Quantitative epigenetics and evolution

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Abstract

Epigenetics refers to chemical modifications of chromatin or transcribed DNA that can influence gene activity and expression without changes in DNA sequence. The last 20 years have yielded breakthroughs in our understanding of epigenetic processes that impact many fields of biology. In this review, we discuss how epigenetics relates to quantitative genetics and evolution. We argue that epigenetics is important for quantitative genetics because: (1) quantitative genetics is increasingly being combined with genomics, and therefore we should expand our thinking to include cellular-level mechanisms that can account for phenotypic variance and heritability besides just those that are hard-coded in the DNA sequence; and (2) epigenetic mechanisms change how phenotypic variance is partitioned, and can thereby change the heritability of traits and how those traits are inherited. To explicate these points, we show that epigenetics can influence all aspects of the phenotypic variance formula:

\[ V_P = V_G + V_E + V_{GxE} + 2COV_{GE} + V_\varepsilon \]

requiring new strategies to account for different potential sources of epigenetic effects on phenotypic variance. We also demonstrate how each of the components of phenotypic variance not only can be influenced by epigenetics, but can also have evolutionary consequences. We argue that no sources of epigenetic effects on phenotypic variance can be easily cast aside in a quantitative genetic research program that seeks to understand evolutionary processes.

Impacts of epigenetics research

Epigenetics refers to chemical modifications of chromatin or transcribed DNA that can influence gene activity and expression without changes in DNA sequence (Jablonka and Raz 2009; Kilvitis et al. 2014). The most well-studied epigenetic mechanism is the methylation of cytosines within DNA sequences (Rapp and Wendel 2005; Allis and Jenuwein 2016), although other mechanisms like histone modifications (Berger et al. 2009) and a heterogeneous assortment of RNA regulatory systems (Chen et al. 2011) have also been uncovered and studied intensively (reviewed in Allis and Jenuwein 2016). Methylation of cytosines is an important component of silencing transposable elements (Gibney and Nolan 2010; Huff and Zilberman 2014; Fultz et al. 2015; Ikeda and Nishimura 2015), but is also found in complex patterns across other genomic contexts (Niederhuth et al. 2016). For example, dense gene promoter methylation is associated with silencing of genes, but methylation in gene bodies—which tends to be polymorphic between individuals—is poorly associated with gene expression and varies widely across taxa (Becker et al. 2011; Nätt et al. 2012; Schmitz et al. 2013; Niederhuth and Schmitz 2017).

Epigenetics research has surged over the last 20 years, thanks to advances in our knowledge about chromatin modifications and their modifiers, and the often-complex connections between these modifications and gene expression patterns (reviewed in Allis and Jenuwein 2016). In particular, “the modern era of epigenetic research” is transforming our understanding of the molecular basis of development and disease (Allis and Jenuwein 2016). But the integration of epigenetics into evolutionary thinking is still gathering momentum (Laland et al. 2015; Burggren 2016; Richards et al. 2017). Epigenetics is one of several new research topics that are emerging in evolutionary biology that go beyond the original tenets of the Modern Synthesis (Huxley 1942), the major integration of disparate biological and mathematical concepts into evolutionary biology.
theory in the mid-20th century (Pigliucci 2007; Plutynski 2009; Pigliucci and Müller 2010; Robertson and Richards 2015).

Proponents of a so-called “Extended Evolutionary Synthesis” are dissatisfied with a definition of evolution that is limited to changing DNA sequence based on allele frequencies over time. Instead, the evolutionary process would be more completely described by investigating non-genetic mechanisms like epigenetics that were not initially included in the Modern Synthesis, and that might be able to overcome some of the limitations of strictly DNA sequence-based inheritance (Pigliucci 2007; Bonduriansky and Day 2009; Day and Bonduriansky 2011; Robertson and Richards 2015). In particular, because epigenetic mechanisms are more dynamic and reversible than DNA sequence, epimutations have been implicated as a faster source of adaptation than genetic mutations (Jablonka and Lamb 1989, Jablonka et al. 1998). Epimutations can provide higher phenotypic variance at equilibrium due to different epigenetic states, and enhance the adaptive possibilities of asexual or low-diversity taxa (Jablonka and Lamb 1989; Geoghegan and Spencer 2012, 2013). Organisms may use the additional variation afforded by epigenetic mechanisms as a bet-hedging strategy in unknown environments (Jablonka and Lamb 1989, Jablonka et al. 1998; Pál and Miklós 1999). Epigenetic modifications could also "hold" a potentially advantageous phenotype for multiple generations, allowing time for more stable genetic variants to stabilize the phenotype (i.e. canalization or genetic assimilation, Waddington 1942, 1953; West-Eberhard 2005; Klironomos et al. 2013; Kronholm and Collins 2016).

Epigenetics should be incorporated into evolutionary theory, because the molecular underpinnings of phenotypic variation are the subject of many evolutionary research programs (McNiven et al. 2014). We simply need to expand our thinking to include epigenetic mechanisms, which are another class of molecular mechanisms. Evolution relies on inheritable phenotypic variation, which is provided by alleles (which may be DNA-based alleles or epigenetic-based alleles, as we will describe) that confer different patterns of gene expression/function that ultimately result in different phenotypes in different individuals (Falconer and Mackay 1996). Roff (2007) and Hill (2012) took stock of evolutionary biology in the genomics era, where the advent of high throughput genome sequencing technologies of the last few decades (e.g., Liberles 2001; Goodwin et al. 2016) has been revolutionary across the life sciences. They point out that even quantitative genetics, a branch of evolutionary biology that statistically analyzes phenotypes and has treated molecular mechanisms as a black box, has been greatly influenced by the genomic era, with many research programs now focusing on the number and distribution of effect sizes of loci affecting a trait in a population through the use of quantitative trait locus (QTL) or genome-wide association mapping (GWAM) (Li et al. 2016). Genomics approaches have also provided new resolution to understanding the interactions among loci, and the change in the phenotypic effects of loci under different environments (Mackay 2001). High-throughput genome sequencing technologies have enabled the evolutionary biology community to blend quantitative genetics and genomic approaches to address the enduring question about whether phenotypic variation is controlled by many loci of small effect, fewer loci with a wider distributions of effect sizes, or just a few loci of large effect on the phenotype (Roff 2007; Kalisz and Kramer 2008; Stinchcombe and Hoekstra 2008; Hill 2012). Further, we now have more power to ask whether loci are behaving in an additive or more complex fashion in their phenotypic effects (Ruppell et al. 2004; Wolf et al. 2005; Mackay 2014).

