Invited Review

Genomics-assisted breeding in minor and pseudo-cereals

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Minor and pseudo-cereals, which can grow with lower input and often produce specific nutrients compared to major cereal crops, are attracting worldwide attention. Since these crops generally have a large genetic diversity in a breeding population, rapid genetic improvement can be possible by the application of genomics-assisted breeding methods. In this review, we discuss studies related to biparental quantitative trait locus (QTL) mapping, genome-wide association study, and genomic selection for minor and pseudo-cereals. Especially, we focus on the current progress in a pseudo-cereal, buckwheat. Prospects for the practical utilization of genomics-assisted breeding in minor and pseudo-cereals are discussed including the issues to overcome especially for these crops.

Key Words: QTL analysis, genome-wide association study, marker-assisted selection, genomic selection, minor cereal, pseudocereal, buckwheat.

Introduction

The world population is predicted to exceed 9.7 billion by 2050 (FAOSTAT 2018), and as a result, the global food crisis will be exacerbated over the next few decades. The crisis will also be worsened by the ongoing progression of global-scale climate change and the limitation of resources for agricultural production, such as water, fertilizer, and suitable land. Therefore, in order to circumvent negative consequences, multifaceted efforts that incorporate multiple strategies, such as the improvement of cultivation management technologies (e.g., smart farming systems) and the reduction of food loss during cultivation and distribution, will be needed.

The world’s current food production is mainly supported by a small number of major crops, and, thus is vulnerable to major threats, such as pests, diseases, and, now, climate change (Massawe et al. 2016). Therefore, the diversification of food crops, including underutilized crops, is an important strategy for distributing risk of major threats in food production and for stabilizing global food production. Importantly, some of the crops that are currently underutilized, like minor and pseudo-cereals, are also suitable for low-input agriculture and exhibit high resistance to a variety of environmental stressors (Zhang et al. 2018). Because minor and pseudo-cereals are at least as nutritious as major cereals and often produce special nutrients that are not produced by major cereal crop, they are also important in supporting the dietary health of human populations (Cernansky 2015, Cheng et al. 2017, Kaur et al. 2014, Mir et al. 2018, Nowak et al. 2016, Sytar et al. 2016, Vega-Gálvez et al. 2010). As a result, minor and pseudo-cereals are currently attracting the attention of various stakeholders and breeders (Chirurgwi et al. 2019, D’Amelia et al. 2018, Joshi et al. 2019, Upadhyaya and Vettriventhal 2018, Zurita-Silva et al. 2014).

However, because minor and pseudo-cereals have a short history of modern breeding, their yields are often lower than those of major cereals. For example, minor and pseudo-cereals often have characteristics, such as lodging and seed shattering, that cause low yield (Cheng et al. 2017, Hinterthuer 2017). Minor and pseudo-cereals also possess other disadvantages, such as the presence of anti-nutrients, poor digestibility, and low palatability, all of which greatly affect their utilization as food sources (Kaur et al. 2014, Zurita-Silva et al. 2014). Therefore, it is necessary to establish breeding programs to build upon the crops’ advantages and to compensate for their disadvantages. One of the most important means of addressing this is to improve the genetic quality of crop species through efficient breeding programs (Abberton et al. 2016, Tester and Langridge 2010).

During the last few decades, three major DNA marker-based methods aiming at efficient genomics-assisted breeding, namely biparental QTL mapping, genome-wide association studies (GWAS), and genomic selection (GS), have been developed and applied to the breeding of crops...
Indeed, both researchers and breeders of major crops have developed and used these methods as front runners for genomics-assisted breeding. However, recent advances in molecular genetic technology and the commoditization of such technologies have improved the availability of genomic methods, even for the breeding of minor crops, like minor and pseudo-cereals. Crucial disadvantages of minor and pseudo-cereals, such as lodging, seed shattering, and other characteristics affecting their utility, show their lack of refinement as domestic food crops. The success of genetic improvement depends on the efficient use of genetic resource. Major crops that have experienced intense modern breeding may be exposed to a loss of genetic diversity in a breeding population (Esquinas-Alcázar 2005, Fu 2015). On the other hand, breeding populations of minor crops are expected to retain genetic variation held in their founder populations due to their shorter history of modern breeding. Thus, it should be noted that both minor and pseudo-cereals are expected to have great potential to be improved through modern breeding utilizing genomics information (Armstead et al. 2009, Varshney et al. 2012).

