Alternative Modes of Introgression-Mediated Selection Shaped Crop Adaptation to Novel Climates

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Abstract

Recent plant genomic studies provide fine-grained details on the evolutionary consequences of adaptive introgression during crop domestication. Modern genomic approaches and analytical methods now make it possible to better separate the introgression signal from the demographic signal thus providing a more comprehensive and complex picture of the role of introgression in local adaptation. Adaptive introgression has been fundamental for crop expansion and has involved complex patterns of gene flow. In addition to providing new and more favorable alleles of large effect, introgression during the early stages of domestication also increased allelic diversity at adaptive loci. Previous studies have largely underestimated the effect of such increased diversity following introgression. Recent genomic studies in wheat, potato, maize, grapevine, and ryegrass show that introgression of multiple genes, of as yet unknown effect, increased the effectiveness of purifying selection, and promoted disruptive or fluctuating selection in early cultivars and landraces. Historical selection processes associated with introgression from crop wild relatives provide an instructive analog for adaptation to current climate change and offer new avenues for crop breeding research that are expected to be instrumental for strengthening food security in the coming years.

Key words: crop wild relatives, purifying selection, fluctuating selection, hybridization, polygenic adaptation, climate change.

Significance

Climate change threats on crops are increasing while the genetic base of crops is shrinking. This, coupled with increasing global demand for food, compels us to investigate the genetic diversity associated with adaptation and productivity in unfavorable climates. Results from recent studies show that some crops have benefited from increased genetic diversity following introgression with wild relatives. Such introgression compensated for the effect of unfavorable variants fixed during domestication through various modes of selection on introgressed variants. Characterizing how selection has impacted introgressed genomic variation may show the way forward for adapting current crops to climate change.

The Adaptive Significance of Introgression

The concept of Introgression (see Glossary) was originally coined by Anderson and Hubricht (1938). In their seminal paper, introgression was already defined as a pervasive process with a marked adaptive significance. The theoretical framework illustrating the evolutionary significance of introgression was developed 70 years later (Barton 2001). Since the publication of this theoretical paper, and thereafter in empirical studies, the adaptive significance has been said to be mainly driven by the incorporation of favorable mutations that are rare in the initial stages of introgression. A deeper understanding of its adaptive value led to a consistent increase in publications aiming to identify the adaptive component of introgression in multiple plant groups (Suarez-Gonzalez, Lexer, et al. 2018), a trend that continues today in the genomic era (e.g., Menon et al.)
hybridization and repeated backcrossing. The purpose of introgression in crop breeding is to incorporate alleles from one species to the gene pool of another.

- **Linkage drag**—Undesirable effect of nontargeted QTLs in linkage disequilibrium with the QTLs of interest that are being introgressed.

- **Mutation/genetic load**—The decrease in fitness of the average individual in a population relative to the fittest genotype due to the presence of deleterious genes in the gene pool.

- **Purifying selection**—Selection against deleterious alleles.

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**Glossary**

- **Disruptive selection**—A mode of natural selection in which extreme values of a trait are favored over intermediate values. Under disruptive selection (also called diversifying selection), the variance of the trait increases and the population splits into two distinct groups.

- **Fluctuating selection**—A mode of natural selection characterized by the fluctuation of the direction of selection on a given phenotype over a relatively brief period of evolutionary time (e.g., across years).

- **Hybridization**—The process of interbreeding between two dissimilar genetic entities (taxa, varieties, or lines) to produce a hybrid with new allele combinations.

- **Incomplete lineage sorting**—A phenomenon in population genetics in which gene copies do not coalesce (looking backwards in time) into a common ancestral copy until beyond previous speciation events. This phenomenon causes discordance between the gene tree and the population or species tree. Incomplete lineage sorting is also termed as deep coalescence or retention of ancestral polymorphisms.

- **Introgression**—The transfer of genetic variation between species through hybridization and repeated backcrossing. The purpose of introgression from CWR has the potential to increase the genome-wide genetic diversity which has several adaptive consequences: (1) it can promote new allelic combinations that are favorable for particular environmental conditions, (2) it can improve the efficiency of purifying selection to eliminate deleterious alleles, and (3) it can also promote disruptive or fluctuating selection in geographical areas with a spatially or temporally highly variable climate.

