

Annual Review of Plant Biology Crop Biodiversity: An Unfinished Magnum Opus of Nature

Matthew B. Hufford,¹ Jorge C. Berny Mier y Teran,² and Paul Gepts²

¹Department of Ecology, Evolution, and Organismal Biology, Iowa State University, Ames, Iowa 50011–1020, USA; email: mhufford@iastate.edu

2Department of Plant Sciences, University of California, Davis, California 95616–8780, USA; email: [jcberny@ucdavis.edu,](mailto:jcberny@ucdavis.edu) plgepts@ucdavis.edu

Annu. Rev. Plant Biol. 2019. 70:727–51

The *Annual Review of Plant Biology* is online at plant.annualreviews.org

[https://doi.org/10.1146/annurev-arplant-042817-](https://doi.org/10.1146/annurev-arplant-042817-040240) 040240

Copyright © 2019 by Annual Reviews. All rights reserved

ANNUAL CONNECT

- www.annualreviews.org
- · Download figures
- Navigate cited references
- Keyword search
- · Explore related articles
- · Share via email or social media

Keywords

farmer selection, gene flow, genetic diversity, parallel and convergent evolution, phenotypic plasticity, seed systems

Abstract

Crop biodiversity is one of the major inventions of humanity through the process of domestication. It is also an essential resource for crop improvement to adapt agriculture to ever-changing conditions like global climate change and consumer preferences. Domestication and the subsequent evolution under cultivation have profoundly shaped the genetic architecture of this biodiversity. In this review, we highlight recent advances in our understanding of crop biodiversity. Topics include the reduction of genetic diversity during domestication and counteracting factors, a discussion of the relationship between parallel phenotypic and genotypic evolution, the role of plasticity in genotype \times environment interactions, and the important role subsistence farmers play in actively maintaining crop biodiversity and in participatory breeding. Linking genotype and phenotype remains the holy grail of crop biodiversity studies.

1. INTRODUCTION

Some of the most colorful sights around the world are vegetable markets and the stalls of grocery stores. The grains, vegetables, fruits, and spices of innumerable colors, sizes, shapes, and aromas offer a kaleidoscopic view of—mostly domesticated—crop biodiversity (**Figure 1)**.

Figure 1

The kaleidoscope of fruits and vegetables sold in this market are the result of millennia of farmer selection during and after domestication around the world. Crop biodiversity is represented by both different crops and varieties within crops. From **<https://www.publicdomainpictures.net>**, license CC0 Public Domain.

Underlying these everyday, down-to-earth sights are important considerations about diversity. First, what qualifies as crop biodiversity, and which species fit this definition? We define crop plants here as any plant being cultivated, mainly those that have been or are being domesticated. Of the 390,000 to 420,000 plant species that are estimated to exist [\(76,](#page-20-0) [126\)](#page-22-0), some 2,500 species may have been domesticated [\(30,](#page-18-0) [59\)](#page-19-0). This is a fairly generous estimate, which encompasses not only domesticated species but also wild species that are cultivated in some way. We must consider the nature and degree of domestication of these species [\(28\)](#page-18-0), the extensiveness of their distribution, and the level of their production. For example, some 80% of plant-derived foods originate in just 17 botanical families (of a total of 416 families); several of these are major contributors of domesticated species, including the Poaceae, Fabaceae, Apiaceae, Brassicaceae, Rosaceae, Arecaceae, and Zingiberaceae. About 100 species contribute 90% of all calories in the human diet [\(76\)](#page-20-0); however, other domesticated species fulfill important functions as sources of essential nutrients, such as proteins, vitamins, and minerals and as medicines. This collective crop biodiversity can be considered at several levels: (*a*) populations or lines within species, (*b*) assemblages of crops in time and space (i.e., agroecosystems), and (*c*) several agroecosystems composing an agricultural landscape. All organisms populating agroecosystems and agricultural landscapes can be considered components of agrobiodiversity, including crop and noncrop plants, insects, and microorganisms, both beneficial and detrimental [\(22,](#page-17-0) [68\)](#page-19-0). In this review, we specifically focus on domesticated crop species, regardless of their degree of domestication, and their wild relatives, especially their progenitors. Genetic diversity within these species is not only the raw material for plant breeding, but also a key factor in assuring the productivity and stability of agriculture (e.g., [8,](#page-17-0) [89,](#page-20-0) [133, 136\)](#page-22-0).

Second, how is crop biodiversity produced? Crop biodiversity is the result of a triad of interacting factors, which include the plant itself, its environment, and the actions of humans, as illustrated by the domestication triangle (**Figure 2**). Whereas natural evolution is confined to organismal characteristics and interactions with the environment, crop evolution involves the allimportant contributions of humans, who are responsible for selection and dissemination of crop plants, not only in the first stages of transition from gathering to cultivation but in all subsequent stages as well. Today, this process includes both modern plant breeding and management of seed stocks by subsistence farmers, activities that continue to adapt crops to the needs of their respective users. Thus, our domesticated crop plants should be considered not merely as biological organisms, each with their genetic, life history, and production characteristics, but more significantly as biocultural artifacts. We use the word artifact in the anthropological sense, namely as a product of human ingenuity and skill within their cultural contexts. Crops are one of the most important inventions of humanity, and humans and crops are now symbiotically related because of their mutual dependence for survival. Seven billion humans cannot be fed without agriculture because plants contribute 80% of our food; conversely, humans assure the survival of some of our major crops, given that domestication has eliminated their ability to survive on their own. Thus, crop cultivation represents a case of mutually assured survival of two classes of organisms further underscoring the need to understand selection during domestication and contemporary plant breeding and the continued involvement of farmers in shaping crop biodiversity to fit their own uses.

Third, to what extent has evolution during domestication shaped crop biodiversity? The evolutionary processes that shaped diversity, regardless of their nature and origin (see the previous paragraph), have operated from before the initiation of cultivation of wild progenitors some 10,000– 12,000 years ago, through the initial selection for adaptation to cultivation and subsequent worldwide dissemination [\(77\)](#page-20-0), up until today. For example, the general reduction in diversity of crops, especially observable at the molecular level, traces back to the domestication process, or in some cases even before this process. Understanding the contemporary biodiversity of crops, therefore, requires an analysis of its evolutionary history [\(50,](#page-19-0) [46, 47,](#page-18-0) [74, 85,](#page-20-0) [98, 106–108, 115,](#page-21-0) [132\)](#page-22-0). Studies **Domestication:** the process of heritable genetic adaptation to human cultivation and consumption conditions

Figure 2

Domestication and subsequent crop evolution involve interactions of plant, environmental, and human factors. All three factors are required for domestication to generally take place, but specifics vary among domestication regions. The figure lists examples of specific factors that have been implicated in domestication. Modified from Reference [43](#page-18-0) with permission.

of crop evolution are very active not only because the transition from hunting-gathering lifestyles to those of farming is one of the most important evolutionary steps of humanity but also because wild relatives represent a significant aspect of sound conservation due to their utility in breeding by cross hybridization.

The importance of crop biodiversity will rise in the near future because of additional qualitative or quantitative demands on agricultural production. The human population is expected to increase from 7.5 billion in 2017 to 9.8 billion in 2050 and 11.2 billion in 2100 [\(166\)](#page-23-0). The increase in demand for agricultural products due to a larger population will be compounded by increased consumption accompanying greater affluence. Because expecting a sharp reduction in the human population is illusory [\(15\)](#page-17-0), humanity is faced with a long-term need for increased agricultural production. There is no single magical solution that will achieve this goal [\(79\)](#page-20-0). However, genetic crop improvement has been responsible for about half of on-farm yield increases [\(154\)](#page-23-0); the other has been contributed by cultivation practices. Furthermore, there is increasing pressure for agricultural production to evolve to a more sustainable model that is resource efficient and reduces environmental impact (e.g., [35,](#page-18-0) [123\)](#page-22-0). There is no doubt that other factors like education, infrastructure, and policy, among others, also play a role, but these are topics beyond the scope of this review.

Biodiversity in general and crop biodiversity, in particular, are threatened. Paradoxically, agricultural activity is the second most prevalent threat in this regard [\(104\)](#page-21-0). Agricultural biodiversity is harmed by habitat destruction through agricultural conversion [\(174\)](#page-24-0); urbanization [\(80\)](#page-20-0); road construction [\(86\)](#page-20-0); increasing global cultural homogenization [\(77\)](#page-20-0); and introduction of modern, elite varieties with limited conservation of traditional varieties (i.e., genetic erosion) [\(168, 169\)](#page-24-0).

Although the existence of human-induced global climate change is not in doubt, the future intensity of this change remains uncertain given various emission profiles of greenhouse gases. Uncertainties relate mainly to the political will, both nationally and internationally, to control emissions. What is certain is that global average temperatures will increase, and rainfall will become more unpredictable and erratic. What is uncertain are the magnitude and interactions of these changes and their effects on Earth and its biota. Climate change leads to shifts in climate zones and biome distributions [\(140,](#page-22-0) [145\)](#page-23-0). In this regard, changes in crop distribution to higher altitudes or latitudes are likely [\(54,](#page-19-0) [152\)](#page-23-0). Mitigation of these changes requires a multifaceted response in which crop biodiversity will play a role by, for example, providing variety or crop substitution or an earlier phenology [\(73, 93,](#page-20-0) [119\)](#page-22-0). Not surprisingly, a current emphasis in biodiversity utilization includes screening for heat and drought tolerance to identify new sources of adaptive genetic diversity to mitigate yield reductions caused by global climate change (e.g., [39\)](#page-18-0).

Additional pressures on crop biodiversity are the current diversification of human diet and crop nutritional quality to improve human well-being, especially that of children [\(2,](#page-17-0) [72\)](#page-20-0). Although this is a complex issue involving mainly sociopolitical and economic factors affecting food security, agricultural biodiversity plays roles through crop species diversification in agroecosystems and biofortification of crops (e.g., [56,](#page-19-0) [112\)](#page-21-0).

In this review, we focus on the genetic diversity of domesticated plants from the perspective of crop evolution and domestication. Understanding this diversity, including how it has been shaped and maintained, is fundamental to all steps of genetic conservation (whether in gene banks or in situ) and the utilization of genetic resources in plant breeding. We concentrate on specific issues that have recently been studied, spanning the full range of genomic studies of crop diversity and gene flow to parallel and convergent evolution to plasticity and ending with the farmers' ongoing roles in shaping their seed stocks. We end with a series of future issues or recommendations regarding the study, management, and utilization of crop biodiversity.

2. ORIGIN OF VARIATION AND THE COST OF DOMESTICATION

2.1. Origin of Variation

During domestication and after domestication, crop biodiversity was shaped by both genomewide demographic effects and locus-specific selection on favorable alleles (**Figure 3**). In most crop-wild relative pairs evaluated to date, reduced genetic diversity is observed in the crop due to what is known as a domestication bottleneck. Ancient cultivators likely focused their efforts on particular wild subpopulations and, through both unconscious and conscious selection on specific alleles, increased the prevalence of desirable traits such as reduced branching and shattering that facilitated harvest [\(31\)](#page-18-0). A focus on a few wild relative subpopulations would result in a subset of genome-wide diversity being passed to domesticates. Continued selection on loci underlying domestication traits would further reduce diversity in these specific chromosomal regions relative to the genomic background. For example, maize retains approximately 83% of genome-wide diversity found in its wild relative teosinte [\(66\)](#page-19-0), whereas near domestication loci, such as *teosinte branched1* (*tb1*), essentially all diversity in maize has been lost [\(160\)](#page-23-0). The same general trend has been observed in many of the world's major crops, including rice [\(18\)](#page-17-0), wheat [\(58\)](#page-19-0), soybean [\(84\)](#page-20-0), and common bean [\(11\)](#page-17-0), with weaker bottlenecks typically observed in perennials relative to annual crops [\(41\)](#page-18-0).