Accordingly, the aim of the present paper was to review the progress achieved to date, in regards to the use of biparental QTL mapping, GWAS, and GS for the genomics-assisted breeding of minor and pseudo-cereals, and to discuss future prospects for the application of genomics-assisted breeding.

**Biparental QTL mapping**

One strategy, namely biparental QTL mapping, has become routinely performed in many crop species owing to the appearance of versatile DNA markers and the development of statistical models and methods (Kearsey and Farquhar 1998). Indeed, QTLs for a variety of traits have been reported for multiple crop species. In fact, Bernardo (2008) reported that 1,200 QTL mapping studies, which involved 10,000 QTLs and 12 major crop species, had been published, and the number of such studies has increased since then (Bernardo 2016).

On December 26th, 2018, we searched the Web of Science database (http://www.webofknowledge.com/) for titles that contained terms “QTL”, “QTLs”, “quantitative trait locus”, “quantitative trait loci”, or “markers” + “associated”. The searching query was the same as that described by Bernardo (2008), except that we focused on different crop species (see below). As a result, 220 QTL mapping studies involving 12 words meaning minor and pseudo-cereal species, we identified: millet, spelt (Triticum spelta L.), emmer (Triticum turgidum L.), einkorn (Triticum monococcum L.), sorghum (Sorghum bicolor L. Moench), quinoa (Chenopodium quinoa Willd), amaranth (Amaranthus L.), buckwheat (common buckwheat (Fagopyrum esculentum Moench) and tartary buckwheat (Fagopyrum tataricum Gaertn.), chia (Salvia hispanica L.), oat (Avena sativa L.), rye (Secale cereale L.), and teff [Eragrostis tef (Zuccagni) Trotter]. Although the number is far smaller than that reported for major crops, the number of the studies published per year has been increasing steadily, especially during the last decade (Fig. 1).

Most of the QTL mapping studies of minor and pseudo-cereals have focused on sorghum, which is the forth largest calorie-supplying cereals in the world (FAOSTAT 2013). Interestingly, because sorghum is important both as a food and feed crop and as a bioenergy crop, QTL mapping studies of sorghum increased most dramatically after the late 2000s (e.g., Murray et al. 2008). In addition, the...
development of QTL mapping in sorghum may have been easier than that in other minor and pseudo-cereals because the researchers could use DNA markers developed for other species, such as maize (Zea mays L.) (e.g., Hulbert et al. 1990, Lin et al. 1995). This was a great advantage in the 1990s, when marker development was costly, and the application of markers from other species helped to identify orthologous QTLs. Another advantage of sorghum as a target of QTL mapping is its autogamous reproductive system, which has facilitated the development of experimental populations, such as recombinant inbred lines (RILs) and near-isogenic lines (NILs) (e.g., Harris et al. 2007, Sanchez et al. 2002). Recently, a nested association mapping (NAM) population, which includes 2,214 RILs derived from 10 diverse global lines crossed with an elite reference line, was developed (Bouchet et al. 2017) and is being used by the sorghum research community.

Studies involving the QTL mapping of other minor and pseudo-cereals, however, are relatively limited and mostly include studies in rye (Erath et al. 2016, 2017, Miedaner et al. 2018, Myśkowski et al. 2018, Wang et al. 2015), oat (Admassu-Yimer et al. 2019, Babiker et al. 2015, Pellizzaro et al. 2016, Schneider et al. 2015, Sunstrum et al. 2018, Zimmer et al. 2018), and millet. Of the various types of millet, pearl millet (Ambawat et al. 2016, Aparna et al. 2015, Kumar et al. 2016b, 2017, 2018, Pucher et al. 2018, Punnuri et al. 2016, Taunk et al. 2018) and foxtail millet (Fang et al. 2016, Jia et al. 2017, Mauro-Herrera and Doust 2016, Ni et al. 2017, Odonkor et al. 2018, Wang et al. 2017a, 2017b, 2019) have received the majority of research attention, although one study also focused on proso millet (Rajput et al. 2016). Notably, very few QTL mapping studies have focused on pseudocereals. For wheat’s close relatives, triticale (a hybrid of wheat and rye) and progeny populations derived from crossing wheat and its wild relatives have been analyzed with QTL mapping (Dhariwal et al. 2018, Kalih et al. 2015, Miedaner et al. 2016, Wen et al. 2018). For example, emmer wheat has been used as a donor of disease resistance, drought resistance, productivity, and grain nutrient to durum wheat (Triticum turgidum ssp. durum) and bread wheat (Merchunk-Ovnat et al. 2016a, 2016b, Zhang et al. 2014).

