# Spring 2023 – Epigenetics and Systems Biology Lecture Outline (Epigenetics and Evolution) Michael K. Skinner – Biol 476/576 Weeks 15 and 16 (April 18 & 25)

# **Epigenetics and Evolution**

- Darwinian Evolution
- Lamarck's Environment and Evolutionary Biology
- History Environment and Evolutionary Biology
- Waddington Environment and Evolutionary Biology
- Molecular and Genetic Aspects of Evolutionary Biology
- Hopeful Monsters and Evolutionary Biology
- Epigenetics and Evolutionary Biology
- Sociobiology and Evolutionary Biology
- Sexual Selection and Evolutionary Biology
- Epigenetic Transgenerational Inheritance and Evolutionary Biology
- Summary Epigenetics and Evolutionary Biology

# **Required Reading**

Laland, et al. (2014) Does evolutionary theory need a rethink? Nature 54:161-4

Skinner MK (2015) Environmental Epigenetics and a Unified Theory of the Molecular Aspects of Evolution: A Neo-Lamarckian Concept that Facilitates Neo-Darwinian Evolution. Genome Biol Evol. 26;7(5):1296-302

# Books (Reserve in Library)

Jablonka, E. & Lamb, M.J. (2014). Evolution in Four Dimensions: Genetic, Epigenetic, Behavioral and Symbolic Variation in the History of Life. MIT Press, Cambridge.

# <u>Literature</u>

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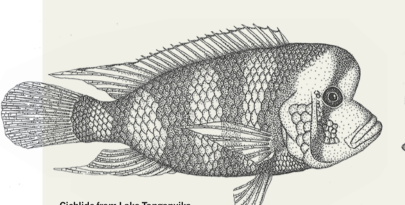
# COMMENT

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Cichlids from Lake Tanganyika (left) and from Lake Malawi (right) evolved similar body shapes.

# Does evolutionary theory need a rethink?

Researchers are divided over what processes should be considered fundamental.

# POINT Yes, urgently

Without an extended evolutionary framework, the theory neglects key processes, say Kevin Laland and colleagues.

Charles Darwin conceived of evolution by natural selection without knowing that genes exist. Now mainstream evolutionary theory has come to focus almost exclusively on genetic inheritance and processes that change gene frequencies.

Yet new data pouring out of adjacent fields are starting to undermine this narrow stance. An alternative vision of evolution is beginning to crystallize, in which the processes by which organisms grow and develop are recognized as causes of evolution.

Some of us first met to discuss these advances six years ago. In the time since, as members of an interdisciplinary team, we have worked intensively to develop a broader framework, termed the extended evolutionary synthesis<sup>1</sup> (EES), and to flesh out its structure, assumptions and predictions. In essence, this synthesis maintains that important drivers of evolution, ones that cannot be reduced to genes, must be woven into the very fabric of evolutionary theory.

We believe that the EES will shed new light on how PAGE 162 >

# COUNTERPOINT No, all is well

Theory accommodates evidence through relentless synthesis, say Gregory A. Wray, Hopi E. Hoekstra and colleagues.

In October 1881, just six months before he died, Charles Darwin published his final book. *The Formation of Vegetable Mould*, *Through the Actions of Worms*<sup>11</sup> sold briskly: Darwin's earlier publications had secured his reputation. He devoted an entire book to these humble creatures in part because they exemplify an interesting feedback process: earthworms are adapted to thrive in an environment that they modify through their own activities.

Darwin learned about earthworms from conversations with gardeners and his own simple experiments. He had a genius for distilling penetrating insights about evolutionary processes — often after amassing years of observational and experimental data — and he drew on such disparate topics as agriculture, geology, embryology and behaviour. Evolutionary thinking ever since has followed Darwin's lead in its emphasis on evidence and in synthesizing information from other fields.

A profound shift in evolutionary thinking began PAGE 163 >

**POINT: YES, URGENTLY** be evolution works. We hold that organisms are constructed in development, not simply 'programmed' to develop by genes. Living things do not evolve to fit into pre-existing environments, but co-construct and coevolve with their environments, in the process changing the structure of ecosystems.

The number of biologists calling for change in how evolution is conceptualized is growing rapidly. Strong support comes from allied disciplines, particularly developmental biology, but also genomics, epigenetics, ecology and social science<sup>1,2</sup>. We contend that evolutionary biology needs revision if it is to benefit fully from these other disciplines. The data supporting our position gets stronger every day.

Yet the mere mention of the EES often evokes an emotional, even hostile, reaction among evolutionary biologists. Too often, vital discussions descend into acrimony, with accusations of muddle or misrepresentation. Perhaps haunted by the spectre of intelligent design, evolution-

ary biologists wish to show a united front to those hostile to science. Some might fear that they will receive less funding and recognition if outsiders — such as physiologists or developmental biologists — flood into their field.

However, another factor is more important: many conventional evolutionary biologists study the processes that we claim are neglected, but they comprehend them very differently (see 'No, all is well'). This is no storm in an academic tearoom, it is a struggle for the very soul of the discipline.

Here we articulate the logic of the EES in the hope of taking some heat out of this debate and

encouraging open discussion of the fundamental causes of evolutionary change (see Supplementary Information; go.nature.com/boffk7).

# **CORE VALUES**

The core of current evolutionary theory was forged in the 1930s and 1940s. It combined natural selection, genetics and other fields into a consensus about how evolution occurs. This 'modern synthesis' allowed the evolutionary process to be described mathematically as frequencies of genetic variants in a population change over time — as, for instance, in the spread of genetic resistance to the myxoma virus in rabbits.

In the decades since, evolutionary biology has incorporated developments consistent with the tenets of the modern synthesis. One such is 'neutral theory', which emphasizes random events in evolution. However, standard evolutionary theory (SET) largely retains the same assumptions as the original modern synthesis, which continues to channel how people think about evolution.

The story that SET tells is simple: new variation arises through random genetic mutation; inheritance occurs through DNA; and natural selection is the sole cause of adaptation, the process by which organisms become well-suited to their environments. In this view, the complexity of biological development — the changes that occur as an organism grows and ages — are of secondary, even minor, importance.

In our view, this 'gene-centric' focus fails to capture the full gamut of processes that direct evolution. Missing pieces include how physical development influences the generation of variation (developmental bias); how the environment directly shapes organisms' traits (plasticity); how organisms modify environments (niche construction); and how organisms transmit more than genes across generations (extragenetic inheritance). For SET, these phenomena are just outcomes of evolution. For the EES, they are also causes.

Valuable insight into the causes of adaptation and the appearance of new traits comes from the field of evolutionary developmental biology ('evo-devo'). Some of its experimental findings are proving tricky to assimilate into SET. Particularly thorny is the observation that much variation is not random because developmental processes generate certain forms more readily than others<sup>3</sup>. For example, among one group of centipedes, each of the more than 1,000 species has an odd number of leg-bearing segments, because of the mechanisms of segment development<sup>3</sup>.

In our view, this concept — developmental bias — helps to explain how organisms adapt to their environments and diversify into many different species. For example, cichlid fishes in Lake Malawi are more closely related to other cichlids in Lake Malawi than to those in Lake Tanganyika, but species in both lakes have strikingly similar body shapes<sup>4</sup>. In each case, some fish have large fleshy lips, others protruding foreheads, and still others short, robust lower jaws.

SET explains such parallels as convergent evolution: similar environmental conditions select for random genetic variation with equivalent results. This account requires extraordinary coincidence to explain the multiple parallel forms that evolved independently in each lake. A

more succinct hypothesis is that developmental bias and natural selection work together<sup>4,5</sup>. Rather than selection being free to traverse across any physical possibility, it is

guided along specific routes opened up by the processes of development<sup>5,6</sup>.

Another kind of developmental bias occurs when individuals respond to their environment by changing their form — a phenomenon called plasticity. For instance, leaf shape changes with soil water and chemistry. SET views this plasticity as merely fine-tuning, or even noise. The EES sees it as a plausible first step in adaptive evolution. The key finding here is that plasticity not only allows organisms to cope in new environmental conditions but to generate traits

that are well-suited to them. If selection preserves genetic variants that respond effectively when conditions change, then adaptation largely occurs by accumulation of genetic variations that stabilize a trait after its first appearance<sup>5,6</sup>. In other words, often it is the trait that comes first; genes that cement it follow, sometimes several generations later<sup>5</sup>.

Studies of fish, birds, amphibians and insects suggest that adaptations that were, initially, environmentally induced may promote colonization of new environments and facilitate speciation<sup>5,6</sup>. Some of the best-studied examples of this are in fishes, such as sticklebacks and Arctic char. Differences in the diets and conditions of fish living at the bottom and in open water have induced distinct body forms, which seem to be evolving reproductive isolation, a stage in forming new species. The number of species in a lineage does not depend solely on how random genetic variation is winnowed through different environmental sieves. It also hangs on developmental properties that contribute to the lineage's 'evolvability'.

In essence, SET treats the environment as a 'background condition', which may trigger or modify selection, but is not itself part of the evolutionary process. It does not differentiate between how termites become adapted to mounds that they construct and, say, how organisms adapt to volcanic eruptions. We view these cases as fundamentally different<sup>7</sup>.

Volcanic eruptions are idiosyncratic events, independent of organisms' actions. By contrast, termites construct and regulate their homes in a repeatable, directional manner that is shaped by past selection and that instigates future selection. Similarly, mammals, birds and insects defend, maintain and improve their nests — adaptive responses to nest building that have evolved again and again<sup>7</sup>. This 'niche construction', like developmental bias, means that organisms co-direct their own evolution by systematically changing environments and thereby biasing selection<sup>7</sup>.

# **INHERITANCE BEYOND GENES**

SET has long regarded inheritance mechanisms outside genes as special cases; human culture being the prime example. The EES explicitly recognizes that parent–offspring similarities result in part from parents reconstructing their own developmental environments for their offspring. 'Extra-genetic inheritance' includes PAGE164 ▶

Plasticity: commodore butterflies emerge with

different colours in dry (left) and wet seasons.

# **COUNTERPOINT: NO, ALL IS WELL** during the 1920s, when

a handful of statisticians and geneticists began quietly laying the foundations for a dramatic transformation. Their work between 1936 and 1947 culminated in the 'modern synthesis', which united Darwin's concept of natural selection with the nascent field of genetics and, to a lesser extent, palaeontology and systematics. Most importantly, it laid the theoretical foundations for a quantitative and rigorous understanding of adaptation and speciation, two of the most fundamental evolutionary processes.

In the decades since, generations of evolutionary biologists have modified, corrected and extended the framework of the modern synthesis in countless ways. Like Darwin, they have drawn heavily from other fields. When molecular biologists identified DNA as the material basis for heredity and trait variation, for instance, their discoveries catalysed fundamental extensions to evolutionary theory. For example, the realization that many genetic changes have no fitness consequences led to major theoretical advances in population genetics. The discovery of 'selfish' DNA prompted discussions about selection at the level of genes rather than traits. Kin selection theory, which describes how traits affecting relatives are selected, represents another extension<sup>12</sup>.

Nonetheless there are evolutionary biologists (see 'Yes, urgently') who argue that theory has since ossified around genetic concepts. More specifically, they contend that four phenomena are important evolutionary processes: phenotypic plasticity, niche construction, inclusive inheritance and developmental bias. We could not agree more. We study them ourselves.

But we do not think that these processes deserve such special attention as to merit a new name such as 'extended evolutionary synthesis'. Below we outline three reasons why we believe that these topics already receive their due in current evolutionary theory.

# **NEW WORDS, OLD CONCEPTS**

The evolutionary phenomena championed by Laland and colleagues are already well integrated into evolutionary biology, where they have long provided useful insights. Indeed, all of these concepts date back to Darwin himself, as exemplified by his analysis of the feedback that occurred as earthworms became adapted to their life in soil.

Today we call such a process niche construction, but the new name does not alter the fact that evolutionary biologists have been studying feedback between organisms and the environment for well over a century<sup>13</sup>. Stunning adaptations such as termite mounds, beaver dams, and bowerbird displays have long been a staple of evolutionary studies. No less spectacular are cases that can only be appreciated at the microscopic or molecular scale, such as viruses that hijack host cells to reproduce and 'quorum sensing', a sort of group think by bacteria.

Another process, phenotypic plasticity, has drawn considerable attention from evolutionary biologists. Countless cases in which the environment influences trait variation have been documented — from the jaws of cichlid fishes that change shape when food sources alter,

A worm cast pictured in Charles Darwin's final book.

to leaf-mimicking insects that are brown if born in the dry season and green in the wet. Technological advances in the past decade have revealed an incredible degree of plasticity in gene expression in response to diverse environmental conditions, opening the door to understanding its material basis. Much discussed, too, was a book<sup>5</sup> by behavioural scientist Mary Jane West-Eberhard that explored how plasticity might precede genetic changes during adaptation.

So, none of the phenomena championed by Laland and colleagues are neglected in evolutionary biology. Like all ideas, however, they need to prove their value in the marketplace of rigorous theory, empirical results and critical discussion. The prominence that these four phenomena command in the discourse of contemporary evolutionary theory reflects their proven explanatory power, not a lack of attention.

# **MODERN EXPANSION**

Furthermore, the phenomena that interest Laland and colleagues are just four among many that offer promise for future advances in evolutionary biology. Most evolutionary biologists have a list of topics that they would like to see given more attention. Some would argue that epistasis — complex interactions among genetic variants — has long been under-appreciated. Others would advocate for cryptic genetic variation (mutations that affect only traits under specific genetic or environmental conditions). Still others would stress the importance of extinction, or adaptation to climate change, or the evolution of behaviour. The list goes on.

We could stop and argue about whether 'enough' attention is being paid to any of these. Or we could roll up our sleeves, get to work, and find out by laying the theoretical foundations and building a solid casebook of empirical studies. Advocacy can take an idea only so far.

What Laland and colleagues term the standard evolutionary theory is a caricature that views the field as static and monolithic. They see today's evolutionary biologists as unwilling to consider ideas that challenge convention.

We see a very different world. We consider ourselves fortunate to live and work in the most exciting, inclusive and progressive period of evolutionary research since the modern synthesis. Far from being stuck in the past, current evolutionary theory is vibrantly creative and rapidly growing in scope. Evolutionary biologists today draw inspiration from fields as diverse as genomics, medicine, ecology, artificial intelligence and robotics. We think Darwin would approve.

# **GENES ARE CENTRAL**

Finally, diluting what Laland and colleagues deride as a 'gene-centric' view would de-emphasize the most powerfully predictive, broadly applicable and empirically validated component of evolutionary theory. Changes in the hereditary material are an essential part of adaptation and speciation. The precise genetic basis for countless adaptations has been documented in detail, ranging from antibiotic resistance in bacteria to camouflage coloration in deer mice, to lactose tolerance in humans.

Although genetic changes are required for adaptation, non-genetic processes can sometimes play a part in how organisms evolve. Laland and colleagues are correct that phenotypic plasticity, for instance, may contribute to the adaptedness of an individual. A

seedling might bend towards brighter light, growing into a tree with a different shape from its siblings'. Many studies have shown that this kind of plasticity is beneficial, and that it can readily evolve if there PAGE 164 > **POINT: YES, URGENTLY** Ithe transmission of epigenetic marks (chemical changes that alter DNA expression but not the underlying sequence) that influence fertility, longevity and disease resistance across taxa<sup>8</sup>. In addition, extra-genetic inheritance includes socially transmitted behaviour in animals, such as nut cracking in chimpanzees or the migratory patterns of reef fishes<sup>8,9</sup>. It also encompasses those structures and altered conditions that organisms leave to their descendants through their niche construction - from beavers' dams to wormprocessed soils<sup>7,10</sup>. Research over the past decade has established such inheritance to be so widespread that it should be part of general theory.

Mathematical models of evolutionary dynamics that incorporate extra-genetic inheritance make different predictions from those that do not<sup>7-9</sup>. Inclusive models help to explain a wide range of puzzling phenomena,

# "There is more to inheritance than genes."

such as the rapid colonization of North America by the house finch, the adaptive potential of invasive plants with low genetic diversity, and how reproductive isolation is established.

Such legacies can even generate macro-evolutionary patterns. For instance, evidence suggests that sponges oxygenated the ocean and by doing so created opportunities for other organisms to live on the seabed<sup>10</sup>. Accumulating fossil data indicate that inherited modifications of the environment by species has repeatedly facilitated, sometimes after millions of years, the evolution of new species and ecosystems<sup>10</sup>.

# **BETTER TOGETHER**

The above insights derive from different fields, but fit together with surprising coherence. They show that variation is not random, that there is more to inheritance than genes, and that there are multiple routes to the fit between organisms and environments. Importantly, they demonstrate that development is a direct cause of why and how adaptation and speciation occur, and of the rates and patterns of evolutionary change.

SET consistently frames these phenomena in a way that undermines their significance. For instance, developmental bias is generally taken to impose 'constraints' on what selection can achieve — a hindrance that explains only the absence of adaptation. By contrast, the EES recognizes developmental processes as a creative element, demarcating which forms and features evolve, and hence accounting for why organisms possess the characters that they do.

Researchers in fields from physiology and ecology to anthropology are running up against the limiting assumptions of the standard evolutionary framework without realizing that others are doing the same. We believe that a plurality of perspectives in science encourages development of alternative hypotheses, and stimulates empirical work. No longer a protest movement, the EES is now a credible framework inspiring useful work by bringing diverse researchers under one theoretical roof to effect conceptual change in evolutionary biology.

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**COUNTERPOINT: NO, ALL IS WELL is** genetic variation in the response<sup>14</sup>. This role for plasticity in evolutionary change is so well documented that there is no need for special advocacy.

Much less clear is whether plasticity can 'lead' genetic variation during adaptation. More than half a century ago, developmental biologist Conrad Waddington described a process that he called genetic assimilation<sup>15</sup>. Here, new mutations can sometimes convert a plastic trait into one that develops even without the specific environmental condition that originally induced it. Few cases have been documented outside of the laboratory, however. Whether this is owing to a lack of serious attention or whether it reflects a genuine rarity in nature can be answered only by further study.

Lack of evidence also makes it difficult to evaluate the role that developmental bias may have in the evolution (or lack of evolution) of adaptive traits. Developmental processes, based on features of the genome that may be specific to a particular group of organisms, certainly can influence the range of traits that natural selection can act on. However, what matters ultimately is not the extent of trait variation, nor even its precise mechanistic causes. What matters is the heritable differences in traits, especially those that bestow some selective advantage. Likewise, there is little evidence for the role of inherited epigenetic modification (part of what was termed 'inclusive inheritance') in adaptation: we know of no case in which a new trait has been shown to have a strictly epigenetic basis divorced from gene sequence. On both topics, further research will be valuable.

All four phenomena that Laland and colleagues promote are 'addons' to the basic processes that produce evolutionary change: natural

# **"What matters** is the heritable differences in traits, especially those that bestow some selective advantage."

selection, drift, mutation, recombination and gene flow. None of these additions is essential for evolution, but they can alter the process under certain circumstances. For this reason they are eminently worthy of study.

We invite Laland and colleagues to join us in a more expansive extension, rather than imagining divisions that do not exist.

We appreciate their ideas as an important part of what evolutionary theory might become in the future. We, too, want an extended evolutionary synthesis, but for us, these words are lowercase because this is how our field has always advanced<sup>16</sup>.

The best way to elevate the prominence of genuinely interesting phenomena such as phenotypic plasticity, inclusive inheritance, niche construction and developmental bias (and many, many others) is to strengthen the evidence for their importance.

Before claiming that earthworms "have played a more important part in the history of the world than most persons would at first suppose"<sup>11</sup>, Darwin collected more than 40 years of data. Even then, he published only for fear that he would soon be "joining them"<sup>17</sup>.

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# Environmental Epigenetics and a Unified Theory of the Molecular Aspects of Evolution: A Neo-Lamarckian Concept that Facilitates Neo-Darwinian Evolution

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# Abstract

Environment has a critical role in the natural selection process for Darwinian evolution. The primary molecular component currently considered for neo-Darwinian evolution involves genetic alterations and random mutations that generate the phenotypic variation required for natural selection to act. The vast majority of environmental factors cannot directly alter DNA sequence. Epigenetic mechanisms directly regulate genetic processes and can be dramatically altered by environmental factors. Therefore, environmental epigenetics provides a molecular mechanism to directly alter phenotypic variation generationally. Lamarck proposed in 1802 the concept that environment can directly alter phenotype in a heritable manner. Environmental epigenetics and epigenetic transgenerational inheritance provide molecular mechanisms for this process. Therefore, environment can on a molecular level influence the phenotypic variation directly. The ability of environmental epigenetics to alter phenotypic and genotypic variation directly can significantly impact natural selection. Neo-Lamarckian concept can facilitate neo-Darwinian evolution. A unified theory of evolution is presented to describe the integration of environmental epigenetic and genetic aspects of evolution.

Key words: epigenetics, Lamarck, Darwin, natural selection, environment, review.