The knowledge of the genetic architecture of traits can be used to study many aspects of evolution. For instance, genetic architecture can provide evidence for the degree to which negative selection, stabilizing selection, and balancing selection shaped the evolution of a trait, as reflected in the distributions of the minor allele frequencies of the QTLs/SNPs, as well as in the distribution of effect sizes of the QTLs/SNPs (Erickson et al. 2004; Josephs et al. 2017). Knowledge of the genetic architecture can also be leveraged in artificial selection. Agricultural breeding programs currently make prodigious use of a technique known as genomic selection to improve plant and animal breeding (Meuwissen et al. 2001; Koopae and Koshkoiyeh 2014). The approach first uses GWAM to find associations between molecular markers and the phenotype(s) of interest in crops or livestock. Then, individuals with unknown phenotype or pedigree are genotyped at the same SNP sites (using a low-cost SNP chip) to forecast their phenotype(s) from the effect sizes associated with each allele from the GWAM study (Koopae and Koshkoiyeh 2014). In this way, superior stock can be selected for breeding and development without further measuring of phenotype. In addition to this type of application, genomic selection has been used in model systems to make more precise predictions about adaptive evolution in response to selection (e.g., Edwards et al. 2016; Kooke et al. 2016).

**Epigenetics and the missing heritability problem**

A well-known problem with genetic mapping is that not all of the genetic variance (V_G) in the phenotype can be accounted for by variance in the genome, a phenomenon known as the “missing heritability” problem (Brachi et al. 2011; Caballero et al. 2015). In QTL studies of recombinant
inbred lines (RIL) or GWAM studies of natural variants, some differences in phenotypes cannot be accounted for by the sum of the significantly correlated genomic regions. Multiple explanations have been put forth to account for the missing heritability (Dickson et al. 2010; Rockman 2012; Hemani et al. 2013; Yang et al. 2015). Epigenetic changes are another possible source of some of the missing heritability, because epigenetic changes can be inherited and act in an additive fashion on phenotypic variance (Morgan et al. 1999; Henderson and Jacobsen 2007; Slatkin 2009; Bell and Spector 2011; González-Recio 2011; Migicovsky and Kovalchuk 2011; Goddard and Whitelaw 2014; Triantaphyllopoulos et al. 2016). Epigenetic changes that have phenotypic effects (mediated through their effects on gene expression) and that are also inherited could generate associations between individuals and phenotypes that may not be explained based on variation in DNA sequence-based markers.

There are many examples of epigenetically inherited effects that could account for missing heritability. One particularly illustrative example comes from the agouti locus in mice (Morgan et al. 1999; Waterland and Jirtle 2003). Differences in silencing by methylation of a retrotransposon inserted upstream of the AY allele at the agouti locus is correlated with ectopic agouti expression and disease risk. Low methylation of the transposable element is correlated with yellow coat color and diseases such as obesity, diabetes, and tumors, while high methylation of the transposable element is associated with brown coat color and lessened disease risk (Morgan et al. 1999; Waterland and Jirtle 2003). Importantly, specific agouti epialleles in female mice can be induced by a high methyl donor diet, and the epigenetic state of the agouti allele is incompletely reset by meiosis in the female germ line (Morgan et al. 1999; Waterland and Jirtle 2003). Thus, coat color, along with susceptibility to cancer and metabolic disorders, can be induced in the mother and inherited by the offspring, illustrating that epigenetic changes can be inherited in a manner like DNA sequence-based mutations, but can also be induced in a specific, directional way by the environment.

Several other studies in ecological epigenetics provide examples of epigenetic effects that are inherited (Sano and Kim 2013; Robertson and Richards 2015; Richards et al. 2017). Verhoeven et al. (2010) reared apomictic dandelions (Taraxacum officinale) under several different stresses (nutrient, salt, and chemical stresses). They found that the stresses triggered considerable methylation variation throughout the genome, and many of these were faithfully transmitted to the offspring. These results demonstrated that epigenetic variation can be readily generated by environmental challenges and that those changes can be inherited. In another example, Richards et al. (2008, 2012) demonstrated the persistence of epigenetic differences in plants from the Fallopia species complex (referred to as Japanese knotweed sensu lato) with little DNA sequence variation. The Fallopia species complex occupies a wide range of habitats in Europe and has colonized an even wider array of habitats in the northeastern US (including marshes and beaches). Richards et al. (2012) found that the invasive Fallopia at sites on Long Island in New York, USA had only four variable AFLP positions out of 200 positions genome-wide. They found diversity was five times higher at epigenetic loci, and this diversity varied among sites. Importantly, these data were collected from leaf tissue that was grown from rhizomes in a common garden. Therefore, the DNA methylation patterns were not merely induced by different sites, but persisted in tissue that was created in a common environment. Perhaps not all of this epigenetic variation would be inherited through meiosis, but some of it could be as has been found in other studies (Cubas et al. 1999; Feng et al. 2010; Herrera et al. 2013). More importantly, for the biology of this species, clonal propagation is a common mode of expansion and therefore the persistence of differences in DNA methylation through clonal propagation is highly relevant (Verhoeven and Prete 2014; Douhovnikoff and Dodd 2015; Rendina González et al. 2016; Spens and Douhovnikoff 2016). Neither the Verhoeven et al. (2010) study nor the Richards et al. (2012) study correlated inherited epigenetic variation with phenotypic variation, but such linkages have been made in other studies (reviewed in Richards et al. 2017). Cubas et al. (1999), for instance, characterized a naturally occurring mutant of toadflax (Linaria vulgaris) that has a different floral symmetry pattern. They found that methylation changes, rather than DNA sequence changes, to a floral symmetry gene explained the phenotypic change. The changes in methylation of the gene segregate and are inherited much like DNA sequence variants, but spontaneous resetting of methylation sometimes occurs in nature that recovers the non-mutant phenotype.