In terms of materials and methods, RILs are most commonly used for QTL mapping, followed by F₂ populations. Therefore, even in minor cereals, RIL populations have been developed for QTL mapping and are shared among researchers. In triticale, a doubled haploid (DH) population have been created and used (Dhariwal et al. 2018, Kalih et al. 2015, Miedaner et al. 2016, Wen et al. 2018). For example, emmer wheat has been used as a donor of disease resistance, drought resistance, productivity, and grain nutrient to durum wheat (Triticum turgidum ssp. durum) and bread wheat (Merchunk-Ovnat et al. 2016a, 2016b, Zhang et al. 2014).

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In contrast, there are few reports of shared materials for the QTL mapping of pseudocereals. This is partly due to the fact that the major pseudocereals, such as common buckwheat, quinoa, and amaranth, are all allohexaploids. In common buckwheat, an inbred population (F₁) have been derived from two interspecific hybrids between common buckwheat (Fagopyrum esculentum) and its autogamous wild relative (Fagopyrum homotropicum) (Hara et al. 2011). The development of immortal populations, such as RILs and DH populations, will facilitate collaborative phenotyping and the aggregation of phenotype data from multiple institutions.

Among the strategies used for QTL mapping, the use of single nucleotide polymorphism (SNP) markers is the most common. Such markers can be investigated using either DNA array technologies (e.g., Babiker et al. 2015, Dhariwal et al. 2018, Erath et al. 2017, Schneider et al. 2015) or next-generation sequencing (NGS) technologies, such as genotyping-by-sequencing (GBS; Elshire et al. 2011) and restriction site-associated DNA sequencing (RAD-Seq; Baird et al. 2008, Miller et al. 2007), (e.g., Punnuri et al. 2016, Rajput et al. 2016, Sunstrum et al. 2018, Wang et al. 2017a, Wen et al. 2018). Not a few studies have used simple sequence repeats (SSRs) as markers (e.g., Aparna et al. 2015, Fang et al. 2016, Taunk et al. 2018).

Furthermore, QTL mapping has been used to investigate a variety of target traits. Studies frequently focused on biotic and biotic stress resistance, including disease resistance, drought tolerance, frost tolerance, aluminum tolerance, etc. (e.g., Aparna et al. 2015, Erath et al. 2017, Schneider et al. 2015, Sunstrum et al. 2018), as well as a variety of general agronomical and morphological traits. QTL mapping has also been performed in minor cereals to identify loci associated with nutrient content (e.g., Kumar et al. 2018). Many traits, such as disease resistance and seed shattering, are often controlled by primary genes, so the strategy of identifying QTLs and performing subsequent marker-assisted selection (MAS) is promising for the improvement of crop traits.

The major disadvantage of using QTL mapping in minor and pseudo-cereals is that experimental populations are needed. Indeed, when compared to the resources available for major cereals, relatively few experimental populations of minor and pseudo-cereals have been developed, and even though genotyping of materials in populations is no longer a bottleneck, the construction of experimental populations will require both time and money. It is worth noting that other strategies, such as GWAS, do not require the construction of experimental populations. However, QTL mapping is still superior in its estimation of allele location and effect, especially for low-frequency alleles (Bernardo 2016). Alleles that are useful but underutilized in breeding history may be rare in a population of breeding materials or a population of germplasm collections.
More recently, the advent of NGS technology encouraged researchers to conduct GWAS. The purpose of GWAS is to detect markers associated with target traits from a huge number of genome-wide markers. GWAS can be performed using natural genetic resources or breeding populations because the use of high-density marker sets permits the utilization of linkage disequilibrium (LD) between QTLs and markers without creating linkage blocks, as in biparental QTL mapping. Therefore, GWAS can capture a substantial proportion of the variation found in a species-wide population (Hamblin et al. 2011).