# Introduction

Charles Darwin's concept of evolution by natural selection is the unifying theme for much of modern biology (Darwin 1859). Remarkably, Darwin had no understanding of the molecular mechanisms involved in this process. Integration of Darwin's thinking with advances in genetic and molecular sciences over the past century facilitated the development of a well supported neo-Darwinian theory of evolution (Olson-Manning et al. 2012). The current primary concept for the molecular basis of evolution involves genetics and mutations, such that random DNA sequence and chromosomal alterations create a genetic variation that directly impacts phenotype and phenotypic variation. The majority of models in evolutionary biology involves DNA sequence mutations as the primary molecular mechanism underlying heritable phenotypic variation (Laland et al. 2014). A conundrum in evolutionary theory is that the frequency of potentially advantageous genetic mutations is extremely low (Jablonka and Raz 2009; Day and Bonduriansky 2011; Kuzawa and Thayer 2011; Nei and Nozawa 2011; Laland et al. 2014). Although recent studies with organisms such as microbes demonstrate genotypic variation are sufficient (Levy and Siegal 2008; Avelar et al. 2013; Ho and Zhang 2014) and additional mechanisms such as random genetic drift, genetic assimilation, directed mutations and epistasis also play important roles, genetic theory alone has difficulty explaining some aspects of evolution (Laland et al. 2014). For example, phenotypic mutation rates and genotypic mutation rates are dramatically different and genetics has been the primary molecular mechanism considered (Burger et al. 2006), but the inclusion of an additional mechanism such as epigenetics can help explain this discordance. Understanding the origins of genotypic variation and rapid evolutionary phenomenon under environmental pressure is difficult to explain with only classic genetics considered. Opposing groups of evolutionary biologists are now debating the need to "rethink" the theory (Laland et al. 2014). Genetics is the primary molecular mechanism considered in classic neo-Darwinian evolution theory (Olson-Manning et al. 2012) (table 1 and fig. 1).

In addition to evolution considerations, a large number of biological phenomena have been observed that cannot be

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# Table 1

**Evolution Theory Components** 

Neo-Lamarckian concept

Environment directly alters phenotype generationally

Darwinian evolution theory

Natural selection acts on phenotypic variation

Neo-Darwinian evolution theory

Genetic mutations promote phenotypic variation on which natural selection acts

Unified evolution theory

Environmental epigenetic alterations promote genetic mutations to alter genotypic variation Environmental epigenetics and genetic mutations both promote phenotypic variation on which natural selection acts

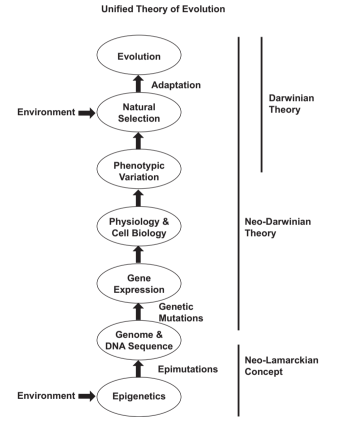


Fig. 1.—Schematic of the unified theory of evolution. No dominance is suggested by the appearance of specific circles (e.g., epimutations vs. genetics) such that all are equally important components.

easily explained by genetics alone. These include the fact that identical twins with similar genetics generally have discordant disease (Zwijnenburg et al. 2010; Kratz et al. 2014; Tan et al. 2015), or the fact that generally only a small percentage of a disease population has been found to have a correlated genetic mutation, or the fact that many diseases have increased

in frequency an order of magnitude in only a couple decades, or the fact that hundreds of environmental contaminants not able to alter DNA sequence have been shown to alter disease or phenotype later in life (Skinner 2014a). Many biological observations do not follow normal Mendelian genetic rules and are difficult to explain with classic genetic processes or mechanisms (McClintock 1984). An example in evolution is that the rates of molecular and morphological evolution are largely decoupled and these patterns of phenotypic divergence are regulatory and not classic genetic mutations (Janecka et al. 2012). Epigenetic resolution of the "curse of complexity" in adaptive evolution of complex traits has been suggested (Badyaev 2014).

Recently documented molecular mechanisms that can dramatically influence genome activity and contribute to phenotypic variation involve epigenetics (Skinner et al. 2010). Many of the above phenomenon when epigenetics is considered as an additional molecular mechanism can be more easily understood, such as the discordance of identical twins (Zwijnenburg et al. 2010; Kratz et al. 2014; Tan et al. 2015). Waddington (1953) coined the term epigenetics and the classic epigenetic definitions of Waddington (1953) and others (Skinner 2011) are descriptive, without an understanding of the molecular elements (Skinner 2011). Considering our current molecular understanding, epigenetics is defined as "molecular processes around DNA that regulate genome activity independent of DNA sequence and are mitotically stable" (Skinner et al. 2010). These epigenetic mechanisms include DNA methylation, histone modifications, chromatin structure, and selected noncoding RNA (ncRNA) (Skinner 2014a). Epigenetic processes such as DNA methylation can become programmed (e.g., imprinted) and be inherited over generations (Skinner 2014a). Environmental factors have been shown to promote the epigenetic transgenerational inheritance of phenotypic variation. Several examples of environmentally induced epigenetic transgenerational inheritance of phenotypic change have been shown to be inherited for hundreds of generations (Cubas et al. 1999). Therefore, like genetic changes, epigenetic changes can have an important role in short-term microevolution (Day and Bonduriansky 2011) and contribute to macroevolutionary (i.e., at or above the level of species) processes, such as speciation and adaptive radiation (Rebollo et al. 2010; Flatscher et al. 2012). A number of insightful reviews have proposed a role for epigenetics in evolution, primarily as a responsive molecular mechanism in natural selection (Jablonka et al. 1998; Pigliucci 2007; Laland et al. 2014).

# **Environment and Evolution**

A variety of environmental factors can influence evolution and general biology. These range from ecological parameters such as temperature and light to nutritional parameters such as caloric restriction or high fat diets. A host of environmental chemicals from phytochemicals to toxicants can also influence phenotype and health (Skinner 2014a). Environment has a critical role in natural selection and Darwinian evolution (Darwin 1859). Natural selection is a process in which environmental factors influence the survival or reproductive success of individuals bearing different phenotypes. The current paradigm in evolutionary biology holds that changes in DNA sequence underlie the variation that can evolve in response to natural selection (Laland et al. 2014) (table 1). Although James Baldwin in 1896 suggested environment through sociobiology type mechanisms (i.e., behavior) could alter phenotypic variation, these are thought to be due to genetic changes and considered a neo-Darwinian process (Baldwin 1896; Paenke et al. 2007). Therefore, in neo-Darwinian evolution the primary link between the environment and evolution is to mediate the natural selection process (Olson-Manning et al. 2012; Laland et al. 2014).

In contrast, Lamarck proposed one of the early evolutionary theories in 1802 in that environment promotes the phenotypic alterations associated with evolution (Lamarck 1802; Calabi 2001). This is distinct to the role of environment providing selective pressure in natural selection, such that environment directly alters the phenotype to influence evolution. This theory was seen as conflicting with Darwin's natural selection evolutionary theory and so was discounted and today is not seriously considered in modern evolutionary theory or neo-Darwinian evolution (Day and Bonduriansky 2011). However, if there was a molecular mechanism that generationally could facilitate the ability of the environment to alter genotypic and phenotypic variation, such a neo-Lamarckian concept may facilitate evolution (table 1 and fig. 1).

Interestingly, Darwin (1868) himself was a strong proponent of the inheritance of acquired characteristics. The blending of inheritance and evolution by natural selection appeared to be a fundamentally flawed concept that would require an untenably high mutation rate in order to maintain the trait variation required for selection (Jenkins 1867). To address this, Darwin (1868) proposed pangenesis, a complex theory of environmentally responsive somatic cell transmittance to offspring. Therefore, Darwin conceptually supported Lamarck's theory of the inheritance of acquired characteristics, but until the last 30 years the potential molecular mechanism was unclear.

# **Environmental Epigenetics**

Epigenetics provides molecular mechanisms for the environment to directly alter phenotypic variation and its subsequent inheritance (Crews et al. 2007; Skinner, Gurerrero-Bosagna, Haque, et al. 2014). A variety of epigenetic mechanisms have been identified including DNA methylation, histone modifications, chromatin structure, and selected ncRNA. All these mechanisms have the ability to program and alter gene expression and have been shown to have a critical role in normal development and biological processes (Skinner et al. 2010; Skinner 2014a). For example, the ability to generate an embryonic stem cell requires the erasure of DNA methylation such that the cell becomes pluripotent (Seisenberger et al. 2013). Although the vast majority of environmental factors cannot alter DNA sequence, epigenetic processes can be dramatically altered in response to environmental factors from nutrition to temperature (Skinner 2014a). All organisms that have been investigated contain highly conserved epigenetic processes (e.g., DNA methylation) that can be environmentally modified (Skinner 2014a). Epigenetics provides an additional molecular mechanism, integrated with genetics, to regulate biology.

The ability of environment to directly alter the development and function of cells and tissues is critical for the health and phenotype of the individual. This direct environmental epigenetic effect on the individual would likely have a limited impact on evolution, unless the epigenetic changes could be transmitted between generations. A large number of environmental factors from nutrition to toxicants have been shown to induce the epigenetic transgenerational inheritance of disease and phenotypic variation (Skinner 2014a). Epigenetic transgenerational inheritance is defined as the germline transmission of epigenetic information between generations in the absence of direct exposure (Skinner et al. 2010). Environmental exposures during a critical period of germline development, fetal gonadal sex determination or gametogenesis, have been shown to permanently program epigenetic marks such as DNA methylation (Skinner 2014a). Nutrition (Pembrey et al. 2006; Burdge et al. 2011), temperature (Song et al. 2013), stress (Skinner 2014b), and toxicants (Anway et al. 2005; Skinner 2014a) have all been shown to promote the epigenetic transgenerational inheritance of phenotypic variation (Skinner 2014a). The phenomenon has been observed in plants, insects, fish, rodents, pigs, and humans (Skinner 2014a). In mammals the altered transgenerational phenotypes have been observed for generations (Skinner 2014a), with environmentally induced epigenetic transgenerational inheritance of phenotypic variation in plants being transmitted for hundreds of generations (Cubas et al. 1999). Therefore, environment can promote the epigenetic transgenerational inheritance of phenotypic variation. The ability of environment to alter phenotype and alter phenotypic variation, independent of genetics, through this epigenetic mechanism is proposed to be important for evolution (Anway et al. 2005; Jablonka and Raz 2009; Day and Bonduriansky 2011; Kuzawa and Thayer 2011; Skinner 2014a).

Darwin proposed that one of the critical determinants of evolution was sexual selection (Darwin 1859). A previous study investigated the ability of an environmental factor (toxicant) to promote the epigenetic transgenerational inheritance of an alteration in mate preference associated with sexual selection (Crews et al. 2007). An FO generation gestating female rat was exposed to the agricultural fungicide vinclozolin transiently and then the F3 generation animals (great-grand-offspring) were obtained to assess alterations in mate preference behavior (Anway et al. 2005). A dramatic alteration in mate preference was observed (Crews et al. 2007) along with epigenetic alterations (termed epimutations) in the germline (sperm) (Guerrero-Bosagna et al. 2010). Transgenerational transcriptome changes in the brain regions correlated with the alterations in mate preference behavior (Skinner et al. 2008). Therefore, an environmental factor that altered sexual selection was found to promote a permanent alteration in the sperm epigenome in an imprinted-like manner that was inherited for multiple generations (Crews et al. 2007; Skinner et al. 2010). These studies suggest that environmental epigenetics may play an important role in evolutionary change. The role of epigenetics in mate choice and evolution has been further discussed (Zeh JA and Zeh DW 2008; Bonduriansky and Day 2013). Indeed, several recent reviews have suggested a role for epigenetics in microevolution and macroevolution (Jablonka and Raz 2009; Rebollo et al. 2010; Skinner et al. 2010; Day and Bonduriansky 2011; Kuzawa and Thayer 2011; Flatscher et al. 2012; Klironomos et al. 2013; Badyaev 2014; Jaeger and Monk 2014; Skinner 2014a).

# **Unified Theory**

Environmental epigenetics and epigenetic transgenerational inheritance provide a molecular mechanism for the neo-Lamarckian concept that environmental factors directly alter phenotype (table 1). The ability of environmental epigenetics to alter phenotypic variation provides an initial element for evolution where environment can directly establish the variation and phenotype in a population (fig. 1). Although aspects of the original Lamarckian evolution theory were not accurate (Lamarck 1802), such as having "directed" phenotypes within a generation (Koonin and Wolf 2009; Koonin 2014), the concept that environment can directly impact phenotype is supported by environmental and transgenerational epigenetic studies (Crews et al. 2007; Koonin and Wolf 2009; Koonin 2014; Skinner, Gurerrero-Bosagna, Haque, et al. 2014). Therefore, the first aspect of the unified theory involves the ability of environment to impact epigenetic programming generationally to alter phenotypic variation (fig. 1).

The well-established aspect of Darwinian evolution is the ability of environment through natural selection to act on phenotypic variation within an evolutionary event (Darwin 1859; Olson-Manning et al. 2012). The classic neo-Darwinian view is that genetic mutations and genetic variation are the primary molecular mechanism involved in generating the phenotypic variation (Nei and Nozawa 2011; Olson-Manning et al. 2012) (table 1). Although epigenetics can also have a critical role in the establishment and maintenance of phenotypic variation, the genetic mutations and genotype of the phenotype will be critical. This neo-Darwinian natural selection event for evolution is the other component of the unified theory (fig. 1).

A combination of environmental epigenetic impacts on phenotypic variation and the ability of environment to mediate natural selection will both be important for evolution. Therefore, this neo-Lamarckian concept facilitates neo-Darwinian evolution (fig. 1). This unified theory provides an expanded understanding of the molecular aspects of evolution and solutions for issues such as the mechanisms for rapid evolutionary phenomenon. The mechanisms that environment can impact evolution are also expanded. An integration of epigenetics and genetics will be essential to consider in our future understanding of the molecular aspects of evolution (Jablonka and Raz 2009; Day and Bonduriansky 2011; Laland et al. 2014; Skinner 2014a).

An additional important consideration involves the ability of epigenetic processes to promote genetic mutations (table 1). In cancer biology, altered epigenetics has been shown to promote genome instability and formation of genetic mutations (Feinberg 2004). Nearly all genetic mutations can be directly influenced by epigenetic processes. The most frequent point mutation (single nucleotide polymorphism) is a C to T conversion that is facilitated by CpG DNA methylation (Jones et al. 1992). Repeat elements in the genome when expanded create copy number variations (CNV) that are controlled by hypermethylation of DNA (Liu et al. 2012). Transposable elements are also silenced by hypermethylation of DNA (Yagi et al. 2012). Translocation events and inversions are also influenced by histone modifications, DNA methylation, and ncRNA (Solary et al. 2014). Therefore, epigenetics can directly influence genetic mutations and the origin of genotypic variation is influenced by environmental epigenetic alterations (table 1). In contrast, genetic mutations have been shown to influence epigenetics (Furey and Sethupathy 2013). Recently, we have found that environmentally induced epigenetic transgenerational inheritance of disease and phenotypic variation can promote genetic mutations (i.e., CNV) in later generations (Skinner MK, Guerrero-Bosagna C, Hague MM, unpublished data). Therefore, environmental epigenetics may not only promote increased phenotypic variation, but epigenetics can also drive genetic change and increase genotypic variation. This also needs to be considered in the unified evolution theory (fig. 1).

# Discussion

Environmental epigenetics and epigenetic transgenerational inheritance alter phenotypic variation which can be acted on by natural selection. Therefore, environmental epigenetics can directly influence phenotype and this neo-Lamarckian concept can facilitate natural selection and neo-Darwinian evolution. These different aspects of evolution should not be seen as conflicting, but instead can form a unified theory for evolution (fig. 1). This expanded understanding of the molecular aspects of evolution provides novel insights into the mechanism for rapid evolutionary events. An expanded understanding of how environment impacts evolution is also provided. The unified theory provides novel considerations that environment can both act to directly influence phenotypic variation and directly facilitate natural selection (fig. 1). Previous evolutionary models have primarily considered genetics and mutations as the primary molecular driver for evolution (Nei and Nozawa 2011; Olson-Manning et al. 2012; Laland et al. 2014). More recently, a number of models have started to consider epigenetics in these evolution models as well (Rebollo et al. 2010; Skinner et al. 2010; Day and Bonduriansky 2011; Kuzawa and Thayer 2011; Flatscher et al. 2012; Klironomos et al. 2013; Badyaev 2014; Jablonka and Lamb 2014; Jaeger and Monk 2014). For example, consideration of epigenetics as an additional molecular mechanism has assisted in the understanding of genetic drift (Gordon et al. 2012), genetic assimilation (Zuckerkandl and Cavalli 2007), and directed mutation (Jablonka and Lamb 2007; Kryazhimskiy et al. 2014). The consideration of epigenetics can also be used to better understand neutral evolution (Kimura 1989) through mechanisms, such as robustness (Ohta 2011). The unified theory suggests additional variables that should be considered are the multiple roles of environment and the integration of epigenetics into future evolution models.

Epigenetic transgenerational inheritance of phenotypic variation will have an important role in microevolutionary and macroevolutionary changes, including speciation. A recent study was designed to investigate the epigenetic changes associated with phylogenetic distance in Darwin's finches (Skinner, Gurerrero-Bosagna, Hague, et al. 2014), a wellknown example of adaptive radiation (Darwin 1859; Lack 1947; Burns et al. 2002; Grant and Grant 2008; Huber et al. 2010; Donohue 2011). Erythrocyte DNA was obtained from five species of sympatric Darwin's finches that vary in phylogenetic relatedness. Genome-wide alterations in genetic mutations, using CNV, were compared with epigenetic alterations associated with differential DNA methylation regions (epimutations) (Skinner, Gurerrero-Bosagna, Haque, et al. 2014). A greater number of epimutations than genetic mutations were observed among the different species, with the number of epimutations increasing with phylogenetic distance. The number, chromosomal locations, regional clustering, and overlap of epimutations suggest that epigenetic change has likely had a role in the speciation and evolution of Darwin's finches (Skinner, Gurerrero-Bosagna, Hague, et al. 2014). A number of additional observations also support a role of epigenetics and speciation. Using Drosophila and maternally inherited ncRNA silencing of transposons a role for epigenetics and speciation was discussed (Brennecke et al. 2008). The role of epigenetics and a punctuated equilibrium in the mobilization of transposable elements was also suggested (Zeh et al. 2009). An interesting study comparing Neanderthal and human DNA methylation maps also supports a role for epigenetics in speciation (Gokhman et al. 2014) and evolution.

Although the causal role of epimutations was not established in the Darwin's finch adaptive radiation (Skinner, Gurerrero-Bosagna, Haque, et al. 2014) or other models (Brennecke et al. 2008; Zeh et al. 2009; Gokhman et al. 2014), the causal role of genome-wide genetic mutations has also not been established (Laland et al. 2011). Future studies need to focus on the causal relationship of epigenetic alterations in relation to phenotypic variation that is acted on by natural selection. Genetics and genetic mutations are critical for evolution, but they are not the only molecular factors to consider. Although the major paradigm in the biological sciences is genetic determinism, this paradigm is limited in its ability to explain biological phenomenon ranging from the molecular basis of disease etiology (Skinner 2014a) to certain aspects of evolution by natural selection (Skinner et al. 2010; Day and Bonduriansky 2011; Longo et al. 2012). As Thomas Kuhn suggested during a scientific revolution when the current paradigm reveals anomalies then new science needs to be considered (Kuhn 1962). This type of challenge to current paradigms is also supported by other scientific philosophy, such as Popper (Rieppel 2008) and Macintyre (MacIntyre 1977). A paradigm shift is required to explain how genetics and epigenetics integrate to regulate genome activity and evolution, and these advances will need to be incorporated into future evolutionary biology modeling (Rebollo et al. 2010; Skinner et al. 2010; Day and Bonduriansky 2011; Kuzawa and Thayer 2011; Flatscher et al. 2012; Klironomos et al. 2013; Badyaev 2014; Jablonka and Lamb 2014; Jaeger and Monk 2014; Skinner 2014a) and theory.

# Summary

The integration of environmental epigenetics into the molecular aspects of evolution theory suggests a neo-Lamarckian concept that facilitates neo-Darwinian evolution. Several of the novel factors to be considered are summarized below. In regards to the neo-Lamarckian concept:

- 1. Environmental epigenetics provides a molecular mechanism for Lamarck's proposal that environment can directly alter phenotype in a heritable manner.
- 2. Environmental exposures at critical developmental windows promote the epigenetic transgenerational inheritance of germline (e.g., sperm) epimutations that alter phenotypic variation.
- 3. Direct environmental exposures of developing somatic tissue can alter somatic epigenomes and phenotype in the individual exposed, but this will not be heritable and the phenotypes will often be distinct to transgenerational phenotypes.
- 4. In regards to novel aspects of neo-Darwinian evolution:
- 5. Transgenerational germline epimutations alter genome stability to promote genetic mutations and genotypic variation in subsequent generations.