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- **Purifying selection**—Selection against deleterious alleles.
Intraspecific diversity: Species have a wide intraspecific diversity that sometimes spans wide geographic regions. It is relatively common to find diverse adaptive introgression events in different parts of the geographic range of the study species (Blanco-Pastor, Manel, et al. 2019; Zhou et al. 2020; Lovell et al. 2021). This means that a broad intraspecific sampling may be required to uncover a complete picture of introgression.

2. Intraspecific diversity: Species have a wide intraspecific diversity that sometimes spans wide geographic regions. It is relatively common to find diverse adaptive introgression events in different parts of the geographic range of the study species (Blanco-Pastor, Manel, et al. 2019; Zhou et al. 2020; Lovell et al. 2021). This means that a broad intraspecific sampling may be required to uncover a complete picture of introgression.

3. Genetic structure analyses and genome-wide introgressions: Genetic clusters should be used as operational units to identify gene flow at the intraspecific level, but neutral and putative adaptive genetic structure usually differ. These differences can be observed at the chromosome level with different signature of selection in different individuals from the same gene pool or genetic cluster (e.g., Freitas et al. 2021; Lovell et al. 2021). Local ancestry blocks across the different chromosomes can be inferred using sophisticated probabilistic hidden Markov models such as those implemented in HAPMIX (Price et al. 2009) or Ancestry_HMM (Corbett-Detig and Nielsen 2017).

4. Confounding signals:
   - The adaptive hybridization/introgression signal can be easily confounded with incomplete lineage sorting (ILS; Pamilo and Nei 1988; Doyle 1992; Maddison 1997; Rosenberg and Nordborg 2002; Blanco-Pastor et al. 2011; Tang et al. 2022). Analyses that distinguish both patterns (such as the four taxon $f_4$ [Patterson et al. 2012] or $f_d$ [Martin et al. 2015] statistics or the five-taxon $D_{FST}$ [Pease and Hahn 2015] statistic, which are all related to the genome-wide summary ABBA-BABA test [Green et al. 2010]) are essential tools for detecting introgression. Like the local ancestry block identification mentioned above, we could also perform these introgression analyses across the genome in sliding windows to detect the origin of the introgressed genomic blocks. With $f_4$ or $f_d$ values across the genome, it is also possible to calculate, for each introgressed individual, the genome-wide proportion of genomic introgression coming from each of the potential donor species (Zhou et al. 2020).

5. Investigating the timing of introgression: The temporal pattern of introgression among potential donors can be calculated by the size of the introgression tracts also using hidden Markov models (Guan 2014). This can be achieved under the expectation that tract size negatively correlates with the age of introgression assuming that recombination events keep breaking off donor haplotypes after the initial hybridization (Zhou et al. 2020). Recent extensions to the hidden Markov model of Corbett-Detig and Nielsen (Ancestry_HMM) (Corbett-Detig and Nielsen 2017; Medina et al. 2018) allow to directly infer the timing of the introgression events.

6. Identifying the selective value of introgressed regions: Putative selection signals can be identified in introgressed regions through the use of $F_{ST}$ and $d_{xy}$ scans, the XP-CLR statistic (Chen et al. 2010) or more sophisticated models (Günther and Coop 2013) in sliding windows (e.g., He et al. 2019). Other possibilities include checking the overlap between introgressed regions and significant GWAS loci (e.g., Cheng et al. 2019). To check for the effect of introgressed regions on polygenic traits one option is to partition the phenotypic variation for a trait into components explained by introgressed and nonintrogressed regions using, for example, the GCTA-GREML method (Yang et al. 2011). It is also possible to investigate if introgression has influenced nucleotide diversity patterns, this can be done by calculating nucleotide diversity values in the same sliding windows that were previously associated with introgression by $f_{ST}$-type statistics. Higher allelic diversity in introgressed regions could point to an effect of either disruptive or fluctuating selection on adaptive alleles (see also Box 2).
Box 2: Effect of (1) mean climate across years and (2) interannual variability (std) of climate on natural selection processes