Another advantage of GWAS is that they could solve some of the disadvantages of biparental QTL analysis. For example, GWAS does not require the construction of mapping populations, thereby shortening the time necessary for analysis. Because mapping populations of less-studied minor or pseudo-cereals would sometimes need to be constructed from scratch, the possibility of using natural or breeding populations is a major advantage. Another disadvantage of QTL analysis is the bias in detectable QTLs, which depend on parents used to generate the mapping population, potentially identifying useless markers or QTLs in the target breeding population. For example, when one of the QTL mapping parents is even a wildtype of the target population. In contrast, GWAS is not biased by the selection of cases, e.g., when an allele is fixed in the cultivated breeding analysis. Because mapping populations of less-studied population, potentially identifying useless markers or QTLs populations’ high heterogeneity is expected, GWAS may be minor or pseudo-cereals, in which the use of the breeding genetic diversity. Morris et al. (2013a), for example, found that their global sorghum accessions exhibited strong population structure, according to both morphological type and geographic origin. Additionally, evaluation of the GWAS accuracy for a known sorghum gene Tannin1 revealed difficulty in treating strong population structure (Morris et al. 2013b). Strong population structure has also been reported in foxtail millet (Jia et al. 2013). However, Newell et al. (2011) analyzed the LD and population structure of an oat population to evaluate the feasibility of GWAS and found relatively weak population structure, despite high diversity. In GWAS of plant species, especially for species with large genetic diversity, the existence of strong population structure sometimes worries researchers. It is recommended that researchers evaluate the population structure of candidate populations and select GWAS methods that are optimal for the situation.

Several methods have been proposed for circumventing the effect of population structure in GWAS. Using principal components or eigenanalysis (implemented in the R package rrBLUP, one of the familiar GWAS tools; Endelman 2011, Endelman and Jannink 2012) is one of the typical ways. On the other hand, when the total number of genotypes in an experimental population is quite large, it may also be possible to conduct separate GWAS (Crowell et al. 2016). Bayesian methods or other variable selection methods that are used for genomic prediction can have a possibility of GWAS with circumventing the effect of population structure.
structure. For the application of GWAS to genomics-assisted breeding, the analysis of population genetics and choice of optimal method would work be key to identifying truly useful QTLs.

Another disadvantage of GWAS is that rare and low-frequency alleles can sometimes have relatively large effects on target traits (Manolio et al. 2009). Indeed in the field of human genetics, researchers have tried to solve this issue by collecting data from hundreds of thousands of people (e.g., Marouli et al. 2017). In GWAS of plant species, however, collecting and evaluating such a large number of genotypes may be impossible. Recently developed high-throughput phenotyping systems will help us collect phenotype data from large sample populations (e.g., Spindel et al. 2018). For example, it is reported that image analysis can be used for phenotyping in GWAS of sorghum (Zhou et al. 2019). If the problems of size of experimental field, cost, and phenotyping can be solved, the collection and utilization of genotypes that encompass the diversity of the population or species is a key factor in GWAS, in order to incorporate minor allele into the population. Therefore, the collaboration of researchers, gene banks, and breeders will be necessary.

The use of GWAS is expected to promote the genomics-assisted breeding of less-studied minor and pseudo-cereals. However, it is important to note that GWAS can only detect major genes, as in QTL analysis. MAS may be one strategy for using the QTL information produced by GWAS, as mentioned in the section of biparental QTL mapping. Spindel et al. (2016) used markers detected by GWAS as fixed effects in a genomic prediction model for rice, which may be another useful strategy.

### Genomic selection

Meuwissen et al. (2001) proposed the idea of GS for accelerating the genetic improvement of quantitative traits controlled by a number of genes. When using this strategy, a prediction model is built based on the relationship between phenotypic values and genome-wide marker genotypes in a training population and then used to calculate the genotypic values of selection candidates. GS models utilize the LD between QTLs and high-density markers, as in GWAS. Because the purpose of GS is to predict the breeding values of selection candidates, causal loci are not necessarily identified (Hamblin et al. 2011). By combining GS with accelerating generations using off-season nursery, like conventional MAS based on QTL mapping, breeders can conduct more than one cycle of GS per cycle of phenotypic selection, thereby achieving high-speed breeding.