- 6. Phenotypic variation is derived from a combination of integrated genetic and epigenetic processes on which natural selection acts.
- 7. Environment has a critical role in natural selection, as well as in the induction of heritable adaptive phenotypic variation.

As shown in figure 1, these concepts and components contribute to a unified theory that integrates environmental epigenetics into the molecular aspects of evolution. It is important to note that there is not a dominance of genetics or epigenetics, but the two molecular processes integrate to regulate biology.

Previously, an environmental exposure was found to promote the epigenetic transgenerational inheritance of phenotypic traits such as mate preference, which can play an important role in evolution (Crews et al. 2007; Skinner 2014a). Several reviews have subsequently suggested a role for epigenetics in evolution (Jablonka and Raz 2009; Rebollo et al. 2010; Skinner et al. 2010; Day and Bonduriansky 2011; Kuzawa and Thayer 2011; Flatscher et al. 2012) and experimental models have shown the importance of epigenetic associated genes (Mihola et al. 2009) and molecular elements (Long et al. 2013; Skinner, Gurerrero-Bosagna, Hague, et al. 2014) in evolution. The current report extends these studies to present a unified theory that combines both neo-Lamarckian and neo-Darwinian aspects and expands our understanding of how environment impacts evolution. The integration of epigenetics and genetics will be critical for all areas of biology including evolution.

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Associate editor: Dan Graur

### Spring 2023 - Epigenetics and Systems Biology Lecture Outline (Epigenetics and Evolution) Michael K. Skinner - Biol 476/576 Weeks 15 and 16 (April 18 & 25)

### Epigenetics and Evolution

Darwinian Evolution

- Lamarck's Environment and Evolutionary Biology
- History Environment and Evolutionary Biology
   Waddington Environment and Evolutionary Biology
- Molecular and Genetic Aspects of Evolutionary Biology
- Hopeful Monsters and Evolutionary Biology
- Epigenetics and Evolutionary Biology
- Sociobiology and Evolutionary Biology
- Sexual Selection and Evolutionary Biology
- Epigenetic Transgenerational Inheritance and Evolutionary Biology
- Summary Epigenetics and Evolutionary Biology

### Required Reading

Laland, et al. (2014) Does evolutionary theory need a rethink? Nature 54:161-4

Skinner MK (2015) Environmental Epigenetics and a Unified Theory of the Molecular Aspects of Evolution: A Neo-Lamarckian Concept that Facilitates Neo-Darwinian Evolution. Genome Biol Evol. 26;7(5):1296-302

### Books (Reserve in Library)

Jablonka, E. & Lamb, M.J. (2014). Evolution in Four Dimensions: Genetic, Epigenetic, Behavioral and Symbolic Variation in the History of Life. MIT Press, Cambridge. Spring 2023 – Epigenetics and Systems Biology Discussion Session (Evolutionary Biology) Michael K. Skinner – Biol 476/576 Week 15 (April 20)

### Epigenetics and Evolutionary Biology

### Primary Papers

- 1. Skinner, et al. (2014) Genome Biology and Evolution 6:1972-1989. (PMID: 25062919)
- 2. Anastasiadi D, et al. (2021) Trends Ecol Evol. 36(12):1124-1140. (PMID: 34489118)
- 3. Sadler KC. (2022) Bioessays. 20:e2200036. (PMID: 36403219)

### Discussion

Student 2 – Ref #1 above

- What was the model system and experimental design?
- What epigenetic observations were provided and how might environmental epigenetics impact evolution?
- Is this a Lamarckian contribution to evolution?

### Student 3 - Ref #2 above

- · What was the role of epigenetics in phenotypic variation?
- · What epigenetic differences were observed between the species?
- What is the integration of genetics, epigenetics and evolution suggested?

### Student 4 - Ref #3 above

- What are the model systems and experimental data considered?
- What phylogeny associations were observed?
- How could epigenetics be involved in the potential adaptive response?

Spring 2023 – Epigenetics and Systems Biology Discussion Session (Epigenetics and Evolutionary Biology) Michael K. Skinner – Biol 476/576 Week 16 (April 27)

### **Epigenetics and Evolutionary Biology**

Primary Papers

- 1. Luo, et al. (2020) Cell Reports 33:108306. (PMID: 33113358)
- 2. Aagaard, et al. (2020) Mol Ecol. (22):5765-5783. (PMID 36112081)
- 3. McNew, et al. (2017) BMC Evolution 17(1):183. (PMID: 28835203)

### Discussion

### Student 5 – Ref #1 above

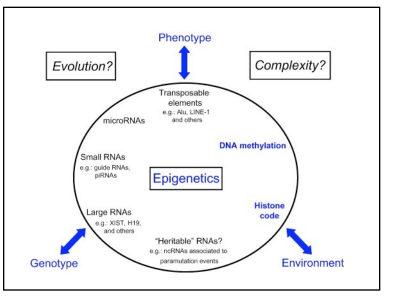
- What was the experimental design and model system?
- What epigenetic process and gene network effects were observed?
- Does this provide evidence for environmental induction of epigenetic alterations in a gene network for evolutionary adaptation?

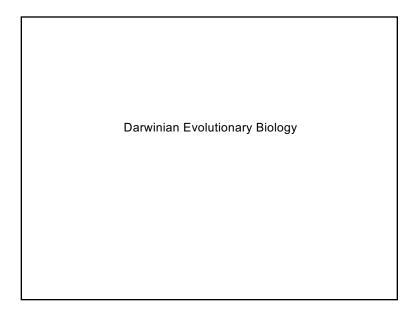
Student 6 – Ref #2 above

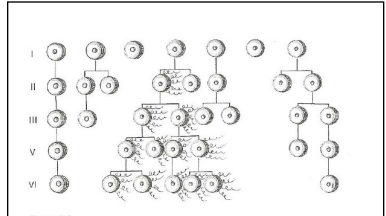
- · What is the model system, phenotypic change, and environmental factor?
- What epigenetic change was observed?
- · How did the environment, epigenetics and genetics integrate?

### Student 7 - Ref #3 above

- What was the experimental design and approach?
- What molecular alterations were observed in what cell types?
- What molecular mechanism can promote rapid evolutionary events?

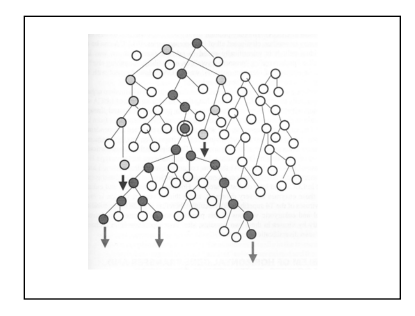


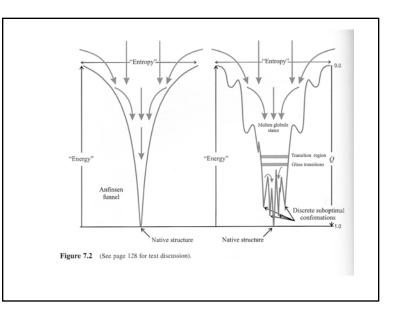


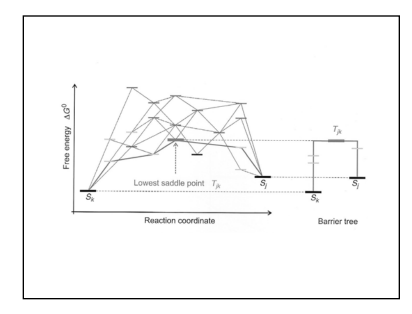


# Figure 1.1

Universal Darwinism: the frequency of the hairy entity, which first appears in generation II, increases in subsequent generations because it survives better and multiplies more than its competitors.



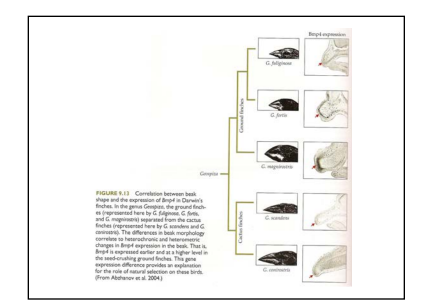


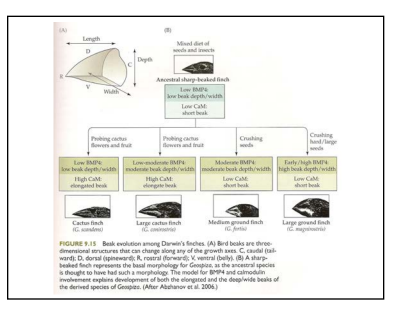


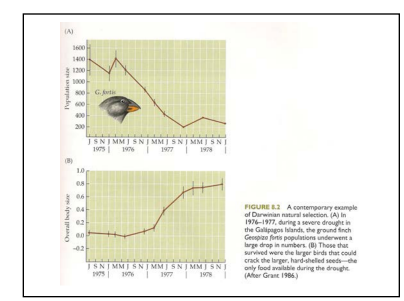
### **Classical Darwinism: Natural selection**

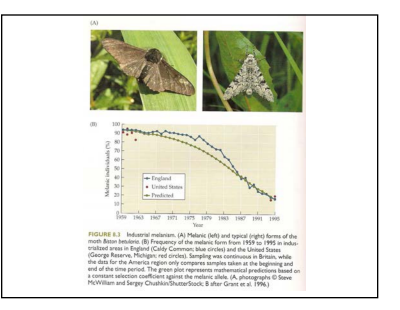
Classical Darwinian emphasis on natural selection can be summarized in a few sentences:

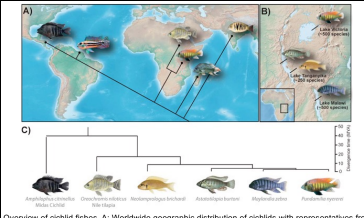
- There is variation among the individual organisms that make up a population of a species.
- There is an enormous amount of death, and most individuals will not survive to reproduce.
- Death is selective. Those individuals that best fit into the environment they encounter are more likely to survive; those that do not fit the environment well are usually eliminated.
- 4. When those individuals that survive reproduce, their progeny have a high likelihood of inheriting the variations that allowed their parents to survive. If individuals who carry those variations continue to be favored (selected), over time this natural selection will alter the overall characteristics of the population.
- 5. When populations of a species become reproductively isolated (i.e., separated in such a way that members of one population cannot mate with members of another\*), each population can randomly acquire a distinct and separate suite of variations. If the environmental conditions faced by the isolated populations are different, different variations will be selected. Anatomical and physiological











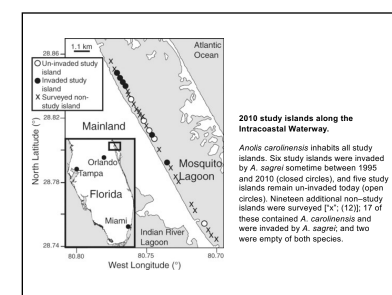
Overview of cichlid fishes. A: Worldwide geographic distribution of cichlids with representatives from India and Madagascar forming the most basal lineages and the monophyletic African and South and Central American lineages as sister-groups. B: The cichlids' center of biodiversity is East Africa, where more than 1,500 cichlid species are recognized. Estimated species numbers for the big lakes Victoria, Tanganyika and Malawi are given in brackets. C: Phylogenetic tree of the six cichlid species from Central America (A. citrinellus) and Africa with high-quality genome drafts. Divergence times are based on the lower timescale of [51].

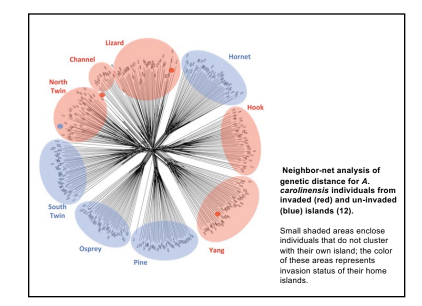
# Rapid evolution of a native species following invasion by a congener.

Science. 2014 Oct 24;346(6208):463-6. Stuart YE, Campbell TS, Hohenlohe PA, Reynolds RG, Revell LJ, Losos JB.

### Abstract

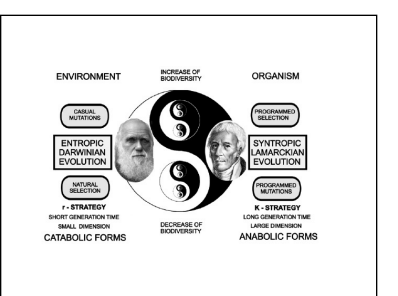
In recent years, biologists have increasingly recognized that evolutionary change can occur rapidly when natural selection is strong; thus, real-time studies of evolution can be used to test classic evolutionary hypotheses directly. One such hypothesis is that negative interactions between closely related species can drive phenotypic divergence. Such divergence is thought to be ubiquitous, though well-documented cases are surprisingly rare. On small islands in Florida, we found that the lizard Anolis carolinensis moved to higher perches following invasion by Anolis sagrei and, in response, adaptively evolved larger toepads after only 20 generations. These results illustrate that interspecific interactions between closely related species can drive evolutionary change on observable time scales.

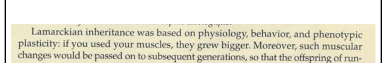




Lamarck' s

Environment and Evolutionary Biology





### Lamarck, evolution, and the inheritance of acquired characters. Genetics. 2013 Aug;194(4):793-805. Burkhardt RW Jr.



Scientists are not always remembered for the ideas they cherished most. In the case of the French biologist Jean-Baptiste Lamarck, his name since the end of the nineteenth century has been tightly linked to the idea of the inheritance of acquired characters. This was indeed an idea that he endorsed, but he did not claim it as his own nor did he give it much thought. He took pride instead in advancing the ideas that (1) nature produced successively all the different forms of life on earth, and (2) environmentally induced behavioral changes lead the way in species change. This article surveys Lamarck's ideas about organic change, identifies several ironies with respect to how his name is commonly remembered, and suggests that some historical justice might be done by using the adjective "Lamarckian" to denote something more (or other) than a belief in the inheritance of acquired characters.

## Lamarck concludes:

Nature has produced all the species of animals in succession, beginning with the most imperfect or simplest, and ending her work with the most perfect, so as to create a gradually increasing complexity in their organisation; these animals have spread at large throughout all the habitable regions of the globe, and every species has derived from its environment the habits that we find in it and the structural modifications which observation shows us.

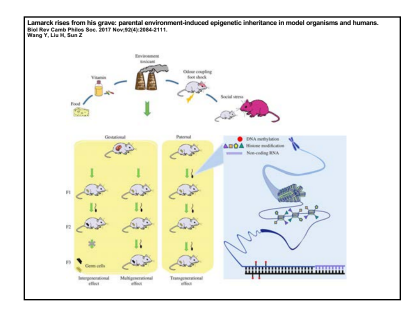
### Is Lamarckian evolution relevant to medicine? Handel AE, Ramagopalan SV. BMC Med Genet. 2010 May 13;11:73

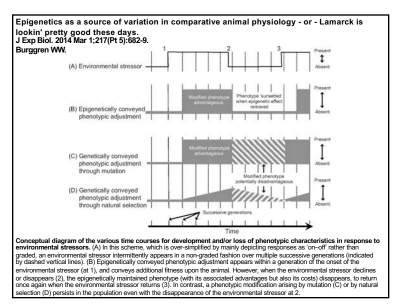
### Abstract

**BACKGROUND:** 200 years have now passed since Darwin was born and scientists around the world are celebrating this important anniversary of the birth of an evolutionary visionary. However, the theories of his colleague Lamarck are treated with considerably less acclaim. These theories centre on the tendency for complexity to increase in organisms over time and the direct transmission of phenotypic traits from parents to offspring.

DISCUSSION: Lamarckian concepts, long thought of no relevance to modern evolutionary theory, are enjoying a quiet resurgence with the increasing complexity of epigenetic theories of inheritance. There is evidence that epigenetic alterations, including DNA methylation and histone modifications, are transmitted transgenerationally, thus providing a potential mechanism for environmental influences to be passed from parents to offspring: Lamarckian evolution. Furthermore, evidence is accumulating that epigenetics plays an important role in many common medical conditions.

SUMMARY: Epigenetics allows the peaceful co-existence of Darwinian and Lamarckian evolution. Further efforts should be exerted on studying the mechanisms by which this occurs so that public health measures can be undertaken to reverse or prevent epigenetic changes important in disease susceptibility. Perhaps in 2059 we will be celebrating the anniversary of both Darwin and Lamarck.

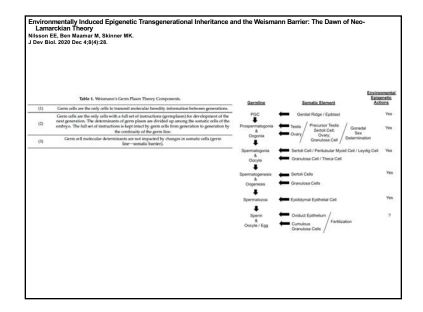


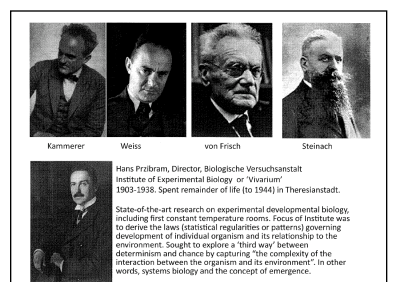


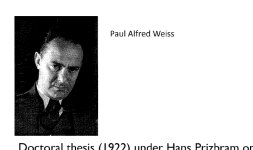
History

Environment and Evolutionary Biology

In the 16<sup>th</sup>-17<sup>th</sup> centuries the central question was how a fully integrated multicellular organism develops from a single cell (the fertilized egg). *Preformationism* believed that adult features were present fully formed in the egg and simply unfolded during growth. *Epigenesis* held that traits emerge as a consequence of the progressive interaction of the constituent parts of the zygote.

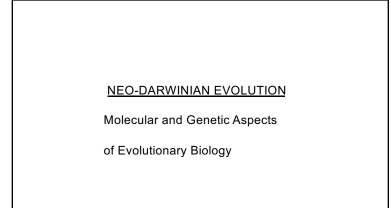




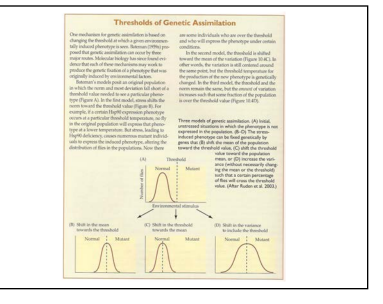


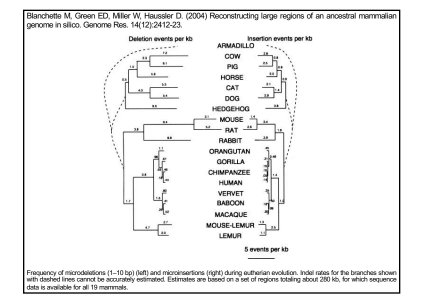
Doctoral thesis (1922) under Hans Prizbram on the responses of butterflies to light and gravity. Became Assistant Director of the Vivarium. Studied cell differentiation and the transplanting and reforming of connections in the nerves of limbs; used newts and frogs. Emphasized concept emergence and the idea of "plastic reactions" or the ability to change as a result of experience. Moved to the USA in 1931, published *Principles of Development* in 1939, and in 1954 he became one of the founding professors at the Rockefeller University; awarded the National Medal of Science in 1979.