In another example linking epigenetic variation and phenotypic variation, Johannes et al. (2009) developed a recombinant inbred line (RIL) population of the model plant Arabidopsis thaliana that segregates for differentially methylated positions (DMPs)—differences in whether individual cytosines are methylated or not. The epigenetic recombinant inbred lines (epiRILs) are nearly isogenic in terms of the DNA sequences, but show variation and high heritability for many traits relating to growth and morphology, including plant height, flowering time, and primary root length (Zhang et al. 2013; Cortijo et al. 2014; Kooke and Keurentjes 2015). The epiRILs were derived from crosses between the Columbia wild-type genotype and a mutant line derived from the Columbia wild-type with a mutation in the DECREASED DNA METHYLATION 1

SPRINGER NATURE
(DDM1) locus. Mutant plants (ddm1/ddm1) exhibit 70% less methylation genome-wide than wild-type plants (Vongs et al. 1993). From this initial cross, Johannes et al. (2009) selected backcross progeny that were homozygous for the wild-type (DDM1/DDM1) at the DDM1 locus to propagate 505 epiRILs through six rounds of single-seed descent. Thus, the epiRILs are nearly isogenic in terms of DNA sequences, but segregate for stably inherited methylation polymorphisms (DMPs), which are homozygous within each epiRIL (Johannes et al. 2009). The epiRILs show phenotypic variation and high heritability for plant height, flowering time, and primary root length (Johannes et al. 2009; Cortijo et al. 2014; Kooke and Keurentjes 2015). Of particular interest, Cortijo et al. (2014) found multiple epiQTLs accounting for 60–90% of the heritability of flowering time and primary root length. Thirty percent of the heritable DMPs identified in the epiRIL population also overlapped with naturally occurring epipolymorphic regions among 138 natural A. thaliana accessions, suggesting that epigenetic variation could be important in the wild. These epiQTLs have all the necessary properties (stability, inheritance, variance, phenotypic effects) to be targets of natural or artificial selection.

Differences between epigenetic effects and genetic effects

Parental effects refer to traits expressed in parents that influence traits expressed in offspring (Hadfield 2012). Under this umbrella are both paternal effects and maternal effects, allowing for fathers’ and mothers’ phenotypes to have different effects on offsprings’ phenotypes. Maternal effects have generally received the most theoretical attention, presumably because paternal maternal provisioning and postnatal maternal care are the most significant sources of parental effects (Kirkpatrick and Lande 1989; Wolf and Wade 2016). Epigenetic and parental effects differ in that epigenetic effects are a more specific class of phenomena than parental effects; epigenetic effects result from specific types of cellular mechanisms, whereas parental effects can include any mechanism by which a trait expressed in the parents is also expressed in the offspring. Parental effects can be due to inherited epigenetic effects, but they can also be due to non-epigenetic mechanisms, like cultural transmission, prenatal nutrient provisioning, postnatal care, and maternal transmission of mitochondria, chloroplasts, and other cytoplasmic factors to the offspring (Kirkpatrick and Lande 1989; Wolf and Wade 2016). Epigenetics also differs from parental effects because epigenetic effects are not necessarily inherited, whereas parental effects are, by definition, inherited. Consider, for instance, a change in the methylation state at a particular region of a chromosome of an organism in response to some environmental stimulus, where the methylation changes are reset in the germ line. The methylation changes in this example would be epigenetic, but they would not be a parental effect. This is because, in this example, the methylation changes in the parents would not be transmitted to the offspring, and therefore the methylation changes in the parents would not serve as the vehicle through which “information” about the parental traits are passed to the offspring. In summary, parental effects and epigenetics are overlapping but also distinct concepts that should be modeled separately and can have different effects on the phenotype and on inheritance (Santure and Spencer 2006).

Differences between epigenetic effects and genetic effects

Epigenetic mechanisms can contribute to phenotypic variation and could be playing a role in the missing heritability of quantitative traits. But epigenetic contributions to heritability are different from DNA sequence based contributions to heritability, because epigenetic states can be reset, either before reaching the germ line or after transmission to the offspring (Saze et al. 2003; Feng et al. 2010; Hackett et al. 2013), and because epigenetic differences among individuals can arise at different rates from DNA sequence-based differences (Kronholm and Collins 2016). Modeling efforts have demonstrated that epigenetic effects have different ramifications for evolution than standard DNA sequence based processes. Furrow (2014), for instance, used a population genetic model to understand the response of epigenetically induced phenotypic variation to selection over multiple generations. He found that, because epimutation rates are high, the response to selection decays more rapidly across generations than would be expected based on the observed heritability of a trait alone. Similarly, Santure and Spencer (2006) found a smaller than expected response to selection when they included epigenetic differences that depended on the parent from whom a chromosomal region was inherited (genomic imprinting; Macdonald 2012). Thus, traits may be less responsive to natural selection than expected due to the effect of epigenetic phenomena on observed heritabilities (but see Klironomos et al. 2013). In addition to the rate of epimutation, Kronholm and Collins (2016) found that the effect size of the epimutations influences the rate of adaptation, where small-effect epimutations generally speed up adaptation, and large-effect epimutations slow down adaptation. In contrast to models that found that epigenetic effects may slow down adaptation, a model by Uller et al. (2015) showed that epigenetic effects can speed up evolution in heterogeneous but predictable environments, due to the environmental induction.
of some epigenetic states and their transmissibility. Another model by Leimar and McNamara (2015) showed that incompletely-resetting epigenetic systems are ideally suited for actively transmitting environmental information from previous generations, and can evolve readily. Thus, selection may be actively maintaining that the cellular machinery does not completely reset certain environmentally induced epigenetic states across generations, something that Shea et al. (2011) refer to as a “selection-based effect” of epigenetics.