Exceptions to this general pattern have been found. Domesticated carrot, for instance, shows no reduction in overall diversity compared to wild populations potentially due to its outcrossing **Plasticity:** the capacity of a genotype to produce different phenotypes under different environmental conditions

Figure 3

Fate of genetic diversity during and after domestication. The blue line indicates neutrally evolving loci subject to genetic drift. The red line depicts loci subject to selection, leading to either reduction of diversity (or even monomorphism) during (*a*) the initial domestication step and (*b*) subsequent steps. (*c*) An increase of diversity may occur during the domestication process or subsequently during dissemination or breeding for certain genes involved in agronomic adaptation or consumer preference. Adapted from Reference [47.](#page-18-0)

mating system and ongoing history of gene flow with wild relatives [\(69\)](#page-19-0). Likewise, high diversity is found at the *PvTFL1y* locus in common bean, which underlies the determinacy domestication phenotype [\(82\)](#page-20-0). Determinacy in common bean is achieved through loss of function of *PvTFL1y*, a factor that may explain the diversity of putatively causal alleles because a greater proportion of mutations will break a gene than will create a novel function (see more about the *PvTFL1y* locus in Section 4). A more general analysis of genic sequences in common bean identified a higher diversity in nonsynonymous mutations in domesticated types, suggesting an increase in functional diversity selected during and after domestication [\(12\)](#page-17-0) (although an alternative explanation is provided by an insufficient purging of deleterious mutations). Traits that were particularly affected include tolerance of abiotic stress, flowering time, and morphology [\(11, 12\)](#page-17-0).

An additional, important factor determining diversity levels at domestication loci is whether causal alleles are sourced from standing variation in the wild or de novo mutations in protodomesticates. Many domesticated traits are deleterious in the wild. For example, decreased shattering and reduced dormancy are appealing to cultivators, but these characteristics would dramatically reduce survival and reproduction in the wild. Given the loss of fitness conferred by domestication traits, some have speculated that their causal mutations must have occurred de novo in cultivated crops. Others have countered, however, that standing variation has likely played an important role given the long waiting time for novel beneficial mutations and the fact that many domestication

Standing variation:

preexisting genetic variation acted upon domestication or selection postdomestication

De novo mutation:

change in DNA sequence appearing during or after the domestication process (in contrast with standing variation)

traits are polygenic, which would require multiple de novo mutations per trait [\(158\)](#page-23-0). Given the relatively rapid time frame of domestication, it is likely that standing variation has been a source of adaptation [\(9\)](#page-17-0). In fact, the causal polymorphism at *tb1,* for example, which leads to an unbranched phenotype in maize has been shown as a transposon insertion that is segregating in teosinte [\(160\)](#page-23-0). Because standing variation is older and has had more opportunity to recombine onto multiple genomic backgrounds, the selection signature of reduced diversity can be more attenuated (i.e., a soft sweep) than selection on de novo mutations [\(9\)](#page-17-0). Selection scans based on sweep detection in domesticates or increased differentiation between domesticated types and wild relatives may therefore be biased toward identification of selection on de novo mutations. However, methods have recently been developed using approximate Bayesian computation [\(121\)](#page-22-0) and machine learning frameworks [\(144, 147\)](#page-23-0), which appear to be capable of detecting soft sweeps and distinguishing these from hard sweeps. Future application of these methods may help clarify the extent to which domestication relied on standing variation versus de novo mutations.

2.2. Cost of Domestication

The general decline in crop diversity due to demographic bottlenecks and targeted selection during domestication also has implications for the prevalence of deleterious alleles in crop genomes. The reduction in effective population size that is associated with declines in diversity lowers the efficiency of purifying selection, permitting the accumulation of deleterious alleles. In the context of crops, this phenomenon has been termed the cost of domestication (reviewed in [110\)](#page-21-0) and was first proposed by Lu and colleagues [\(95\)](#page-20-0) as an explanation for the excess of deleterious alleles in domesticated rice. The trend of increased genetic load in crops has since been observed in maize [\(176\)](#page-24-0), cassava [\(135\)](#page-22-0), grape [\(182\)](#page-24-0), sunflower [\(137\)](#page-22-0), and other species, suggesting this is a general phenomenon tied to domestication. Several computational methods have been developed to identify deleterious alleles, primarily based on constraint across a phylogeny (e.g., [25,](#page-18-0) [159\)](#page-23-0). In the future, detection of particularly damaging alleles in genes of agronomic importance could facilitate the amelioration of load in crop germplasm through recombination or targeted gene editing.

3. POSTDOMESTICATION EXPANSION, GENE FLOW, AND ADAPTATION

Following domestication, many crops spread beyond their initial centers of origin to near-global distributions. During this period of expansion, crops often encounter new populations of their progenitor and closely related wild species. In multiple instances, hybridization between crops and these newly sympatric wild relatives has been documented. At times, this hybridization provided crops with adaptations to local conditions. In other new environments where wild relatives did not occur, crops have acquired adaptations de novo as they colonized these new regions. As we describe below, the processes of expansion, gene flow, and adaptation postdomestication have each meaningfully shaped crop biodiversity (**Figure 4**).

Patterns of diversity and genetic similarity of crops to specific populations of extant wild relatives suggest that, in many instances, crops have arisen from one or a few centers of origin. For example, Matsuoka and colleagues [\(103\)](#page-21-0) determined that maize was likely domesticated a single time in the Balsas River basin of southwest Mexico, a finding that has been corroborated by additional genetic analyses [\(170\)](#page-24-0) and archaeological data [\(128\)](#page-22-0). The number of domestication events for other crops like Asian rice remains in question, with some investigations suggesting a single origin potentially in the Pearl River valley of southern China [\(64\)](#page-19-0), and others suggesting

Cost of

domestication:

accumulation of deleterious mutants during and subsequent to domestication due to demographic effects and strong selection limiting recombination

Figure 4

Migration and gene flow history of a fictional domesticate originating in central Mexico. Plots to the right of each population place diversity of the local domesticated population (*red solid line*) in the context of the ancestral diversity of the crop's progenitor (*green dashed line*). Black arrows represent repeated migration and founding events. Although gene flow with wild relatives outside the crop's center of origin can help restore diversity lost during the domestication bottleneck, repeated founding events further erode diversity. Original figure by M. Hufford.

multiple origins in both China and south Asia [\(21\)](#page-17-0). In still other cases, such as the clear independent domestications of common bean in the Lerma River valley of Mexico and the southern Andean highlands of South America [\(48,](#page-19-0) [82\)](#page-20-0), multiple domestication events are clearly supported. However, few examples of diffuse, noncentric domestications have been conclusively proven, and most crops likely share a history of local domestication followed by expansion.

Geographic expansion of a species is known to affect patterns of genetic diversity. For example, as humans migrated out of Africa, diversity was subsampled repeatedly through a series of colonization and/or founder events. In fact, a survey of genetic diversity in modern human populations has demonstrated the effects of this subsampling persist even today, with decreasing diversity observed in humans with increasing distance from Africa [\(134\)](#page-22-0). Given their similar history of expansion, it is not surprising to find the same geographic trend in the genetic diversity of crops. For example, Wang and colleagues [\(176\)](#page-24-0) recently demonstrated a decline in genetic diversity in modern maize landraces with increasing distance from the Balsas River valley. Although not extensively studied, this pattern is likely repeated in many of the world's crops, particularly those such as rice, wheat, common bean, and barley, which experienced sustained, postdomestication expansion over several millennia. Additionally, serial founder effects during expansion have likely shaped patterns in the local prevalence of deleterious alleles. Similar to the cost of domestication described above, declining effective population size due to repeated expansion bottlenecks has likely increased the prevalence of deleterious alleles as crops spread away from their site of origin. This has been clearly demonstrated in maize [\(176\)](#page-24-0) and may well be a more common trend.

In some crops, gene flow with newly encountered populations of wild relatives has likely been an important countervailing force to genetic erosion experienced due to expansion. A recent review [\(70\)](#page-19-0), for example, summarized evidence for substantial wild-to-crop gene flow during postdomestication expansion of many of the world's most important crops including potato, barley, and maize. Gene flow has been so substantial between domesticated potato and several wild relative species, for instance, that it long obscured the identity of the initial progenitor and likely contributed to the high level of diversity currently found in this domesticate [\(55\)](#page-19-0). In crops such as barley, postdomestication introgression from wild relatives has not remained geographically constrained because it was carried by the crop throughout its period of expansion, leading to an ever-widening genetic base of the crop [\(130\)](#page-22-0). However, in other cases like common bean [\(117,](#page-21-0) [118\)](#page-22-0), gene flow has predominantly occurred from domesticate to wild relative, doing little to ameliorate the diversity lost in the crop owing to expansion. In fact, substantial crop-to-wild gene flow has the potential to rapidly diminish wild diversity or even genetically swamp wild relatives to extinction [\(60,](#page-19-0) [164\)](#page-23-0).

Beyond the neutral, demographic effects of serial founding events and gene flow, the process of adaptation required when colonizing new environments has shaped crop diversity. Some common adaptive challenges for crops during expansion include abiotic gradients of elevation, precipitation, temperature, and latitude, as well as exposure to novel biotic stressors [\(65,](#page-19-0) [97\)](#page-21-0). Sources of adaptive variants for crops encountering new environments include the standing variation with which they arrived, gene flow from locally adapted wild relatives, and de novo mutations. Although the former two sources of variation allow for rapid adaptation, the latter requires a waiting time until an adaptive mutation randomly occurs. Perhaps, for this reason, the few studies that have investigated the source of adaptation in expanding crops suggest standing variation and gene flow from wild relatives predominate [\(161\)](#page-23-0). In fact, some have hypothesized that, given the constraints associated with de novo mutations, adaptation may commonly occur across a broad range of species through standing variation and hybridization [\(9,](#page-17-0) [122\)](#page-22-0). Importantly, the winnowing effects of selection on diversity during adaptation would not be as pronounced if adaptation were to occur from standing variation [\(9\)](#page-17-0) or repeated introgression rather than from de novo mutations.

Most recently, modern plant breeding, with its awareness of crop biodiversity, has added to the genetic diversity through a thorough mixing of available germplasm, including introgression from wild types and landraces (e.g., tomato) [\(93\)](#page-20-0), interspecific crosses (e.g., cotton) [\(181\)](#page-24-0), and reconstruction of polyploids (e.g., strawberry) [\(96\)](#page-21-0).