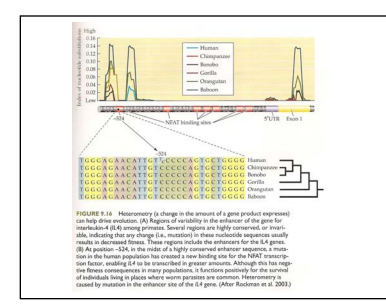
Table 1.1					
Type of theory	Hereditary transmission	Unit of variation	Origin of variation	Target of selection	Unit of evolution
Darwin's Darwinism	Gemmules transferred from the soma to sex cells	Gemmule	Random + induced in the soma	Individual (sometimes also the group)	The population of individuals
Weismann's neo-Darwinism	Transfer of determinants through the germ line	Determinant	Random + induced in the germ line	Individual (mainly) + determinants, cells, organs	The population of individuals, cells, or determinants
Modern Synthesis neo-Darwinism	Transfer of genes in the germ line	Genes in the germ line	Random mutation	Individual .	The population of individuals
Molecular neo-Darwinism	DNA replication	DNA sequence	Random DNA changes; rarely also directed changes (see chapter 3)	Mainly the individual (also the gene, the group, lineage, and species)	Mainly the population of individuals
Selfish gene neo-Darwinism	DNA replication	DNA sequence	Random DNA changes	The genc, the individual, the group	The population of alleles of the gene

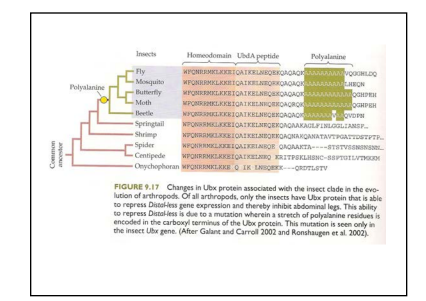


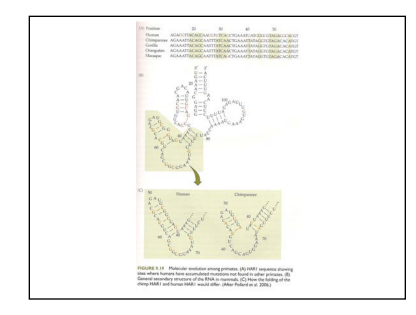
Operation	Original Target								
(a) Operations affecting gene orders									
Reversal	12345	1	-	3	-2	2 4	5		
Translocation	12 <u>345</u> 67 <u>89</u>	1 6	2 7		8 5	9			
Fusion	12345	1	2	3	4	5	6	7	
Fission	1234567	1 6	2 7	3	4	5			
Transposition	12345678	1	4	5	6	2	3	7	8
Block interchange (special)	12345678	1	4	5	6	2	3	7	8
Block interchange (general)	12345678	1	6	7	4	5	2	3	8
(b) Operations affecting gene contents									
Duplication	12345	1	2	3	2'	3'	4	5	5
Insertion	12345	1	2	3	6	4	5		
Deletion	12345	1	4	5					

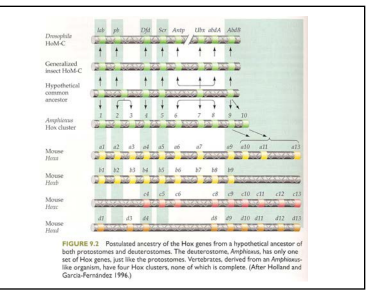


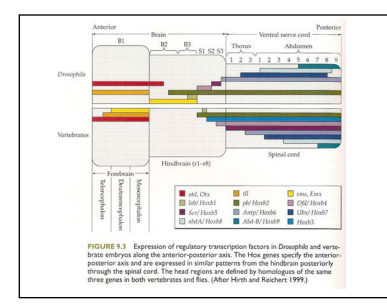


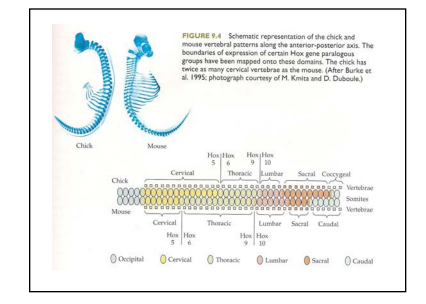


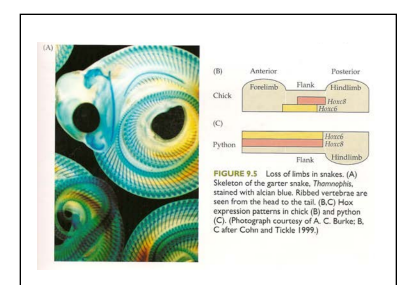


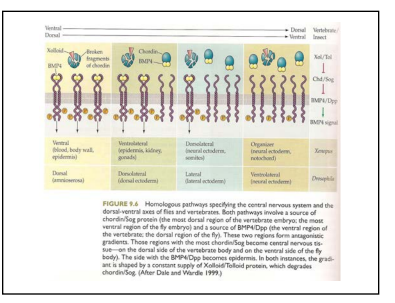




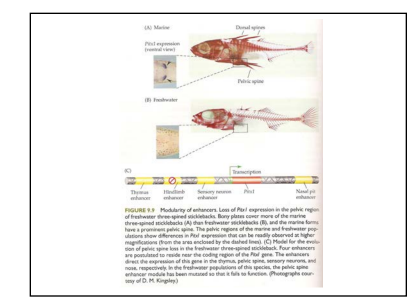


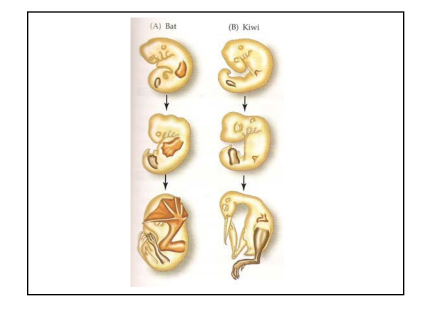


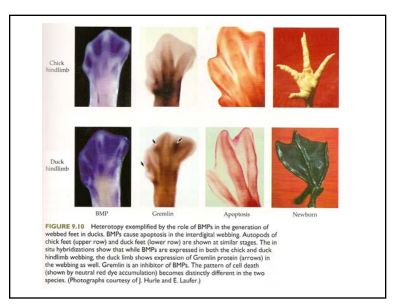


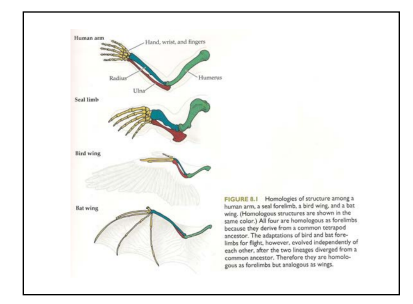


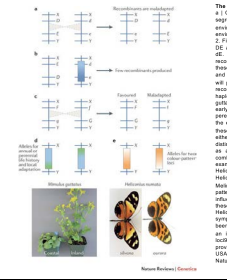
Gene	Function	protostomes and deuterostomes				
		Distribution				
achaete-scute group	Cell fate specification	Cnidarians, Drosophila, vertebrates				
Bcl2/Drob-1/ced9	Programmed cell death	Drosophila, nematodes, vertebrates				
Caudal	Posterior differentiation	Drosophila, vertebrates				
delta/Xdelta-1	Primary neurogenesis	Drosophila, Xenopus				
Distal-less/DLX	Appendage formation (proximal-distal axis)	Numerous phyla of protostomes and deuterostomes				
Dorsal/NFxB	Immune response	Drosophila, vertebrates				
forkhead/Fox	Terminal differentiation	Drosophila, vertebrates				
Fringe/radical fringe	Formation of limb margin (apical ectodermal ridge in vertebrates)	Drosophila, chick				
Hac-1/Apaf/ced 4	Programmed cell death	Drosophila, nematodes, vertebrates				
Hox complex	Anterior-posterior patterning	Widespread among metazoans				
in-12/Notch	Cell fate specification	C. elegans, Drosophila, vertebrates				
Otx-1, Otx-2/Otd, Emx-1, Emx-2/ems	Anterior patterning, cephalization	Drosophila, vertebrates				
Pax6/eyeless; Eyes absent/eya	Anterior CNS/eye regulation	Drosophila, vertebrates				
Polycomb group	Hox expression/cell differentiation control	Drosophila, vertebrates				
Netrins, Split proteins, and their receptors	Axon guidance	Drosophila, vertebrates				
RAS	Signal transduction	Drosophila, vertebrates				
ine occulus / Six3	Anterior CNS/eye pattern formation	Drosophila, vertebrates				
og/chordin, dpp/BMP4	Dorsal-ventral patterning, neurogenesis	Drosophila, Xenopus				
inman/Nkx 2-5	Heart/blood vascular system	Drosophila, mouse				
nd, msh	Neural tube patterning	Drosophila, vertebrates				





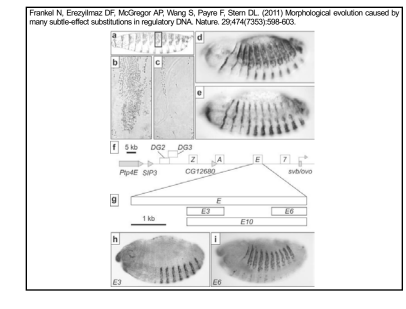


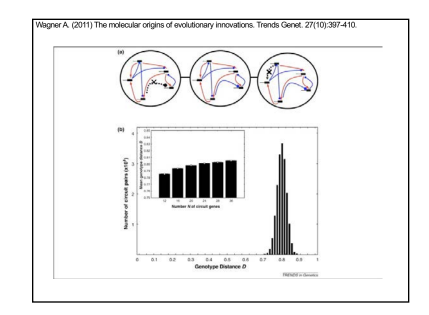


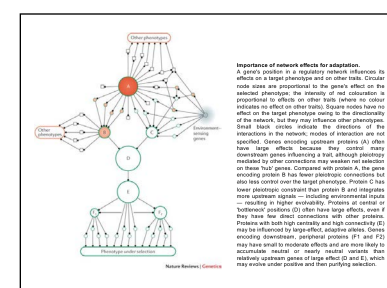


#### The dual nature of recombination.

a | Consider two ecologically important genes, D and E, segregating for alleles that are adapted to different environments. Alleles D and E are best suited to environment 1, and alleles e and d are best in environment 2. Finally, genes X and Y are neutral loci. Mating between DE and de can produce maladaptive haplotypes De and dE. b | An inversion on the DE haplotype will repress recombination between these loci and increase fitness o these alleles in their favoured environment. c | If alleles and g are maladapted, then recombination between them will produce the high fitness FG haplotype. In this case, recombination aids in the emergence of adaptive haplotype FG. d | The inland, annual ecotype of Mimulus guttatus occurs in seasonally dry habitats and flowers early in the spring, whereas the sympatric coastal, perennial form is found in wetter areas and is domant in the early spring and flowers later. Hybridization between these ecotypes would produce offspring that are less fit in either habitat. Traits that confer local adaptation to these distinct environments are located on an inversion (showr as a long rectangle) that preserves these phenotypic combinations81. e | Heliconius butterflies are a classic example of Mullerian mimicry. Many species of the genus Heliconius (for example, Heliconius numata silvana and Heliconius numata aurora) mimic the wing patterns o Melinae spp. to avoid predators. Each of these wind patterns requires a distinct combination of alleles that influence colour and shape, and recombinants between these distinct types are maladapted. The differen Heliconius mimics are closely related and occur sympatrically, yet hybrids are rarely found in nature. It has been shown that two phenotypically distinct mimics have an inversion that harbours at least two colour-pattern loci97. Photographic images in panels d and e were provided by David Lowry (University of Texas at Austin, USA) and Mathieu Joron (Museum National d'Histoire Naturelle, Paris, France), respectively.







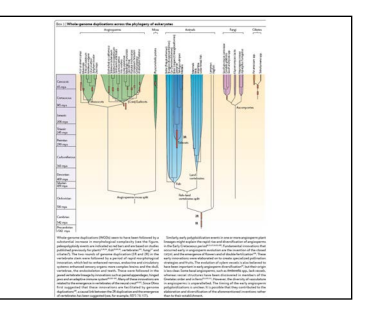
#### Box 2 | Key issues in network evolution

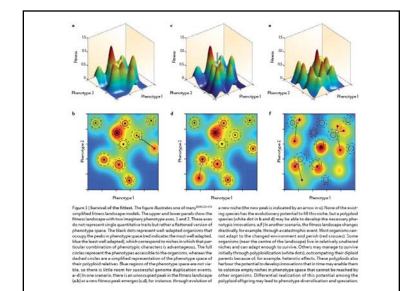
Integrating targeted genome engineering with laboratory evolution and computational modelling could considerably increase our understanding of the following open issues in network evolution. Impact of network rewiring on metabolic functioning What is the adaptive value of introducing new enzymatic reactions or rewiring regulatory links in particular environments? Systematic network modifications by means of genome engineering<sup>90</sup> will allow researchers to map the fitness landscape of metabolic networks and also explore the space of plausible alternative molecular circuits. Neutral evolution and emergence of key innovations How does the neutral evolution of metabolic networks influence the emergence of evolutionary innovations91? A computational study showed that the presence of alternative metabolic circuits with the same phenotype is a key facilitator of evolutionary novelty (that is, the ability to utilize new nutrients)92. In principle, this prediction can be tested experimentally by measuring the fitness of alternative network circuits under various environmental conditio Role of promiscuous enzyme activities in network evolution Promiscuous functions - weak activities for which the enzyme is not directly selected - have been suggested to have important roles as raw materials for future adaptive evolution93,94. Generating large pools of mutations in numerous targeted promis enzymes and exposing the mutant strains to repeated rounds of selection will shed light on how novel promiscuous pathways evolve. Importance of regulatory versus structural mutations in adaptive evolution Phenotypic changes could arise through mutations in cis-regulatory sequences or coding regions, but their relative importance remains intensely debated<sup>95</sup>. This issue could be addressed by directed evolution in vitro<sup>81</sup> by modifying the targets of available genetic variation. Convergent evolution of network structure and function How frequent is convergent evolution at the network level%? Replaying adaptive network evolution in the laboratory would allow the prevalence of convergence to be estimated and computational predictions on the availability of alternative evolutionary trajectories to be tested.

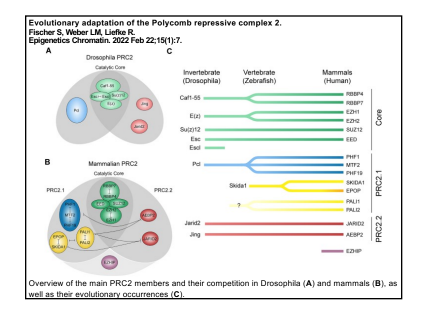
The evolutionary significance of ancient genome duplications.

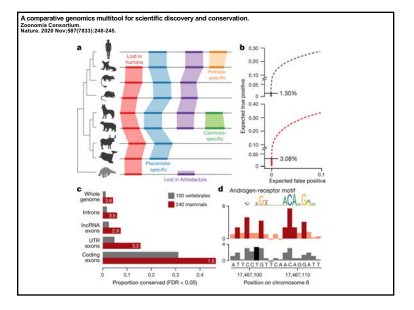
Van de Peer Y, Maere S, Meyer A.

Nat Rev Genet. 2009 Oct;10(10):725-32.



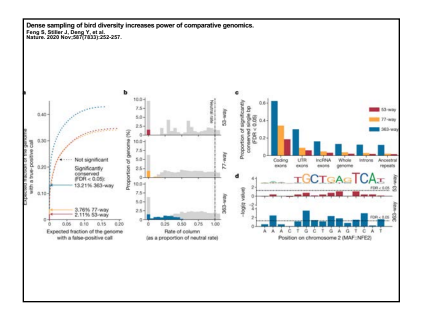






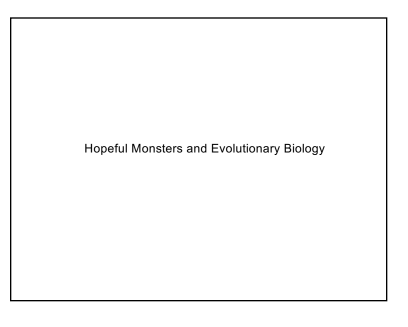
(a)	Common ancestor	(b) Timescale since polyploidyzation	(c) Source of phenotypic changes	(d) Materials	(e) Methods
	apoind 1 Actopation Standing variation				
	polypoid Environmental robustness Generalist niche	Polyploidization event	<ul> <li>Merge and attenuation of inherited adaptation</li> <li>Genome shock?</li> <li>Chromosome doubling</li> <li>Hybrid vigor</li> </ul>	Synthetic & natural polyploids	RNA-seq Epigenome
	Mutational robustness-	Microevolution	Inherited variations & new mutations	Natural variation	Population genetics
	Diploidization ·····	Macroevolution	New mutations	Paleo- polyploids	Phylogeny

#### Robustness and the generalist niche of polyploid species: Genome shock or gradual evolution? Shimizu KK.



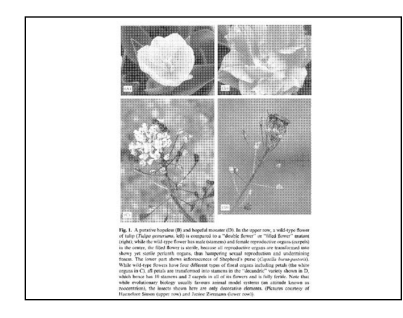
# The proper place of hopeful monsters in evolutionary biology.

Theissen G. Theory Biosci. 2006 Mar;124(3-4):349-69.



#### Abstract

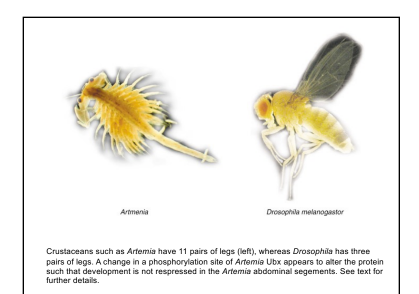
Hopeful monsters are organisms with a profound mutant phenotype that have the potential to establish a new evolutionary lineage. The Synthetic Theory of evolutionary biology has rejected the evolutionary relevance of hopeful monsters, but could not fully explain the mechanism and mode of macroevolution. On the other hand, several lines of evidence suggest that hopeful monsters played an important role during the origin of key innovations and novel body plans by saltational rather than gradual evolution. Homeotic mutants are identified as an especially promising class of hopeful monsters. Examples for animal and plant lineages that may have originated as hopeful monsters are given. Nevertheless, a brief review of the history of the concept of hopeful monsters reveals that it needs refinements and empirical tests if it is to be a useful addition to evolutionary biology. While evolutionary biology is traditionally zoocentric, hopeful monsters might be more relevant for plant than for animal evolution. Even though during recent years developmental genetics has provided detailed knowledge about how hopeful monsters can originate in the first place, we know almost nothing about their performance in natural populations and thus the ultimate difference between hopeful and hopeless. Studying the fitness of candidate hopeful monsters (suitable mutants with profound phenotype) in natural habitats thus remains a considerable challenge for the future.



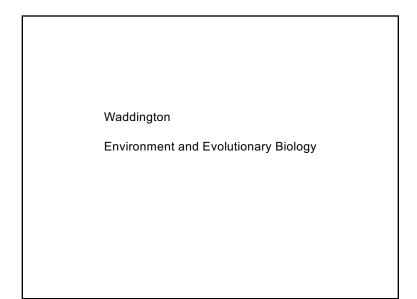
# Hopeful monsters and morphogens at the beach.

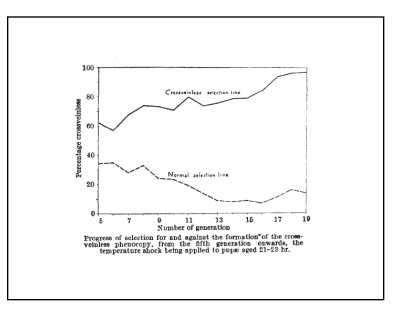
Niswander L, Anderson KV.

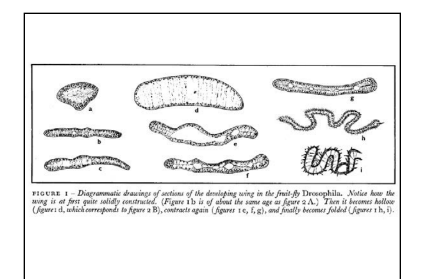
Nat Cell Biol. 2002 Nov;4(11):E259-62.