Several authors have emphasized how the decoupling of phenotypic change from genotypic change enabled by epigenetic mechanisms can allow populations to change in ways that might not otherwise be possible. For instance, epigenetic changes that happen at faster rates than DNA sequence changes can drive more rapid evolutionary change by serving as a bridge to allow phenotypic changes to occur first, and genetic changes later (Bonduriansky and Day 2009; Geoghegan and Spencer 2013; Klironomos et al. 2013; Kronholm and Collins 2016). In addition, epimutations may allow for a higher mutational load and lower mean fitness of adapted populations than would otherwise be expected (due to the higher epimutation rate; Klironomos et al. 2013), but recent studies found that epigenetic variation may compensate for the negative effects of inbreeding (Vergeer et al. 2012; Liebl et al. 2013). When van der Graaf et al. (2015) combined theoretical modeling with high-resolution epigenetic analysis of multiple independent Arabidopsis thaliana mutation accumulation lines, they found that the mean fitness due to epimutations was not lower than expected. The epimutation rates they reported were high enough to rapidly uncouple genetic from epigenetic variation, but low enough for new epialleles to sustain long-term selection responses. We will need more information before we can make conclusions about the specific ramifications of epigenetics on evolutionary processes. Much of the differences among the conflicting evolutionary models for partitioning phenotypic variance can be accounted for by quantifying the distributions and phenotypic effects of epimutations in multiple different species (Kronholm and Collins 2016). Yet those variables are presently largely unknown and we must wait for the empirical data to parameterize those variables (sensu van der Graaf et al. 2015).

**Epigenetic effects on phenotypic variance**

Quantitative genetic studies are based on understanding components of variance within the framework $V_P = V_G + V_E$, where $V_P$ is the total phenotypic variance in a trait in a population, $V_G$ is the phenotypic variance attributed to genetic variance, and $V_E$ refers to all other sources of phenotypic variance that are not the genetic variance (Falconer and Mackay 1996). In this simple, dichotomous partitioning, $V_E$ is often misleadingly termed the “environmental variance”, even though it contains more than just environmental variance. But $V_E$ can also be defined sensu stricto as the phenotypic variance attributed only to environmental variance (Scheiner and Goodnight 1984). When the definition of $V_E$ is limited to the phenotypic variance attributed to truly environmental variance, two additional terms are partitioned out in the phenotypic variance equation: $V_{GxE}$, the phenotypic variance attributed to genotype-by-environment interaction, and $V_I$, the residual or error variance that is not explained by any of the other sources of variation (i.e., not explained by $V_G$, $V_E$, or $V_{GxE}$) (Scheiner and Goodnight 1984). We argue that each component of the equation $V_P = V_G + V_E + V_{GxE} + V_I$ can be influenced by epigenetics, and an additional partition, COVGE, can also be influenced by epigenetics. We also show how each of the variance components from the expanded phenotypic variance equation can influence evolution. Therefore, the influences of epigenetics on any of the variance components cannot be safely set aside and ignored when thinking about evolutionary processes and evolutionary outcomes.

**Epigenetics, $V_G$, and evolution**

The genetic variance portion of phenotypic variance, $V_G$, can be dissected into: $V_A + V_D + V_I + V_{Ge}$, where $V_A$ is the additive genetic variance, meaning the portion of the total genetic variance that is inherited in an additive fashion. $V_D$ (dominance variance), $V_I$ (gene–gene interaction), and $V_{Ge}$ (residual genetic variance) are other portions of the total genetic variance (Falconer and Mackay 1996). $V_A$ is the portion of phenotypic variance that is necessary for evolutionary change, and when standardized by the total phenotypic variance, $V_P$, it is defined as the narrow sense heritability ($h^2$). Narrow-sense heritability is important in both agricultural breeding and in evolutionary theory, because it describes variation that is necessary for traits to evolve (Lopez-Fanjul and Garcia-Dorado 2011). The importance of narrow-sense heritability to evolution is represented by the “breeder’s equation”, $R = h^2 S$, where $R$ is the response to selection on a trait, $h^2$ is the heritability of the trait, and $S$ is the strength of selection on the trait (Falconer and Mackay 1996). In this equation, evolution is the response to selection.

To the extent that epigenetic mechanisms contribute additively to the phenotype, then epigenetic variance may be found in the additive genetic variance component, $V_A$, of the total genetic variance. It may be possible to partition epigenetic variance from $V_A$ (see the section below titled, Incorporating epigenetic effects into quantitative genetics studies; Spencer 2002; Santure and Spencer 2006, 2011; Tal et al. 2010; Macdonald 2012; Varona et al. 2015). But
epigenetic mechanisms can also contribute to dominance deviations, because there is no reason to assume that epigenetic re-patterning of chromosomes should have purely additive effects on the phenotype. Therefore, epigenetics can also contribute to the variance in dominance deviations (\(V_D\)) portion of the total genetic variance. The contribution of epigenetics to dominance deviations has not been thoroughly investigated, to our knowledge, but it would helpful to learn more about how much epigenetic mechanisms contribute to \(V_D\).

