PERSPECTIVES

Rethinking inheritance, yet again: inheritomes, contextomes and dynamic phenotypes

N. G. Prasad¹, Sutirth Dey², Amitabh Joshi³ and T. N. C. Vidya⁴

¹ Department of Biological Sciences, Indian Institute of Science Education and Research Mohali, Knowledge City, Sector 81, SAS Nagar, P.O. Manauli, Mohali, Punjab 140 306, India.

² Population Biology Laboratory, Biology Division, Indian Institute of Science Education and Research Pune, Dr. Homi Bhabha Road, Pune 411 008, India.

³ Evolutionary Biology Laboratory, Evolutionary and Organismal Biology Unit, Jawaharlal Nehru Centre for Advanced Scientific Research, Jakkur P.O., Bengaluru 560 064, India.

⁴ Animal Behaviour and Sociogenetics Laboratory, Evolutionary and Organismal Biology Unit, Jawaharlal Nehru Centre for Advanced Scientific Research, Jakkur P.O., Bengaluru 560 064, India.

For correspondence: Email: tncvidya@jncasr.ac.in; tncvidya@gmail.com

© All authors contributed equally to this work. This is contribution no. 1 from FOGEG (see Acknowledgments for details).

Running title: An extended view of inheritance

Key-words: Extended evolutionary synthesis; non-genetic inheritance; phenotypic variance partitioning; cultural evolution; epigenetics; parental effects.
Abstract

In recent years, there have been many calls for an extended evolutionary synthesis, based in part upon growing evidence for non-genetic mechanisms of inheritance i.e. similarities in phenotype between parents and offspring that are not due to shared genes. While there has been an impressive marshalling of evidence for diverse forms of non-genetic inheritance (epigenetic, ecological, behavioural, symbolic), there have been relatively few studies trying to weld the different forms of inheritance into a common conceptual structure, a development that would be important to formalizing elements of the extended evolutionary synthesis. Here, we develop a framework for an extended view of inheritance and introduce some conceptual distinctions that we believe are important to this issue. In this framework, the phenotype is conceived of as a dynamic entity, its state at any point in time resulting from intertwined effects of previous phenotypic state, and of hereditary materials (DNA and otherwise) and environment. We contrast our framework with the standard gene-based view of inheritance, and also discuss our framework in the specific context of recent attempts to accommodate non-genetic inheritance within the framework of classical quantitative genetics and the Price equation. In particular, we believe that the extended view of inheritance and effects on the phenotype developed here is particularly well suited to algorithmic modeling for simulation studies of evolutionary dynamics.
Introduction

In recent decades, increasing attention within evolutionary biology is being focused on the issue of non-genetic (i.e. not based on gene sequence variation) inheritance (reviewed at length in Jablonka and Lamb 2005; Pigliucci and Müller 2010). There is increasing evidence for the inheritance of environmentally induced epigenetic states from parents (reviewed in Bonduriansky and Day 2009; Jablonka and Raz 2009), as well as for the passing on of culturally acquired (i.e. learnt) behaviours to offspring and, indeed, other peers, referred to as cultural inheritance or transmission (Cavalli-Sforza and Feldman 1981; Richerson and Boyd 2005; Jablonka and Lamb 2005; El Mouden et al. 2014). The literature in these areas is growing but, as a result of coming from diverse subdisciplinary backgrounds of inquiry, it is often confusing with regard to terminology and underlying concepts. In this brief essay, we delineate some concepts and conceptual distinctions, and attendant terminology, that we believe may help ameliorate some of the confusion in attempts to bring together various forms of inheritance into a coherent and expanded view of evolutionary dynamics. Much of what we are drawing upon has been said before: what we hope to accomplish is to (a) highlight some specific aspects of present attempts to extend the evolutionary synthesis with regard to extra-genic inheritance where we believe that conceptual clarity is still lacking, and (b) make some suggestions as to how that conceptual clarity can be attained. Given the numerous recent reviews of different aspects on non-genetic inheritance and its evolutionary implications, the list of references is not comprehensive and is somewhat biased towards broad reviews or books wherein leads to most of the primary literature can be found.
Following the rediscovery of Mendel's laws, the study of heredity gradually narrowed to an almost exclusive focus on genic inheritance, especially after Johannsen's (1911) delineation of the concepts of genotype and phenotype, mirroring the hard dichotomy of germline and soma, laid down by Weissmann (1904), that excluded any possibility of the inheritance of acquired characteristics (discussed in Schwarz 2008). Subsequently, this narrower, Mendelian, version of heredity was what was reconciled with a neo-Darwinian (sensu Romanes 1888) emphasis on natural selection as the principal mechanism of adaptive evolution, largely by the work of Fisher (1930), Haldane (1932) and Wright (1932), becoming a major foundation of the Modern Synthesis (Mayr 1992). The importance of genes as explanatory factors in biology got further strengthened following the advent of molecular genetic understanding of DNA structure, replication, expression and regulation. Thus, as heredity was central to the mechanism of natural selection, the narrowing of heredity to genic inheritance naturally led to the gene-centric bias of much modern evolutionary thinking, at least insofar as it pertained to adaptive evolution within populations (i.e. microevolutionary change) (discussed in Gould 2002; Amundson 2005; Bonduriansky 2012).