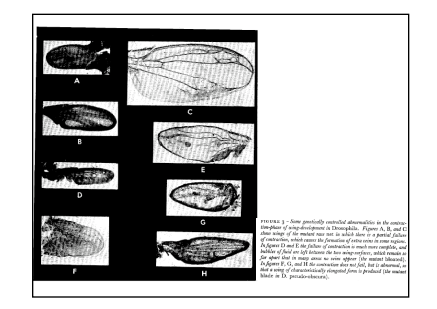


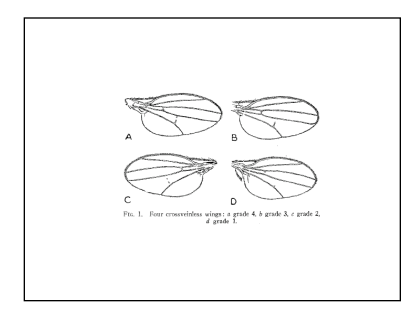
"If it could be demonstrated that any complex organ existed, which could not possibly have been formed by numerous, successive, slight modifications, my theory would absolutely break down." (Darwin 1859, p. 189)

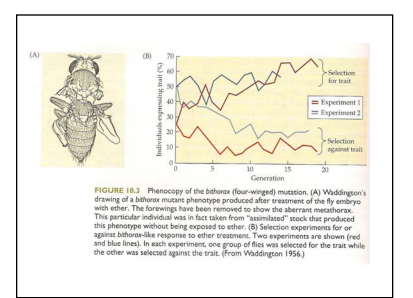


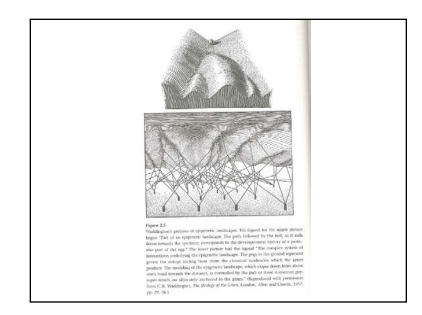


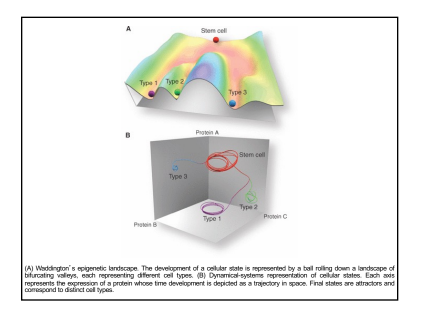


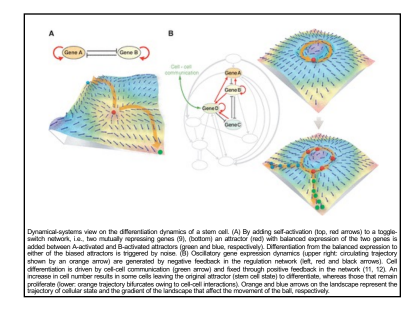




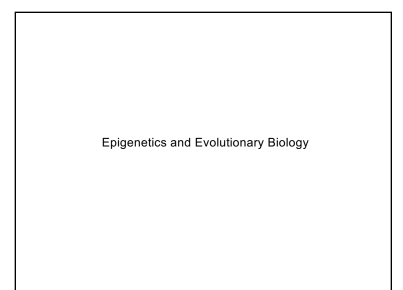


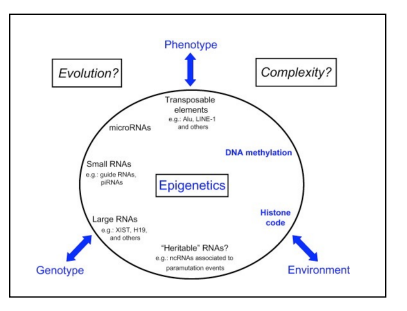


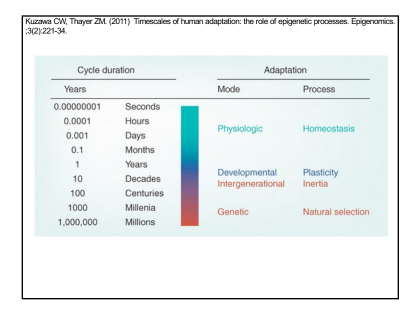


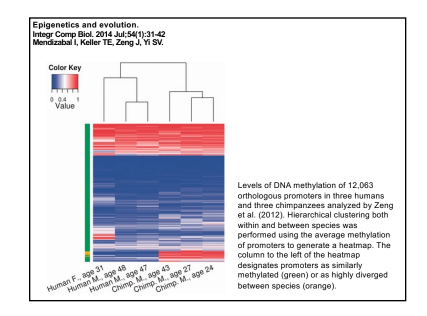


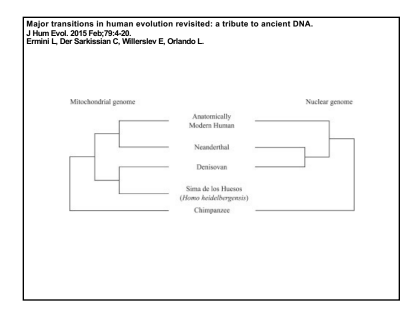
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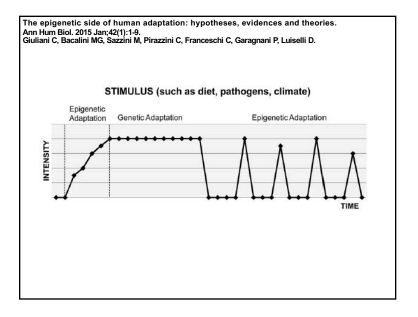


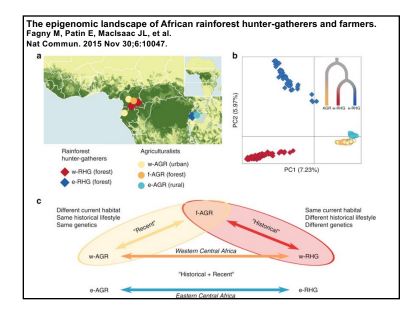


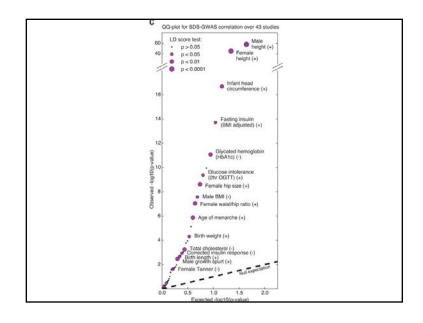


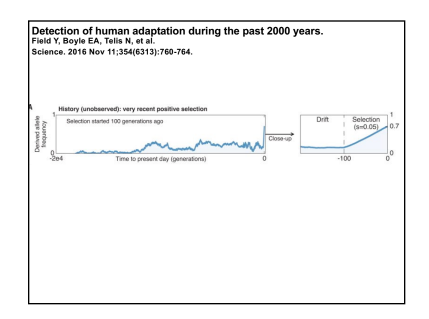


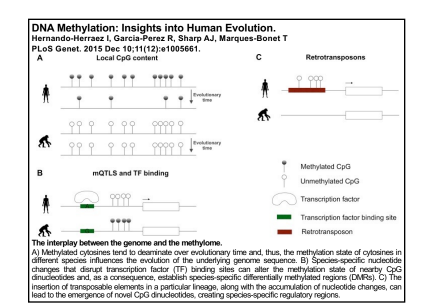




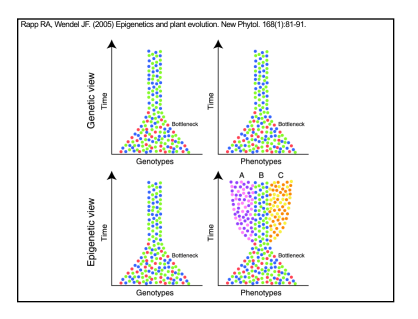


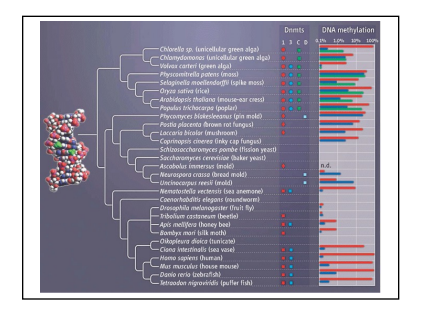


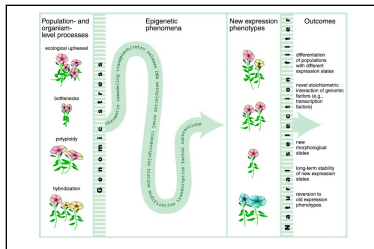




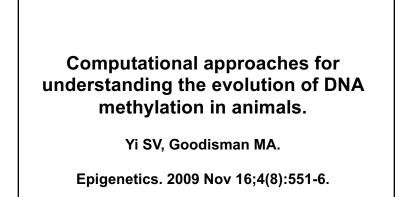
Reference	Species	Methodology	Tissue	Highlights
Wang, J. (2012)	Human, Macaque	MeDIP-chip and SEQUENOM MassARIPAY	Prefrontal cortex	>100 differentially methylated regions; Validated DMRs associated with genes with neural functions and with schizophrenia and Alzheimer's disease
Pai, A. (2011)	Human, Chimpanzee	Bumina 27K array	Liver, heart, and kidney	14.5% of promoter CpG sites are differentially methylates between tissues; 8.6% of promoter CpG sites are differentially methylated between species, Interspecies differences in promoter methylation underlife 12%-18% of gene expression differences
Molaro, A. (2011)	Human, Chimpanzee	Whole-genome bisulfite sequence	Sperm	70% of genes are hypomethylated in both chimpanzee and human sperm; 6% and 35% of orthologous SVAs had a methylation level below 50% in chimpanzee and human sperm, respectively
Martin, D.I.K. (2011)	Human, Chimpanzee, Orangutan	MethylSeq	Neutrophils	10% of CpG Islands-like regions present different methylation states between chimpanzees and humans; Regions with differential methylation might have diverge in gene regulatory function
Fukuda, K. (2013)	Human, Chimpanzee	MeDIP-chip (chromosomes 21 and 22)	Peripheral blood leukocytes	16 sDMRs between chimpanzees and humans in chromosomes 21 and 22; Genetic changes underlying these differences in methylation include gainloss of CTCF-binding sites and changes in CpG density
Hernando- Herraez, I. (2013)	Human, Chimpanzee, Bonobo, Gorilla, Orangutan	Illumina 450K array	Peripheral blood	-9% of the assayed CpG sites showed significant methylation differences between chimpanzees and humans; 184 genes perfectly conserved at protein level show significant epigenetic differences between chimpanzees and humans
Hernando- Herraez, I. (2015)	Human, Chimpanzee, Gorilla, Orangutan	Whole-genome bisulfite sequence	Peripheral blood	72% of the hypomethylated regions (HMRs) were shared among all four species; 42.6% of HMRs were on humar CpG islands; 52.6% of HMRs were on human CpG shores
Goldman, D. (2014)	Neandertal, Denisovan	Deamination rate as a proxy for DNA methylation	Femur, costae, and tibia bones	>2,000 DMRs between archaic and present-day humans Substantial changes in methylation in the HOXD cluster
Fraser, H. B (2012)	Human	Ilumina 27K array	Lymphoblastoid cell Ines	21.4% of CpG sites differed in methylation between populations; 5.4% of these CpG sites were strongly associated with local SNPs
Heyn, H. 2013	Human	Illumina 450K array	Lymphoblastoid cell lines	439 population specific differentially methylated CpG sites (pop-CpG); Significantly decreased gene expression associated to promoter hypermethylation in 12.9% (13 out of 101) of pop-CpG; Significantly increased gene expression associated to gene body methylation in 23.9% (27 out of 113) of pop-CpG

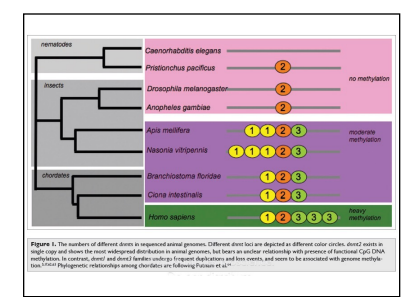


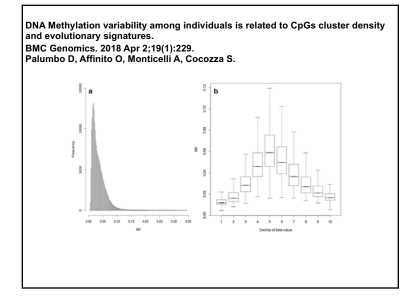


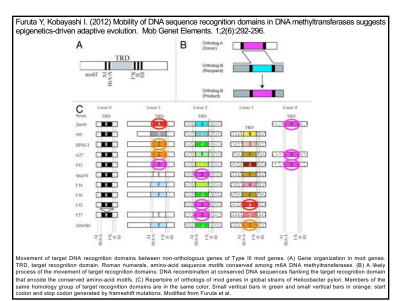


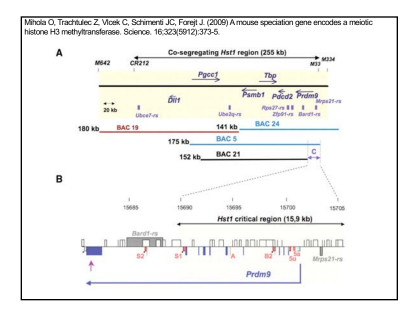
Conceptualization of the interaction between epigenetics and evolutionary change. A number of population-level processes (left) cause genomic stress, leading to the induction of epigenetic phenomena (large arrow, center). These various phenomena operate in an ecological and evolutionary context to produce novel phenotypes (right center), ranging from nolecular to morphological. These new phenotypes are subjected to the filter of natural selection – those surviving may then undergo longer-term evolutionary processes such as retention or loss of initially epigenetically fixed states.









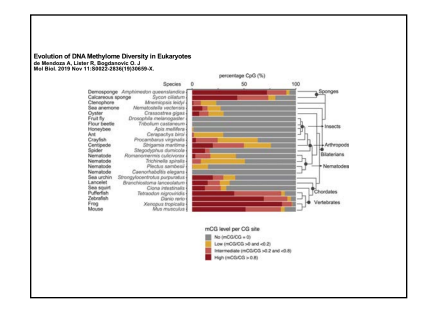


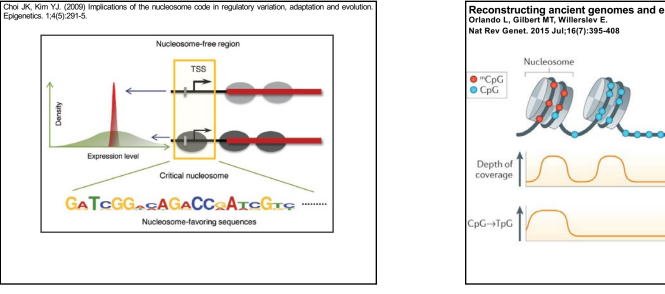
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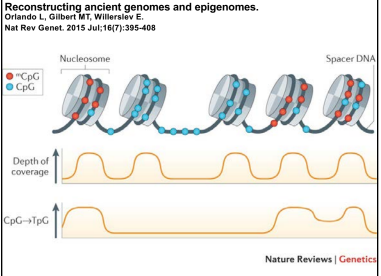
Epigenetics. 1;4(5):291-5.

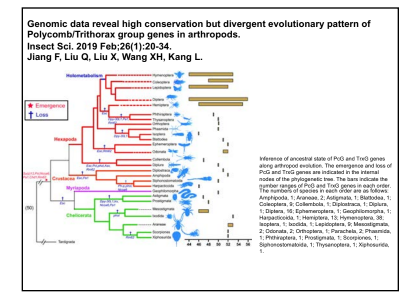
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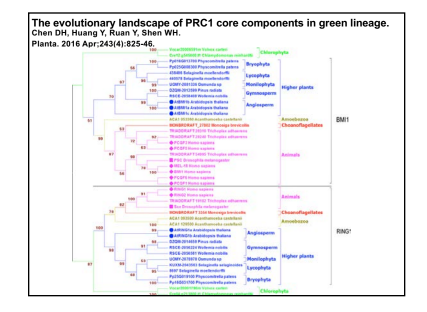
Expression level

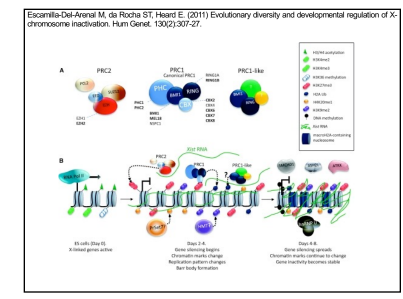


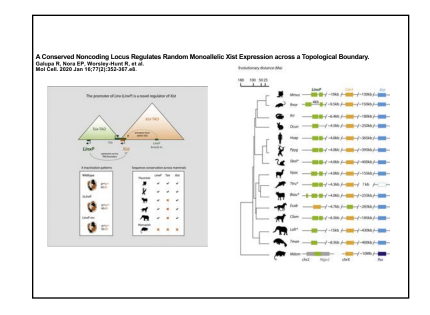




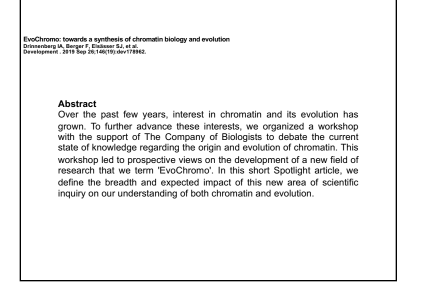


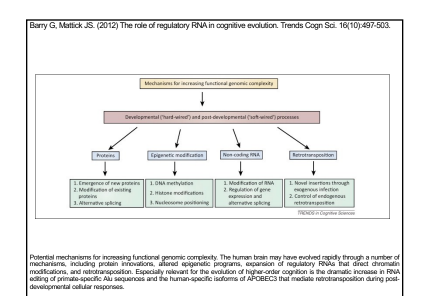


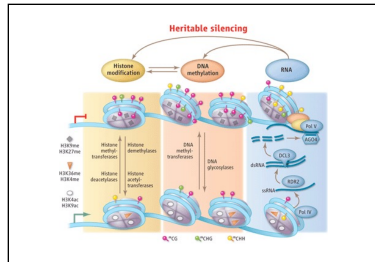




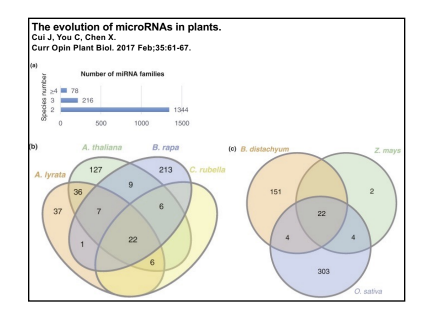
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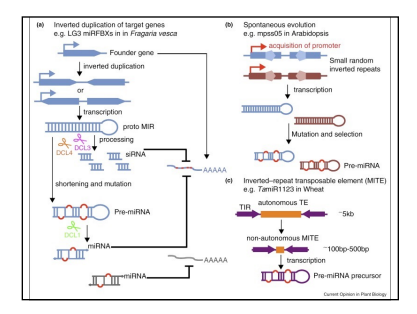


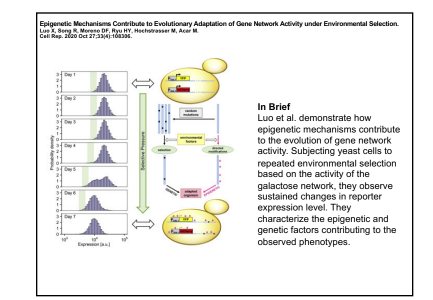




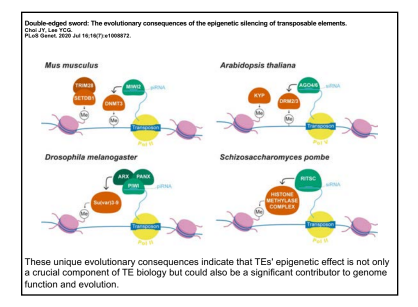
Plant epigenetic mechanisms include DNA methylation, histone modification, and RNA-directed DNA methylation (RdDM). RdDM involves two plant-specific RNA polymerases (Poi IV and Poi V), an RNA-dependent RNA polymerase (RDR2), an enzyme that cleaves double-stranded RNA (DCL3), and an Argonaute-family RNA-binding protein (AgO4). [Adapted with permission from (199)]



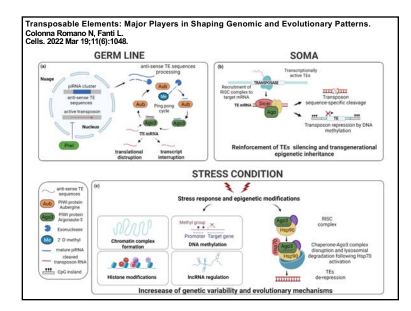


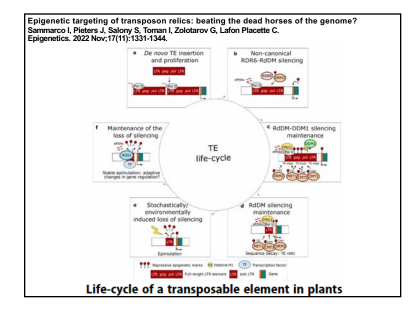


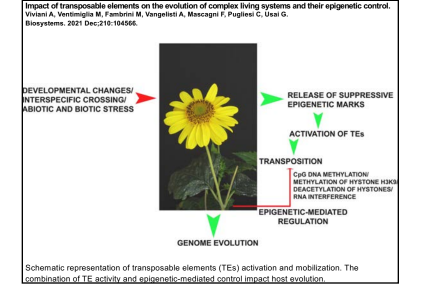
Epigenetic mechanisms			
DNA methylation	Low P Intermediate P High P	Chemical-based Bisulfite-treatment # [96] RRBS [97] Enrichment-based MeDIP # [98] Bio-CAP # [99] Enzyme digest-based MRE # [98]	
Histone PTM	H3K4me1 H3K4me2 H3K4me3 H3K9me3 H3K27ac H3K27me3 H3K36me3	Enrichment-based Histone-specific ChIP # [100]	Fer: Indexes Indexes Francer Frometer Formeter Gene body
Nucleosome occupancy	DNase I hypersensitive sites (DHS)	Enzyme digest-based DNase I # [101]	Manager Promoter Gene body

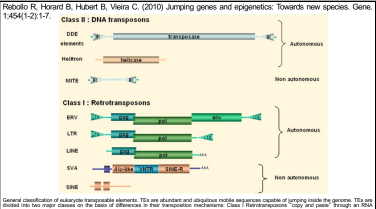


## 









divided into two major classes on the basis of differences in their transposition mechanisms: Class / Retrotransposons "copy" and paste" "through an RNA intermediate, whereas Class II DNA transposons just: "Cut and paste" their own molecule. Autonomous retrotransposons harbor tong terminal repeats in their ends (LTR) or not (LINE-Ikke), and can be infectious agents (endogenous retrovinuses). Non-autonomous retrotransposons harbor bas SINEs, are dependent on autonomous elements to be "copied and pasted" in trans. The same dependency is observed among DNA transposons, where MITEs need a full-length transposase coded by autonomous DNA transposons to be "cut and pasted" in trans. Fur a recent classification of eukaryote TEs, please refer to Wicker et al., 2007. Boxes represent open reading frames, transpis are either inverted repeats (R) in blue, or long terminal repeats (LTR) in green, and small bue arrows correspond to duplicated insertion site representations. DDE elements: transposases carrying the aspartate (D), agutamate and (E) molti. MITE (an interserve denoted repeated element, SVA: composte element, SEV. Scotter of parts of SINE (short interspersed nuclear element), VHTR (variable number of landem repeats) and Alu repeats.—the first box represents CCCTCT hexamer repeats; SINE red boxes: Indicate a adjagrostic feature; Gag. P. (Env. retrovinal: the proteins coded by TE open reading frames.