Epigenetically re-patterned genes may influence the expression of other genes and their effects on the phenotype, and thus epigenetics can influence the epistatic variance, \(V_I\), portion of the total genetic variance as well. Shivaprasad et al. (2012) studied gene expression patterns in the \(F_1\) progeny of a cross between cultivated tomato (\(Solanum lycopersicum\)) and a wild relative (\(Solanum pennellii\)) and found that micro or small interfering (si) RNAs were more abundant in hybrids than in either parent, and that accumulation of such transgressive siRNAs correlated with suppression of the corresponding target genes. In one instance, this effect was associated with hypermethylation of the corresponding genomic DNA. Smith and Weigel (2012) pointed out that many siRNAs are produced from non-coding regions or transposable elements, which diverge more quickly than protein-coding genes and thus provide more opportunity for unexpected genetic interactions. To the extent that gene expression changes are mediated epistatically by epigenetic changes (as in Shivaprasad et al. 2012), epigenetics can be influencing \(V_I\).

Epigenetic effects on \(V_G\) can most obviously influence evolution via their effects on \(V_A\) (the additive component of the total genetic variance), since the response to selection (evolution) in the breeder’s equation depends, in part, on \(h^2\), which is a standardized measure of \(V_A\). But epigenetic effects on \(V_I\) (the epistatic component of the total genetic variance) can also be converted to additive genetic variance by the action of genetic drift and genetic bottlenecks that cause some loci to become fixed, eliminating the interaction effects among loci (Cheverud and Routman 1995; Hill 2017). The epigenetic effects that may be involved in epistatic interactions could be evolutionarily important during speciation events, when founders are isolated into smaller populations and genetic diversity is reduced. Epigenetic variation is reprogrammed through hybridization among species in the wild and in agricultural contexts (Salmon et al. 2005; Salmon et al. 2008), and may strengthen reproductive barriers among species (Smith and Weigel 2012). In summary, epigenetic influences on \(V_G\) can influence evolution, via their effects on \(V_I\) as well as via their effects on \(V_A\); the nonadditive component of \(V_G\) should not be ignored when accounting for epigenetic influences on phenotypic variance and evolution.

### Epigenetics, \(V_E\) sensu stricto, and evolution

Phenotypic plasticity is the ability of a genotype to produce distinct phenotypes in response to different environmental conditions (Pigliucci 2001). It can be quantified for a single individual, or for multiple replicates of the same genotype, across different environments. A “genotype” in this instance can refer to individuals with specific configurations of alleles at a locus. It can also refer to clonal copies of an individual (e.g., Fallopia spp.; Richards et al. 2008; Parepa et al. 2014), apomictic offspring (e.g., Taraxicum; Verhoeven and van Gurp 2012) or an inbred strain of a species such as the Landsberg erecta strain of Arabidopsis thaliana (Koornneef and Meinke 2010) or the Sprague–Dawley strain of the rat Rattus norvegicus (Hsu and Lai 2007). Plasticity is typically measured by the phenotype in one environment minus the phenotype in the other environment, and is visualized on a graph as a “norm of reaction” (Pigliucci 2001; Fig. 1). The greater the difference in the phenotype between the two environments, the greater the plasticity. Phenotypic plasticity can also be described as the phenotypic variance in a dataset that is due to differences in the phenotype among two or more different environments. Understood in this way, plasticity is represented by the term \(V_P\) sensu stricto in the partitioning of phenotypic variance (Scheiner and Goodnight 1984).

To the extent that changes in the phenotype in different environments arise from changes in gene expression, epigenetic modifications could be contributing to those changes in gene expression (Duncan et al. 2014). There are examples of epigenetically-mediated plasticity in many taxa, including flowering plants (Geng et al. 2013; Zhang

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**Fig. 1** The phenotype (y-axis) of the same individual/inbred strain/genotype in two different environments (x-axis) as a norm of reaction. The phenotype in Environment 2 minus the phenotype in Environment 1 indicates how much phenotypic plasticity (\(V_P\)) there is.
et al. 2013; Herman and Sultan 2016), yeast (Herrera et al. 2012), honeybees (Kucharski et al. 2008), ants (Bonasio et al. 2012; Simola et al. 2013), mammals (Oberlander et al. 2008; Godfrey et al. 2011; Lim et al. 2012; Hunter et al. 2015), and birds (Lindqvist et al. 2007; Natt et al. 2009; Goerlich et al. 2012; Nätt et al. 2012). Generally speaking, epigenetic modifications are likely a common mechanism of phenotypic plasticity (Nicotra et al. 2010; Richards et al. 2010; Geng et al. 2013; Herman et al. 2014; Herman and Sultan 2016), and can account for some of the phenotypic variance via their effects on response to environment (Sultan 2016), and can account for some of the phenotypic variation in asexual freshwater snail populations of the species *Potamopyrgus antipodarum* from contrasting lake and river habitats. They found that populations exhibit habitat specific differences in shell shape that are adaptive given the water current speeds of their environments. These differences were associated with significant genome-wide DNA methylation differences among snails from different habitats, and the different methylation patterns were repeated in replicate habitats. Because these snails contain few molecular genetic differences, Thorson et al. (2017) hypothesize that the differences in the shell morphology among habitats are due to adaptive phenotypic plasticity that is mediated by epigenetic mechanisms, specifically DNA methylation throughout the genome. It is possible that the observed differences in methylation among habitats are not just environmentally induced but also heritable, but this additional possibility was not tested.

