1.1 Introduction

Life histories—describing essential patterns of organismal growth, maturation, reproduction, and survival—show tremendous variation across individuals, populations, species, and environments. Understanding this variation is the goal of life history research. The analytical framework of life history theory focuses on the variation and interaction of different key maturational, reproductive, and other demographic traits, given that natural selection acts to maximize fitness of a life history as a whole (Roff 1992, Stearns 1992). Fitness integrates over the entire reproductive performance of the organism, and life history traits are the major fitness components underlying this integration. However, the investment into alternative life history traits, and thus the possible set of trait combinations, is restricted by genetic, developmental, physiological, and phylogenetic limits. Apart from explaining variation in life history strategies as a result of natural selection, identifying how such trade-offs and constraints shape life histories is the central aim of life history research.

In this chapter we introduce the basic concepts and definitions of life history theory and argue for the importance of integrating a mechanistic perspective into research on life histories. While most traditional life history research is based on mathematical, statistical, and phylogenetic approaches without explicit reference to underlying mechanisms, today’s principal research challenge is to fill this gap through experimental characterization of the proximate basis of life histories. The analysis of genetic, developmental, and physiological factors that shape life history traits will ultimately allow us to determine how evolutionary changes in such mechanisms generate, facilitate, or constrain the diversification of life histories. Integrating mechanistic and evolutionary analyses of life history variation is part of a global quest in biology that seeks a shared understanding of proximate and ultimate causes of phenotypic variation.

1.2 The life history framework

1.2.1 What is a life history?

A life history encompasses the life of an individual from its birth to its death, describing the age- or stage-specific patterns of maturation, reproduction, survival, and death. The major objective of life history research is to understand how evolution, given selection imposed by ecological challenges, shapes organisms to achieve reproductive success. The second objective of life history research is to understand whether and how, given internal trade-offs and constraints, selection can optimize a set of life history traits to maximize reproductive success. Since organisms dispose of limited resources, which must be competitively allocated to differing functions, such as growth, reproduction, survival, and maintenance, resources invested into one function cannot be invested into another, leading to trade-offs. In addition, life history research explores
taxon-specific features of life cycles and life history decisions, including patterns of sex allocation, alternative phenotypes, or larva-to-adult transitions. For in-depth treatments of the evolution of life histories and life history theory see Stearns (1992), Roff (1992, 2002), and Charlesworth (1994).

1.2.2 Life history traits and fitness

Life history traits represent quantitative, demographic properties of organisms that are directly related to the two major components of fitness, i.e., survival and reproduction. Classical life history analysis considers the following to be the principal life history traits (Stearns 1992):

- size at birth
- growth pattern
- age and size at maturity
- number, size, and sex ratio of offspring
- age- and size-specific reproductive investments
- age- and size-specific mortality schedules
- length of life.

These traits essentially represent the demographic parameters required to estimate fitness as defined by the Malthusian parameter (or similar fitness measures). The Malthusian parameter (also called the instantaneous rate of natural increase, \( r \)) is the solution to the Euler–Lotka equation, which describes population growth by summing reproductive events and survival probabilities over the entire lifetime of individuals (Stearns 1992). Thus, life history traits are directly linked to fitness, with fitness being defined by population growth models from demography.

In contrast to classical life history traits, morphological, physiological, or behavioral traits are considered to contribute to fitness only indirectly (e.g., Roff 2007b). However, this distinction is somewhat arbitrary. For example, certain morphological traits such as body size or gonad size may correspond to life history traits (or at least are correlates thereof).

In the literature, the term “life history trait” is often used interchangeably with fitness components, so that many phenotypic characters with major effects on reproduction and survival have been called life history traits.

Because of their complexity and demographic nature, life history traits are usually treated as quantitative, polygenic traits (Falconer and MacKay 1996). The expression of life history traits is also highly contingent on the environment, so that life history research places particular emphasis onto the concept of phenotypic plasticity, i.e., the ability of a single genotype to produce different phenotypes across environments (Stearns 1992). Plasticity is described by “reaction norms”, mathematical functions that relate the phenotypic values adopted by a given genotype to changes in the environment. Selection shapes life history plasticity by acting on genetic variation for plasticity, which is present when the reaction norms that represent different genotypes are non-parallel across the same range of environments (so-called genotype by environment interactions, or \( G \times E \)). Reaction norms (and thus plasticity) are considered to be optimal when they maximize fitness for each of the different environments (Stearns and Koella 1986).

1.2.3 Trade-offs and constraints

A key postulate of life history theory is that the values and combinations of life history traits are limited by factors internal to the organism, namely trade-offs and constraints. These intrinsic factors ultimately limit and direct the evolutionary response to the external force of selection. A life history trade-off occurs when an increased investment in one fitness component causes a reduced investment in another, i.e., a fitness benefit in one trait exacts a fitness cost in another. Examples of classical life history trade-offs are survival versus reproduction, number versus size of offspring, or current reproduction versus future reproduction (Stearns 1992).