#### "Epigenetics and Systems Biology"

#### Spring 2023 (Odd Years) Biol 476/576 Schedule/Lecture Outline -

Week 1	January 10 & 12	Systems Biology (History/ Definitions/ Theory)
Week 2	January 17 & 19	Systems Biology (Networks & Emergence)
Week 3	January 24 & 26	Systems Biology (Components: DNA to Phenotype)
Week 4	Jan 31 & Feb 2	Systems Biology (Genomics / Technology)
Week 5	February 7 & 9	Epigenetics (History / Molecular Processes)
Week 6	February 14 & 16	Epigenetics (Molecular Processes & Integration)
Week 7	February 21 & 23	Epigenetics (Genomics and Technology)
Week 8	Feb 28 & March 2	Cell & Developmental Biology
Week 9	March 7 & 9	Epigenetics of Cell & Developmental Biology (& Midtern Exam)
Week 10	March 13 - 17	Spring Break
Week 11	March 21 & 23	Environmental Impact on Biology
Week 12	March 28 & 30	Environmental Epigenetics
Week 13	April 4 & 6	Disease Etiology
Week 14	April 11 & 13	Epigenetics & Disease Etiology
Week 15	April 18 & 20	Evolutionary Biology & Genetics
Week 16	April 25 & 27	Epigenetics & Evolutionary Biology
Week 17	May 2 & 4	Grant Review/ Study Section Meeting (& Final Exam)

#### Spring 2023 - Epigenetics and Systems Biology Lecture Outline (Epigenetics and Evolution) Michael K. Skinner – Biol 476/576 Weeks 15 and 16 (April 18 & 25)

#### **Epigenetics and Evolution**

- Darwinian Evolution
- Lamarck's Environment and Evolutionary Biology
- History Environment and Evolutionary Biology
- Waddington Environment and Evolutionary Biology
- Molecular and Genetic Aspects of Evolutionary Biology
- Hopeful Monsters and Evolutionary Biology
- Epigenetics and Evolutionary Biology
- Sociobiology and Evolutionary Biology
- Sexual Selection and Evolutionary Biology
- Epigenetic Transgenerational Inheritance and Evolutionary Biology Summary Epigenetics and Evolutionary Biology

#### Required Reading

Laland, et al. (2014) Does evolutionary theory need a rethink? Nature 54:161-4

Skinner MK (2015) Environmental Epigenetics and a Unified Theory of the Molecular Aspects of Evolution: A Neo-Lamarckian Concept that Facilitates Neo-Darwinian Evolution. Genome Biol Evol. 26;7(5):1296-302

#### Books (Reserve in Library)

Jablonka, E. & Lamb, M.J. (2014). Evolution in Four Dimensions: Genetic, Epigenetic, Behavioral and Symbolic Variation in the History of Life. MIT Press, Cambridge.

Spring 2023 - Epigenetics and Systems Biology Discussion Session (Epigenetics and Evolutionary Biology) Michael K. Skinner - Biol 476/576 Week 16 (April 27)

#### Epigenetics and Evolutionary Biology

- Primary Papers 1. Luo, et al. (2020) Cell Reports 33:108306. (PMID: 33113358)
- 2. Aagaard, et al. (2020) Mol Ecol. (22):5765-5783. (PMID 36112081) 3. McNew, et al. (2017) BMC Evolution 17(1):183. (PMID: 28835203)

#### Discussion

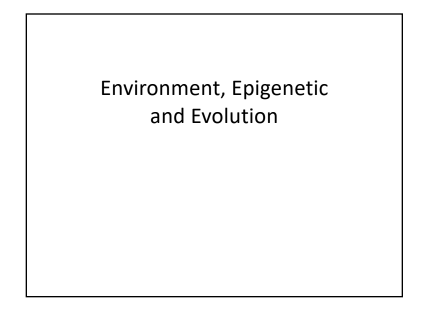
#### Student 5 – Ref #1 above

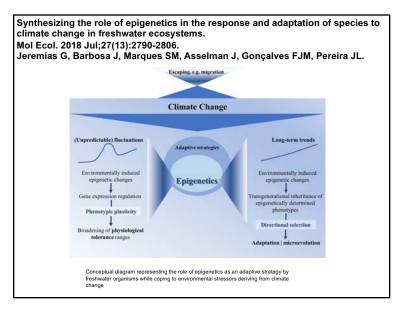
- What was the experimental design and model system?
- What epigenetic process and gene network effects were observed?
  Does this provide evidence for environmental induction of epigenetic alterations in
- a gene network for evolutionary adaptation?

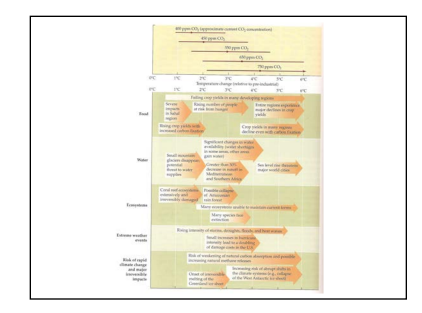
- Student 6 Ref #2 above What is the model system, phenotypic change, and environmental factor?
- · What epigenetic change was observed?
- · How did the environment, epigenetics and genetics integrate?

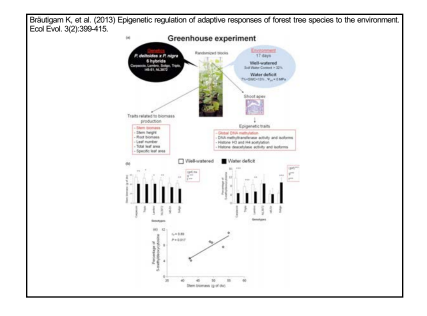
#### Student 7 – Ref #3 above

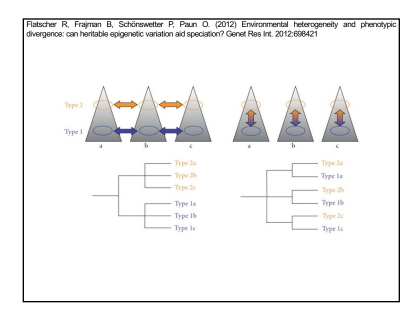
- What was the experimental design and approach?What molecular alterations were observed in what cell types?
- What molecular mechanism can promote rapid evolutionary events?

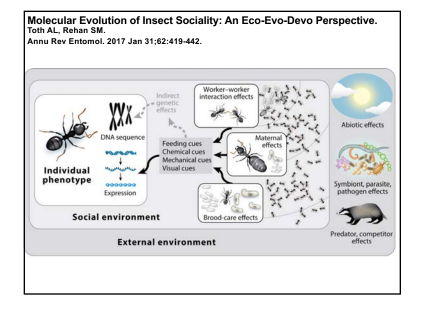


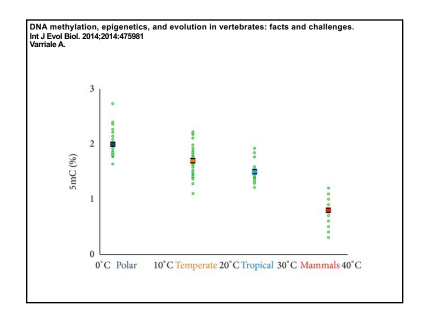


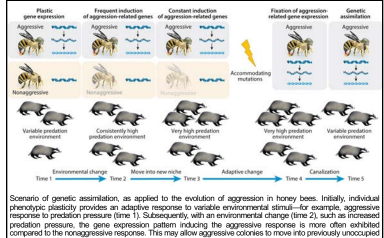




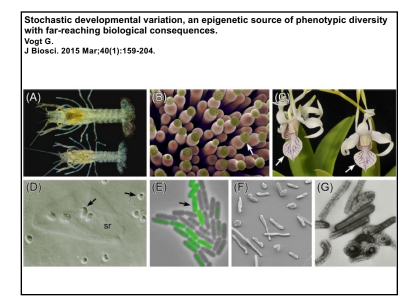


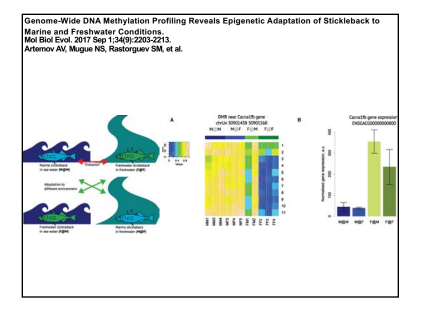


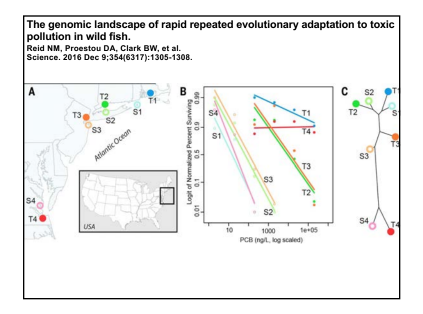


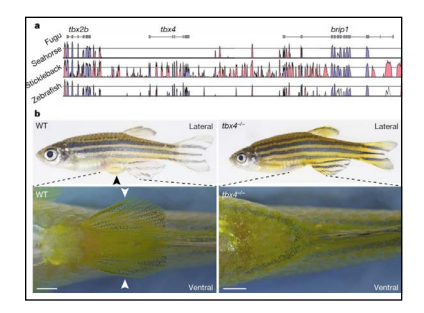


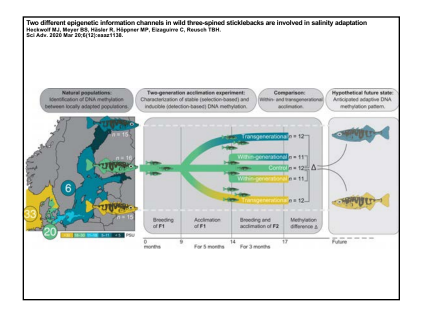
response to predation pressure (time 1). Subsequently, with an environmental change (time 2), such as increased predation pressure, the gene expression pattern inducing the aggressive response is more often exhibited compared to the nonaggressive response. This may allow aggressive colonies to move into previously unoccupied niches in the environment (time 3), such as very high predation environments. Over time, environmentally induced responses in gene expression and aggressive phenotype can become fixed differences as a result of the accumulation of accommodating mutations (time 4). The response then becomes canalized, resulting in a loss of plasticity, and individuals are fixed for the aggressive phenotype, and associated gene expression, even in the absence of the high predation environmental stimulus (time 5).

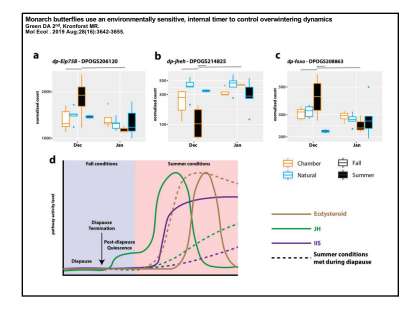


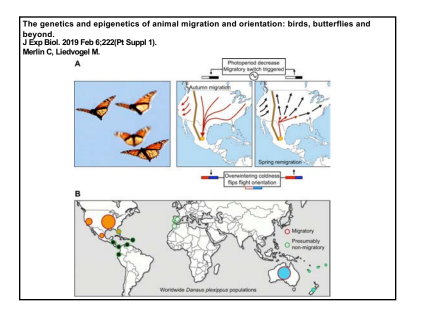


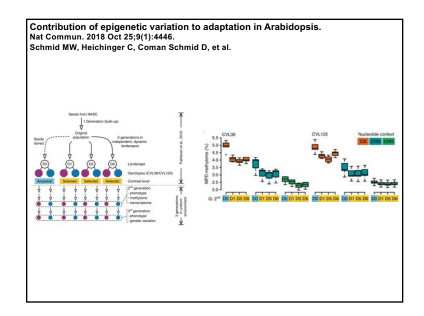


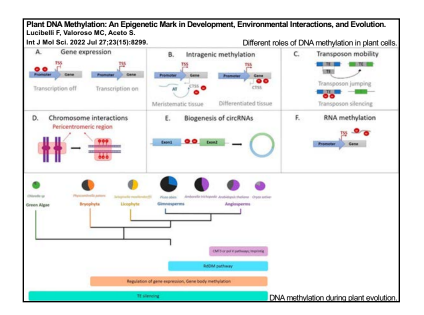


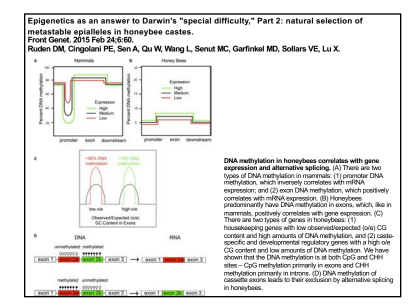


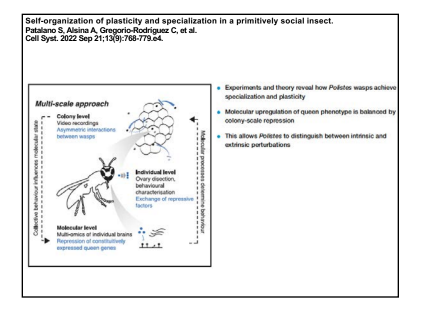




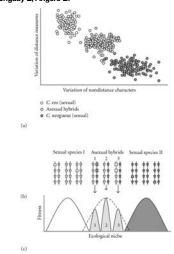




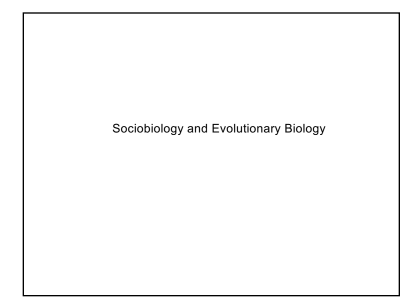


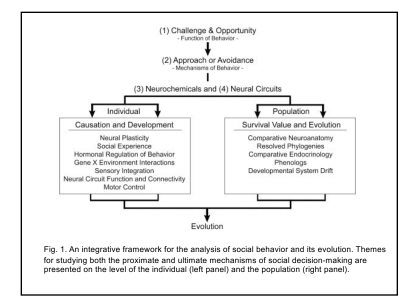


#### The key role of epigenetics in the persistence of asexual lineages. Genet Res Int. 2012;2012:534289. Castonguy E, Angers B.



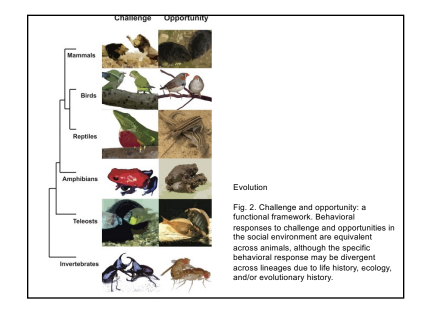
Hypothesis of the epigenetic mechanism underlying the flexibility of a genotype. (a) Phenotypic variation observed in sexual and asexual species. The points represent individual scores of Chrosomus eos, C. neogaeus, and asexual hybrids from two principal component analyses performed on body distance and nondistance measures (modified from [61]). In sexual species, the phenotypic variation among individuals is mostly the result of genetic variation, whereas, in asexual hybrids, it results from differentially expressed alleles of a same genotype. (b) Putative genetic and epigenetic variation at four genes is represented for three individuals per species. Arrows refer to expressed genes. larger arrows to different alleles of an expressed gene (genetic difference), and blocks to silenced genes (epigenetic difference). (c) Under the General Purpose Genotype model, an epigenetically flexible genotype may provide a wide ecological niche for asexual hybrids, where each different epigenetic variant would occupy a narrower niche

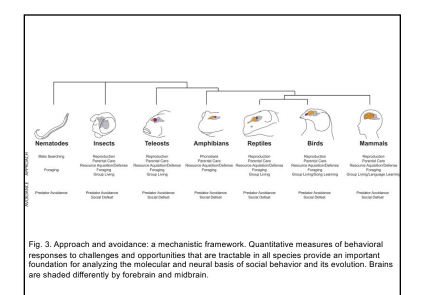


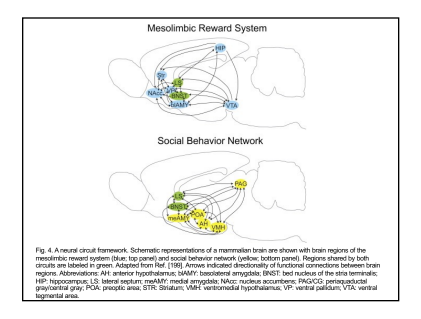


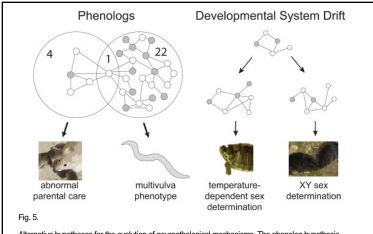
Genes, hormones, and circuits: An integrative approach to study the evolution of social behavior.

O'Connell LA, Hofmann HA. Front Neuroendocrinol. 2010 Dec 14. [Epub ahead of print]

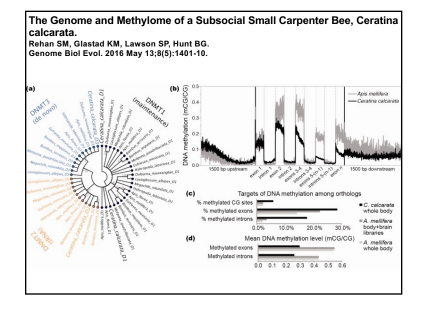


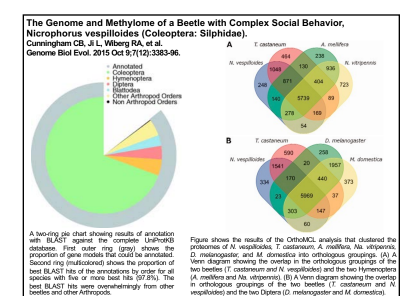






Alternative hypotheses for the evolution of neuroethological mechanisms. The phenolog hypothesis predicts that some gene/protein-interaction networks underlying social behavior and other complex phenotypes can be conserved across animals, even if the phenotypes are completely different. The developmental system drift hypothesis states that the molecular mechanisms underlying homologous phenotypes can diverge substantially during the course of evolution. Nodes and edges represent gene networks involved in a phenotype.

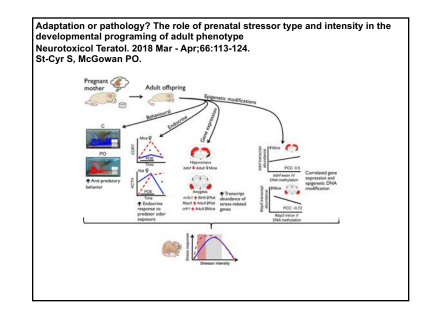


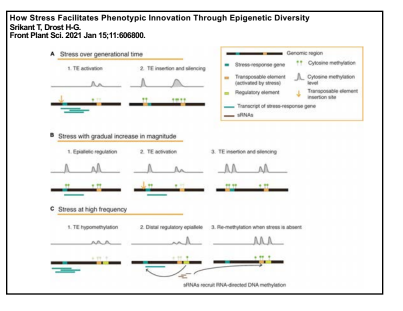


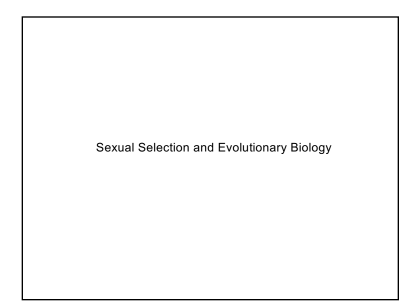
Epigenomics and gene regulation in mammalian social systems Guerrero TP, Fickel J, Benhaiem S, Weyrich A. Curr Zool. 2020 Jun;66(3):307-319.

#### Abstract

Social epigenomics is a new field of research that studies how the social environment shapes the epigenome and how in turn the epigenome modulates behavior. We focus on describing known gene-environment interactions (GEIs) and epigenetic mechanisms in different mammalian social systems. To illustrate how epigenetic mechanisms integrate GEIs, we highlight examples where epigenetic mechanisms are associated with social behaviors and with their maintenance through neuroendocrine, locomotor, and metabolic responses. We discuss future research trajectories and open questions for the emerging field of social epigenomics in nonmodel and naturally occurring social systems. Finally, we outline the technological advances that aid the study of epigenetic mechanisms in the establishment of GEIs and vice versa.