4. PARALLEL EVOLUTION AND CONVERGENCE

Vavilov [\(172\)](#page-24-0) observed that phenotypic variation tended to show parallelism among closely related species, as stated in his article "The Law of Homologous Series in Variation." He even used this observation to predict the occurrence of specific phenotypes in related species. Many of the traits observed by Vavilov were domestication traits, contrasting wild and domesticated types of the same species. The existence of these domestication syndrome traits [\(53, 57\)](#page-19-0) supposes a common set of selection pressures that has acted during and after domestication, leading to parallel phenotypic variation as illustrated by Reference [124](#page-22-0) for domesticates of the Americas. This group of crops provides an excellent model to illustrate the concept of homologous series of phenotypic variation because several genera show multiple domestications within the Americas, notably *Capsicum* and *Phaseolus*, which both contain five domesticated species. In addition, two of the domesticated *Phaseolus* species have been domesticated twice, in Mesoamerica and the Andes [\(48,](#page-19-0) [81\)](#page-20-0).

The determinacy phenotype—a diversification trait, which appeared after domestication—is present in three of the domesticated *Phaseolus*: *P. vulgaris* (common bean), *P. lunatus* (lima bean), and *P. coccineus* (runner bean). In the former two species, the determinacy trait appears in both the Andean and Mesoamerican gene pools. In *Capsicum*, loss of pungency is observed in three domesticated species: *C. annuum*, *C. frutescens*, and *C. chinense* [\(156\)](#page-23-0). In addition to domestication [\(40,](#page-18-0) [92,](#page-20-0) [124\)](#page-22-0), other examples of repeated selection are high-altitude selection (see Section 3), edaphic conditions [\(10,](#page-17-0) [165\)](#page-23-0), and host-plant adaptation [\(42,](#page-18-0) [52,](#page-19-0) [114\)](#page-21-0).

To what extent does parallel phenotypic evolution reflect changes at the same or different loci, and are these structural or, more often, regulatory? An answer to these questions would reveal how natural selection shapes the form and function of living organisms. Moreover, the assumption of convergence is part of the justification of model systems like *Arabidopsis* and rice, which rely on the widespread distribution of shared molecular and genetic mechanisms. It is also of practical importance because it helps identify candidate genes that can be used for gene isolation and marker-assisted selection (e.g., [91,](#page-20-0) [113\)](#page-21-0).

Two important factors affecting the relationship of phenotypic and molecular convergence are the homology of the trait and the phylogenetic relationships among the species being studied. There are two extreme situations: homologous traits in closely related species (same genus or family) versus heterologous traits in distantly related species (different families). In between, all combinations are possible to a degree. In the former situation, the same or similar genes or gene networks are presumably involved; in the latter, different genes or pathways are more likely implicated. These two situations are called, respectively, parallel and convergent (or collateral) evolution [\(124,](#page-22-0) [157\)](#page-23-0).

It has been suggested, however, that data did not support a role of phylogenetic distance in distinguishing between parallelism and convergence [\(5\)](#page-17-0). The common bean provides a case in point. Wild types of this crop show a viny growth habit, allowing them to compete for resources with native vegetation. By contrast, domesticated types display a range of growth habits, from indeterminate, climbing, to determinate, bush. With determinate plants, main stem and branches end in a terminal inflorescence, whereas with indeterminate types the terminal meristems remain vegetative. Thus, the determinacy trait fundamentally changes the growth pattern of bean plants in that it stops its modular growth [\(82,](#page-20-0) [162\)](#page-23-0). Determinacy is controlled by a single recessive allele at the *PvTFL1y* locus, a homolog of the *Arabidopsis TERMINALFLOWER1* (*TFL1*) gene [\(83,](#page-20-0) [138\)](#page-22-0). A sequence analysis of variation at the *PvTF1y* locus showed that indeterminate types—wild or domesticated—showed only synonymous nucleotide substitutions. Different de novo mutations leading to loss of function of the gene (i.e., a determinate phenotype) were only present among domesticated types and not among wild types, consistent with field observations. Among these

Convergence:

independent evolution of the same phenotype in phylogenetically distinct lineages (phenotypic convergence), determined by homologous genes (genotypic convergence) or not

mutations, the most abundant one was a retrotransposon insertion in the fourth exon; others included nonsynonymous substitutions, indels, a putative intron-splicing failure, and a deletion of the entire locus. The multiple mutations leading to the same determinate phenotype provide evidence of parallel evolution under human selection [\(82\)](#page-20-0). Farmers in distinct locations of the centers of origin of the species have selected for a dwarf and early plant type controlled by this determinate phenotype.

Equally remarkable is that other plants, whether they also belong to the Fabaceae family [e.g., soybean [\(163\)](#page-23-0); pea [\(37\)](#page-18-0); pigeon pea [\(142\)](#page-23-0); and others [\(151\)](#page-23-0)] or more distant botanical families, also show phenotypic variation attributable to homologs of *TFL1*. Among these are the Scrophulariaceae (snapdragon, *CENTRORADIALIS*) [\(26\)](#page-18-0), Solanaceae (tomato, *SELF-PRUNING*) [\(129\)](#page-22-0), and Vitaceae (grape, *VvTFL1*) [\(14\)](#page-17-0) in dicots and Poaceae (barley, *HvCEN*) [\(23\)](#page-18-0). Similarities exist also for the seed color gene *P* with both another legume, pea, and a Brassicaceae, *Arabidopsis)* [\(105\)](#page-21-0). Another example of phenotypic variation among closely and distantly related taxa caused by homologous genes are mutations in *replumless* orthologs, affecting fruit opening and seed shattering in the Brassicaceae (*Arabidopsis* and *Brassica*) and in rice (*Sh1* gene), despite their very different phylogenetic position and fruit or seed anatomy [\(6\)](#page-17-0).

Are there features that increase the occurrence of molecular convergence? Lenser & Theißen [\(90\)](#page-20-0) suggest that a nodal position in gene regulatory networks, such as occupied by the *FT*/*TFL1* genes, could be more prone to convergence because they collect regulatory signals from several upstream sources and control several downstream targets. Relatively simple pathways like the anthocyanin pathway could also be a target, especially for a gene situated at the beginning of the pathway such as the *P* color gene in common bean. White seed color controlled by recessive alleles at the *P* locus appeared repeatedly in the Andean and Mesoamerican domesticated gene pools through distinct de novo mutations at the same *P* locus, controlled by a basic helix-loop-helix transcription factor [\(105\)](#page-21-0).

In summary, the determinacy case is one of the most extensive examples to date of phenotypic variation attributable to a homologous gene that is widely distributed phylogenetically. This case provides a counterexample of the prevailing concept of parallel versus convergent evolution. One could also argue that initial identification based on homology between phylogenetically distant species favors molecular convergence in unrelated species. Thus, to date, there is an insufficient number of cases that have been described to quantify the factors involved in molecular convergence. More information is needed from comparisons involving homologous organs or biochemical pathways among a range of species of varying phylogenetic distances.

5. PLASTICITY AND GENOTYPE × ENVIRONMENT INTERACTIONS

Due to the sessile nature of plants, the environment plays an outsized, compelling role in adaptation in these species under both natural and agricultural settings [\(36\)](#page-18-0). Although phenotypic plasticity is the ability of a genotype to express phenotypic variation in different environments, genotype \times environment interaction (G \times E) can be interpreted as differences in plasticity [\(29, 36,](#page-18-0) [125\)](#page-22-0). For plasticity to be under selection, there needs to be environmental variation, genetic differentiation between competing individuals, transgenerational inheritance, and adaptiveness [\(101\)](#page-21-0). Although these four conditions are often met in a natural landscape, they may be less prominent under cultivation, particularly under intensive modern production, which strives to limit genotypic (through monocropping) and environmental variation (through cultivation practices). By contrast, range expansion of domesticated forms, as compared with wild ancestors, has exposed domesticates to novel environments [\(57,](#page-19-0) [173\)](#page-24-0) where plasticity could be adaptive as it may provide general adaptation**.**

Parallel evolution: the development of the same trait in two separate lineages descending from phylogenetically related individuals

Genotype × environment interaction (G×E): the nonlinear response of genotypes to environmental conditions affecting the stability of

genotypes

Quantitative trait locus (QTL): gene controlling the

expression of a trait specifying a quantity or number, which is usually subject to genetic and environmental effects

The detection and study of variation in plasticity can be difficult because environmental variation can encompass both biotic and abiotic factors, agricultural management (for crops), the interactions between these factors, and different temporal and spatial scales. Variation is also found in the degree of plasticity across phenotypic traits [\(1,](#page-17-0) [29\)](#page-18-0). A meta-analysis [\(29\)](#page-18-0) found that natural species are more plastic than domesticates for an array of traits. In addition, more quantitative trait loci (QTLs) per environment were found for wild species compared with crops. Moreover, although the existence of QTLs by environmental interactions is common, differential sensitivity to environmental conditions (change in magnitude of plasticity) is the most frequent type of interaction. A comparison of productivity traits between domesticated types and their wild forms in six herbaceous species found that, even though the domesticates performed better under favorable conditions, plasticity in the wild forms led to higher productivity under water limitation [\(102\)](#page-21-0). Similarly, wild maize (teosinte), when exposed in environmental chambers to simulated early Holocene conditions similar to those at the time of domestication (low carbon dioxide and temperature), showed significant plasticity in its morphology, resembling the morphology of its domesticated descendant [\(127\)](#page-22-0). A higher plasticity was also observed in teosinte relative to maize when comparing gene expression levels under modern and simulated early Holocene conditions [\(94\)](#page-20-0).

By contrast, when considering association with arbuscular mycorrhizae under high and low phosphorus availability, wild plants benefited from the association across environmental conditions (low plasticity), and domesticated forms could modulate the symbiosis based on phosphorus availability [\(99\)](#page-21-0). Modern breeding, although usually evaluated in more stable and uniform conditions, does employ multienvironment testing, which allows the detection and selection of beneficial responses to an array of biotic and abiotic stresses, and when recombined improves the overall productivity and plasticity of crops, as has been found in maize [\(34\)](#page-18-0) and wheat [\(51\)](#page-19-0).

There are different molecular mechanisms associated with plasticity and its variation among genotypes. Although relatively few $G \times E$ genetic factors have been cloned, emerging research suggests that loss-of-function mutations and nonsynonymous changes in receptors play important roles in plasticity [\(29\)](#page-18-0). Polyploidy and epigenetics are increasingly active areas for plasticity research. Domesticated species are more likely to originate from polyploid than diploid wild species [\(141\)](#page-22-0), perhaps because polyploidy can lead to phenotypic novelty and plasticity due to gene dosage effects [\(32\)](#page-18-0). Allopolyploids may show increased plasticity relative to autopolyploids due to their allelic diversity [\(171\)](#page-24-0). Furthermore, allopolyploidy allows gene expression to resemble each of its genome progenitors depending on environmental conditions [\(149\)](#page-23-0). Epigenetics, the heritable change in gene and phenotype expression that is not coded in DNA, is controlled by mechanisms such as DNA methylation, histone modification, small RNA, and enhanced homologous recombination [\(63\)](#page-19-0). The epigenetic profile can be shaped by standing and de novo epimutations and may be environmentally induced, with this latter property having special importance during range expansion or under rapidly changing environments [\(139\)](#page-22-0). The lability of epigenetic profiles may have facilitated plastic changes during crop domestication and adaptation. Although the variation explained by the epigenetic profile might be small compared with the genetic factors [\(3\)](#page-17-0), epigenomic signatures during domestication have been found, for example, in cotton [\(155\)](#page-23-0).