While the gene-centric version of microevolutionary theory has been extensively criticized for being limited in scope, especially in the light of ever-increasing evidence for non-genetic inheritance (e.g. Richerson and Boyd 1985; Odling-Smee et al 2003; Jablonka and Lamb 2005; Pigliucci and Müller 2010), certain aspects of this theory, we believe, have not received as much attention from critics as they should have. First, in addition to the framework of population genetics in the strict sense, there is also the essentially phenotypic framework of quantitative genetics, arising from Fisher's (1918) seminal paper on the correlations between relatives for
polygenic traits following Mendelian inheritance. Interestingly, several critics of the Modern
Synthesis ignore the quantitative genetics tradition altogether (e.g. Amundson 2005; Jablonka
and Lamb 2005; Laland et al. 2014), although it is considerably more relevant than population
genetics is to understanding adaptive evolution. While quantitative genetics did attempt to root
its statistical analysis of trait correlations and evolution in Mendelian genetics, its most important
insights – the notion of breeding value for fitness and its variance as determinants of adaptive
evolutionary responses to fitness differences among individuals – are essentially not dependent
on any underlying mechanistic model of inheritance. Indeed, the statistical approach of
quantitative genetics, which effectively black-boxed the details of how genotypes affect
phenotypes, reached its later, even more generalized, fruition in the Price (1970, 1972) equation
that makes no assumption about the pattern of inheritance at all, subsuming it into a correlation
of phenotypic values between individuals and immediate descendants. What the results from this
statistical tradition clarify is that ultimately what matters in determining the response to a given
selective scenario is not so much the genes that pass from parent and offspring but rather their
statistical effects on offspring phenotype. This important distinction is rooted in the fact that
heredity is ultimately about phenotypic similarity between parents and offspring, even though
one major underlying reason for such similarity is the material genes that pass from parent to
offspring, albeit subject to the vagaries of mutation, recombination, sex and chance. Thus, what
an individual inherits, in terms of a genetic endowment leading to the propensity towards certain
phenotypic values, is usually different from what the same individual transmits to its offspring in
terms of expected effects on the offspring phenotype. This principle is reflected in the clear
distinction between the genotypic value (G) and the breeding value (A) of a trait in quantitative
genetics (Falconer and Mackay 1996). This distinction has implications for the development of
an extended theory of evolutionary change incorporating varied non-genetic forms of
inheritance. There are now some studies that use the framework of quantitative genetics or the
Price equation to incorporate non-genetic inheritance into models of evolutionary change
(Bonduriansky and Day 2009; Helanterä and Uller 2010; Danchin et al. 2011; Santure and
Spencer 2011; El Mouden et al. 2013), and we will discuss them in subsequent sections.

A second aspect of the gene-centric view of the evolutionary process that we believe has
not received much critical attention is the essentially static conception of genotype and
phenotype, at least within an individual's lifetime. Even the notion of the genotype-phenotype
(G-P) map, which was developed as a reaction to the black-boxing of the details of development
in Mendelian genetics (Alberch 1991), is an attempt to define a relationship (a map) linking
genotypic space and phenotypic space. Despite the complexity of this relationship, it is
nevertheless conceived of as a relationship between two spaces that are static within an
individual's lifetime, though not over evolutionary timescales.