Theoretically, under climate-driven directional selection, the allelic diversity in adaptive genes (HeA) is lowest at the two extremes of the climatic gradient. This is because the selective pressure on a particular allele is greatest at the climatic extremes. On the other hand, under fluctuating selection, the HeA reaches the lowest values at the end of the gradient representing the lowest climatic variability and reaches the highest value at the end of the gradient representing the highest climatic variability. This is because high adaptive diversity is required to cope with changing climatic conditions. In the latter case, the selective pressure alternates between the two different alleles over a short period of evolutionary time (see figure). Introgression can play an important role for selection processes occurring across gradients of average climate by introducing newly adaptive alleles previously nonexistent in the population. Such beneficial alleles will increase in frequency in the population by means of selective sweeps or shifts, thus reducing their heterozygosity levels. These climatic gradients have been mainly associated with mean values of climatic parameters (e.g., latitudinal, longitudinal, or altitudinal gradients of mean [daily, monthly, seasonal, yearly, and cross-years] temperature and/or precipitation values), for which numerous examples exist in the literature (e.g., Lasky et al. 2017; Exposito-Alonso et al. 2018; Blanco-Pastor et al. 2021). Another type of climatic gradients is associated with the temporal variability of the climate. Climatic conditions at a given site can fluctuate over different time scales, that is, interseasonal variations or relatively stochastic interannual variations. In this case, the probability of fixation of adaptive alleles will be lower as high heterozygosity would be preferred under such conditions. A good example that associates the diversity of adaptive alleles for a particular trait with the two types of gradients can be found in Keep, Rouet et al. (2021). These authors showed that investment in sexual reproduction during the first year (heading first year, HFY) in perennial ryegrass was partially associated with a gradient of mean summer maximum daily temperature values (SMDT), but the diversity of HFY-associated adaptive loci (which was highest in a geographic region with a signature of introgression, see Fig. 2) was better predicted by a linear combination of both mean SMDT values and the interannual variation on precipitation during the wet season, the latter being the most significant environmental predictor in the model. Perennial ryegrass does not usually flower during the first year and invest in vegetative tissues only, which is an adaptive advantage in optimal rainfall conditions. But first-year flowering is an adaptive strategy in years of low rainfall, given that greater investment in seed formation provides greater fitness under a drought escape strategy (Volaire 2018; Blanco-Pastor et al. 2021), a pattern that was also found at the continental level (Keep, Sampoux et al. 2021).
of domestication and that this introgression facilitated adaptation to novel conditions, particularly during their geographic expansion. The latest genomic studies now provide further details about the evolutionary consequences of adaptive introgression, revealing its pervasiveness, the number of donor species, its geographical scale, the mode of selection that prevails, as well as the genes or set of genes that are subject to selection. Below I present a summary of findings in wheat, maize, potato, grapevine, and forage grasses. In conjunction, these studies reveal a more complex picture of the evolutionary dynamics during crop domestication and highlight the role of introgression in local adaptation through a variety of selection processes beyond directional selection on certain introgressed alleles.

Genomics of Introgressed Crops

Wheat

Bread wheat (*Triticum aestivum*; AABBDD) accounts for 17% of the world’s total cultivated area and is one of the most indispensable staple crops for humans. It originated from hybridization between cultivated tetraploid emmer wheat (*T. turgidum*, AABB) and wild diploid *Aegilops tauschii* (DD) (Marcussen et al. 2014; fig. 1a). The origin of the domestication of bread wheat goes back 10,000 years (Tanno and Willcox 2006). Recent studies have analyzed the role of introgression with wild relatives in shaping the current genetic diversity of the bread wheat genomes (Cheng et al. 2019; He et al. 2019; Zhou et al. 2020). These studies show that cultivated bread wheat had a
single domestication event (Cheng et al. 2019). A polyploidization-related bottleneck caused a substantial loss of genetic diversity in the A, B, and D subgenomes of hexaploid wheat compared with its diploid ancestor, with the loss being much more pronounced in the D subgenome (fig. 1b; Cheng et al. 2019). One explanation for the lack of a similar reduction in diversity in the A and B genomes, with respect to the D genome, is gene flow from different tetraploid wild wheat taxa since the earliest stages of domestication (He et al. 2019; Zhou et al. 2020).