6. FARMER MANAGEMENT OF CROP BIODIVERSITY

It was Darwin [\(27\)](#page-18-0) who remarked that domesticated plants showed more diversity in their harvested organs than in the nonharvested parts, compared to their wild relatives. Numerous examples illustrate Darwin's observation from the multicolored seeds of *Phaseolus* beans to the astonishing diversity among domesticated cabbage (*Brassica* sp.) species. Although several factors can account for this variation, including gene flow (see Section 3), human selection plays a major role in the generation and maintenance of this diversity. This is especially the case in the context of smallholder farming systems in which plants are a primary resource and seeds or vegetative plant materials are part of the capital—instead of being inputs—of the farming enterprises. It is in these systems—often in centers of agricultural origins and domestication [\(173\)](#page-24-0)—that crop evolution and landrace development have taken place for millennia.

6.1. Sociocultural Factors Influencing Crop Biodiversity

Crop biodiversity should be placed in the broader context of plant management of both domesticated and wild or weedy plants, which are managed in parallel. Blancas and colleagues [\(13\)](#page-17-0) studied the way the Náhuatl-speaking people of the village of Coyomeapan managed 122 plant species (including some 35 domesticated species) in different ways and with different intensities within a subsistence system based on the multipurpose use of plant resources and ecosystems. The intensity of plant management appeared to be related to the cultural and economic importance of the plant, its ease of propagation, and a perception of scarcity. On average, some two-thirds of the variation in management intensity could be accounted for by sociocultural factors (e.g., frequency of consumption, number of uses, medicinal use, commercialization), and the remaining variation included 21% by ecological factors (e.g., temporal distribution of harvestable parts, life cycle, ecological dominance) and 12% by the intersection of ecological and sociocultural factors. More intensely managed species are likely subjected to greater selection pressures. Conversely, species that are merely gathered may not be strongly affected unless the gathering involves a selection per se for specific traits (e.g., [87\)](#page-20-0).

The sociocultural organization of farmers plays a role in the structure of crop genetic diversity at the local level. This role is enhanced when crops play an important symbolic role in societies, such as maize in Mesoamerica and pearl millet in Africa. A clear relationship was observed between the spatial limits of genetic groups of pearl millet and boundaries among three ethnolinguistic groups west of Lake Chad [\(71\)](#page-19-0). The authors attributed this situation to limited circulation of pearl millet, a crop that is generally inherited from fathers to sons, whose seeds are obtained generally from relatives and neighbors. Similar patterns were observed in African sorghum populations [\(177\)](#page-24-0). At the level of the African continent, the molecular analysis identified three large geographic sorghum populations (each with their own ecogeographic races) that largely overlapped with local language families. These distributions reflect at least two past farming-language expansions, i.e., the Bantu and Nilo-Saharan expansions.

Although several factors could promote seed exchange across ethnolinguistic boundaries (such as the antiquity of pearl millet or sorghum cultivation, the mixing of landraces and fields, and the ever-increasing density of cultivation), barriers to seed circulation among ethnic groups in these regions seem to have the upper hand (e.g., [116\)](#page-21-0). Correlations between genetic diversity and ethnolinguistic distributions are tied to accumulated local traditional knowledge (TK) and preferences concerning morphological and adaptive characteristics of plants and their use [\(16,](#page-17-0) [183\)](#page-24-0). The importance of such knowledge cannot be overstated. For example, local communities in the Tehuacán valley of Mexico know of nearly 2,000 plant species, including their useful properties, distribution, year-to-year abundance, and interactions with other plants and animals [\(167\)](#page-23-0). Agroforestry systems, such as coffee and cocoa plantations, are probably some of the oldest forms of agriculture. They can be particularly rich in biodiversity and TK associated with each biological and genetic resource they contain [\(89\)](#page-20-0). In turn, this information can be the basis for intensification of new crops, such as indigenous fruit and nut trees [\(88\)](#page-20-0). TK is, therefore, an integral part of crop

Traditional knowledge (TK): information, knowledge, and practices obtained and inherited by local or indigenous communities, generation after generation

biodiversity. Not surprisingly, recent international treaties on biodiversity have included TK as part of their purview.

Within ethnic groups, gender can sometimes be an important factor for farmer preferences. In Ethiopia, both genders ranked earliness as their top priority, most likely because this trait is crucial in assuring food security when faced with depletion of harvests from previous growing seasons. Additional shared preferences, with slightly different rankings, included yield, drought tolerance, and marketability.Women were more preoccupied with addressing fast cooking and taste, whereas men focused more on field problems, including germination and specific adaptation, and market demand [\(7\)](#page-17-0). By contrast, a different study [\(179\)](#page-24-0) did not observe gender differences regarding preferences for high grain yield, earliness, short heads, low kernel weight, and short plants. However, in this study, no postharvest traits like cooking and taste were reported. The diversity of household mixtures of food crops is, therefore, the outcome of a complex set of constraints, which vary across gender preferences and involvement in production, crop location, and time.

6.2. Farmer Practices and Crop Biodiversity

Farmers must make difficult decisions about farming practices that integrate a complex series of biological, climatic, and socioeconomic variables, over some of which they have little control. The practice of planting mixtures is widespread in subsistence agriculture, which is in contrast with industrial agriculture where genetic monocultures are the norm. Farmers in Eritrea and the neighboring province of Tigray in Ethiopia engage in the practice of planting mixtures of barley and wheat, called *hanfets* in the local Tigrigna language, as a means of dealing with the unpredictability of rainfall [\(179\)](#page-24-0). In wet years, the higher yield potential of wheat is favored, whereas in dry years the higher drought tolerance of barley is advantageous. On average across locations and years, the yield of *hanfets* is like that of barley monoculture but markedly higher than that of wheat. Not all *hanfets* mixtures were more stable than pure crops, but generally, they showed fewer or no interactions with location or year within location, unlike pure barley or wheat.

Farmers use several approaches to conserve or enhance crop diversity in their fields. In Santa María Jaltianguis (Oaxaca, Mexico), farmers use three taxa of *Phaseolus* beans in their plantings, including two ecogeographic races of common bean (*P. vulgaris*, races Nueva Granada and Mesoamerica) and runner bean (*P. coccineus*), with distinct altitudinal adaptation [\(180\)](#page-24-0). Farmers consider the altitude of each field to adjust the mixtures planted to increase the proportion of the most adapted component. For example, the most recent introduction of beans—small, black beans—constituted an attempt to introduce marketable beans; it required opening new fields at lower altitudes. Furthermore, farmers adapt these introduced beans to their local environmental conditions by unilateral introgressive hybridization with traditional beans, a process called creolization. Elsewhere, in the state of Guanajuato, Mexico, farmers maintain genetic diversity in the field at a level comparable to those observed in wild beans. They include existing landraces and an improved variety, and tolerate wild beans in their fields [\(184\)](#page-24-0).

The practice of planting mixtures has several potential motivations. It is a risk-mitigating strategy when faced with a variable and uncertain biotic and abiotic environment; it is also a way of dealing with diverse topography spanning a range of microenvironments. Diverse plantings also cater to various postharvest needs, including different cooking characteristics and consumer tastes. Both bananas and beans in smallholder farms of Uganda showed high richness and evenness in frequency of traditional varieties [\(111\)](#page-21-0). In locations with higher disease incidence of anthracnose (*Colletotrichum lindemuthianum*) and angular leaf spot (*Pseudocercospora griseola*), a negative correlation between richness and the disease index was observed, suggesting that increased within-field bean diversity can act as a partial disease control measure. In the Mulumba et al. [\(111\)](#page-21-0) study, richness was determined on the basis of the number of traditional varieties regardless of the genetic relatedness of the varieties. However, traditional bean varieties grown by households in Uganda include both Andean and Mesoamerican domesticates of *Phaseolus vulgaris* [\(178\)](#page-24-0). For both diseases, there is a parallel organization of genetic diversity with Andean and Mesoamerican strains, which are more virulent on Andean and Mesoamerican hosts [\(52,](#page-19-0) [150\)](#page-23-0), respectively. A household or community mixture consisting of both Andean and Mesoamerican varieties is inherently more resistant to diseases than varietal portfolios consisting of one or the other gene pool. Furthermore, a greater evenness in frequencies is more conducive to disease epidemic control, as is planting variety mixtures in spatial patterns within fields [\(111\)](#page-21-0).

During the crop cycle, weeding plays an important role in maintaining the yield potential by removing damaging competition with crop plants. Cassava farmers practice selective weeding of volunteer plants resulting from outcrossing among clonal plants. Because there is a positive correlation between plant size and multilocus heterozygosity, the resulting volunteer population after weeding had a lower level of inbreeding [\(131\)](#page-22-0). This example illustrates how farming practices exert an action on the genetic diversity of crops since domestication and up to today.

Farmers express preferences for certain agronomic, culinary, and consumption traits with varying degrees of heritability. These traits are important because they provide the basis for local adaptation addressing the needs of farmers and local consumers. In two Ethiopian communities, the inheritance of farmer preferences in their durum wheat crops was investigated [\(78\)](#page-20-0). They combined quantitative evaluations of farmer preferences (for flowering time, spike morphology, tilling capacity, and overall quality) and measurements by researchers of agronomic and phenology traits in a collection of 400 Ethiopian wheat varieties, genotyped with single-nucleotide polymorphisms. For farmer traits, 124 putative QTLs were identified, whereas 30 putative QTLs were identified for researcher traits. Twenty of these were shared. This research showed that farmer evaluations were highly correlated across environments and, thus, reproducible and heritable.

6.3. Seed Systems

The previous examples illustrate that farmers in a subsistence context, i.e., often in centers of crop domestication, play an active role in managing their seed stocks, including their biodiversity, in their respective households. These farmer seed systems are often called informal in contrast with the commercial or state-led formal seed sector. Farmer seed networks can be described as systems through which plant materials may come from a range of sources, including farmers' saved seeds, other farmers, markets, nongovernmental organizations, national and international research institutions, and commercial seed suppliers [\(24,](#page-18-0) [178\)](#page-24-0). These farmer networks disseminate not only seeds of wild plants but also new crops and new varieties created by on-farm management, including creolized varieties, increasing crop diversity and providing an avenue for resiliency in the case of biotic, abiotic, or economic shocks. Furthermore, informal systems may be the only system for crops overlooked by commercial systems and research, such as vegetables and grain legumes. An important part of crop biodiversity is, therefore, maintained by these informal networks. Maintaining this biodiversity requires further research and a legal and political system that supports both formal and informal seed systems [\(120\)](#page-22-0).

6.4. Application to Participatory Breeding

The results of Kidane and coauthors [\(78\)](#page-20-0), mentioned in Section 6.2, suggest that farmer knowledge can be integrated into modern plant breeding, using QTL mapping, marker-assisted selection, and genomic selection. However, can farmers themselves be engaged in plant breeding? This is a question that has been asked repeatedly in participatory research focusing on variety selection or breeding (reviewed in [20,](#page-17-0) [67,](#page-19-0) [148\)](#page-23-0) in the hope of obtaining improved varieties that satisfy the

Seed system: a set of biological, social, and economic conditions that affect the production, storage, and distribution of cultivar seeds

Participatory research:

collaborative research between scientists and farmers (or users) leading to the development of a research product or outcome

needs and preferences of farmers and consumers who live in diverse and variable environments. The example of beans in Honduras provides a strong confirmation of the success of participatory variety breeding [\(67\)](#page-19-0) and reflects the use of local crop biodiversity, combined with advanced breeding lines in the development of locally released improved varieties. Additional examples of decentralized selection activities using local crop biodiversity are the on-farm stratified recurrent mass selection experiments conducted in maize. In these experiments, heterogeneous local maize populations planted in the field are divided into plots; in each plot, the best individuals are selected, and their seeds are pooled for the next generation. This process is then repeated several times until the necessary breeding objectives are achieved [\(4,](#page-17-0) [109,](#page-21-0) [153\)](#page-23-0).