The advantage of GS over QTL-based MAS is the involvement of a large number of QTLs with small effects at the selection step (Hayes et al. 2013, Heffner et al. 2010). In biparental QTL analysis and GWAS, only major genes can be detected, so it is difficult to improve the traits controlled by a number of genes. In addition, GS could potentially solve some of the other disadvantages of MAS based on known QTLs, such as the overestimation of QTL effects (Strauss et al. 1992) and disagreement of effective QTLs between mapping and breeding population (Hoeschele and VanRaden 1993, Lande and Thompson 1990, Melchinger et al. 1998). This also mentioned in the ‘Genome-wide association studies’ section as the bias of detectable QTLs in biparental QTL mapping. Because of the advantages of GS over the conventional MAS, GS is a promising alternative for improving the efficiency of plant breeding for both major and minor crops.

In addition, GS should work especially well for under-utilized minor crops with high genetic diversity in a breeding population. Indeed, GS requires neither marker position information nor QTL information. Therefore, GS can even be used for less-studied minor or orphan crops if genotyping and phenotyping systems are built. Moreover, the genetic diversity harbored by breeding populations of minor crops is ideal for GS, which can include a number of QTLs with small effects. Together, both the possibility of immediate application and the high genetic diversity of breeding population give reason to apply GS to minor and pseudo-cereals.

Several attempts for performing GS in minor and pseudo-cereals have been reported. Previous studies have reported empirical GS for the selection of β-glucan concentrations in oat (Asoro et al. 2013). Asoro et al. (2013) conducted GS and compared its efficiency with phenotypic selection based on BLUP (Henderson 1984) and MAS based on GWAS in the GS training population, in which the advantage of GS (i.e., acceleration of breeding cycles and avoiding bias in detectable QTLs) were not hold. Even though they did not demonstrate the advantage of GS over other methods in the level of genotypic improvement, they suggested that greater genetic variation was maintained by GS than by BLUP-based phenotypic selection by accounting for Mendelian segregation and thus avoiding the selection of related individuals. In the ‘Genomics-assisted breeding of buckwheat’ section, we will present studies relating to GS breeding in common buckwheat, as an example of successful GS in an allogamous pseudo-cereal (Yabe et al. 2018).

To conduct GS breeding, breeding strategy must be optimized. GS breeding includes marker genotyping and generally the acceleration of generations, thereby requiring more cost and labor than ordinary phenotypic selection. Especially when the GS of minor or pseudo-cereals is performed from scratch, strategy optimization is crucial. Several studies have addressed GS breeding in minor cereals. In many cases, prediction accuracy has been evaluated to determine the suitability of using GS, optimal prediction method, optimal training population set, and necessary number of DNA markers (e.g., in rye (Auinger et al. 2016, Bernal-Vasquez et al. 2014, 2017, Wang et al. 2014), oat (Asoro et al. 2011), pearl millet (Li et al. 2018) and sorghum (Fernandes et al. 2018, Hunt et al. 2018)). To optimize
breeding strategy for several members of the Poaceae, both the prediction accuracy and estimated response to selection have been calculated (Marulanda et al. 2016). On the other hand, Yabe et al. (2013, 2014b) conducted simulation studies to optimize breeding strategy for the GS breeding of common buckwheat. Such preparation for actual breeding programs may enable researchers to utilize the genetic variation of breeding populations more efficiently and is necessary for GS programs, especially in minor crop populations with high genetic variation.

Recently developed genotyping systems can help us perform GS breeding for minor and pseudo-cereals, for which the allowable cost or scale of breeding is sometimes limited. The utilization of high-throughput phenotyping technology, such as unmanned aerial vehicles (UAVs), could also be used for genomic prediction modeling, even in large-scale fields of sorghum plants (Watanabe et al. 2017). The rapidly developing technology will also help GS breeding in minor cereals with great diversity in a large field.