In the next section, we briefly review the standard gene-based view of inheritance,
emphasizing some points that we will draw upon in order to develop an extended view of
inheritance. The subsequent section will be devoted to incorporating non-genetic inheritance into
a conceptual framework of heredity, and the shaping of the phenotype, that will stress the
dynamic nature of phenotypes even within the lifetime of an individual. In the final section, we
will discuss other studies that have addressed the issue of non-genetic inheritance and its
evolutionary implications and highlight similarities and differences between our view and theirs,
in particular highlighting conceptual distinctions that we believe are important and have not been
drawn earlier. We will also discuss some of the broader implications of the extended view of inheritance developed here.

**The standard gene-based view of inheritance**

Before developing an extended view of inheritance, we briefly reiterate what the standard gene-based model of inheritance is, and how quantitative genetics distinguishes between the genotypic value and breeding value of a trait in an individual. The fundamental components of this view are that information regarding the specification of phenotypes (whether as blueprint or developmental program) is transmitted to offspring from parent(s) via the inheritance of genomes, and that the specification of the phenotype by the genome can be affected by the environment which includes biotic and abiotic components (Fig. 1). Direct environmental effects on the phenotype of an individual (solid dark blue arrow from E to the octagon around P), however, cannot be passed on to that individual’s offspring, as only genes are transmitted between generations. We note that, in this view, environment in a broad sense can affect the genome (solid dark orange arrow from E to the circle around G in Fig. 1) by inducing mutations or transpositions, or via horizontal gene transfer from conspecifics or heterospecifics. We will not treat acquisition of genetic material by an individual via horizontal gene transfer as constituting inheritance from the viewpoint of that individual, as it does not involve inherited material acquired at conception via transmission from parent(s). Of course, once acquired by horizontal gene transfer, such genetic material can subsequently be transmitted by that individual to its own offspring, becoming part of their inheritance. We next stress a few specific points about this gene-based view of inheritance that we believe will be relevant to the development of an extended view of inheritance in the next section.
Although the organism and its phenotype are central in ecology and evolution because Darwinian fitness typically accrues to the individual as a result of its phenotype and how well it functions in a given environmental context, evolutionary dynamics are often tracked at the genetic level. This is because, at least for typical Mendelian traits, the G-P mapping is simple, and the transmission of genes is amenable to mathematical representation. For more complex polygenic phenotypes, a phenotypic approach (quantitative genetics) is taken, although here too there is often an underlying genetic model (see (iv, vi) below).

The genome plays two distinct roles here: that of a transmissible genome and an expressible genome. During transitions between generations, the genome acts as the transmissible material of heredity, whereas during an individual's lifetime the genome acts as an expressible material that directs the generation of the phenotype. While there is a degree of materialistic continuity to the genome between generations, functionally the expressed genome can vary temporally and spatially within the lifetime and the body of the individual, respectively (see also (v) below).

When an individual reproduces, it typically does not transmit the genome it inherited from its parents. This difference between the inherited and transmitted genomes arises due to (a) changes in the genome due to mutations, transpositions and horizontal gene transfer during the individual's lifetime, and (b) the effects of recombination and meiosis during sexual reproduction.

In the quantitative genetics approach, an individual's phenotypic value for a trait is conceptualized as being made up of a genotypic value and an environmental effect, with the mean environmental effect assumed to be zero. The genotypic value is the expected phenotypic value of individuals of a given genotype exposed to all possible environments (Falconer and Mackay 1996). Thus, there is a material partitioning of causes of phenotypic value into a
genotypic value and an environmental deviation from it. The genotypic value of an individual is, however, not what the individual transmits in terms of phenotypic value propensity to its offspring. Essentially, in a statistical partitioning of effects, the genotypic value is further divided into a transmissible component (the breeding value or additive component) and a non-transmissible component traditionally ascribed to genotype-by-genotype interactions. The breeding value of an individual is a measure of how much the mean phenotypic value of that individual's offspring is expected to differ from the population mean phenotypic value, were that individual to mate at random within the population. In terms of phenotype, the breeding value is what an individual passes on to its offspring, on an average, and it is a function of the individual's genotype as well as the genotypic composition of the population (Falconer and Mackay 1996).

(v) Thus, in terms of the phenotype, the transmissible genome of an individual is reflected in its breeding value (which is dependent also on the genotypic composition of the population) whereas its expressible genome is reflected in its genotypic value, which is independent of the populational context.