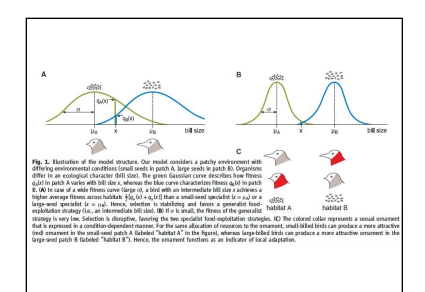


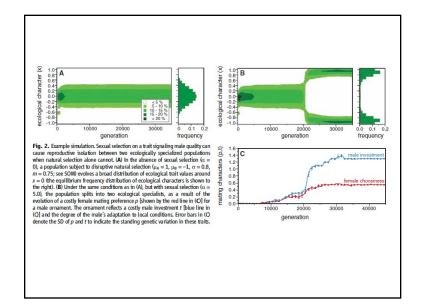


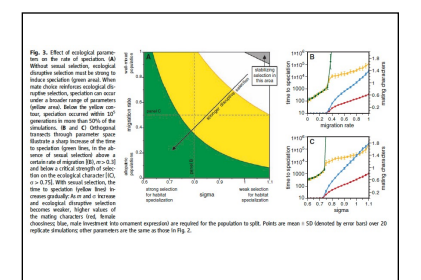


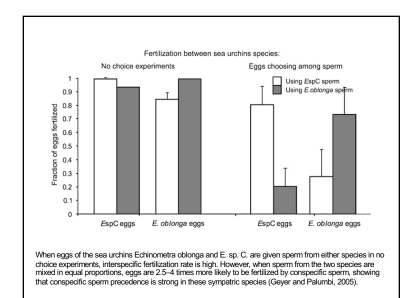
# On the origin of species by natural and sexual selection.

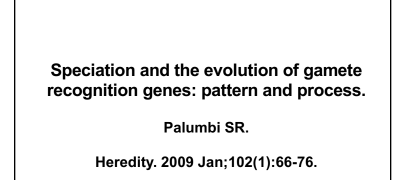
van Doorn GS, Edelaar P, Weissing FJ. Science. 2009 Dec 18;326(5960):1704-7.

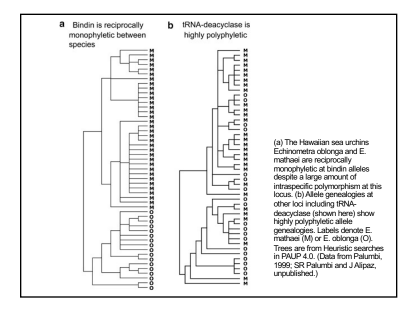


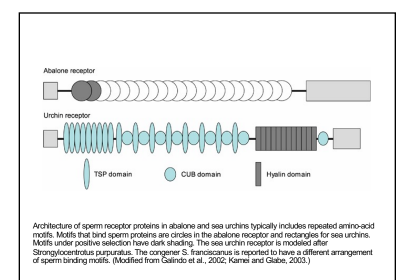






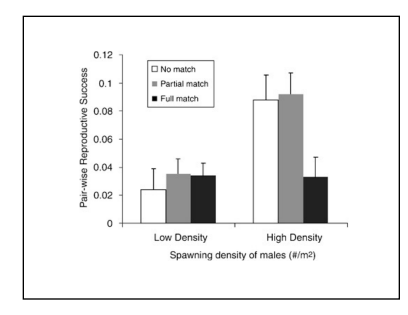


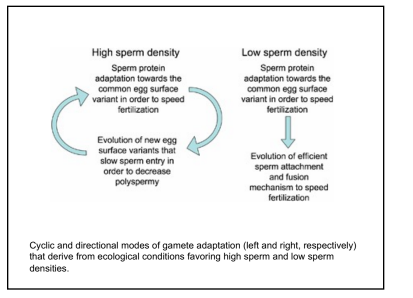


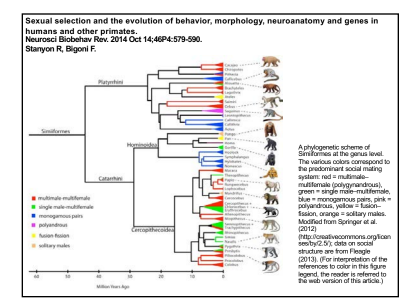


SpurpEBR1	PVISGCPSDQNVTTDIGNATAVVIWPPTATDNSGSQTLTSTNNPGDDFPIGNNTVTY
SpurpEBR1	
SpurpRepeat 1	FN
SpurpRepeat 3	F
SpurpRepeat2	N
SpurpRepeat3	F
SpurpRepeat4	N
SpurpRepeat5	
SpurpRepeat6	F
SpurpRepeat7	N
SpurpRRepeat8	
Em16	FVLTFNP.KVI.T.I.LNDHSH
Em16	FS.GST.A.NDSYTN.
Em16	.GT.N.FLRKPV.Q.TNA.NSH.STTN.
Em16	FS.NPATS.MTN
Em16	FS.G.GST.A.NDSYTN.
Eob27	F
Eob27	FS.GSA.SNN.IASY
Eob27	FS.NPTNIV.VSS
een species. The t	the hyalin-like repeats motifs of the sea urchin sperm receptor within and top 11 sequences are from different repeats of the EBR1 gene sequenced fro puratus (Kamei and Glabe, 2003), showing variation at six amino-acid positi
uences from one in	dividual Echinometra mathaei (Em16) and one E. oblonga (Eo27) show stro atus at about half of the amino-acid positions but are highly variable among

individual clones. (Data from Kamei and Glabe, 2003; SR Palumbi and J Alipaz, unpublished.)







Unravelling the role of epigenetics in reproductive adaptations to early-life environment. Bar-Sadeh B, Rudnizky S, Pnueli L, Bentley GR, Stöger R, Kaplan A, Melamed P. Nat Rev Endocrinol. 2020 Sep;16(9):519-533.

#### Key points

- Human reproductive function adjusts to changing environmental conditions.
- Key 'windows of susceptibility' during various stages of early development are the most sensitive to events or exposures that can impart long-term reprogramming of adult reproductive function.
- Epigenetic modifications have a role in regulating the central control of reproduction and pubertal onset and likely mediate much of the adaptive response.
- Human cohort data are useful for identifying methylation in proxy tissues that correlates with phenotypic variation, but determining cause and effect is challenging because hormones affect the epigenome and epigenetic ageing.
- Understanding which of the modifications are functional and responsible for the phenotype requires integrating the study of human tissues, animal and cell models and molecular approaches.
- Characterization and elucidation of these adaptive mechanisms are needed to inform the clinician of alternative reproductive strategies, and the implications for fertility treatment and healthy ageing.

Evolutionary history of sexual selection affects microRNA profiles in Drosophila sperm. Hotzy C, Fowler E, Kiehl B, Francis R, Mason J, Moxon S, Rostant W, Chapman T, Immler S. Evolution. 2022 Feb;76(2):310-319.

Our findings suggest that long-term adaptation may affect miRNA profiles in sperm and that these may show varied interactions with short-term environmental changes. Epigenetic Transgenerational Inheritance and Evolutionary Biology

# **Epigenetic Inheritance Systems**

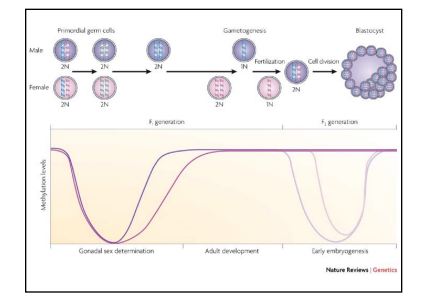
The Inheritance of Environmentally Induced Traits

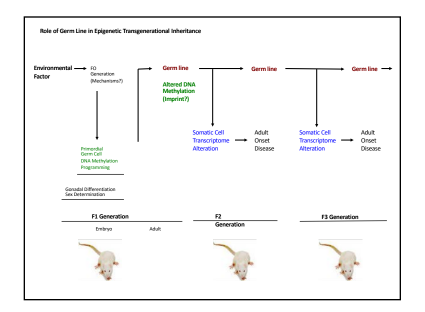
Ecological evolutionary developmental biology, or "eco-evo-devo," has the data to bring two controversial alternative inheritance systems back into the discussion of evolutionary biology. The first idea concerns the inheritance of environmentally acquired traits, an ancient idea usually associated with Jean-Baptiste Lamarck (1744–1829), but which was also used by Charles Darwin and many other Victorian naturalists. The second controversial idea usually goes by the name "genetic assimilation," and it concerns the genetic fixation of an adaptive, plastic response into the genome. In this hypothesis, a response that was once part of a phenotypically plastic repertoire is now part of the normative genetic "program."

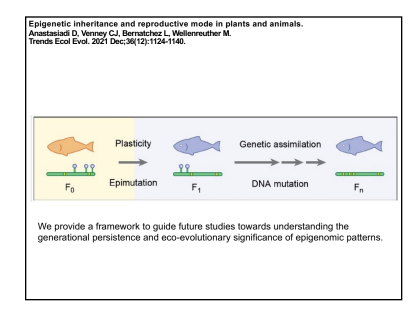
# Weismann proposed that only the germline counted in heredity, and that the germline was separate from the somatic lineages of cells that formed the body. Therefore, anything that affected the individual could not influence heredity if the germline was not affected. Weismann cut off the tails of mice for nineteen generations and showed that a tailless race did not develop.

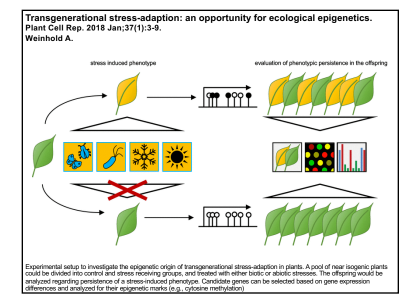
### The Ghost of Lamarck

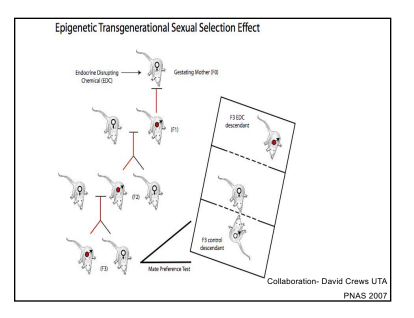
Epigenetic inheritance systems recall the specter of a banished ghost—Lamarckian inheritance. The year 2009 is not only the bicentenary of Darwin's birth and the centenary of the Woltereck and the Johannsen papers described in Chapter 1, it is also the bicentenary of Lamarck's *Philosophie Zoologique*.











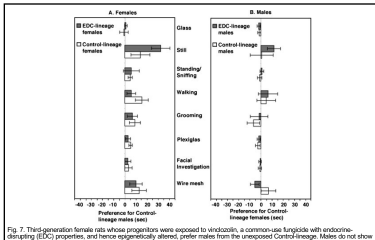
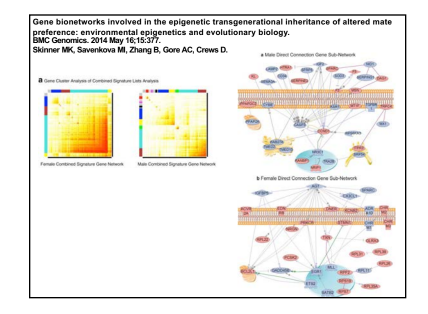
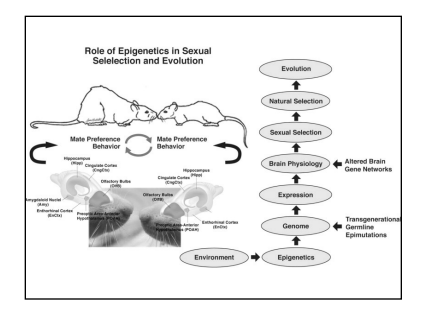
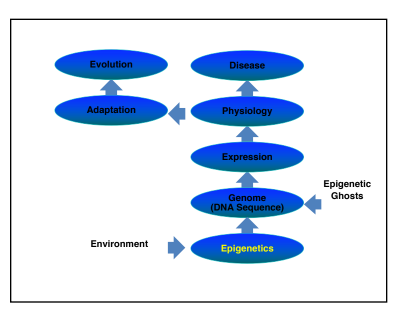
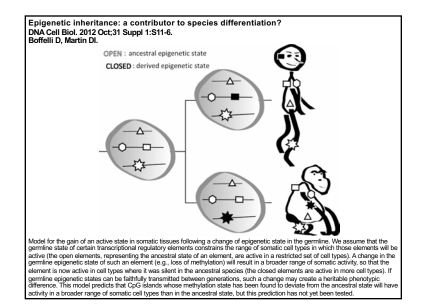


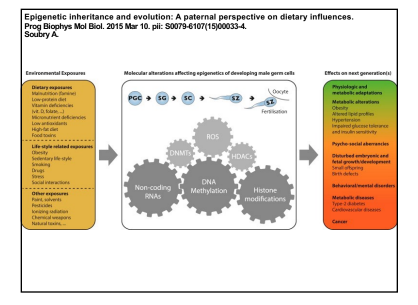
Fig. 7. Third-generation female rats whose progenitors were exposed to vinclozolin, a common-use fungicide with endocrinedisrupting (EDC) properties, and hence epigenetically altered, prefer males from the unexposed Control-lineage. Males do not show this preference. See Fig. 6 for further details. Both females and males from Control - and EDC-lineages three tested with pairs of Control- and EDC-lineage stimulus partners. Presented are the meen (+1 standard error) differences in the time spent in each behavior. Left panel: Behaviors exhibited by females from Control- and EDC-lineages towards males from Control-landes (positive, right side) and EDC-lineage (negative, left side). Right panel: Behaviors exhibited by males from Control- and EDC-lineages towards females from Control-lineage (positive, right side) and EDC-lineage (regative, left side). The various behavioral measures and test are described in Crews et al. (2007). Reprinted by permission from Crews et al. (24).

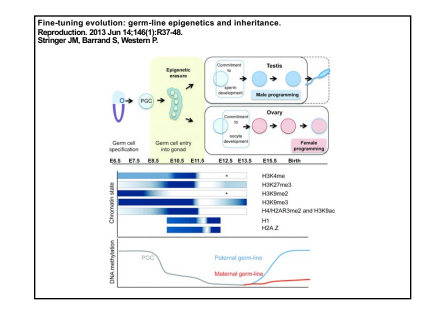




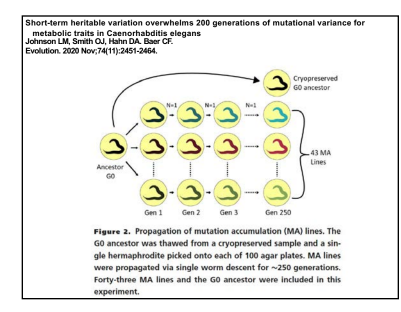




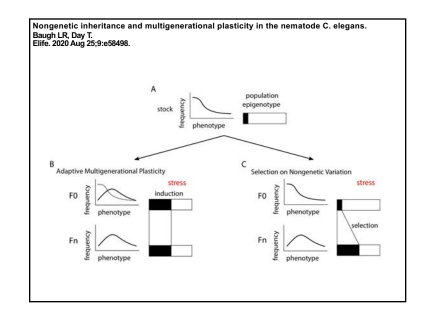


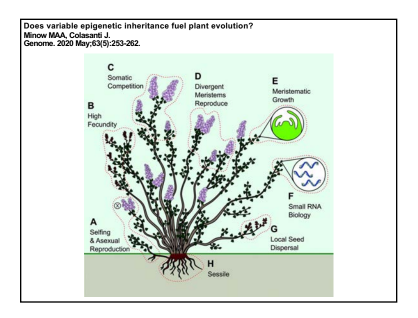


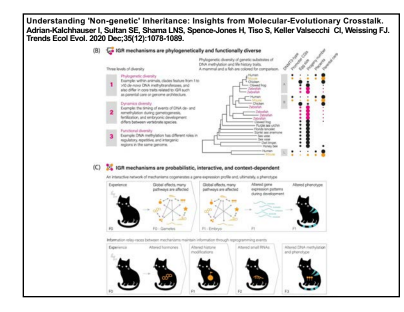
,	Acquisition (induction) of epigenet	g., marks such as methylation patte Ic variant	inis/aca particular locas
Inheritance of epigenetic variant	Obligatory	Facilitated	Independent of DNA sequence variation
Obligatory	The variant and its inheritance are fully determined by the specific DNA sequence	The DNA sequence affects the likelihood of acquiring particular variants; their inheritance is dependent on the DNA sequence	The DNA sequence does not determine which variant is acquired, but its inheritance is dependent on the DNA sequence
Facilitated	The variant is determined by the DNA sequence; some variants are more likely to be inherited than others	The DNA sequence affects both the likelihood of acquiring particular variants and the likelihood of their inheritance	The DNA sequence does not determine which variant is acquired but does affect the likelihood of it being inherited
Independent of DNA variation	The variant is determined by the DNA sequence, but the likelihood of it being inherited is not	The DNA sequence affects the likelihood of acquiring particular variants but not the likelihood of their being inherited	The DNA sequence determines neithe which variant is acquired nor the likelihood of it being inherited

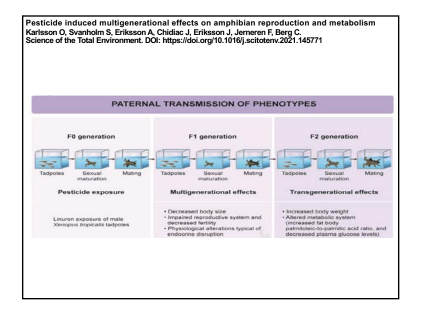


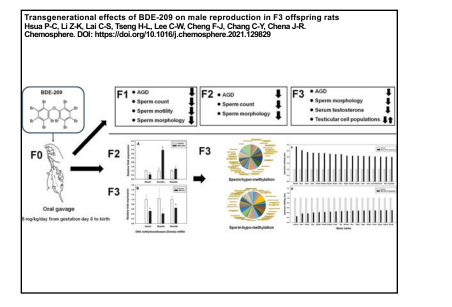
A	000	000	000	00	000	
Epimutation first	000-	→ 0 00- 0 0 0 00	→0 <u>0</u> 0-	→ <b>○</b>	→ 0 0 0 0 0 0	
B Epimutation first, provides survival advantage	000 000- 000	→ 000- 000- 000	→ <mark>○</mark> ○ ○ _	→ <sup>0</sup> 0 −	000 000 000	
C Epimutation as a programmed respons passive acceleration of genetic takeover	000 *000_ 000	→ 0 0 0 0 0 0 0 0 0	+ 000 000	000 000	0	Key Individual with active gene
D Epimutation as a programmed respons active acceleration of genetic takeover	000 •000 000	+ 0 0 0 0 0 0 0	→ 0 0 0 0 0 0		• -==	Individual with silenced gene

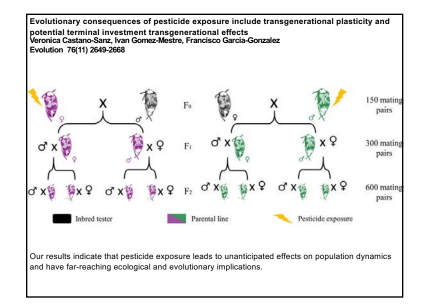


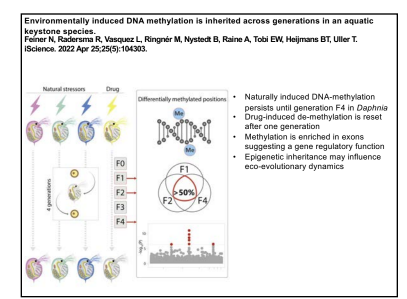


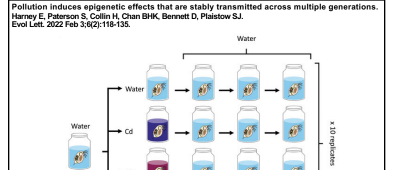












F1

Persistent effects are likely to influence phenotypic development, which could contribute to the rapid adaptation, or extinction, of populations confronted by anthropogenic stressors.