While epigenetic modifications may provide insight into the molecular details of phenotypic plasticity (Nicotra et al. 2010), the two are not entirely interchangeable. As described above, epigenetics is also found in the *V* _G_ part of the dissection of phenotypic variance, which is not a plastic response to environment. Furthermore, phenotypic plasticity can be due to many biochemical mechanisms (reviewed in Kelly et al. 2012), and not all of these mechanisms involve epigenetics. For instance, phenotypic plasticity can be due to the kinetic constraints of enzyme functions in response to temperature. This type of plasticity can be seen in plants, which have optimal temperature ranges for photosynthesis, driven by the kinetic properties of the enzymes and other mechanisms involved (Yamasaki et al. 2002). Therefore, photosynthetic reactions can go slower or faster depending on the temperature regime experienced by the plant, and enhanced growth of a plant under moderately higher temperatures could be due to increased rates of enzymatic reactions and not necessarily to epigenetic mechanisms. Optimal environmental ranges for enzymes are a fundamental aspect of cellular physiology across the domains of life (Geueke and Kohler 2010), so this example is generalizable to any taxon. Similarly, to the extent that an organism grows larger in response to nutrients simply because it has more materials to build more cells and more structures, this does not necessarily result from an epigenetic mechanism mediating the plastic response of organism size to nutrient levels. In summary, some phenotypic variance can be attributed to epigenetic modifications that are not plasticity (like epigenetics influencing *V* _G_ ) and some phenotypic variance is due to plasticity that is not the result of epigenetic modifications.

Epigenetic effects on *V* _E_ *sensu stricto* may not have obvious ramifications for evolution, given that environmental variation does not have a heritable component to it. But epigenetic states are different from DNA sequence variation in that epigenetic states can be environmentally induced and then inherited in the germ line (Morgan et al. 1999; Waterland and Jirtle 2003; Verhoeven et al. 2010; Richards et al. 2012; Frévard et al. 2013; and reviewed in Richards et al. 2017). Thus, epigenetic mechanisms can account for some of the plastic response of a trait to the environment (which is *V* _E_ *sensu stricto*) that is then converted into additive genetic variance. Surprising as it may first seem, epigenetic influences on *V* _E_ *sensu stricto* have evolutionary implications for the calculation of *V* _A_ , and therefore epigenetic influences on *V* _E_ *sensu stricto* may potentially influence evolution.

### Epigenetics, *V* _GE_ , and evolution

The effects of the environment on phenotype can vary by genotype (Fig. 2). In other words, the expression and function of genes can change depending on the specific
alleles comprising a genotype (Gillespie and Turelli 1989). This widespread phenomenon (Baye et al. 2011; De Marais et al. 2013; Rauw and Gomez-Raya 2015), called the “genotype-by-environment interaction”, is partitioned from the total phenotypic variance (Scheiner and Goodnight 1984). Any genotypes in Fig. 2 that have non-zero slopes are displaying plasticity (Pigliucci 2001), and the fact that different genotypes have different slopes indicates that there is genetic variation for plasticity (Scheiner and Lyman 1989). Thus, there are environmental and genetic components in $V_{GxE}$.

Just as there can be epigenetic contributions to the environmental ($V_e$) and genetic ($V_G$) components of phenotypic variance, there can also be epigenetic contributions to the genotype-by-environment interaction ($V_{GxE}$). Take, for instance, the genotypes in Fig. 2 in environment 1. Within that environment, there is genetic variance in the phenotype—the three genotypes have different genotypic means. As described above, epigenetics can be contributing to the genetic variance. Similarly, take the same genotypes in Fig. 2 in environment 2. Within that environment, there is also genetic variation in the phenotype and, therefore, epigenetic mechanisms can be contributing to the genetic variance within that environment. Now take genotype 1 and examine how its phenotype changes across environments. This is phenotypic plasticity, and epigenetic mechanisms can contribute to phenotypic plasticity, as described above.

The signature of epigenetic effects on $V_{GxE}$ has been documented in a few studies. Bossdorf et al. (2010) treated a set of natural inbred lines of Arabidopsis thaliana with the demethylating agent 5-azacytidine and examined the consequences of this treatment for plasticity of plant traits to nutrient levels. By looking at how demethylation influenced the plastic responses of the inbred lines as compared to their control counterparts, they found that the effect of epigenetic re-patterning on plastic responses varies among inbred lines (Fig. 3). But the changes in the plastic responses only weakly matched what would be expected by the relatedness of the lines to one another. This suggests that the inbred lines differed in their methylation patterns in ways that are at least partly decoupled from their DNA sequence similarity. Similarly, Tatra et al. (2000) investigated the response of DNA methylation to irradiance-mediated plasticity of stem elongation in two ecotypes of Stellaria longipes, using a multifactorial design of light and 5-azacytidine. They found that the plastic responses to light were altered differently by demethylation among the two different ecotypes. In another study, Herman and Sultan (2016) used demethylation in Polygonum persicaria to show that the inheritance of adaptive response in offspring of drought-exposed plants was mediated by DNA methylation. The effect of demethylation on inheritance of the adaptive response varied among inbred lines. These studies demonstrate empirically that epigenetic modifications can account for some of the differences in plastic responses among genotypes. To the extent that epigenetic differences are inherited, epigenetic mechanisms may account for some of the inherited differences in plasticity among different genotypes.

The genotype-by-environment interaction can be thought of as genetic variance for plasticity, and can be broken down into additive and dominance variance components. This allows for calculating the heritability (in the narrow-sense) of plasticity, which can evolve (Scheiner and Lyman 1989). There are many examples of the evolution of plastic responses to the environment (Dudley and Schmitt 1995; Gotthard and Nylin 1995; Pigliucci 2001). As such, epigenetic influences on $V_{GxE}$ can influence evolution, because $V_{GxE}$ contains heritable genetic variation for plasticity and some of this heritable variation can be due to epigenetic mechanisms.