(vi) The partitioning of phenotypic value into breeding value and a non-transmissible component, and the analogous partitioning of phenotypic variance in a population into a variance of breeding values (the so-called additive genetic variance) and a non-transmissible variance are purely phenotypic, reflecting the degree to which phenotypic differences among individuals are transmitted to the next generation. In this sense, quantitative genetics is essentially a phenotypic theory. Under the assumption of a large number of Mendelian genes contributing to phenotypic value, however, corresponding genetic models can also be developed and have been used in quantitative genetics (Falconer and Mackay 1996).
In this standard gene-based view, the phenotype is conceived of as a largely static entity, at least within the individual's lifetime.

One last point we wish to make before moving on to developing an extended view of inheritance is that the framework of quantitative genetics, simple though it is, can incorporate many biological phenomena that are often mentioned in the extended evolutionary synthesis literature as being beyond the gene-based view. For example, at least from the perspective of effects on phenotypic value, an environmental effect and genotype-by-environment interaction reflect phenotypic plasticity and genetic variation for it. Similarly, many types of adaptive niche construction can be conceptualized as a subset of positive genotype-environment covariance.

An extended view of inheritance

We now develop an extended view of inheritance that takes cognizance of phenomena like transgenerational epigenetic inheritance, parental effects, ecological inheritance and cultural inheritance. We follow the categorization of Danchin et al. (2011); types of non-genetic inheritance have been categorized differently (e.g. Bonduriansky and Day 2009; Helanterä and Uller 2010), but these differences do not affect the development of our view and we defer their discussion to the next section. We first describe some terms and conceptual distinctions we believe to be helpful in discussing such an extended view of inheritance. Our primary focus throughout remains the phenotype, and the acquisition from, and contribution to, other individuals of effects on the phenotype. Rather than the distinction between vertical, horizontal and oblique transmission (e.g. Bonduriansky and Day 2009; Helanterä and Uller 2010), we believe it is important to distinguish between heritable phenotypic effects acquired as a material
endowment at conception and those acquired subsequently in an individual's lifetime. Consequently, we will distinguish, in a manner similar to the gene-based view, between inheritome, phenotype and environment (Fig. 2). In Fig. 2, time runs from top to bottom and, at conception, the focal individual has an inheritome $I_1$, inherited from its parent(s) (dashed dark orange arrow at top). The inheritome includes DNA sequence, as well as any epigenetic modifications of DNA/chromatin and cytoplasmic mediators of parental effects. The focal individual also receives a phenotype $P_1$ as a parental endowment (dashed dark blue arrow at top), comprising, for example, cytoplasmic components and cellular structure. The environment is conceptualized as a dynamic entity encompassing all individuals and the physical and cultural (if applicable) backdrop within which they live. Clearly, though individuals live and die, the environment has a continuity, while subject to changes, that transcends individual lifetimes: it is a superset to the other categories depicted in Fig. 2. When examining inheritance/transmission of phenotypic variations from the point of view of a focal individual (Fig. 2), we refer to its local environment at a given time step $i$ as its contextome at that time step ($C_i$). We treat the initial phenotype as distinct from the inheritome because, over the course of a lifetime, the phenotype of an individual undergoes vastly more change than its inheritome. For example, the phenotype of a multicellular organism changes much more than the inheritome over the course of developing from a zygote to an adult. The inheritome, on the other hand, does have considerable integrity across the lifetime. We propose that the terms *inherit* and *transmit* be used solely in the context of an inheritome being received from parent(s) and inheritomic components being passed on to offspring, respectively. We propose using the terms *acquire* and *contribute* to refer to the receiving and giving of changes to the inheritome and/or phenotype that occur after conception. Of course, changes to the inheritome that are acquired, in this sense, can subsequently be
transmitted and therefore inherited by offspring. We further propose a distinction between parents and peers in terms of this distinction between inheritance and acquisition. Parents transmit phenotypic effects via the inheritome and, transiently, through the initial phenotype, whereas peers contribute phenotypic effects via the inheritome, phenotype and contextome. Biological parents, therefore, can affect offspring phenotypes both as parents, at conception, and as peers, subsequently. This distinction is similar to Bonduriansky and Day's (2009) use of the term inheritance solely for vertical transmission, and differs from other, broader, usages (e.g. Odling-Smee et al. 2003; Jablonka and Lamb 2005; Helanterä and Uller 2010; Danchin et al. 2011). However, our usage is narrower than that of Bonduriansky and Day (2009) because we restrict inheritance to heritable phenotypic effects acquired as a material endowment at conception.