Genomics-assisted breeding of buckwheat

It is likely that genomics-assisted breeding will be used to help meet the growing demand for minor and pseudo-cereals, especially for underutilized pseudo-cereals with high genetic diversity in breeding populations. However, the situation is more serious for pseudo-cereals than minor graminaceous cereals, owing to the difficulty of applying genetic information or markers from major graminaceous cereals, as indicated by the small number of studies that have reported performing genetic analysis in pseudo-cereals. The goal of this section is to summarize previous reports of genetic analysis and genomics-assisted breeding in common buckwheat, as an example of an allogamous pseudo-cereal.

Common buckwheat is an annual crop that is grown widely in temperate zones and that is used in both bakery products (e.g., bread, biscuits, and noodles) and non-bakery products (e.g., honey and tea), as a great source of nutrients (Giménez-Bastida et al. 2015). In addition, the perception of common buckwheat as a healthy and functional food is growing (Giménez-Bastida and Zielinski 2015), and efforts have been made to incorporate buckwheat’s beneficial nutrients into our everyday eating habits (e.g., Baljeet et al. 2010).

The breeding of common buckwheat, however, has had a difficult history. Most types of common buckwheat are characterized by complete outcrossing, owing to heteromorphic self-incompatibility (Campbell 1997) controlled by the S-locus (Lewis and Jones 1992), and as in breeding programs of other cross-pollinated species, mass selection for population improvement is performed conventionally, in which the lack of pollen control and inbreeding depression hinder efficient genetic improvement (Acquaah 2009). The difficulty of evaluating single plants reduces the efficiency of mass selection, as well (Yano et al. 2002). Therefore, genomics-assisted breeding is expected to provide more rapid results than conventional phenotypic selection methods, especially for crops like common buckwheat, for which selection has been hindered by the disadvantages of phenotype-based mass selection.

Biparental QTL mapping has also been reported in common buckwheat, for several growth and morphological traits. However, common buckwheat’s allogamous reproductive system works as a barrier because the allogamous reproductive system has hindered the development of a suitable mapping population (e.g., RIL, NIL, or NAM population). Hara et al. (2011) used QTL analysis to investigate the photoperiod sensitivity of common buckwheat using an F4 population that was derived from a cross between autogamous lines, in order to circumvent the complex genetic patterns caused by the cross-pollinating reproductive system. In their study, Hara et al. (2011) constructed a linkage map, performed QTL analysis using 63 expressed sequence tag (EST) markers and three candidate genes, which were homologs of Arabidopsis genes, and identified a single candidate gene and two ESTs that were associated with the target trait. Besides, linkage maps for allogamous common buckwheat populations can also be constructed using pseudo-testcross mapping strategy (Grattapaglia and Sederoff 1994), in which a map was constructed for each parent and then two maps were bridged (Konishi and Ohnishi 2006, Yasui et al. 2004). Yabe et al. (2014a), for example, used this strategy and high-density markers, with NGS technology, to construct a high-density linkage map for a common buckwheat population, and, then conducted QTL analysis for main stem length. This study verified the usefulness of linkage maps, even for allogamous populations. Such studies suggested that QTL analysis and other QTL detection methods (e.g., GWAS) will contribute to allogamous crop breeding via conventional MAS for major genes.