Another way in which $V_{GxE}$ can influence evolution is through genetic assimilation. Genetic assimilation is the process where a phenotype originally produced in response to an environmental stimulus later becomes genetically encoded via selection, and the plastic response is lost, due to a directional shift in the environment (Waddington 1956; Pigliucci and Murren 2003; Crispo 2007). Phenotypic plasticity can thus serve as a bridge for populations to adapt to changing environments, and evolution via genetic assimilation can later developmentally hard-wire the induced plastic responses. Under a scenario of genetic assimilation, selection first acts on genetic variation for plasticity ($V_{GxE}$), favoring individuals that are able to alter...
their phenotype in ways that better match the new environment; if the ancestral environment is later lost, selection will subsequently favor constitutive induction of the novel phenotype, due to costs associated with maintaining the metabolic/developmental machinery for a plastic response to the environment (Pigliucci and Murren 2003). To the extent that epigenetic variation among individuals account for some of the extent that epigenetic variation among individuals accounts for the environments from their parents (Oyama et al. 2001), the inheritance of parental environments, means that the environmental induction of epigenotypes, combined with some of the genetic variance in plasticity that selection can act on in the new environment. This plasticity can later become genetically encoded and constitutive via evolution by genetic assimilation.

**Epigenetics, COVGE, and evolution**

When the occurrence of specific environmental conditions depends on an organism’s genotype, or vice versa, there is a genotype–environment covariance (COVGE; Falconer and Mackay 1996). The genotype–environment covariance combines the genetic variance together with the environmental variance (Falconer and Mackay 1996). A standardized version of this covariance is often presented in the literature as a genotype–environment correlation (r_{GE}; Jaffe and Price 2007). Because epigenetic mechanisms can play a role in genetic variance, epigenetic mechanisms can also play a role in COVGE, which is based, in part, on genetic variance. As with each of the other components of the phenotypic variance formula, the epigenetic contribution to COVGE must be accounted for, either by cross-factorial experimentation where each genotype experiences each environment equally, or by explicitly estimating COVGE as part of partitioning of phenotypic variance. If COVGE is not accounted for, then the phenotypic variance it introduces can confound estimates of \( V_G \) or \( V_{GxE} \) (Falconer and Mackay 1996; Jaffe and Price 2007). A covariance between genotype and environment is particularly likely to arise when considering epigenetic mechanisms of phenotypic variance, because epigenetic changes can be environmentally induced and then inherited (Morgan et al. 1999; Waterland and Jirtle 2003; Verhoven et al. 2010; Richards et al. 2012; Frésard et al. 2013; and reviewed in Richards et al. 2017). Given that organisms tend to inherit their environments from their parents (Oyama et al. 2001), the environmental induction of epigenotypes, combined with the inheritance of parental environments, means that the prevalence of different epigenotypes should be particularly nonrandom with respect to the environments in which the epigenotypes are found. Without the COVGE term in the phenotypic variance formula, the mathematical assumption is that the (epi)genetic variance and environmental variance are unrelated, so the addition of this term is important for properly partitioning phenotypic variance when the asumption of (epi)genotype-environment independence is violated.

Incorrect estimation of \( V_G \) and \( V_{GxE} \) could lead to an incorrect heritability estimate of a trait or trait plasticity, and the response of a trait to selection (i.e., evolution) could be miscalculated. The genotype–environment covariance is often ignored in quantitative genetic studies, because it is assumed to be of negligible importance (Falconer and Mackay 1996). However, COVGE could be a large portion of the total phenotypic variance when epigenetic effects are involved, due to induction of epigenetic changes by the environment and the subsequent inheritance of those epigenetic changes (Bonduriansky and Day 2009; Slatkin 2009; Geoghegan and Spencer 2012, 2013; Richards et al. 2017). We propose that accounting for the epigenotype–environment covariance could be accomplished by first clustering individuals, based on their molecular epigenetic marker similarities (using a Bayesian clustering algorithm for example; Pritchard et al. 2000), and then calculating the epigenotype–environmental covariance by combining the information about the environments that the individuals experienced, the individuals’ phenotypes, and the epigenetic clusters that the individuals were assigned to. Development of this or similar approaches is beyond the scope of this paper, but methods for parsing out the epigenotype–environment covariance are needed to properly parse phenotypic variance into its various constituents. It should be possible to calculate the epigenetic contribution to COVGE, but the methods for doing so have not yet been fleshed out.

**Epigenetics, \( V_{\epsilon} \), and evolution**

The residual or error variance, \( V_{\epsilon} \), as defined by Scheiner and Goodnight (1984), includes only measurement error, microenvironmental variance, and developmental stochasticity. Microenvironmental variance refers to environmental variance that was not explicitly accounted for in the study design. If there are a large of amount of environmental effects that are not explicitly accounted for by \( V_E \ sensu stricto \) in the study design, then the microenvironmental variance can actually be larger than \( V_E \ sensu stricto \) (Mulder et al. 2013). Yet the microenvironmental variance is an invisible constituent of \( V_{\epsilon} \) in the phenotypic variance partitioning, because by definition it was not measured. Given that epigenetic changes can influence the environmental variance of the phenotype, as described above, epigenetic changes can also influence microenvironmental variance of the phenotype, because microenvironmental variance is just phenotypic variance attributable to environmental axes that were unmeasured. This also means that epigenetic effects on microenvironmental variance can influence evolution in the same was that epigenetic effects
on environmental variance can influence evolution, as described above.