In Fig. 2, the set of arrows between $I_1$, $I_2$, $P_1$, $P_2$, $C_1$ and $C_2$ encompasses the various ways in which inheritome, phenotype and contextome affect phenotypes, either directly (arrows impinge upon $I$, $P$ or $C$) or in an interacting manner (arrows impinge upon other arrows linking $I$, $P$ or $C$). This pattern of potential phenotypic effects of $I$, $P$ and $C$ is iterated at every time step, as indicated by the dashed grey arrow labeled 'time' connecting $I_2$, $P_2$, and $C_2$ to $I_n$, $P_n$, and $C_n$. Within a given time step, the specific expressed inheritome at that time step directs the generation of the phenotype at that time step, also subject to interactions with the contextome at that time step. In contrast to the gene-based view (fig. 1), however, the phenotype is also directly affected by the phenotype in the previous time step in a manner also subject to interactions with the contextome in the previous time step. Moreover, the phenotype can also be affected directly by the contextome in the previous time step. Thus, in this view, the phenotype during the lifetime
of an individual is a dynamic flow from $P_i$ to $P_{i+1}$, mediated by inputs from the inheritome ($I_{i+1}$) and the contextome ($C_i$ and $C_{i+1}$), which themselves are dynamic flows. Thus, the phenotype initiated at conception undergoes transformation throughout the lifetime of the individual under the joint inputs from its previous state and the also changing inheritome and contextome, and interactions between them, in a manner echoing the 'triple helix' metaphor of Lewontin (2002).

The inheritome and contextome of the focal individual, in this view, are also conceptualized as flows through time. The inheritome may be altered as a result of its own previous state, and the previous state of the phenotype and contextome, as well as interactions among them (solid dark orange arrows between entities in time steps 1 and 2 in Fig. 2). These changes to the inheritome encompass mutations/transpositions (directed or otherwise), resetting of epigenetic marks on DNA and chromatin, or acquisition of cytoplasmic constituents that may affect offspring phenotypes, mediated by either the physical aspects of the experienced environment, or biological interactions with other individuals. This includes a subset of phenomena included under the label of non-genetic inheritance in the all-encompassing formulations (e.g. Odling-Smee et al. 2003; Jablonka and Lamb 2005; Helanterä and Uller 2010; Danchin et al. 2011). We prefer to treat the remaining mechanisms of non-genetic inheritance, broadly construed, as being changes in the contextome, which is also acquired at conception as a local reflection of the environmental state at that point of time. The contextome of an individual is changing as part of the dynamic change in environment (solid green arrow from $C_1$ to $C_2$ in Fig. 2). We note that these changes include modifications to the physical or cultural aspects of the environment due to the effects of phenotypes of other individuals, conspecific or heterospecific (e.g. social interactions, competition, niche construction). In addition, an
individual's contextome can be affected by the same individual's phenotype, subject to interactions with the contextome, and the altered contextome, in turn, can exert effects on subsequent phenotypic states (solid green arrow from $P_1$ to $C_2$ in Fig. 2). In our view, therefore, we are treating ecological and cultural inheritance (\textit{sensu} Danchin et al. 2011) as being encompassed by the changes to an individual's contextome and not as inheritance in our narrower usage. Of course, changes made to an individual's phenotype by its contextome can lead to phenotypic correlations between parent(s) and offspring, as does inheritance, provided the contextome remains the same for the offspring. This is analogous to genotype-environment covariance in quantitative genetics and we do not believe that it is helpful to label this phenomenon as inheritance, especially since these effects are typically mediated by the phenotypes of many peers that are part of the individual's contextome, rather than by one or two parents.

During its lifetime, an individual can, in principle, reproduce at any time step. When it does so, it transmits to its offspring inheritomic materials which, in the case of sexual reproduction, together with inheritomic materials transmitted by the other parent, form the inheritome of the newly conceived offspring. The parents also pass on an initial phenotype to offspring which will then undergo changes through the offspring's lifetime. As was the case for genomes (Fig. 1), what is transmitted/passed on is dependent upon the reproducing individual's inheritome/phenotype at that time step but is also different from it (distinction between $I_2$ and $I'_2$, and $P_2$ and $P'_2$ in Fig. 2). One point of departure from the gene-based view of inheritance here is that the inheritome is more plastic than genomes typically are, since the former are subject to a greater variety of genetic, epigenetic and cytoplasmic changes. Consequently, the source of
inheritomic materials to be transmitted to offspring by an individual during reproduction at any
time step is likely to vary from time step to time step, in contrast to the germline genome which
is assumed to be relatively unchanged through life, barring rare mutations.