SUMMARY POINTS

- 1. Crop biodiversity is both one of the most important inventions of humanity and a crucial resource for plant breeding to develop crop varieties.
- 2. The overall evolutionary pattern of genetic diversity is one of gradual reduction due to demographic (genome-wide) and selective (gene-specific) causes from the initial domestication process until today. This reduction is tempered, however, by several factors including gene flow in wild \times domesticated and domesticated \times domesticated hybridizations, selection for functional diversity and broader adaptation, and, more recently and to a certain extent, modern plant breeding.
- 3. There is a cost to domestication owing to the accumulation of deleterious mutations in domesticated populations caused by limited recombination in populations with reduced effective size. A future focus of plant breeding will be the elimination of these deleterious alleles through recombination or targeted genome editing.
- 4. It remains to be determined to what extent the domestication syndrome and homologous series of crop variations are caused by genetic variation in similar (parallel evolution) or different genes or gene networks (convergent evolution). An answer to this question would help quantify the role of model systems and predictability across the taxa of candidate genes.
- 5. Plasticity is a form of genotype \times environment interaction. Its magnitude relative to total genetic variation, its inheritance and heritability, and its evolution remain to be fully understood.
- 6. Farmers are active agents of the conservation and development of genetic diversity, reflecting their membership in different ethnic and socioeconomic groups, their gender, and their farming practices. Their abilities to select and conserve genetic diversity are being utilized in participatory breeding, leading to the release of locally adapted varieties.

FUTURE ISSUES

1. To promote their utilization, gene banks will need in-depth characterization, including systematic DNA sequencing (e.g., [175\)](#page-24-0), high-throughput phenotyping [\(146\)](#page-23-0), and biochemical [\(19\)](#page-17-0) and physiological observations [\(49\)](#page-19-0), while maintaining current activities aimed at maintenance of the viability and disease-free status of collections [\(38\)](#page-18-0).

- 2. Historical demography coupled with selection during domestication and adaptation have shaped crop diversity over millennia [\(176\)](#page-24-0). We must therefore better understand how modern in situ diversity and the evolutionary potential of a crop have been determined by historical diversity in its progenitor, the strength of its domestication and expansion bottlenecks, and the historical opportunities for gene flow with locally adapted wild relatives.
- 3. A full accounting of temporal and spatial patterns of crop biodiversity require an understanding of how plants function at different levels from the genome to the whole-plant levels so that the effects of genes and gene networks can be translated into a population genetic perspective (e.g., [47,](#page-18-0) [75\)](#page-20-0).
- 4. There is a need to assess the variation of plastic responses among wild ancestors to determine whether this variation was lost during domestication and whether it is advantageous in a domesticated genetic background and under agricultural conditions [\(29,](#page-18-0) [102\)](#page-21-0).
- 5. The current emphasis on individual plants or crops should be complemented with a more comprehensive view of the crop's interactions with other plants, animals, and microorganisms in its (agro)ecosystem with potential benefits for plant breeding, integrated pest management, and crop husbandry [\(17,](#page-17-0) [61,](#page-19-0) [143\)](#page-23-0).
- 6. The focus on crop wild relatives must be maintained, as these relatives have proven to be a source of additional genetic diversity, especially considering the overall reduction in genetic diversity of the domesticated gene pool and changes in pest and disease pressures, global climatic conditions, and market and consumer demands [\(28\)](#page-18-0).
- 7. Local informal seed systems and participatory breeding involving farmers and scientists are crucial to promote utilization of crop biodiversity and maximize local adaptation, including consumer preference [\(67,](#page-19-0) [100\)](#page-21-0).
- 8. Genetic conservation efforts, including gene banks, are chronically underfunded and deserve better support commensurate with their importance as stewards of biodiversity as an insurance policy for the adaptability and resilience of agro- and managed ecosystems [\(45,](#page-18-0) [62\)](#page-19-0).
- 9. Because efforts such as the international Convention on Biological Diversity (and its agreements like the Nagoya Protocol), as well as the national Intellectual Property Law, have had counterproductive effects, limiting the use and conservation of this diversity [\(33, 44\)](#page-18-0), a significant legal effort needs to be developed to promote research and education in biodiversity conservation.

DISCLOSURE STATEMENT

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

ACKNOWLEDGMENTS

Recent research by M.B.H. in this area has been supported by National Science Foundation (NSF) grant IOS-1546719. P.G. received recent funding from the US Department of Agriculture, National Institute of Food and Agriculture (2013–67013–21224 and 2016–67013–24460) and NSF ERA-CAPS (European Research Area Network for Coordinating Action in Plant Sciences) (1539848). We thank J. Dubcovsky for a helpful discussion of the parallel and convergent evolution and genetic diversity of wheat.

LITERATURE CITED

- 1. Abley K, Locke JCW, Leyser HMO. 2016. Developmental mechanisms underlying variable, invariant and plastic phenotypes. *Ann. Bot.* 117:733–48
- 2. Allen T, Prosperi P, Cogill B, Flichman G. 2014. Agricultural biodiversity, social–ecological systems and sustainable diets. *Proc. Nutr. Soc.* 73:498–508
- 3. Aller EST, Jagd LM, Kliebenstein DJ, Burow M. 2018. Comparison of the relative potential for epigenetic and genetic variation to contribute to trait stability. *G3* 8:1733–46
- 4. Alves ML, Belo M, Carbas B, Brites C, Paulo M, et al. 2018. Long-term on-farm participatory maize breeding by stratified mass selection retains molecular diversity while improving agronomic performance. *Evol. Appl.* 11:254–70
- 5. Arendt J, Reznick D. 2008. Convergence and parallelism reconsidered: What have we learned about the genetics of adaptation? *Trends Ecol. Evol.* 23:26–32
- 6. Arnaud N, Lawrenson T, Østergaard L, Sablowski R. 2011. The same regulatory point mutation changed seed-dispersal structures in evolution and domestication. *Curr. Biol.* 21:1215–19
- 7. Assefa T, Sperling L, Dagne B, Argaw W, Tessema D, Beebe S. 2014. Participatory plant breeding with traders and farmers for white pea bean in Ethiopia. *J. Agric. Educ. Ext.* 20:497–512
- 8. Barot S, Allard V, Cantarel A, Enjalbert J, Gauffreteau A, et al. 2017. Designing mixtures of varieties for multifunctional agriculture with the help of ecology. A review. *Agron. Sustain. Dev.* 37:13
- 9. Barrett RDH, Schluter D. 2008. Adaptation from standing genetic variation. *Trends Ecol. Evol.* 23:38–44
- 10. Beebe S, Lynch J, Galwey N, Tohme J, Ochoa I. 1997. A geographical approach to identify phosphorusefficient genotypes among landraces and wild ancestors of common bean. *Euphytica* 95:325–36
- 11. Bellucci E, Bitocchi E, Ferrarini A, Benazzo A, Biagetti E, et al. 2014. Decreased nucleotide and expression diversity and modified coexpression patterns characterize domestication in the common bean. *Plant Cell* 26:1901–12
- **12. Bitocchi E, Rau D, Benazzo A, Bellucci E, Goretti D, et al. 2016. High level of nonsynonymous changes in common bean suggests that selection under domestication increased functional diversity at target traits.** *Front. Plant Sci.* **7:2005**
- 13. Blancas J, Casas A, Pérez-Salicrup D, Caballero J, Vega E. 2013. Ecological and socio-cultural factors influencing plant management in Náhuatl communities of the Tehuacán Valley, Mexico. *J. Ethnobiol. Ethnomed.* 9:39
- 14. Boss PK, Sreekantan L, Thomas MR. 2006. A grapevine TFL1 homologue can delay flowering and alter floral development when overexpressed in heterologous species. *Funct. Plant Biol.* 33:31–41
- 15. Bradshaw CJA, Brook BW. 2014. Human population reduction is not a quick fix for environmental problems. *PNAS* 111:16610–15
- 16. Brush SB. 2004. *Farmers' Bounty*. New Haven, CT: Yale Univ. Press
- 17. Cadotte MW. 2017. Functional traits explain ecosystem function through opposing mechanisms. *Ecol. Lett.* 20:989–96
- 18. Caicedo AL, Williamson SH, Hernandez RD, Boyko A, Fledel-Alon A, et al. 2007. Genome-wide patterns of nucleotide polymorphism in domesticated rice. *PLOS Genet*. 3:1745–56
- 19. Cañas RA, Yesbergenova-Cuny Z, Simons M, Chardon F, Armengaud P, et al. 2017. Exploiting the genetic diversity of maize using a combined metabolomic, enzyme activity profiling, and metabolic modeling approach to link leaf physiology to kernel yield. *Plant Cell* 29:919–43
- 20. Ceccarelli S, Guimaraes EP, Weltzien E. 2009. *Plant Breeding and Farmer Participation*. Rome: FAO
- 21. Civáň P, Craig H, Cox CJ, Brown TA. 2015. Three geographically separate domestications of Asian rice. *Nat. Plants* 1:15164
- 22. Collins WW, Qualset CO. 1999. *Biodiversity in Agroecosystems*. Boca Raton, FL: CRC

for an excess of nonsynonymous mutations in coding regions of domesticated genotypes, potentially due to selection for tolerance to abiotic and biotic stresses.

