of 0.082 GB.

| Parents | Focal individuals | LD array | Time (s) | Memory (GB) |
|---------|------------------|----------|----------|-------------|
| $F_{20}$ | $F_2$            | 3        | 37.41    | 0.079       |
| $F_{20}$ | $F_2$            | 50       | 8.14     | 0.080       |
| $F_{20}$ | $F_2$            | 400      | 7.95     | 0.082       |
| $F_{20}$ | $F_{10}$         | 3        | 49.33    | 0.079       |
| $F_{20}$ | $F_{10}$         | 50       | 8.48     | 0.080       |
| $F_{20}$ | $F_{10}$         | 400      | 9.40     | 0.082       |
| $F_1$   | $F_2$            | 3        | 26.70    | 0.080       |
| $F_1$   | $F_2$            | 50       | 35.10    | 0.080       |
| $F_1$   | $F_{10}$         | 3        | 24.66    | 0.079       |
| $F_1$   | $F_{10}$         | 50       | 35.41    | 0.080       |
| $F_1$   | $F_{10}$         | 400      | 10.35    | 0.082       |
| Average |                  |          | 22.13    | 0.080       |

### Discussion

Our results highlight three points for discussion: (1) the performance of AlphaPlantImpute; (2) the performance of AlphaPlantImpute compared to PlantImpute; and (3) future development of AlphaPlantImpute.

**Performance of AlphaPlantImpute**

This paper presents a new heuristic method, called AlphaPlantImpute, for phasing and imputation of SNP array data in diploid plant species. AlphaPlantImpute explicitly leverages features of plant breeding programmes to impute LD focal individuals to HD. The explicit utilisation of pedigree information and heuristics developed specifically to track the inheritance of parental haplotypes using the LD genotypes of focal individuals are likely to be the reasons for AlphaPlantImpute’s robust and consistent performance across all tested scenarios. AlphaPlantImpute achieves high imputation accuracy of between 0.8 and 1.0 for the majority of scenarios. For scenarios where the imputation accuracy was below 0.8, increasing the number of LD markers increased the imputation accuracy.

Increasing number of selfing events separating parents and focal individuals from $F_2$ to $F_{10}$ only slightly decreases the imputation accuracy. Decreasing the level of inbreeding in the parents or increasing the chromosome size decreases the imputation accuracy when the number of LD markers is 10 or less. However, in both cases, the decrease in the imputation accuracy could be mitigated by increasing the number of LD markers to 20 SNP or more.

Decreasing the number of focal individuals in the bi-parental population slightly decreases the imputation accuracy. This was most evident when the number of LD markers
was 10 SNP or less. The likely cause of this is that inferring the most likely linkage between alleles for two markers is difficult with fewer focal individuals, since fewer individuals will be homozygous at the markers. In this case, the algorithm defaults to the linkage pattern of alleles in the parents. This may be sub-optimal for imputing markers in regions with elevated recombination rates, i.e. hotspots. When there was a single focal individual in the focal family, the accuracy of imputation for that individual varied. The likely cause of this is whether an individual had a recombination or whether it had inherited the parental haplotypes without recombination. One solution to this situation could be to utilise the most likely linkage from related families with more genotyped focal individuals (see section: Future work and developments).

Overall, the results suggest that for a given population, high imputation accuracy can be achieved even when the number of LD markers is low, and small increases in the number of markers can achieve high accuracies depending on the biology of the species (i.e. recombination rate, obligate outcrossing) and the pedigree design (outbred, inbred, level of selfing).

In this study, we tested a maximum LD marker array size of 400 SNP. With declining genotyping and sequencing costs, the number of markers on an LD array may increase to a few thousand. To test the scalability of AlphaPlantImpute with denser SNP panels, we performed an additional simulation of a bi-parental population with two inbred parents and 100 $F_2$ focal individuals for imputation. Parents had HD genotypes for 25,000 SNP on a single chromosome, and $F_2$ focal individuals had genotypes at LD for 1000, 3000 and 5000 SNP. The time and memory requirements for AlphaPlantImpute for imputing the $F_2$ focal individuals to HD was 17 s and 0.094 GB for 1000 SNP, 28 s and 0.114 GB for 3000 SNP and 644 s and 0.137 GB for 5000 SNP. We are working on further optimisation of the time and memory requirements of AlphaPlantImpute for larger SNP panels.