Discussion

In this final section, we discuss some implications of the extended view of inheritance developed
in the preceding section, especially in the context of earlier attempts to formalize theories of non-
genetic inheritance. There are two aspects in which our formulation differs from much of the
previous work on this theme. The first is our separation of inheritomic and contextomic
mediation of phenotypic correlations between parents and offspring, and our restriction of the
term inheritance to describe only the former. The second is our emphasis on the inheritome, and
especially the phenotype and contextome, as dynamic flows rather than static entities. We will
now discuss some implications of these two emphases.

There are two approaches that have been taken to formalizing the evolutionary
consequences of the various types of non-genetic inheritance mechanisms. One is to treat each
category – epigenetic inheritance, parental effects, ecological inheritance and cultural inheritance
– separately, as is done by Jablonka and Lamb (2005). The other approach is to seek a combined
framework within which all forms of non-genetic inheritance can be formalized with regard to
their evolutionary implications. This approach has been taken from a quantitative genetics
perspective by Bonduriansky and Day (2009) and Danchin et al. (2011), and using the
framework of the Price equation by Helanterä and Uller (2010). We note that the important
elements of quantitative genetics framework for explaining adaptive evolution can be derived as
special cases assuming Mendelian inheritance from the Price equation (Frank 1997).
Bonduriansky and Day (2009) treat the change in mean phenotypic value in a population between subsequent generations as being divisible into components due to the change in mean additive genotypic value and in the mean non-genetic component of inheritance. Danchin et al. (2011: Box 4) use a partitioning of variance approach to partition phenotypic variance in the population, $V_P$, into transmitted genetic variance ($V_G$ in their notation, $V_A$ in standard quantitative genetic notation: Falconer and Mackay 1996), various kinds of transmitted non-genetic variance ($V_{TNG}$) combining phenotypic variance components variance ascribable to epigenetic, parental, ecological and socio-cultural inheritance, and a non-transmitted component. They propose to call the fraction of phenotypic variance that is transmissible $[(V_A + V_{TNG})/V_P]$ the 'inclusive heritability' of a trait (Danchin and Wagner 2010) and suggest that it “quantifies the whole evolutionary potential of a trait and can be seen as the result of both genetic and non-genetic heritability” (Danchin et al. 2011). They further suggest that it may be possible to estimate different components of the transmissible phenotypic variance through extensions to classic breeding experiments that also include multiple environmental contexts thought to play a role in non-genetic inheritance, and by tracking epigenetic changes. It is not clear to us that the kinds of partitionings envisaged by Bonduriansky and Day (2009) and Danchin et al. (2011) will actually be experimentally feasible in the context of ecological and cultural inheritance.

Helanterä and Uller (2010) fit various types of non-genetic inheritance mechanisms into the Price equation and try to categorize them into clusters based on the effect each type of non-genetic inheritance has on the different terms in the Price equation (Table 1 in Helanterä and Uller 2010). They find that their approach suggests a clustering different from that into genetic,
epigenetic, behavioural and symbolic inheritance (Jablonka and Lamb 2005). They find three clusters that group mechanisms of non-genetic inheritance that will be similarly incorporated into a Price equation framework, and these three clusters are differentiated by whether “inheritance” is by “vertical transmission” (parent to offspring), “induction (environmentally determined changes between parents and offspring) or “acquisition” (phenotypes affected by peers or other sources). These three clusters differ in the way they mediate adaptive evolutionary change of phenotypes across generations, with and without classic responses to selection in the sense of the breeder’s equation in quantitative genetics (Helanterä and Uller 2010).