12. Provides evidence

- 23. Comadran J, Kilian B, Russell J, Ramsay L, Stein N, et al. 2012. Natural variation in a homolog of *Antirrhinum CENTRORADIALIS* contributed to spring growth habit and environmental adaptation in cultivated barley. *Nat. Genet.* 44:1388–92
- 24. Coomes OT, McGuire SJ, Garine E, Caillon S, McKey D, et al. 2015. Farmer seed networks make a limited contribution to agriculture? Four common misconceptions. *Food Policy* 56:41–50
- 25. Cooper GM, Stone EA, Asimenos G, Green ED, Batzoglou S, Sidow A. 2005. Distribution and intensity of constraint in mammalian genomic sequence. *Genome Res*. 15:901–13
- 26. Cremer F, Lönnig W-E, Saedler H, Huijser P. 2001. The delayed terminal flower phenotype is caused by a conditional mutation in the *CENTRORADIALIS* gene of snapdragon. *Plant Physiol*. 126:1031–41
- 27. Darwin C. 1859. *On the Origin of Species by Means of Natural Selection*. London: Murray
- 28. Dempewolf H, Baute G, Anderson J, Kilian B, Smith C, Guarino L. 2017. Past and future use of wild relatives in crop breeding. *Crop Sci*. 57:1070–82
- 29. Des Marais DL, Hernandez KM, Juenger TE. 2013. Genotype-by-environment interaction and plasticity: exploring genomic responses of plants to the abiotic environment. *Annu. Rev. Ecol. Evol. Syst.* 44:5– 29
- 30. Dirzo R, Raven PH. 2003. Global state of biodiversity and loss. *Annu. Rev. Environ. Resour.* 28:137–67
- 31. Doebley JF, Gaut BS, Smith BD. 2006. The molecular genetics of crop domestication. *Cell* 127:1309–21
- 32. Dubcovsky J, Dvorak J. 2007. Genome plasticity a key factor in the success of polyploid wheat under domestication. *Science* 316:1862–66
- 33. Dutfield G. 2017. Traditional knowledge, intellectual property and pharmaceutical innovation: What's left to discuss? In *The Sage Handbook of Intellectual Property*, ed.M David, D Halbert, pp. 649–64. London: Sage
- 34. Duvick DN, Smith JSC, Cooper M. 2004. Long-term selection in a commercial hybrid maize breeding program. *Plant Breed. Rev.* 24 (Part 2):109–51
- 35. Dwivedi SL, Lammerts van Bueren ET, Ceccarelli S, Grando S, Upadhyaya HD, Ortiz R. 2017. Diversifying food systems in the pursuit of sustainable food production and healthy diets. *Trends Plant Sci*. 22:842–56
- 36. El-Soda M, Malosetti M, Zwaan BJ, Koornneef M, Aarts MGM. 2014. Genotype × environment interaction QTL mapping in plants: lessons from *Arabidopsis. Trends Plant Sci*. 19:390–98
- 37. Foucher F, Morin J, Courtiade J, Cadioux S, Ellis N, et al. 2003. *DETERMINATE* and *LATE FLOWER-ING* are two *TERMINAL FLOWER1/CENTRORADIALIS* homologs that control two distinct phases of flowering initiation and development in pea. *Plant Cell* 15:2742–54
- 38. Fu Y-B. 2017. The vulnerability of plant genetic resources conserved ex situ. *Crop Sci*. 57:2314–28
- 39. Gammans M, Mérel P, Ortiz-Bobea A. 2017. Negative impacts of climate change on cereal yields: statistical evidence from France. *Environ. Res. Lett.* 12:054007
- 40. Gaut BS. 2015. Evolution is an experiment: assessing parallelism in crop domestication and experimental evolution. *Mol. Biol. Evol.* 32:1661–71
- 41. Gaut BS, Díez CM, Morrell PL. 2015. Genomics and the contrasting dynamics of annual and perennial domestication. *Trends Genet*. 31:709–19
- 42. Geffroy V, Sicard D, de Oliveira J, Sévignac M, Cohen S, et al. 1999. Identification of an ancestral resistance gene cluster involved in the coevolution process between *Phaseolus vulgaris* and its fungal pathogen *Colletotrichum lindemuthianum*. *Mol. Plant Microbe Interact*. 12:774–84
- 43. Gepts P. 2004. Crop domestication as a long-term selection experiment. *Plant Breed. Rev.* 24(Part 2):1–44
- 44. Gepts P. 2004. Who owns biodiversity and how should the owners be compensated? *Plant Physiol*. 134:1295–307
- 45. Gepts P. 2006. Plant genetic resources conservation and utilization: the accomplishments and future of a societal insurance policy. *Crop Sci*. 46:2278–92
- 46. Gepts P. 2014. The contribution of genetic and genomic approaches to plant domestication studies.*Curr. Opin. Plant Biol.* 18:51–59
- 47. Gepts P. 2017. Genetic aspects of crop domestication. In *Routledge Handbook of Agricultural Biodiversity*, ed. D Hunter, L Guarino, C Spillane, PC McKeown, pp. 147–67. New York: Routledge

65. Describes the genome-wide extent of introgressive hybridization between sympatric maize and wild *Zea mays* **ssp.** *mexicana* **leading to highland adaptation in Mexico.**

67. Provides a successful example of participatory research involving local farmers, agricultural researchers, and nongovernmental organizations in using biodiversity for crop breeding.

68. Provides a broad overview of agricultural biodiversity, including its nature, valuation, effect on human health, socioeconomic and legal aspects, and conservation.

- 48. Gepts P, Osborn TC, Rashka K, Bliss FA. 1986. Phaseolin-protein variability in wild forms and landraces of the common bean (*Phaseolus vulgaris*): evidence for multiple centers of domestication. *Econ. Bot.* 40:451–68
- 49. Gilbert ME, Medina V. 2016. Drought adaptation mechanisms should guide experimental design. *Trends Plant Sci*. 21:639–47
- 50. Glémin S, Bataillon T. 2009. A comparative view of the evolution of grasses under domestication. *New Phytol*. 183:273–90
- 51. Grogan SM, Anderson J, Baenziger PS, Frels K, Guttieri MJ, et al. 2016. Phenotypic plasticity of winter wheat heading date and grain yield across the US Great Plains. *Crop Sci*. 56:2223–36
- 52. Guzmán P, Gilbertson RL, Nodari R, Johnson WC, Temple SR, et al. 1995. Characterization of variability in the fungus *Phaeoisariopsis griseola* suggests coevolution with the common bean (*Phaseolus vulgaris*). *Phytopathology* 85:600–7
- 53. Hammer K. 1984. Das Domestikationssyndrom. *Kulturpflanze* 32:11–34
- 54. Hannah L, Roehrdanz PR, Ikegami M, Shepard AV, Shaw MR, et al. 2013. Climate change, wine, and conservation. *PNAS* 110:6907–12
- 55. Hardigan MA, Laimbeer FPE, Newton L, Crisovan E, Hamilton JP, et al. 2017. Genome diversity of tuber-bearing *Solanum* uncovers complex evolutionary history and targets of domestication in the cultivated potato. *PNAS* 114:E9999–10008
- 56. Harjes CE, Rocheford TR, Bai L, Brutnell TP, Kandianis CB, Sowinski SG. 2008. Natural genetic variation in *lycopene epsilon cyclase* tapped for maize biofortification. *Science* 319:330–33
- 57. Harlan JR. 1992. *Crops and Man*. Madison, WI: Am. Soc. Agron.
- 58. Haudry A, Cenci A, Ravel C, Bataillon T, Brunel D, et al. 2007. Grinding up wheat: a massive loss of nucleotide diversity since domestication. *Mol. Biol. Evol.* 24:1506–17
- 59. Hawksworth D, Kalin-Arroyo MT, Hammond PM, Ricklefs RE, Cowling R, et al. 1995. Magnitude and distribution of biodiversity. In *Global Biodiversity Assessment*, ed. VH Heywood, RT Watson, pp. 107–92. Cambridge, UK: Cambridge Univ. Press
- 60. Hegde SG, Nason JD, Clegg JM, Ellstrand NC. 2006. The evolution of California's wild radish has resulted in the extinction of its progenitors. *Evolution* 60:1187–97
- 61. Heil M. 2014. Herbivore-induced plant volatiles: targets, perception and unanswered questions. *New Phytol*. 204:297–306
- 62. Heywood VH. 2017. Plant conservation in the Anthropocene—challenges and future prospects. *Plant Divers*. 39:314–30
- 63. Holeski LM, Jander G, Agrawal AA. 2012. Transgenerational defense induction and epigenetic inheritance in plants. *Trends Ecol. Evol.* 27:618–26
- 64. Huang X, Zhao Y, Wei X, Li C, Wang A, et al. 2012. Genome-wide association study of flowering time and grain yield traits in a worldwide collection of rice germplasm. *Nat. Genet.* 44:32–39
- **65. Hufford MB, Lubinksy P, Pyhäjärvi T, Devengenzo MT, Ellstrand NC, Ross-Ibarra J. 2013. The genomic signature of crop-wild introgression in maize.** *PLOS Genet.* **9:e1003477**
- 66. Hufford MB, Xu X, van Heerwaarden J, Pyhajarvi T, Chia JM, et al. 2012. Comparative population genomics of maize domestication and improvement. *Nat. Genet.* 44:808–11
- **67. Humphries S, Rosas JC, Gómez M, Jiménez J, Sierra F, et al. 2015. Synergies at the interface of farmer–scientist partnerships: agricultural innovation through participatory research and plant breeding in Honduras.** *Agric. Food Secur.* **4:27**
- **68. Hunter D, Guarino L, Spillane C, McKeown PC, eds. 2017.** *Routledge Handbook of Agricultural Biodiversity***. New York: Taylor & Francis**
- 69. Iorizzo M, Senalik DA, Ellison SL, Grzebelus D, Cavagnaro PF, et al. 2013. Genetic structure and domestication of carrot (*Daucus carota* subsp. *sativus*) (Apiaceae). *Am. J. Bot.* 100:930–38
- 70. Janzen GM,Wang L, Hufford MB. 2019. The extent of adaptive wild introgression in crops. *New Phytol*. 221:1279–88
- 71. Jika AKN, Dussert Y, Raimond C, Garine E, Luxereau A, et al. 2017. Unexpected pattern of pearl millet genetic diversity among ethno-linguistic groups in the Lake Chad basin. *Heredity* 118:491–502
- 72. Jones AD. 2016. On-farm crop species richness is associated with household diet diversity and quality in subsistence- and market-oriented farming households in Malawi. *J. Nutr.* 147:86–96
- 73. Jump AS,Marchant R, Peñuelas J. 2009. Environmental change and the option value of genetic diversity. *Trends Plant Sci*. 14:51–58
- 74. Kantar MB, Nashoba AR, Anderson JE, Blackman BK, Rieseberg LH. 2017. The genetics and genomics of plant domestication. *BioScience* 67:971–82
- 75. Kessler A, Kalske A. 2018. Plant secondary metabolite diversity and species interactions. *Annu. Rev. Ecol. Evol. Syst.* 49:115–38
- 76. Kew R. Bot. Gard. 2016. *State of the World's Plants 2016*. Kew, UK: Kew R. Bot. Gard. **https:// stateoftheworldsplants.org/2016/**
- **77. Khoury CK, Achicanoy HA, Bjorkman AD, Navarro-Racines C, Guarino L, et al. 2016. Origins of food crops connect countries worldwide.** *Proc. R. Soc. London Ser. B* **283:20160792**
- 78. Kidane YG, Mancini C, Mengistu DK, Frascaroli E, Fadda C, et al. 2017. Genome wide association study to identify the genetic base of smallholder farmer preferences of durum wheat traits. *Front. Plant Sci.* 8:1230
- 79. Kiers ET, Leakey RRB, Izac A-M, Heinemann JA, Rosenthal E, et al. 2008. Agriculture at a crossroads. *Science* 320:320–21
- 80. Knapp S, Winter M, Klotz S, Bennett J. 2017. Increasing species richness but decreasing phylogenetic richness and divergence over a 320-year period of urbanization. *J. Appl. Ecol.* 54:1152–60
- 81. Kwak M, Kami JA, Gepts P. 2009. The putative Mesoamerican domestication center of *Phaseolus vulgaris* is located in the Lerma-Santiago basin of Mexico. *Crop Sci*. 49:554–63
- **82. Kwak M, Toro O, Debouck D, Gepts P. 2012. Multiple origins of the determinate growth habit in domesticated common bean (***Phaseolus vulgaris* **L.).** *Ann. Bot.* **110:1573–80**
- 83. Kwak M, Velasco DM,Gepts P. 2008.Mapping homologous sequences for determinacy and photoperiod sensitivity in common bean (*Phaseolus vulgaris*). *J. Hered.* 99:283–91
- 84. Lam H-M, Xu X, Liu X, Chen W, Yang G, et al. 2010. Resequencing of 31 wild and cultivated soybean genomes identifies patterns of genetic diversity and selection. *Nat. Genet.* 42:1053–59
- 85. Larson G, Piperno DR, Allaby RG, Purugganan MD, Andersson L, et al. 2014. Current perspectives and the future of domestication studies. *PNAS* 111:6139–46
- 86. Laurance WF, Clements GR, Sloan S, O'Connell CS, Mueller ND, et al. 2014. A global strategy for road building. *Nature* 513:229–32
- 87. Law W, Salick J. 2005. Human-induced dwarfing of Himalayan snow lotus, *Saussurea laniceps* (Asteraceae). *PNAS* 102:10218–20
- 88. Leakey RRB. 2012. Participatory domestication of indigenous fruit and nut trees: new crops for sustainable agriculture in developing countries. In *Biodiversity in Agriculture: Domestication, Evolution, and Sustainability*, ed. P Gepts, TR Famula, RL Bettinger, SB Brush, AB Damania, et al., pp. 479–501. Cambridge, UK: Cambridge Univ. Press
- 89. Leakey RRB. 2017. *Multifunctional Agricultura: Achieving Sustainable Development in Africa*. London: Academic
- 90. Lenser T, Theißen G. 2013. Molecular mechanisms involved in convergent crop domestication. *Trends Plant Sci*. 18:704–14
- 91. Li L, Tacke E, Hofferbert H-R, Lübeck J, Strahwald J, et al. 2013. Validation of candidate gene markers for marker-assisted selection of potato cultivars with improved tuber quality. *Theor. Appl. Genet.* 126:1039–52
- 92. Li W, Gill BS. 2006. Multiple genetic pathways for seed shattering in the grasses. *Funct. Integr. Genomics* 6:300–9
- 93. Lin T, Zhu G, Zhang J, Xu X, Yu Q, et al. 2014. Genomic analyses provide insights into the history of tomato breeding. *Nat. Genet.* 46:1220–26
- 94. Lorant A, Pedersen S, Holst I, Hufford MB, Winter K, et al. 2017. The potential role of genetic assimilation during maize domestication. *PLOS ONE* 12:e0184202
- **95. Lu J, Tang T, Tang H, Huang J, Shi S, Wu C-I. 2006. The accumulation of deleterious mutations in rice genomes: a hypothesis on the cost of domestication.** *Trends Genet***. 22:126–31**