**Methods for incorporating epigenetic effects into quantitative genetics studies**

With the growing evidence that epigenetic changes can have different ramifications for evolution than DNA sequence changes, methods are being developed to actually incorporate epigenetic effects into models of quantitative genetics. Models of genomic imprinting have found that the estimation of the total genetic variance is not affected by imprinting, but the partitioning of the genetic variance in the phenotype changes and the response to selection for a highly imprinted trait cannot be predicted by the breeder’s equation (Spencer 2002; Santure and Spencer 2006, 2011). Day and Bonduriansky (2011) propose a model for non-genetic inheritance more generally based on the Price equation (Price 1972) that emphasizes the interactions between genetic and epigenetic effects with the following parameters for both (1) the effects of selection, (2) changes that occur in transmission from the parent to offspring generation (“reproductive transmission”) and (3) changes that occur in the parent generation (“survival transmission”). This model allows for changes in DNA sequence, the fact that some genomic contexts are more likely to acquire epigenetic effects than others, and the change in the epigenetic component is dependent on genotype. They found that epigenetic variation can determine the pattern of phenotypic variation and that phenotypic change can be decoupled from the dynamics of the genotype. Slatkin (2009) modeled disease risk as a function of diallelic molecular epigenetic processes, and particularly the rate of environmental induction (the probability of the epigenetic state being induced) by adding a term, $V_C$, to the equation $V_P = V_G + V_E$, that reapportions some of the phenotypic variance from $V_G$ and $V_E$. Calculating $V_C$ requires determining the phenotypic covariances between parents and offspring, between sibs, and between uncles and nephews. The design allows for estimation of the heritable epigenetic variance and epigenetic transmissibility using information about the number of opportunities for epigenetic reset between generations, and assumptions about environmental induction. They introduce a new term, $r^2$, which is the epigenetic heritability, or the proportion of phenotypic variance attributable to potentially heritable epigenetic variance. Different from standard heritability ($h^2$), the contribution of epigenetic heritability ($r^2$) to parent-offspring similarity depends on the transmissibility coefficient: if the epigenetic states tend to be reset frequently, then the impact of $r^2$ on parent-offspring similarity is low; if the epigenetic states tend to be transgenerationally stable, then the impact of $r^2$ on parent-offspring similarity is high. The introduction of a new type of heritability with different ramifications for similarity among relatives could enhance our understanding of evolutionary processes by describing how different components of heritable phenotypic variation respond to selection.

The Tal et al. (2010) study provides a practical way forward for parsing out epigenetic influences in quantitative genetic studies. But like in other studies, Tal et al. (2010) do not account for the influence of epigenetic differences on $V_G + V_E$. Perhaps the Tal et al. (2010) formula could be reformulated to account for epigenetic influences on $V_G + V_E$ by partitioning $V_G + V_E$ and $V_G$ in a similar manner to what they describe for partitioning $V_C$ from $V_G$ and $V_E$ (sensu lato). The development of such a method lies outside the scope of this paper, but it would be a fruitful area of future research. Similarly, Richards et al. (2017) recommend that future studies on phenotypic variance should parse out a higher-order interaction term involving epigenetic effects, which could be something like $V_G + V_{E, C}$. This variance component would account for the ways in which DNA-encoded differences in plasticity change depending on the epigenetic effects.

It is important to note that Tal et al. (2010) define epigenetic inheritance broadly to include the transmission of phenotypic variation by any means besides transmission of DNA sequence variation. Their definition of epigenetics is broader than the more mechanistic, cellular definition of epigenetics that we have described (namely, chemical modifications of chromatin or transcribed DNA that can influence gene activity and expression without changes in DNA sequence; Jablonka and Raz 2009; Kilvitis et al. 2014), and their definition includes additional parental effects (Kirkpatrick and Lande 1989; Wolf and Wade...
It may be that there is no way in principle to separate epigenetics effects *sensu stricto* from other non-genetic forms of inheritance using a purely phenotypic dataset and a pedigree. Yet, despite potentially overestimating the epigenetic effects *sensu stricto*, the Tal et al. (2010) approach can be used to set upper bounds on the amount epigenetic influence on phenotypic variance and provide data to generate hypotheses for future research. For instance, they give the example of rapidly increasing heritability in inbred lines (Grewal 1962; Hoi-Sen 1972; Lande 1975), where one may expect $V_A$ (and therefore $h^2$) to be small due to a lack of DNA sequence diversity. The classic interpretation is that the inbred lines must have a DNA mutation rate that is three orders of magnitude higher than is normally expected in order to increase the heritability and increase it that quickly. But if the estimates of $V_C$, $r^2$, and the transmissibility coefficient are greater than zero, then the heritability may be partially generated by epigenetic mechanisms. The specific epigenetic mechanisms that contribute to heritability in the inbred lines could be examined in follow-up studies that use epigenomic mapping techniques (e.g., Zhang et al. 2013; Cortijo et al. 2014; Kooke and Keurentjes 2015). Overall, the Tal et al. (2010) study provides a pragmatic approach for teasing out epigenetic influences on phenotypic variance and heritability, but the methods developed so far are imperfect and require further development to accurately quantify epigenetic influences.

**Conclusions**

The modern era of epigenetics research presents opportunities for advancing our understanding of evolutionary processes, just as the genomics era more generally has greatly influenced evolutionary disciplines. We have demonstrated how epigenetic phenomena can contribute to each of the components of phenotypic variance in the expanded formula: $V_G$, $V_E$, $V_{GE}$, $V_f$, and $COV_{GE}$. It is important to consider the influence of epigenetic phenomena on phenotypic variance when hypothesizing about the mechanisms behind phenotypic variance observed in a quantitative genetic study. One cannot assume that only DNA sequence-based phenomena account for patterns when epigenetic and other non-genetic phenomena could be playing a role as well; and accounting for parental effects will not necessarily capture epigenetic phenomena. The contribution of epigenetic mechanisms cannot be easily cast aside, especially given the increasing reliance on molecular mechanisms as a part of our understanding of evolutionary processes, and the fact that epigenetic mechanisms can have different ramifications for evolution from DNA sequence-based mechanisms. Some existing studies have developed methods for partitioning epigenetic variance from phenotypic variance, which represents a step in the right direction, but we need more methods that carefully consider all of the areas where variance due to epigenetic factors may need to be accounted for. The challenge of developing methods to parse out all of the ways that epigenetics can contribute to evolutionary models will require significant effort, but the difficulty of deciphering the role of epigenetic mechanisms does not preclude its importance.

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**Compliance with ethical standards**

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

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