Our position is intermediate between the two approaches described above. We note that, of the three clusters of Helanterä and Uller (2010), the “vertical transmission” case corresponds to inheritomic transmission, while their other two clusters are encompassed in contextomic continuity, coupled with contextome effects on phenotype, in our framework. Similarly, the three terms $V_G$, $V_{\text{TEpi}}$ and $V_{\text{PNGE}}$ in the formulation of Danchin et al. (2011: Box 4), encompassing phenotypic variance due to additive genetic variance, epigenetic variance and parental effect variance, respectively, together correspond to variance due to inheritomic transmission in our framework. What Danchin et al. (2011: Box 4) term transmitted ecological and socio-cultural variation is encompassed in contextomic continuity, coupled with contextome effects on phenotype, in our framework. We believe there is a point to separating inheritomic and contextomic effects on the phenotype, for two reasons. First, their effects on adaptive evolutionary dynamics differ, as shown by Helanterä and Uller (2010). Second, following classical quantitative genetics, if we define phenotypic value as being made up of an inheritomic value plus a contextomic value, then it is clear that we need to separately partition each into a
transmissible and non-transmissible component. This is exactly analogous to the partitioning of Bonduriansky and Day (2009: Eqns. 2a,b). The partitioning into a transmissible and non-transmissible component of genotypic value is standard in quantitative genetics. Moreover, there is already considerable work on incorporating epigenetic effects, genomic imprinting and parental effects into quantitative genetics models (Kirkpatrick and Lande 1989; Spencer 2002, 2009; Johannes et al. 2008; Tal et al. 2010; Santure and Spencer 2011), suggesting that the partitioning of inheritomic value into transmissible and non-transmissible components is likely to be feasible. It is not clear to us at this time whether a partitioning of contextomic value that is similar in structure to the partitioning for inheritomic value will be possible and, therefore, we believe it might be best to treat it separately. Experimentally, too, we speculate that estimation of the partitioned phenotypic variances due to contextomic and inheritomic effects may require fairly different kinds of designs. A framework analogous to classical quantitative genetics directly suggests possible experimental designs to estimate transmitted versus non-transmitted components on inheritomic variance. For contextomic effects, a completely mechanism free approach like the Price equation may be more suited, with the drawback that it does not lend itself to suggesting designs for experiments.

Our explicit conceptualization of inheritome, contextome and phenotype as flows in time rather than static entities during an individual's lifetime is another point of departure from many earlier treatments. In particular, this conceptualization suggests that the G-P map metaphor is fundamentally misplaced, as a map is a relation between static entities. In our framework, the inheritome-phenotype map is itself a flow in time and we believe further work is needed to model this relationship more accurately as a dynamic one. One implication of this view of
dynamic phenotype is for genome wide association studies (GWAS). It has already been pointed out that the problem of “missing heritability” (Maher 2008) is potentially explained if there is a substantial non-genetic component to inheritance in the sense of parent-offspring similarity in phenotype (Danchin et al. 2011). If phenotypes are viewed as changing in time, based largely on previous phenotypic state, modulated by inheritomic and contextomic effects on the phenotype, then it is very likely that, at least for complex traits, the best predictors of phenotypic state might be previous phenotypic states rather than genotypes. If this speculation is correct, it is a potential explanation for the efficacy of traditional holistic systems of medicine (like Ayurveda in India) in dealing with complex so-called life-style diseases, as these systems try to correlate diseased phenotypic states with a constellation of previous phenotypic states along different trait-axes.

Overall, we believe that the framework for an extended view of inheritance that we have developed here provides a good basis for thinking about specific lines of inquiry relating to the conceptual partitioning of inheritomic and contextomic values, as well as related experimental approaches for obtaining estimates of such partitioned effects and variances. We believe this framework can be fruitfully extended to issues like how non-genetic effects on phenotypes, and their inheritance or acquisition, can affect fitness surfaces; this is an issue we have not touched upon here. Finally, we believe that the extended view of inheritance and effects on the phenotype developed here is particularly well suited to algorithmic modeling for simulation studies of evolutionary dynamics along the lines of the extended evolutionary synthesis (Pigliucci and Müller 2010).

Acknowledgments
This is contribution no. 1 from the Foundations of Genetics and Evolution Group (FOGEG).

FOGEG is an informal association of SD, AJ, NGP and TNCV getting together periodically to work on conceptual issues at the foundations of genetics and evolutionary biology. All authors contribute equally to the manuscripts and the sequence of authors is chosen at random for each submission. The last author acts as corresponding author for that submission. We thank Sachit Daniel and K. P. Mohanan for useful discussions, especially during the early phases of the crystallization of these ideas. We also thank the Indian Academy of Sciences, Bengaluru, for supporting a discussion meeting on 'Foundations of Evolutionary Theory' at Orange County, Coorg, in February 2014, at which many discussions on the extended evolutionary synthesis took place. AJ thanks the Department of Science and Technology, Government of India, for support via a J. C. Bose Fellowship. SD and NGP thank IISER Pune and IISER Mohali, respectively, for in-house funding.
References


Frank S. A. 1997 The Price equation, Fisher's fundamental theorem, kin selection and causal

MA, USA.