F2

F3

Unexposed

F0 Direct Exposure

#### **WO**FUNDAMENTAL CONCEPTS IN GENETICS

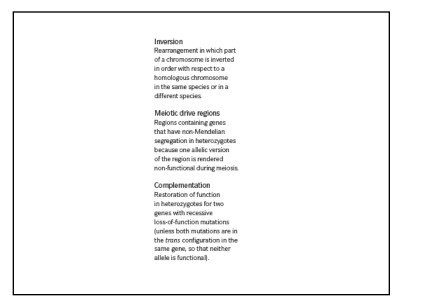
3 generations conditioning

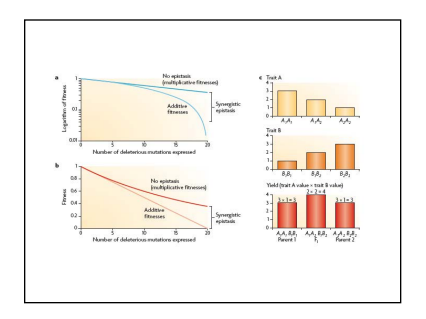
## The genetics of inbreeding depression

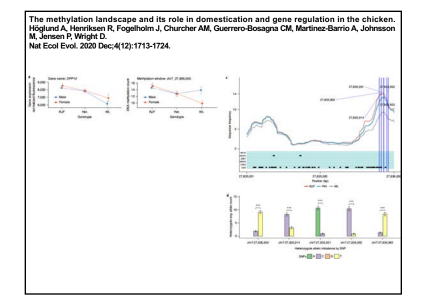
#### Deborah Charlesworth\* and John H. Willis\*

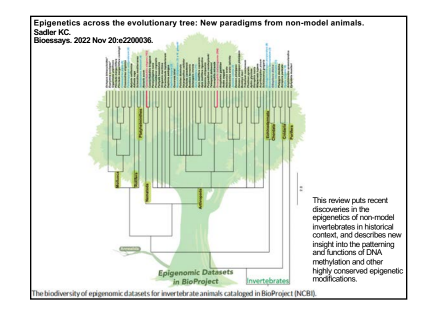
Abstract | Inbreeding depression — the reduced survival and fertility of offspring of related individuals — occurs in wild animal and plant populations as well as in humans, indicating that genetic variation in fitness traits exists in natural populations. Inbreeding depression is important in the evolution of outcrossing mating systems and, because intercrossing inbred strains improves yield (heterosis), which is important in crop breeding, the genetic basis of these effects has been debated since the early twentieth century. Classical genetic studies and modern molecular evolutionary approaches now suggest that inbreeding depression and heterosis are predominantly caused by the presence of recessive deleterious mutations in populations.

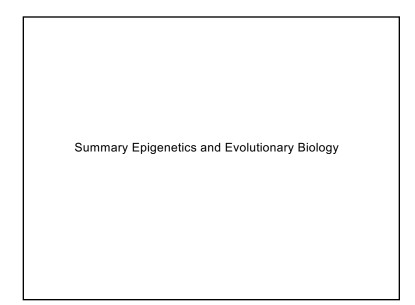
Inbreeding, Epigenetics and Evolutionary Biology

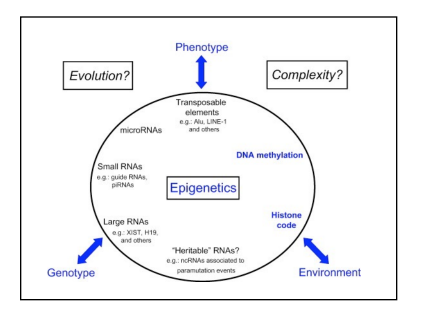


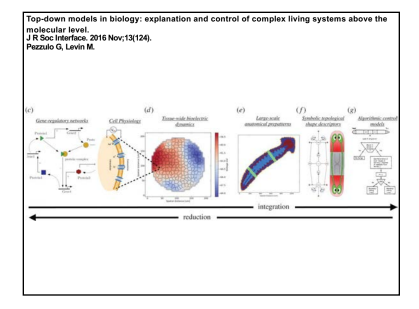


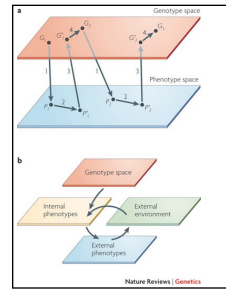








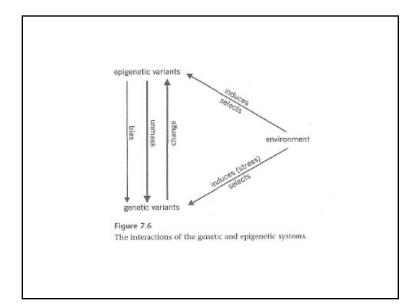


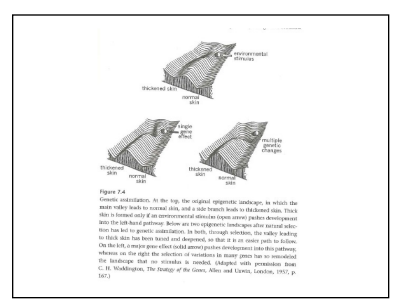


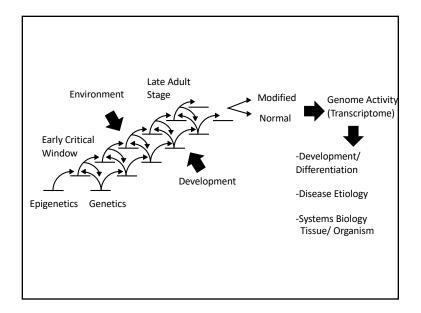
The concept of a genotype-phenotype (G-P) map is a widely used metaphor for the multiple ways in which genotypic information influences the phenotype of an organism. The term dates at least to 1970 when Jim Burns proposed linking population genetic and bichemical variation116, but the importance of the relationship between genotype and phenotype has long been apparent. Two early versions of the G-P map concept are the epigenetic landscape of Conrad Hal Waddington117 and Richard Lewonth's concept of evolution as taking place in the space of all possible genotypes (c space) and the space of all

#### possible phenotypes (P space)118.

This relationship is shown in part a of the figure, which indicates the mean position of a population in G and P spaces over two generations. There are four key parts to the evolutionary process, shown as numbered arrows: (1) the epigenetic process creates the phenotype using genotypic information; (2) natural selection acts in P space to change the average phenotype or parents away from the average phenotype or parents away from the average phenotype of parents average northe average phenotype or parents average phenotype of a parents attraction and the average phenotype or parents away from the average phenotype of parents average phenotype of an adverage phenotype of parents average phenotype of parents average and the space an attraction and recombination after position in G space. An attractive concept of the G-P map at the level of the individual is shown in part b of the figure. An individual can be conceptualized as occupying a single point in G space, and this position plus the environment (including other individuals; such as parents) combine to create the internal phenotypic state of the organism throughout its life. These internal phenotypes inturn shape there mytomment that individual occupies, creating complex feedback relationships between genes, environment sungests that we should explicitly toraden the G-P map to the genotype-environment-phenotype (G-E-P) map.

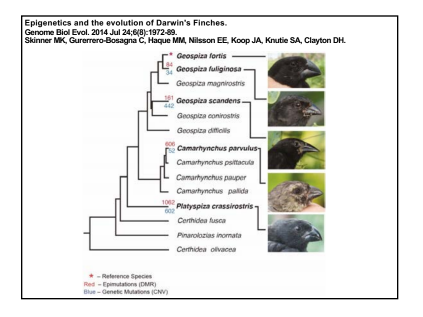






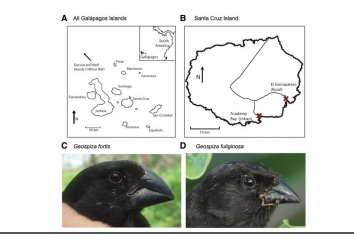
Inheritance system	Variation is-targeted (biased generation)?	Variation subject to developmental filtering and modification?	Variation constructed through direct planning?	Variations can change the selective environment?
Genetic	Generally not, except for the directed changes that are part of development and the various types of interpretive mutation	Usually not, although expressed genetic changes may have to survive selection between cells prior to sexual or asexual reproduction	No	Only insolar as genes can affect all aspects of epigenetics, behavior, and culture
Epigenetic	Yes, a lot of epigenetic variations are produced as specific responses to inducing signals	Ves, selection can occur between cells prior to reproduction; epigenetic states can be modified or reversed during meiosis and early embryogenesis	No	Yes, because the products of cellular activities can affect the environment in which a cell, its neighbors, and its descendants live
Behavioral	Yes, because of emotional, cognitive, and perceptual biases	Yes, behavior is selected and modified during the animal's lifetime	No	Yes, new social behavior and traditions alter the social and sometimes also the physical conditions in which an animal lives
Symbolic	Yes, because of emotional, cognitive, and perceptual biases	Yes, at many levels, in many ways	Yes, at many levels, in many ways	Yes, very extensively, by affecting many aspects of the social and physical conditions of life

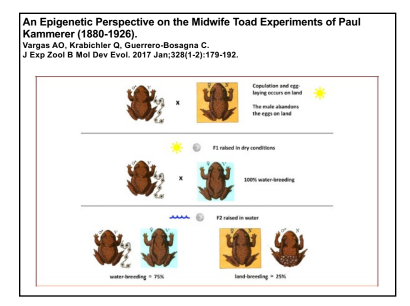
The reproduction of i	nformation				
Inheritance system	Organization of Information	Dedicated copying system?	Transmits latent (nonexpressed) information?	Direction of transmission	Range of variation
Genetic	Modular	Yes	Yes	Mostly vertical	Unlimited
Epigenetic Self-sustaining loops	Holistic	No	No	Mostly vertical	Limited at the loop level
Structural templating	Holistic	No	No	Mostly vertical	unlimited at the cell level Limited at the structure level.
RNA silencing	Holistic	Yes	Sometimes	Vertical and sometimes	unlimited at the cell level Limited at the single transcript level, unlimited at the cell level
Chromatin marks	Modular and holistic	Yes (for	Sometimes	horizontal Vertical	Unlimited
Organism-level developmental legacies	Holistic	methylation) No	No	Mostly vertical	Limited
Behavioral					
Behavior-affecting substances	Holistic	No	No	Both vertical	Limited at the single behavior
Nonimitative social learning	Holistic	No	No	and horizontal Both vertical and horizontal	level, unlimited for lifestyles Limited at the single behavior level, unlimited for
Imitation	Modular	Probably	No	Both vertical and horizontal	lifestyles Unlimited
symbolic	Modular and holistic	Yes, several	Yes	Both vertical and horizontal	Unlimited

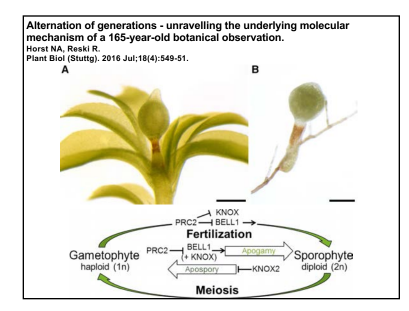


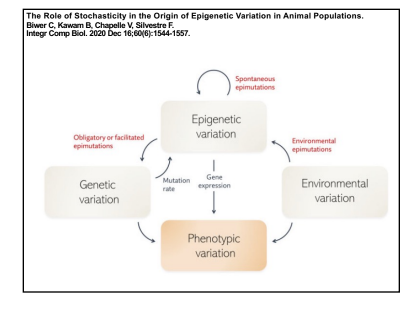
Epigenetic variation between urban and rural populations of Darwin's finches. BMC Evol Biol. 2017 Aug 24;17(1):183.

McNew SM, Beck D, Sadler-Riggleman I, Knutie SA, Koop JAH, Clayton DH, Skinner MK.



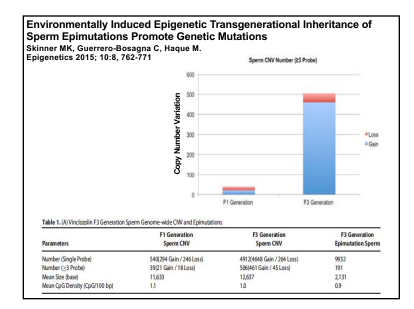


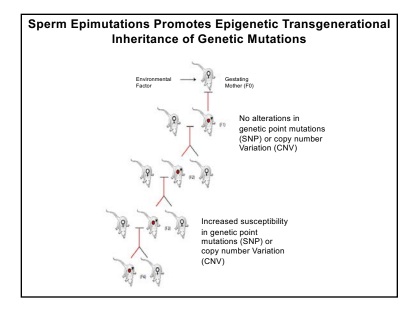


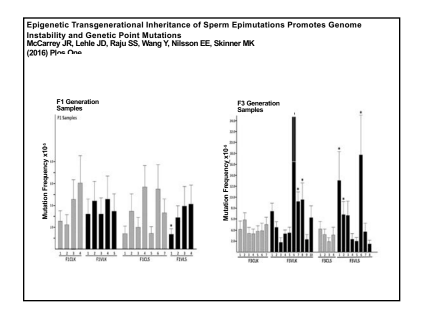


Leaptrog to special Verzijden M. Nature. 2019 Oct;574(	ion boosted by mother's influence. 7776):38-39.
	It has now been found that mothers of a species of frog affect the behaviour of their offspring — influencing female mating preferences and aggression between males. Such behaviours might lead to the formation of new species.

Genetic Mutation	Epigenetic Alteration	DNA Sequence Alteration
Point Mutation (SNP)	DNA Methylation (CpG)	Susceptibility C ➤ T Conversion
Copy Number Variation (CNV)	Hypomethylation (Repeats)	Susceptibility Repeat Element Alteration (CNV)
Transposon Migration	Hypomethylation DNA	Susceptibility Transposon Migratio
Translocation	DNA Methylation and Histone Alterations	Susceptibility Translocation at Break Point
Telomere Length	DNA Methylation Alteration	Alteration in Telomere Length







Environmental Epigenetics and a Unified Theory of the Molecular Aspects of Evolution: A Neo-Lamarckian Concept that Facilitates Neo-Darwinian Evolution. Skinner MK.

Genome Biol Evol. 2015 Apr 26;7(5):1296-302.

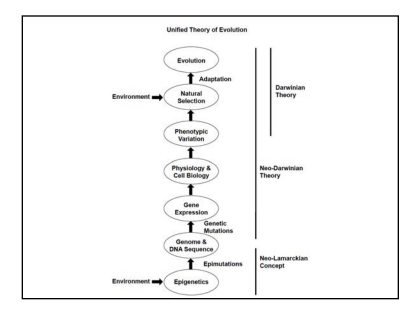
#### Evolution Theory Components

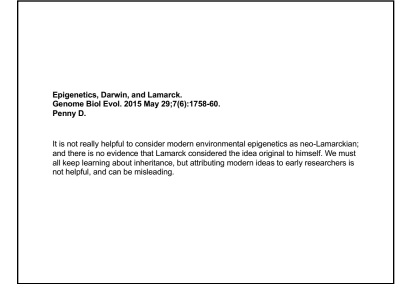
Neo-Lamarckian concept Environment directly alters phenotype generationally Darwinian evolution theory Natural selection acts on phenotypic variation Neo-Darwinian evolution theory

Genetic mutations promote phenotypic variation on which natural selection acts

Unified evolution theory

Environmental epigenetic alterations promote genetic mutations to alter genotypic variation Environmental epigenetics and genetic mutations both promote phenotypic variation on which natural selection acts





Heredity determined by the environment: Lamarckian ideas in modern molecular biology. Tikhodeyev ON. Sci Total Environ. 2020 Mar 25;710:135521. LAMARCKIAN INHERITANCE (PLANT EXAMPLE) prolonged exposure to streptomycin functional plastid ribosomes NO plastid ribosomes X maternal inheritance NO plastid plastid of the gene NO gene acquired expression plastid phenotype translation acquired normal plant albino phenotype

Lamarck and Panspermia - On the Efficient Spread of Living Systems Throughout the Cosmos. Steele EJ, Gorczynski RM, Lindley RA, Liu Y, Temple R, Tokoro G, Wickramasinghe DT, Wickramasinghe NC. Prog Biophys Mol Biol. 2019 Dec;149:10-32.

#### Table 1

2

6

Evidence consistent with Lamarckian evolutionary processes.

- Environmental Stimulation as the Directional Mutational Driver
- Role of Epigenetic Gene Targeting
- **Rapid Genetic Adaptation** 2
- Penetration of the Weismann Barrier
- Horizontal Gene Transfer (HGT)
- Central Role of Reverse Transcription

The summaries of evidence for Horizontal Gene Transfer phenomena are well covered at the Wikipedia site https://en.wikipedia.org/wiki/Horizontal\_gene\_ transfer.

#### Table 2

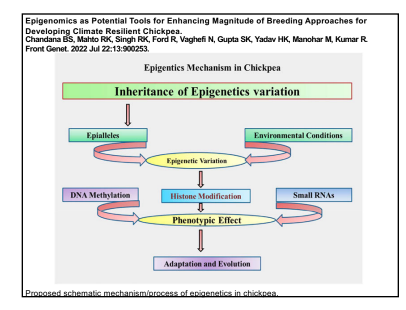
Cosmic distribution and numbers of living systems.

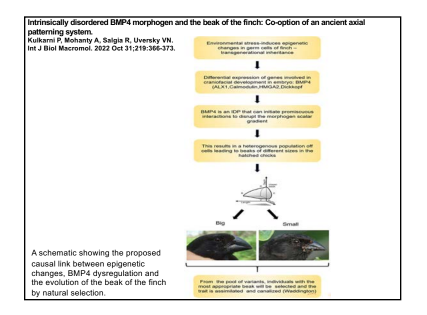
1053 Viruses – terrestrial number 10<sup>31</sup> 1052 • Bacteria/Archaea - terrestrial number  $\geq 10^{30}$ 1032-1052 Single cell eukaryotes - terrestrial number 10<sup>20</sup>-10<sup>30</sup> 1042

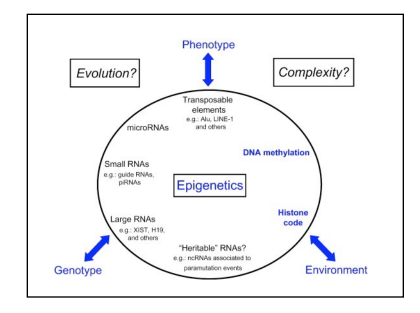
1029

1029

- Complex Metazoans terrestrial number  $\ge 10^{20}$
- Higher plants, terrestrial number  $\geq 10^7$  species
- Higher animals, terrestrial number ≥10<sup>7</sup> species







s Biology	Systems Biology	Revolutiona	ary Systems Biology
Ecosystem Populations	Ecosystem	Ecosystem	Nonlinear
Organisms	<b>↑</b> ↓ Organisms	Organisms	
↑ Physiology	↑↓ Physiology ↑↓	↓ Physiology	Robustness
Organ Systems	Organ Systems ↑↓	Organ Systems ↓	Synergism
Organs Tissues	Organs ↑↓ Tissues	Organs J Tissues	Emergence
Cells	∱↓ Cells	Cells	Epigenetics
Organelles ↑	Organelles ↑ ↓	Organelles	Holism
Macromolecules	Macromolecules ↑↓	Macromolecules	nonsm
	Populations     A     Organisms     Physiology     A     Organ Systems     A     Organs     Tissues     Cells     Organelles     A	↑     ↑     Populations       Populations     ↑     Populations       ↑     Organisms     ↑       Physiology     ↑     ↓       Physiology     ↑     ↓       Organ Systems     Organ Systems     ↓       Organs     ↓     ↓       Organelles     ↓     ↓       Macromolecules     ↓     ↓	↑     ↓     ↓       Populations     Populations     Populations       ↑     Populations     Organisms       ↑     ↓     ↓       Organisms     ↓     ↓       ↑     ↓     ↓       Physiology     ↑↓     ↓       Physiology     ↑↓     ↓       Organ Systems     Organ Systems     Organ Systems       ↑     ↓     ↓       Organs     Organs     Organs       ↑     ↓↓     ↓       Organs     Organs     Organs       ↓     ↓     ↓     ↓       Organelles     ↓↓     ↓       ↓     ↓     ↓       Macromolecules     ↓     ↓

#### "Epigenetics and Systems Biology"

#### Spring 2023 (Odd Years) Biol 476/576 Schedule/Lecture Outline -Week 1 January 10 & 12 Systems Biology (History/ Definitions/ Theory) January 10 & 12 January 17 & 19 January 24 & 26 Systems Biology (Networks & Emergence) Systems Biology (Components: DNA to Phenotype) Week 2 Week 3 Systems Biology (Genomics / Technology) Epigenetics (History / Molecular Processes) Week 4 Jan 31 & Feb 2 February 7 & 9 Week 5 Week 6 February 14 & 16 Epigenetics (Molecular Processes & Integration) February 21 & 23 Feb 28 & March 2 Epigenetics (Genomics and Technology) Week 7 Cell & Developmental Biology Epigenetics of Cell & Developmental Biology (& Midterm Exam) Week 8 Week 9 March 7 & 9 Week 10 March 13 - 17 Spring Break Environmental Impact on Biology March 21 & 23 Week 11 Week 12 March 28 & 30 Environmental Epigenetics Disease Etiology Week 13 April 4 & 6 Epigenetics & Disease Etiology Evolutionary Biology & Genetics Epigenetics & Evolutionary Biology Grant Review/ Study Section Meeting (& Final Exam) Week 14 April 11 & 13 April 18 & 20 Week 15 April 25 & 27 May 2 & 4 Week 16 Week 17