77. Illustrates the origin and worldwide dissemination of our crops and the recent trend in mutual dependency of countries on crop biodiversity.

82. Shows that a given (domesticated) phenotype can have multiple origins through distinct de novo lineage-specific mutations at a single locus.

95. First posited a cost of domestication based on an increased frequency of nonsynonymous mutations in the rice domesticated gene pool compared to the wild one.

- 96. Luby J, Hancock J, Dale A, Serçe S. 2008. Reconstructing *Fragaria* × *ananassa* utilizing wild *F. virginiana* and *F. chiloensis*: inheritance of winter injury, photoperiod sensitivity, fruit size, female fertility and disease resistance in hybrid progenies. *Euphytica* 163:57–65
- 97. Mariac C, Jehin L, Saidou AA, Thuillet AC, Couderc M, et al. 2011. Genetic basis of pearl millet adaptation along an environmental gradient investigated by a combination of genome scan and association mapping. *Mol. Ecol.* 20:80–91
- 98. Martínez-Ainsworth NE, Tenaillon MI. 2016. Superheroes and masterminds of plant domestication. *C. R. Biol.* 339:268–73
- **99. Martín-Robles N, Lehmann A, Seco E, Aroca R, Rillig MC, Milla R. 2018. Impacts of domestication on the arbuscular mycorrhizal symbiosis of 27 crop species.** *New Phytol***. 218:322– 34**
- 100. Mastretta-Yanes A, Acevedo Gasman F, Burgeff C, Cano Ramírez M, Piñero D, Sarukhán J. 2018. An initiative for the study and use of genetic diversity of domesticated plants and their wild relatives. *Front. Plant Sci.* 9:209
- 101. Matesanz S, Gianoli E, Valladares F. 2010. Global change and the evolution of phenotypic plasticity in plants. *Ann. NY Acad. Sci.* 1206:35–55
- 102. Matesanz S, Milla R. 2018. Differential plasticity to water and nutrients between crops and their wild progenitors. *Environ. Exp. Bot.* 145:54–63
- 103. Matsuoka Y, Vigouroux Y, Goodman MM, Sanchez GJ, Buckler E, Doebley J. 2002. A single domestication for maize shown by multilocus microsatellite genotyping. *PNAS* 99:6080–84
- 104. Maxwell SL, Fuller RA, Brooks TM, Watson JEM. 2016. Biodiversity: the ravages of guns, nets and bulldozers. *Nature* 536:143–45
- 105. McClean PE, Bett KE, Stonehouse R, Lee R, Pflieger S, et al. 2018. White seed color in common bean (*Phaseolus vulgaris*) results from convergent evolution in the *P* (pigment) gene. *New Phytol*. 219:1112– 23
- 106. Meyer RS, DuVal AE, Jensen HR. 2012. Patterns and processes in crop domestication: an historical review and quantitative analysis of 203 global food crops. *New Phytol*. 196:29–48
- 107. Meyer RS, Purugganan MD. 2013. Evolution of crop species: genetics of domestication and diversification. *Nat. Rev. Genet.* 14:840–52
- 108. Miller AJ, Gross BL. 2011. From forest to field: perennial fruit crop domestication. *Am. J. Bot.* 98:1389– 414
- 109. Moreira PA, Mariac C, Zekraoui L, Couderc M, Rodrigues DP, et al. 2017. Human management and hybridization shape treegourd fruits in the Brazilian Amazon basin. *Evol. Appl.* 10:577–89
- 110. Moyers BT, Morrell PL, McKay JK. 2018. Genetic costs of domestication and improvement. *J. Hered.* 109:103–16
- 111. Mulumba JW, Nankya R, Adokorach J, Kiwuka C, Fadda C, et al. 2012. A risk-minimizing argument for traditional crop varietal diversity use to reduce pest and disease damage in agricultural ecosystems of Uganda. *Agric. Ecosyst. Environ.* 157:70–86
- 112. Murray-Kolb LE, Wenger MJ, Scott SP, Rhoten SE, Lung'aho MG, Haas JD. 2017. Consumption of iron-biofortified beans positively affects cognitive performance in 18- to 27-year-old Rwandan female college students in an 18-week randomized controlled efficacy trial. *J. Nutr.* 147:2109–17
- 113. Naderpour M, Lund OS, Larsen R, Johansen E. 2010. Potyviral resistance derived from cultivars of *Phaseolus vulgaris* carrying *bc-3* is associated with the homozygotic presence of a mutated *eIF4E* allele. *Mol. Plant Pathol.* 11:255–63
- 114. Nosil P, Crespi BJ, Sandoval CP. 2002. Host-plant adaptation drives the parallel evolution of reproductive isolation. *Nature* 417:440–43
- 115. Olsen KM,Wendel JF. 2013. A bountiful harvest: genomic insights into crop domestication phenotypes. *Annu. Rev. Plant Biol.* 64:47–70
- 116. Orozco-Ramirez Q, Ross-Ibarra J, Santacruz-Varela A, Brush S. 2016. Maize diversity associated with social origin and environmental variation in Southern Mexico. *Heredity* 116:477–84
- 117. Papa R, Acosta J, Delgado-Salinas A, Gepts P. 2005. A genome-wide analysis of differentiation between wild and domesticated *Phaseolus vulgaris* from Mesoamerica. *Theor. Appl. Genet.* 111:1147–58