Haldane J. B. S. 1932 The time of action of genes, and its bearing on some evolutionary

 Biol.* **2**, e101.

Jablonka E. and Lamb M. J. 2005 *Evolution in four dimensions: genetic, epigenetic, 
behavioural and symbolic variation in the history of life*. Massachusetts Institute of 
Technology (MIT) Press, Cambridge, MA, USA.

Jablonka E. and Raz G. 2009 Transgenerational epigenetic inheritance: prevalence, 
mechanisms and implications for the study of heredity and evolution. *Quart. Rev. Biol.* **84**, 
131–176.

Johannes F., Colot V. and Jansen R. C. 2008 Epigenome dynamics: a quantitative genetics 


503.


Press, Cambridge, MA, USA.


Figure 1. Schematic representation of the relationships between phenotype, genome and environment in the standard gene-based view of inheritance. The focal individual depicted within the gray dashed box has a genome G (dark orange circle) inherited from its parent(s). This genome, when expressed, directs the generation of the phenotype P (dark blue octagon), and the generation of the phenotype is affected by the environment E (green box), which includes both biotic (conspecifics and heterospecifics) and abiotic components. Eventually, the focal individual transmits genomic material G’ to its offspring. What is transmitted, however, is not necessarily identical to what was inherited (G), as a result of mutation and horizontal gene transfer (hence the dark orange arrow from E to G), as well as recombination and meiosis. For the focal individual, an interaction between P and E determines its Darwinian fitness. Solid arrows indicate the role of one component affecting the other, or interactions between components, when an arrow from one component impinges upon an arrow connecting two others. Thus, E interacts with G in generating P, and also affects P directly.
Figure 2. Schematic representation of the relationships between the phenotype Pᵢ, inheritome Iᵢ and environment E, at different time steps (i = 1..n) during the lifetime of the focal individual depicted within the gray dashed box. The environment (solid dark grey box), which includes all individuals (conspecifics and heterospecifics) as well as the physical and cultural backdrop, is a dynamic superset that transcends the lifespan of individuals of various species. It is the sum of all individuals and together with the physical and, for some species, cultural backdrop within which individuals are transiently embedded during their lifespan. In principle the environment changes on a time-scale that is often similar to the lifespans of individuals. Therefore, the effective local environment of a focal individual is referred to as its contextome (Cᵢ at time step i) which changes during an individual's lifetime. Note that the focal individual is in principle a component of every other individual's contextome. At conception (time i = 1), the focal individual has an inheritome I₁, inherited from its parent(s) (dashed dark orange arrow). The inheritome includes DNA sequence, as well as any epigenetic modifications of DNA and cytoplasmic mediators of parental effects. The focal individual also receives a phenotype P₁ as a parental endowment (dashed dark blue arrow); this includes, for example, cytoplasmic
components and cellular structure. At conception, the focal individual inhabits and experiences a contextome $C_1$. In subsequent time steps ($2..n$), the inheritome can change (e.g. from $I_1$ to $I_2$). This change in inheritome includes changes due to mutation, transposition, horizontal gene transfer and the acquisition of new epigenetic marks due to phenotype, and hence there are solid dark orange arrows from $I_1$, $P_1$ and $C_1$ to $I_2$. Change in the phenotype from $P_1$ to $P_2$ encompasses changes directed at time step 2 by the expressed subset(s) of $I_2$, subject to interaction with $C_2$, and also direct effects of and interactions between $P_1$ and $C_1$ (dark blue solid arrows). The contextome can also change from $C_1$ to $C_2$, subject to direct effects of and interactions between $P_1$ and $C_1$ (green solid arrows). The same set of arrows as between times steps 1 and 2 will apply to every pair of time steps $t$ and $t+1$, till $t = n-1$, indicated by the dotted gray arrow labeled 'time' between time steps 2 and $n$. At any time step $r$ (ranging from 2 to $n$), the focal individual may produce offspring, transmitting to them an inheritome $I_r$ that is typically different from $I_r$ (dashed dark orange arrows leading to offspring), and a phenotype $P_r$', determined by, but different from, $P_r$ (dashed dark blue arrows leading to offspring).