Introgression from bread wheat close relatives contributes to a substantial portion (4–32%) of its genome (Zhou et al. 2020). Initially, this introgression (which explained the relatively high diversity of A and B subgenomes in bread wheat) was thought to come only from wild emmer wheat (Cheng et al. 2019; He et al. 2019). However, introgression from wild emmer wheat constitute only a small proportion of the whole history of gene flow, is gene flow from different tetraploid wild wheat taxa since the earliest stages of domestication (He et al. 2019; Zhou et al. 2020).

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He et al. (2019) showed that gene flow from bread wheat close relatives was associated with an increase in effective population size that improved the efficiency of purifying selection in the AABB subgenome and contributed to a reduced mutation load. Highly recombinant regions also showed signatures of directional selection in this polyploid subgenome, with 8% of locally adaptive alleles introduced by introgression. Rare alleles (derived allele frequency, DAF < 0.1) introduced from the third donor, the wild emmer wheat, explained a substantial proportion of the phenotypic variance for the traits crop weight (up to 30.9%), drought susceptibility (up to 22.5%), and plant height (up to 35%). These rare alleles promoted adaptation in specific geographically restricted habitats (He et al. 2019) following a geographical pattern of introgression and adaptation that can largely be explained by the sympatric distribution of landraces with their donors. Emmer wheat and related tetraploids (donors of the AB subgenomes) are native to the Eastern Mediterranean and West Asia, and A. tauschii ssp. stranulata (introgression donor of the D subgenome) is native to the southwestern Caspian Sea, both places with high constraints in relation to summer drought and where the introgression signal is strongest for the AB and D subgenomes, respectively.

**Potato**

The potato (Solanum tuberosum L.) was domesticated 8,000–10,000 years ago from its wild progenitor native to the Peruvian Andes. Autopolyploidization of early diploid...
ancient $S.\, tuberosum$ cultivars (Stenotomum and Phureja groups) produced tetraploids cultivated in the Andes ($S.\, tuberosum$ Andigena group). The potato was subsequently cultivated in equatorial high altitude, and later on in southern latitudes with longer summer days (Spooner et al. 2005). Migration from the Andes to coastal Chile resulted in a subspecific group adapted to long-day conditions ($S.\, tuberosum$ Chilotanum group) that has provided much of the genetic background for commercial cultivars of $S.\, tuberosum$ worldwide (Spooner et al. 2005; Hardigan et al. 2017). The genetic diversity of diploid and tetraploid $S.\, tuberosum$ is extremely high, due to the wild introgressions following autoploidy and range expansion that captured alleles outside their geographical origin (fig. 1c; Hardigan et al. 2017). A recent study on the potato pan-genome shows that both ILS and hybridization/introgression have contributed to blurring evolutionary relationships between wild potato species and landraces (Tang et al. 2022).

Several wild Solanum species (most notably the Andean $S.\, microdontum$) had a role in the spread of cultivated potatoes across the globe by providing tolerance to long-day conditions (Hardigan et al. 2017). The first European potatoes collected during the period 1650–1750 were closely related to Andean landraces and probably not sufficiently adapted to European conditions. These genotypes were crossed with Chilean genotypes preadapted to long-day requirements like those in Europe. The European varieties became progressively more similar to the Chilean genotypes as they became increasingly distant from their Andean counterparts which paved their adaptation to European climatic conditions (fig. 1d; Gutaker et al. 2019). Analysis of gene functions in introgressed regions supports their adaptive role with enrichment in multiple abiotic stress-associated genes with unknown specific functions and retention of wild alleles in most (70%) tetraploids (Hardigan et al. 2017).