**Performance of AlphaPlantImpute compared to PlantImpute**

The imputation accuracy for AlphaPlantImpute was compared to that for PlantImpute (Nettelblad et al. 2009; Hickey et al. 2015). In the majority of cases, the imputation accuracy was higher for AlphaPlantImpute than for PlantImpute. One exception to this was when the chromosome size was 400 cM and when the number of LD markers was 20 or less (e.g. 0.88 vs. 0.90 when the number of LD markers was 20). One reason for this could be that unless there is enough information in the genotypes of focal individual on the LD array, the heuristic algorithm in AlphaPlantImpute is inherently more conservative in determining recombination regions compared to the probabilistic algorithm in PlantImpute. As such, AlphaPlantImpute is more likely to leave positions as missing and fill them in as the parent average in the final step.

The precision of imputation accuracy (calculated as the log of the inverse of the variance in imputation accuracy within each bi-parental population) was also higher in the majority of cases for AlphaPlantImpute than for PlantImpute. This was most apparent with small number of LD markers. The higher precision of imputation accuracy for AlphaPlantImpute is likely a consequence of directly calling allele phase and parent-of-origin and imputed genotypes in turn. The probabilistic algorithm of PlantImpute is marginalising over the all possible phase and genotype, which is probabilistically correct and handles the uncertainty properly, but it seems this is lowering the imputation accuracy. One exception to this was when the chromosome size was 150 cM, where the precision of imputation accuracy was higher for PlantImpute than for AlphaPlantImpute for all LD arrays.

The biggest advantage of AlphaPlantImpute compared to PlantImpute relates to computational requirements. Hickey et al. (2015) report that to perform imputation within a single bi-parental population of 100 $F_2$ focal individuals, PlantImpute required a minimum of 3 h and in excess of 100 GB of memory. In comparison, AlphaPlantImpute required on average ~22 s and ~0.08 GB of memory for all tested scenarios.

The high and consistent accuracies achieved with very low computational requirements makes AlphaPlantImpute an attractive, reliable and practical tool for routine use in plant breeding programmes that are already using or will include SNP array data to inform selection decisions.

**Future work and developments**

At present, the heuristic method in AlphaPlantImpute works within the most common plant breeding programme design of bi-parental populations and it works best when parents are fully inbred or close to being fully inbred. AlphaPlantImpute could be extended in multiple ways. For example, instead of treating each bi-parental population as an independent unit it could simultaneously work across bi-parental populations that share parents. This could increase the imputation accuracy in three ways: (1) information between bi-parental populations could be shared for imputation of focal individuals that are effectively half-sibs (one common parent); (2) information between bi-parental populations could be used to resolve phase where one or both parents are heterozygous at one or more consecutive markers; and (3) if a common parent has no or LD genotypes available, information from its descendants across half-sib bi-parental populations could be leveraged to phase and impute it to high density.

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AlphaPlantImpute could also be extended to include ancestral pedigree information (such as grandparents and great-grandparents). This could be useful for improving phasing and imputation of parents with missing information or that are highly outbred. More simply, AlphaPlantImpute could also be extended so that it can directly read in and exploit phased information for the fully or partially outbred parents. Such phased information could be generated for parents by running AlphaPlantImpute on the bi-parental family from which the fully or partially outbred parent derived.

AlphaPlantImpute could be extended so that it reads in previously inferred most likely linked alleles at two markers. It is likely that linkage patterns are shared across families, especially if the families are related. Using this information across families would be especially suited to imputation situations in bi-parental populations that have only a few genotyped focal individuals (e.g. one genotyped individual per family).

Finally, although SNP arrays for the many domesticated plant species exist, low-coverage sequencing methods such as genotyping-by-sequencing are also used. The heuristics of AlphaPlantImpute might be extended to enable imputation with such data.

**Software availability**

We implemented our method in a software package called AlphaPlantImpute, which is available for download at http://www.AlphaGenes.roslin.ed.ac.uk/AlphaPlantImpute/ along with a user manual.

**Author contributions statement** SG and JH conceived the method. SG further developed the method, coded the final programme, developed the study design and performed the analysis. VM, RCG, EB and GG contributed to the development of components of the method, to the design and analysis and to the interpretation of the results and provided comments on the manuscript. SG and JH wrote the first draft. All authors read and approved the final manuscript.

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**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.
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