Meanwhile, GS is one of the most promising breeding methods for common buckwheat populations with high genetic diversity, partly owing to the crop’s allogamous reproductive system, which can create new combinations of beneficial genes. Moreover, GS, through evaluating genetic ability based on marker genotype and phenotypic information, can realize a higher accuracy than the relatively low accuracy of phenotypic selection with single plant evaluation, thereby solving the problem of allogamous crop breeding. Yabe et al. (2018) performed empirical GS breeding in common buckwheat and verified that GS was useful for mass selection in allogamous populations with low LD. The authors also demonstrated the importance of both breeding cycle acceleration and frequent model updating, the latter of which may be a unique requirement of GS breeding in allogamous crop populations with low LD. In addition, the marker genotyping system used for the study was created from scratch using NGS technology (Yabe et al. 2014a), simulation studies were used to evaluate the
efficiency of GS mass selection and to optimize GS breeding strategy before empirical breeding was attempted (Yabe et al. 2013, 2014b). It is important to note, however, that, even though common buckwheat population possesses high genetic diversity, which is beneficial for GS, the low LD of allozyme common buckwheat populations might actually hinder GS. Such attempt of the combination of simulations and empirical GS breeding suggests that GS breeding can be used to achieve rapid genetic improvement in common buckwheat and confirm the importance of using optimized breeding strategies. It is essential to figure out an optimal breeding program in advance, especially for common buckwheat breeding, with few research results to date due to the short history of genomics-assisted breeding. In the above-mentioned GS for common buckwheat, the GS model only incorporated additive effects, possibly accumulated via mass selection in a population and, on average, maintained, even during seed multiplication after GS. Due to the species’ completely out-crossing reproductive process, it is difficult to utilize the dominance (or overdominance) and epistasis, including dominance, in mass selection of a common buckwheat population, due to heterogeneity. Meanwhile, it may be possible to fix epistasis that consists of additive effects only, even in an outcrossing population, if the effects could be precisely estimated through a large-scale dataset or biparental QTL mapping. When the hybrid breeding performed in maize breeding is applied to buckwheat breeding in the future, the utilization of dominance effects in genomics-assisted breeding will be required. The GS model should not only be chosen based on selection accuracy, but also on the assumed type of the population to be released as a variety. In addition, optimal genomics-assisted breeding methods should be employed. MAS, based on biparental QTL analysis or GWAS, can drastically improve genetic ability in a common buckwheat population, which has not experienced intense selection, especially when the target trait is controlled by a small number of genes or the detected QTL considerably explains the amount of variation in the population. On the other hand, GS is also promising for the genetic improvement of polygenic traits where a single method could be used to build a GS model incorporating detected QTLs, or MAS and GS could be conducted in series. To create an optimal breeding program for common buckwheat, genetic knowledge of the target traits and population as well as breeding simulations will be needed.

Studies mentioned above were performed without using any reference genome information for common buckwheat, because there were no available resources. However, the recently published draft genome of common buckwheat (1.18 Gbp; Yasui et al. 2016b) enables us to make progress. Using the genome information of common buckwheat, the knowledge of agronomically useful genes in other plant species facilitated the identification of potentially useful homologs (Matsui et al. 2018, Yasui et al. 2016b), which can then be analyzed to estimate function and evolutionary history (Che et al. 2018, Mizuno and Yasui 2019). This rapid growth of genomics technology will enhance the genomics-assisted breeding in common buckwheat.

Genomics-assisted breeding methods, such as GS and MAS, will be useful for the genetic improvement needed to meet the growing demand for minor and pseudo-cereals, as well as for the development of new varieties that can adapt to future climates (Varshney et al. 2018). Genomics-assisted breeding of minor and pseudo-cereals is also expected to contribute to global food safety, by simultaneously increasing crop diversification and improving crop yield and quality.