Maize

Maize ($Z.\, mays$ ssp. $mays$) was domesticated in southwestern Mexico approximately 9,000 years ago from the wild teosinte $Z.\, mays$ ssp. $parviglumis$ (Piperno et al. 2009). Gene flow among multiple teosinte species has occurred during Zea diversification (Ross-Ibarra et al. 2009). Introgression from the wild teosinte taxon $Z.\, mays$ ssp. mexicana (hereafter mexicana) has generated a particular interest due to its adaptive significance to highland areas (Hufford et al. 2013; Wang et al. 2017; Gonzalez-Segovia et al. 2019; Calfee et al. 2021). The subspecies mexicana is endemic to the highland regions of Mexico (1,300–300 m a.s.l.) and diverged from the subspecies parviglumis about 60,000 years ago (Ross-Ibarra et al. 2009). Highland regions of central America present several environmental challenges, including a cooler, drier climate with higher UV intensity and a shorter growing season requiring earlier flowering. The ancestry of mexicana subspecies in upland maize comes from an ancestral introgression that has maintained much of its diversity in the various upland maize populations of Mexico (Calfee et al. 2021). After the introgression from mexicana, introduction of introgressed maize has been documented in Guatemala and the southwestern USA (Wang et al. 2017).

Large genomic regions (at megabase scale) introgressed from mexicana and associated with direction selection have been recently identified on chromosomes 3, 4, and 9 (Hufford et al. 2013; Wang et al. 2017; Gonzalez-Segovia et al. 2019). The introgressed region on chromosome 3 corresponds to a chromosomal inversion associated with flowering time. The region on chromosome 4 contains the $Inv4m$ region, a large 14 Mb inversion (Calfee et al. 2021) which overlaps with QTLs for leaf pigmentation and macrohairs (Lauter et al. 2004), and is associated with adaptation to cold stress and increased maize yield at high elevations (Crow et al. 2020). The region on chromosome 9 overlaps with the locus macrohairless1 ($mhl1$), a large-effect locus also linked to the production of macrohairs (Lauter et al. 2004; Moose et al. 2004; Gonzalez-Segovia et al. 2019). Despite the documented introduction of introgressed maize into the southwestern USA, these populations have lost the signature of introgression in chromosome 4, a possible signal of geographically restricted local adaptation (Wang et al. 2017).

Introgression-affected genomic regions in upland maize showed higher recombination rates (Calfee et al. 2021), and a significantly lower number of deleterious single nucleotide polymorphisms (Wang et al. 2017). This supports an improved effect of purifying selection driven by introgression in upland maize, similarly as shown for wheat (He et al. 2019; see above). It also needs to be noted that most of the outlier loci with high introgression from mexicana do not show signs of reduced genetic diversity by selective sweeps (Calfee et al. 2021). This can be explained by an effect of either disruptive or fluctuating selection (see Box 2) over multiple mexicana haplotypes in those introgressed regions. This maintenance of high haplotype diversity is consistent with the selective pressures caused by microhabitat heterogeneity or interannual variability of highland climates (Blanco-Pastor, Fernández-Mazuecos, et al. 2019). However, pronounced allelic clines are observed with elevation such as in the $Inv4m$ and $mhl1$ loci (Calfee et al. 2021). Altogether, both high haplotype diversity and the presence of specific alleles in key genes under directional selection may be at play in the adaptation of upland maize.

Upland maize from central America flowers earlier than lowland maize. The large introgressed region in upland maize on chromosome 3 corresponds to a chromosomal
inversion associated with flowering time (Wang et al. 2017), but no genes from the core maize flowering time pathway overlap with mexicana introgressed loci (Calfee et al. 2021). This is because not all genes causing flowering time differences in maize are fully annotated, and also because flowering time is a highly polygenic trait in maize (Buckler et al. 2009), which may reduce the strength of selection on most individual genes under detectable levels using standard approaches. These results suggest that many more mexicana genes are beneficial at higher altitudes, but only in conjunction and when present in specific allele combinations.

**Grapevine**

The cultivated grapevine, *Vitis vinifera* ssp. *vinifera*, was domesticated in the southern Caucasus 8,000 years ago from the dioecious taxon *V. vinifera* ssp. *sylvestris* (This et al. 2006). It was initially thought that grapevine domestication included two separate domestication events, one derived from wild Caucasian grapevines and the other from wild Western European grapevines (Grassi et al. 2003; Arroyo-García et al. 2006). However, recent genomic data have confirmed a single origin in West Asia with a strong introgression signal from the subspecies sylvestris as it moved toward the western Mediterranean (Terral et al. 2010; Myles et al. 2011; Bouby et al. 2013; Freitas et al. 2021), which lead to a reduction in its genetic load (Zhou et al. 2017; Freitas et al. 2021). One of the first and strongest evidence of introgression was the increase in genetic diversity toward the west, contrary to expectations given the east-to-west direction of domestication.