99. Illustrates an increase in plasticity under domestication of the symbiotic interaction with arbuscular mycorrhizae.

- 118. Papa R, Gepts P. 2003. Asymmetry of gene flow and differential geographical structure of molecular diversity in wild and domesticated common bean (*Phaseolus vulgaris* L.) from Mesoamerica. *Theor. Appl. Genet.* 106:239–50
- 119. Pathak T,Maskey M, Dahlberg J, Kearns F, Bali K, Zaccaria D. 2018. Climate change trends and impacts on California agriculture: a detailed review. *Agronomy* 8:25
- 120. Pautasso M, Aistara G, Barnaud A, Caillon S, Clouvel P, et al. 2013. Seed exchange networks for agrobiodiversity conservation. A review. *Agron. Sustain. Dev.* 33:151–75
- 121. Peter BM, Huerta-Sanchez E, Nielsen R. 2012. Distinguishing between selective sweeps from standing variation and from a *de novo* mutation. *PLOS Genet*. 8:e1003011
- 122. Pfennig KS, Kelly AL, Pierce AA. 2016. Hybridization as a facilitator of species range expansion. *R. Soc. Proc. B* 283:20161329
- 123. Phalan B, Onial M, Balmford A, Green RE. 2011. Reconciling food production and biodiversity conservation: land sharing and land sparing compared. *Science* 333:1289–91
- 124. Pickersgill B. 2018. Parallel vs. convergent evolution in domestication and diversification of crops in the Americas. *Front. Ecol. Evol.* 6:56
- 125. Pigliucci M. 2005. Evolution of phenotypic plasticity: Where are we going now? *Trends Ecol. Evol.* 20:481–86
- 126. Pimm SL, Raven PH. 2017. The fate of the world's plants. *Trends Ecol. Evol.* 32:317–20
- 127. Piperno DR, Holst I, Winter K, McMillan O. 2015. Teosinte before domestication: experimental study of growth and phenotypic variability in Late Pleistocene and early Holocene environments. *Quat. Int.* 363:65–77
- 128. Piperno DR, Ranere AJ, Holst I, Iriarte J, Dickau R. 2009. Starch grain and phytolith evidence for early ninth millennium B.P. maize from the central Balsas River valley, Mexico. *PNAS* 106:5019–24
- 129. Pnueli L, Carmel-Goren L, Hareven D, Gutfinger T, Alvarez J, et al. 1998. The *SELF-PRUNING* gene of tomato regulates vegetative to reproductive switching of sympodial meristems and is the ortholog of *CEN* and *TFL1. Development* 125:1979–89
- 130. Poets AM, Fang Z, Clegg MT, Morrell PL. 2015. Barley landraces are characterized by geographically heterogeneous genomic origins. *Genome Biol*. 16:173
- 131. Pujol B, David P, McKey D. 2005. Microevolution in agricultural environments: how a traditional Amerindian farming practice favours heterozygosity in cassava (*Manihot esculenta* Crantz, Euphorbiaceae). *Ecol. Lett.* 8:138–47
- 132. Purugganan MD, Fuller DQ. 2009. The nature of selection during plant domestication.*Nature* 457:843– 48
- 133. Qualset CO, McGuire PE, Warburton ML. 1995. In California: 'agrobiodiversity' key to agricultural productivity. *Calif. Agric.* 49:45–49
- 134. Ramachandran S, Deshpande O, Roseman CC, Rosenberg NA, Feldman MW, Cavalli-Sforza LL. 2005. Support from the relationship of genetic and geographic distance in human populations for a serial founder effect originating in Africa. *PNAS* 102:15942–47
- 135. Ramu P, Esuma W, Kawuki R, Rabbi IY, Egesi C, et al. 2017. Cassava haplotype map highlights fixation of deleterious mutations during clonal propagation. *Nat. Genet.* 49:959–63
- 136. Reiss ER, Drinkwater LE. 2018. Cultivar mixtures: a meta-analysis of the effect of intraspecific diversity on crop yield. *Ecol. Appl.* 28:62–77
- 137. Renaut S, Rieseberg LH. 2015. The accumulation of deleterious mutations as a consequence of domestication and improvement in sunflowers and other Compositae crops. *Mol. Biol. Evol.* 32:2273–83
- 138. Repinski SL, Kwak M, Gepts P. 2012. The common bean growth habit gene *PvTFL1y* is a functional homolog of *Arabidopsis TFL1. Theor. Appl. Genet*. 124:1539–47
- 139. Richards CL, Alonso C, Becker C, Bossdorf O, Bucher E, et al. 2017. Ecological plant epigenetics: evidence from model and non-model species, and the way forward. *Ecol. Lett.* 20:1576–90
- 140. Rubel F, Kottek M. 2010. Observed and projected climate shifts 1901–2100 depicted by world maps of the Köppen-Geiger climate classification. *Meteorol. Z.* 19:135–41
- 141. Salman-Minkov A, Sabath N, Mayrose I. 2016. Whole-genome duplication as a key factor in crop domestication. *Nat. Plants* 2:16115
- 142. Saxena RK, Obala J, Sinjushin A, Kumar CVS, Saxena KB, Varshney RK. 2017. Characterization and mapping of *Dt1* locus which co-segregates with *CcTFL1* for growth habit in pigeonpea. *Theor. Appl. Genet.* 130:1773–84
- 143. Schmidt JE, Bowles TM, Gaudin ACM. 2016. Using ancient traits to convert soil health into crop yield: impact of selection on maize root and rhizosphere function. *Front. Plant Sci.* 7:373
- 144. Schrider DR, Kern AD. 2016. S/HIC: robust identification of soft and hard sweeps using machine learning. *PLOS Genet*. 12:e1005928
- 145. Schultz J. 2005. *The Ecozones of the World*. Berlin: Springer
- 146. Shakoor N, Lee S, Mockler TC. 2017. High throughput phenotyping to accelerate crop breeding and monitoring of diseases in the field. *Curr. Opin. Plant Biol.* 38:184–92
- 147. Sheehan S, Song YS. 2016. Deep learning for population genetic inference. *PLOS Comput. Biol.* 12:e1004845
- 148. Shelton A, Tracy W. 2016. Participatory plant breeding and organic agriculture: a synergistic model for organic variety development in the United States. *Sci. Anthr.* 4:143
- 149. Shimizu-Inatsugi R, Terada A, Hirose K, Kudoh H, Sese J, Shimizu KK. 2017. Plant adaptive radiation mediated by polyploid plasticity in transcriptomes. *Mol. Ecol.* 26:193–207
- 150. Sicard D, Michalakis Y, Dron M, Neema C. 1997. Genetic diversity and pathogenic variation of *Colletotrichum lindemuthianum* in the three centers of diversity of its host, *Phaseolus vulgaris*. *Phytopathology* 87:807–13
- 151. Sinjushin AA. 2015. Mutations of determinate growth and their application in legume breeding. *Legum. Perspect.* 6:14–15
- 152. Skarbø K, VanderMolen K. 2015. Maize migration: key crop expands to higher altitudes under climate change in the Andes. *Clim. Dev.* 8:245–55
- 153. Smith M, Castillo F, Gómez F. 2001. Participatory plant breeding with maize in Mexico and Honduras. *Euphytica* 122:551–63
- 154. Smith S, Diers B, Specht J, Carver B, eds. 2014. *Yield Gains in Major U.S. Field Crops*. Madison, WI: Am. Soc. Agron., Inc./Crop Sci. Soc. Am., Inc./Soil Sci. Soc. Am., Inc.
- 155. Song Q, Zhang T, Stelly DM, Chen ZJ. 2017. Epigenomic and functional analyses reveal roles of epialleles in the loss of photoperiod sensitivity during domestication of allotetraploid cottons. *Genome Biol*. 18:99
- 156. Stellari GM, Mazourek M, Jahn MM. 2009. Contrasting modes for loss of pungency between cultivated and wild species of *Capsicum. Heredity* 104:460–71
- 157. Stern DL. 2013. The genetic causes of convergent evolution. *Nat. Rev. Genet.* 14:751–64
- 158. Stetter MG, Gates DJ, Mei W, Ross-Ibarra J. 2017. How to make a domesticate.*Curr. Biol.* 27:R896–900
- 159. Stone EA, Sidow A. 2005. Physicochemical constraint violation by missense substitutions mediates impairment of protein function and disease severity. *Genome Res*. 15:978–86
- **160. Studer A, Zhao Q, Ross-Ibarra J, Doebley J. 2011. Identification of a functional transposon insertion in the maize domestication gene** *tb1. Nat. Genet***. 43:1160–63**
- 161. Takuno S, Ralph P, Swarts K, Elshire RJ, Glaubitz JC, et al. 2015. Independent molecular basis of convergent highland adaptation in maize. *Genetics* 200:1297–312
- 162. Tanaka A, Fujita K. 1979. Photosynthesis and yield components in relation to grain yield of the field beans. *J. Fac. Agric. Hokkaido Univ.* 59:145–238
- 163. Tian Z, Wang X, Lee R, Li Y, Specht JE, et al. 2010. Artificial selection for determinate growth habit in soybean. *PNAS* 107:8563–68
- 164. Todesco M, Pascual MA, Owens GL, Ostevik KL, Moyers BT, et al. 2016. Hybridization and extinction. *Evol. Appl.* 9:892–908
- 165. Turner TL, Bourne EC, Von Wettberg EJ, Hu TT, Nuzhdin SV. 2010. Population resequencing reveals local adaptation of *Arabidopsis lyrata* to serpentine soils. *Nat. Genet.* 42:260–63
- 166. United Nations. 2017. *World Population Prospects 2017*[. New York: UN DESA/Popul. Div.](https://population.un.org/wpp/) **https:// population.un.org/wpp/**
- 167. Vallejo-Ramos M, Moreno-Calles AI, Casas A. 2016. TEK and biodiversity management in agroforestry systems of different socio-ecological contexts of the Tehuacán valley. *J. Ethnobiol. Ethnomed.* 12:31

160. Provides an example of standing variation arising 10,000 years before domestication and resulting from a transposable element insertion in the control region of a gene.

- 168. van de Wouw M, Kik C, van Hintum T, van Treuren R, Visser B. 2010. Genetic erosion in crops: concept, research results and challenges. *Plant Genet. Res.* 8:1–15
- 169. van de Wouw M, van Hintum T, Kik C, van Treuren R, Visser B. 2010. Genetic diversity trends in twentieth century crop cultivars: a meta analysis. *Theor. Appl. Genet.* 120:1241–52
- 170. van Heerwaarden J, Doebley J, Briggs WH, Glaubitz JC, Goodman MM, et al. 2011. Genetic signals of origin, spread, and introgression in a large sample of maize landraces. *PNAS* 108:1088–92
- 171. Vaughan DA, Balázs E, Heslop-Harrison JS. 2007. From crop domestication to super-domestication. *Ann. Bot.* 100:893–901
- 172. Vavilov NI. 1922. The law of homologous series in variation. *J. Genet.* 12:47–89
- 173. Vavilov NI. 1926. Centers of origin of cultivated plants. In *Origin and Geography of Cultivated Plants*, ed. VF Dorofeyev, pp. 22–135. Cambridge, UK: Cambridge Univ. Press
- 174. Vellend M, Baeten L, Becker-Scarpitta A, Boucher-Lalonde V,McCune JL, et al. 2017. Plant biodiversity change across scales during the Anthropocene. *Annu. Rev. Plant Biol.* 68:563–86
- 175. Wambugu PW, Ndjiondjop M-N, Henry RJ. 2018. Role of genomics in promoting the utilization of plant genetic resources in genebanks. *Brief Funct. Genomics* 17:198–206
- 176. Wang L, Beissinger TM, Lorant A, Ross-Ibarra C, Ross-Ibarra J, Hufford MB. 2017. The interplay of demography and selection during maize domestication and expansion. *Genome Biol*. 18:215
- 177. Westengen OT, Okongo MA, Onek L, Berg T, Upadhyaya H, et al. 2014. Ethnolinguistic structuring of sorghum genetic diversity in Africa and the role of local seed systems. *PNAS* 111:14100–5
- **178. Wilkus EL, Berny Mier y Teran JC,Mukankusi CM, Gepts P. 2018. Genetic patterns of commonbean seed acquisition and early-stage adoption among farmer groups in western Uganda.** *Front. Plant Sci.* **9:586**
- **genotypic information.** 179. Woldeamlak A, Grando S, Maatougui M, Ceccarelli S. 2008. *Hanfets*, a barley and wheat mixture in Eritrea: yield, stability and farmer preferences. *Field Crops Res*. 109:50–56
- **180. Worthington M, Soleri D, Aragón-Cuevas F, Gepts P. 2012. Genetic composition and spatial distribution of farmer-managed bean plantings: an example from a village in Oaxaca, Mexico.** *Crop Sci***. 52:1721–35**
- 181. Zhang J, Percy RG, McCarty JC. 2014. Introgression genetics and breeding between Upland and Pima cotton: a review. *Euphytica* 198:1–12
- 182. Zhou Y,Massonnet M, Sanjak JS, Cantu D, Gaut BS. 2017. Evolutionary genomics of grape (*Vitis vinifera* ssp. *vinifera*) domestication. *PNAS* 114:11715–20
- 183. Zimmerer KS. 2010. Biological diversity in agriculture and global change. *Annu. Rev. Environ. Resour.* 35:137–66
- 184. Zizumbo-Villarreal D, Colunga-GarcíaMarín P, Payró de la Cruz E, Delgado-Valerio P, Gepts P. 2005. Population structure and evolutionary dynamics of wild–weedy–domesticated complexes of common bean in a Mesoamerican region. *Crop Sci*. 35:1073–83

RELATED RESOURCES

Crop Trust. **<https://www.croptrust.org/>**

Crop Wild Relatives. **<https://www.cwrdiversity.org/>**

Food Agric. Organ. United Nations. *What Is Agrobiodiversity?* **[http://www.fao.org/docrep/007/y5609e/](http://www.fao.org/docrep/007/y5609e/y5609e01.htm#TopOfPage) y5609e01.htm#TopOfPage**

Francis RC. 2015. *Domesticated: Evolution in a Man-Made World*. New York: W. W. Norton *Genesys: The Global Gateway to Genetic Resources*. **<https://www.genesys-pgr.org/welcome>** Gepts P, Famula TR, Bettinger RL, Brush SB, Damania AB, et al., eds. 2012. *Biodiversity in Agriculture: Domestication, Evolution, and Sustainability*. Cambridge, UK: Cambridge Univ. Press

State of the World's Plants. **<https://stateoftheworldsplants.org>**

178. Illustrates how farmer management and participatory research affect seed stock genetic diversity, based on

180. Shows farmers' awareness of adaptation of species-level adaptation and the directed adaptation to local conditions (creolization) by introgressive hybridization.

Annual Review of Plant Biology

Volume 70, 2019

Contents

Errata

An online log of corrections to *Annual Review of Plant Biology* articles may be found at http://www.annualreviews.org/errata/arplant