Genetic variation is essential for efficient genetic improvement. Because of their relatively short or non-intensive history of breeding, the current collections of cultivars of minor and pseudo-cereals are expected to harbor high genetic diversity (Zhang et al. 2018). However, when compared to the resources available for major crop species, the numbers of minor and pseudo-cereal accessions in gene banks are inferior and insufficient (FAO 2010) and DNA polymorphism information of their genetic resources has not been prepared. For these reasons and to promote the use of current accessions for breeding programs, it is crucial that useful genetic resources be collected before they are lost and that the available genetic resources be characterized and evaluated, possibly including DNA polymorphism information (FAO 2010). One of the main advantages of GS, QTL mapping, and GWAS is that no reference genome is required. For example, no reference genome information was used in the GS breeding experiment of buckwheat (Yabe et al. 2018). However, the availability of reference genomes could further increase the usefulness of genomics-assisted breeding. To detect candidate genes that cause associations between markers and phenotypes, for example, GWAS requires marker position information and an annotate genome. Therefore, draft genomes have recently been developed for common buckwheat (Yasui et al. 2016b), pearl millet (Varshney et al. 2017), grain amaranth (Amaranthus hypochondriacus L.; Clouse et al. 2016), tef (Cannarozzi et al. 2014), and quinoa (Yasui et al. 2016a, Zou et al. 2017), and Varshney et al. (2017) verified the usefulness of the draft pearl millet genome for GWAS and genomic prediction. In addition, databases of genomic-resources have also been developed for a variety of minor and pseudo-cereals, e.g., buckwheat (Yasui et al. 2016b) and pearl millet (Jaiswal et al. 2018). The cost efficiency and high-throughput nature of whole-genome sequence analysis are improved rapidly by the improvement of sequencing technology.
Heffner et al. 2010), the development of generation acceleration methods and protocols becomes very important (Ghosh et al. 2018). However, the development of generation acceleration techniques and protocols for minor and pseudo-cereals is generally lacking (Chiurugwi et al. 2019), with few exceptions (e.g., Ghosh et al. 2018, Stetter et al. 2016). In a recent review, Chiurugwi et al. (2019) reviewed the advantages of speed-breeding capsules and speed-breeding centers. Whereas the equipment required to control cultivation environments (e.g., large growth chambers and greenhouses) is usually expensive, speed-breeding capsules, which consist of shipping containers that have been equipped with hydroponic systems, lighting, air conditioning, and greenhouse benches, can be used as more cost-effective generation acceleration facilities. Meanwhile, speed-breeding centers are used for the intensive improvement of target crops and include both generation acceleration facilities and the integration of the technology needed for genomics-assisted breeding. The construction of such facilities and centers is considered important for effective genomics-assisted breeding technologies in minor and pseudo-cereals.

Both marker selection and efficient phenotyping are important factors for the acceleration of breeding. This is because selection accuracy and selection intensity can be increased by improving the efficiency of trait phenotyping (Araus et al. 2018). In genomics-assisted breeding, it is likely that the improvement of trait phenotyping would improve the power of QTL mapping and GWAS, as well as the prediction accuracy of GS. Furthermore, because high-throughput phenotyping technologies, such as drone remote sensing, are generally versatile and applicable to a wide variety of crop species, it should be easy to apply the technologies developed and used in major crops directly to minor and pseudo-cereals. New technologies for phenotyping enable us to perform QTL mapping/GWAS-based breeding, or GS for characteristics that have been difficult to evaluate so far (e.g., Sivasakthi et al. 2018, Spindel et al. 2018). High-throughput phenotyping will be indispensable in genomics-assisted breeding in the future.

Genome editing, which is one of the most advanced technologies to be developed in recent years, is also likely to be useful for crop development in the near future. The present review focused on the genomics-assisted breeding of minor and pseudo-cereals, mainly based on marker selection. The genetic improvement of these crops can be further accelerated by the use of genome editing, in combination with QTL mapping, GWAS, and GS. In minor and pseudo-cereals, an ancestral state may still remain in agronomic traits (e.g., shattering habit) that have been improved in major crops. Although it is possible to utilize the natural variations present in minor and pseudo-cereals for trait improvement, it might also be possible to edit important genes using genome editing (Osterberg et al. 2017, Varshney et al. 2018). In such cases, biparental QTL mapping and GWAS may play important roles in narrowing down candidate genes.

The advantage of developing breeding strategies for minor and pseudo-cereals is that there is no need to recreate the methodological research required for the advancement of major crops. During the last decade, for example, it has become unnecessary to conduct DNA marker development in order to conduct marker-assisted breeding, and a large number of markers are currently available without special development. In addition, the analysis of whole genome sequences, which has been carried out in global collaborative research, can also be carried out by one research team in some cases. Furthermore, the pros and cons of each method have become apparent in major crop studies. As Bernardo (2016) states, researchers and breeders should choose “waves” to ride on as “surfers”, and the researchers and breeders of minor and pseudo-cereals can assess the potential of “waves” that have previously influenced research and breeding in major crops. This is a major advantage of minor cereal- and pseudocereal-breeding programs that implementing genomics-assisted breeding later than it has been performed in major crops. Currently, the genomic information is available and methodologies are mostly refined, which is owing to the trial-and-error that major crop breeding has experienced over a long period, which means genetic improvements in minor and pseudo-cereals can be realized more expeditiously than they have previously been in major crops.

**Author Contribution Statement**

S.Y. and H.I. contributed to the planning the work, to the writing of the manuscript, and to designing the figure.

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