Environmental adaptation played a key role in both the evolution of wild genotypes and the outcome of introgression with cultivated varieties. Freitas et al. (2021) found significantly enriched climate- and pathogenesis-associated loci in introgressed genomic regions of Iberian cultivated grapevines. The proportion of introgressed regions ranged from 25% to 50% of the grapevine genome, in a set of regrapevines. The proportion of introgressed regions ranged from 25% to 50% of the grapevine genome, in a set of regrapevines. The proportion of introgressed regions ranged from 25% to 50% of the grapevine genome, in a set of regrapevines.

Genomic data of archaeological samples further support gene flow from wild to cultivated grapevines in the western Mediterranean occurring in the early stages of viticulture during the Iron Age (510–475 BC; c.a. 2,500 years ago; Ramos-Madrigal et al. 2019). This coincides with the earliest evidence of vine cultivation in the Iberian Peninsula 2,900 years ago under Phoenician influence (Iriarte-Chiapusso et al. 2017), making it the earliest tentative time of a human-mediated grapevine introgression event in this region.

**Ryegrass**

Perennial ryegrass (*Lolium perenne*) is among the most used grasses for cover crops and forage crops in temperate regions of the world and the most prevalent species in European grasslands grazed by cattle. This species has been recently adopted as a model species for the genetics of temperate forage grasses (Byrne et al. 2015; Blanco-Pastor, Manel, et al. 2019; Blanco-Pastor et al. 2021). *L. perenne* received alleles from the close relatives *L. rigidum* and *L. multiflorum* since >380 kya after initial divergence (Blanco-Pastor, Manel, et al. 2019). Introgression from *L. rigidum* to *L. perenne* has taken place in a genetic cluster located in southeastern Europe and near East regions where the two species coexist. In this region, 15% of alleles present in *L. perenne* originated in *L. rigidum*. Gene flow from *L. multiflorum* to *L. perenne* has taken place in northern Italy. In this region, 5% of alleles present in *L. perenne* originated in *L. multiflorum* since >380 kya after initial divergence (Blanco-Pastor, Manel, et al. 2019). Interestingly, North Italian populations have been recently used as a source for the generation of new ryegrass varieties with increased productivity and water-soluble carbohydrate content (Faville et al. 2004).

In these two regions, Keep, Rouet et al. (2021) found significant relationships between the genetic diversity at major QTLs (adaptive loci) (HeA, sensu Keep, Rouet et al. 2021) and both the mean values of the local climate and its temporal variability (Box 2). Keep, Rouet et al. (2021) modelled the HeA index for several adaptive traits from mean climate and climate variability indicators. For several traits such as heading in the first year (HeA_HFY), spring canopy height (HeA_SPH), and heading date (HeA_HDT), the most significant explanatory variable was a climate variability indicator (standard deviation, std). The region with the highest heterozygosity of genes associated with reproductive investment during the first year (HeA_HFY) was found precisely in northern Italy, a region with high variability in spring rainfall and with strong genomic signature of introgression from the relative *L. multiflorum*, suggesting that fluctuating selection has favored maintenance of introgressed genetic variation in this geographical region (fig. 2, see also Box 2).
Concluding Remarks

The studies reviewed here show that genomic data broaden our understanding of the history of crop domestication and gene flow with their wild or early domesticated relatives. Previous studies showed that the genetic basis of crops was produced through domestication events followed by adaptive gene flow from relatives encountered during post-domestication expansion processes (Janzen et al. 2019). New genomic data reveal that introgressed loci of large effect that are target of directional selection explain a small proportion of the full phenotypic variance in traits associated with altitudinal, latitudinal, and longitudinal climatic gradients in different crop species (phenotypic variance associated with cold stress [maize; Crow et al. 2020], drought stress [wheat, grapevine, and ryegrass; He et al. 2019; Blanco-Pastor et al. 2021; Freitas et al. 2021] and day length [potato] differences [Hardigan et al. 2017]). These studies also show that the evolutionary consequences of introgression from wild relatives are also far more diverse than previously identified (Martin and Jiggins 2017; Warburton et al. 2017; Suarez-Gonzalez, Lexer, et al. 2018; Janzen et al. 2019). Introgression between populations and species can provide new beneficial alleles or allele combinations but can also increase the intrapopulation/intravariety genome-wide diversity, thereby enhancing the effect of purifying (wheat, maize, and grapevine), and disruptive or fluctuating selection (maize and ryegrass).

All cases reviewed here reflect that introgression is local or regional in scope, but that in some cases new introgressed alleles spread more widely (e.g., maize or potato). In all cases, adaptive introgression was clearly linked to specific environmental conditions (eastern Mediterranean and west-central Asia for bread wheat, European region for potato, western Mediterranean for grapevine, and northern Italy for ryegrass). These alleles represent a promising source for further introgressions that could be used for adapting these crops to novel stressful climatic conditions, in these same regions or elsewhere (Dempewolf et al. 2014; Brozynska et al. 2016; Zhang and Batley 2020).

From the earliest stages of crop expansion, farmers moved from place to place, selecting and carrying with them the seeds best adapted to the new local conditions, and eventually crossing these genotypes with local wild species or other varieties. This contributed to the adaptation of crops to new climates and led to the formation of landraces. These landraces, most of which disappeared or have been relegated to germplasm banks, have already contributed to crop improvement in multiple traits, such as increased efficiency in nutrient uptake, protein and mineral content, disease resistance, and abiotic stress tolerance (see Carvalho et al. 2011 for a review), and should be considered for breeding modern varieties with higher priority than wild populations, given their more advanced state of domestication.

Past adaptation to newly colonized areas through introgression in species reviewed here provide an instructive analog for adaptation to current climate change. The genomics of adaptation to climate in the past provides new avenues for adapting crops to future climatic constraints, as well as for improving crop productivity under variable climatic conditions. Although it remains important to focus on identifying the substitution of high-effect alleles for breeding, the way forward in research on adaptive introgression from wild to crop species should also address the more subtle effects of diversity gain following introgression. It is long known that increasing intraspecific diversity is a viable strategy to increase yield and/or yield stability, specially under the effect of biotic or abiotic stressors (Frankel 1939; Allard 1961), but this issue has only recently received especial attention in the context of the agricultural revolution under increasing climate variability (Newton et al. 1997, 2009; Prieto et al. 2015; Reiss and Drinkwater 2018; Litrico and Huyghe 2019).

The recent studies reviewed above raise new outstanding questions, for example: Is purifying selection a common selection mode in crops after introgression with wild relatives? Is it an important element in the elimination of deleterious alleles arising from the mutation load associated with the cost of domestication? Does domestication eliminate alleles that have been subject to fluctuating selection in regions with variable climates, thus favoring phenotypic stability? Can we recover these alleles from wild relatives to favor crop adaptation in regions with increasing climatic instability? How important are traits with polygenic architectures, both additive and nonadditive (epistasis), for climate adaptation in crops? How much of this adaptive genomic variation is present in cultivated varieties and how much in wild populations? What are the best breeding approaches to maximize adaptive diversity while avoiding the negative effects of linkage drag associated with introgression from wild relatives?

We note a general pattern in the studies presented here: geographic expansion of crops was driven primarily by an increase in genetic diversity following introgression with wild relatives that offset the cost of domestication. To a large extent, the success of certain modern cultivars is due to the interaction between multiple introgressed alleles from wild relatives, rather than the introduction of specific alleles with a large effect. The effect of polygenic architectures could have been significant in relation to increasing the efficiency of balancing and disruptive/fluctuating selection but the detailed effect of increased adaptive diversity on the evolution of polygenic traits, remains to be fully determined. Re-analysis of genomic data using models that account for the small but significantly coordinated variation of multiple small-effect alleles involved in phenotypic determination under additive (Zhou et al. 2013; Berg and Cooper 2014; Barghi et al. 2020) and nonadditive
(Exposito-Alonso et al. 2020) polygenic architectures will show the way forward for the introgression-mediated breeding of new, more diverse, and climate-resilient crops.

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