Trade-offs are usually described as phenotypic or genetic covariances or correlations among traits, without reference to their causal relationships. If the relationship can be shown to be genetic, negative genetic covariance among traits is expected to limit the evolution of each of these traits. Such genetic or evolutionary trade-offs are considered at the population level, i.e., as defined by genetic correlations among individuals or correlated phenotypic responses to selection. Genetic trade-offs are traditionally assumed to stem from antagonistic pleiotropy or linkage disequilibrium. These trade-offs
also manifest themselves at the physiological or individual level, for example when an individual with increased reproductive effort in one year exhibits a reduction in reproductive output in the next year. Such physiological trade-offs are thought to be due energy limitations, i.e., the allocation of resources among competing functions. Importantly, trade-offs may exist at population level, but not at individual, physiological level (Stearns 1989, Houle 1991, Stearns 1992).

In contrast to trade-offs, the term “constraint” is often used to described absolute limits to or biases upon trait expression and combination. Constraints may describe physical factors, developmental properties, or historical contingencies that prevent an organism from expressing a certain phenotype or a population from attaining a certain fitness optimum in response to selection (Maynard Smith et al. 1985). The distinction between trade-offs and constraints is not strict, and trade-offs are often regarded as one type of constraint. In the life history context, constraints usually refer to phylogenetic, lineage-specific characteristics that impose absolute limits on trait expression in a given organismal group.

1.2.4 Empirical approaches in life history research

Although classic life history analysis has been largely theory-driven, much empirical research has addressed the questions and predictions raised by life history theory, using both non-genetic and genetic approaches (Stearns 1992, Roff 1992, 2002, 2007b; also see Chapter 2). Non-genetic approaches include phenotypic correlations to examine patterns of life history trait covariation among populations and species, experimental phenotypic manipulations, and statistical tools from comparative analysis to control for phylogenetic history. Genetic approaches to the study of life history variation are predominantly based on the framework of quantitative genetics. Most of this work has concentrated on the detection and analysis of genetic trade-offs, either through the study of covariances and correlations among life history traits between relatives (e.g., pedigree analyses) or through correlated responses of life history traits to artificial selection or experimental evolution. This research framework has generated a substantial body of empirical evidence that has revealed how selection operates on life history traits, contingent on the environment and trade-offs (Stearns 1992, Roff 1992, 2002, 2007a,b). Despite these extensive efforts, very few studies have examined the mechanistic underpinnings of life history traits. For example, inferred interrelationships among life history traits rarely describe more than statistically determined associations. A major limitation common to the classical approaches in life history research is therefore the ignorance of the proximate causes that determine or modulate life histories and their evolution.

1.3 The study of causal mechanisms linking genotype to phenotype

Understanding how a genotype translates into a phenotype is one of the most fundamental problems in biology. In most cases, phenotypes cannot be simply inferred from their underlying genotypes, and vice versa, because the mapping of genotypes onto phenotypes is often a non-linear process, shaped by a multitude of complex genetic and environmental interactions. Moreover, a single genotype may generate multiple phenotypes and, conversely, multiple genotypes may generate a single phenotype. That such properties of the genotype–phenotype map are relevant for our understanding of the evolutionary process has been emphasized for a long time (e.g., Lewontin 1974, Houle 2001), but it is only relatively recently that the causal relationships between genotype and phenotype have received increased attention from evolutionary biologists (e.g., Pigliucci 2010). While research at the interface of development and evolution has begun to tackle the significance of the genotype–phenotype map in morphological evolution, the causal connection between genotypes and phenotypes for fitness components is still extremely rudimentary (e.g., Chapter 2 and Roff 2007b).

Traditionally, attempts to link the genotype with the phenotype have been regarded as the principal task of “reductionist” branches of biology, including molecular, cellular, and developmental biology. Developmental genetics in particular has emerged as the prime discipline in connecting gene function during development with phenotypic outcomes,
primarily by relying on mutational analysis and forward genetics. The great power of this approach lies in the typically high degree of causal inference that can be made through carefully controlled manipulation of isolated genetic factors and their phenotypic effects. The general downside of this approach is that such studies are generally limited to the study of single, highly pleiotropic mutations with large phenotypic effects. In addition, developmental genetic analyses are generally limited to the study of a single or a small number of laboratory populations in highly simplified artificial environments, aiming to reduce variation engendered by genetic background or environmental context as much as possible. This research approach starkly contrasts with that of evolutionary biologists, whose primary concern is the study of quantitative genotypic and phenotypic variation among populations or species. Here, in contrast to developmental genetics, the inferred genotype–phenotype relationships are generally of indirect, associative nature, rarely permitting inferences about the causal connections between genotypic and phenotypic variation.

As advocated in many chapters throughout this book, a better future understanding of many issues in life history evolution will require the integration of evolutionary and organismal biology with molecular and developmental biology (e.g., Dean and Thornton 2007). That unfortunate historical separations between biological disciplines can be overcome is well illustrated by the successful rapprochement of evolutionary and developmental biology (e.g., Raff and Kaufman 1983, Carroll et al. 2000, Stern 2010). Although initially mainly concerned with the description of evolutionary diversification or conservation of developmental mechanisms, the central aim of evolutionary developmental biology (evo-devo) has recently shifted to the experimental analysis of how properties of genetic and developmental architecture impact phenotypic evolution. Evo-devo therefore addresses specific issues directly relevant to the understanding of life history evolution, such as the mechanistic basis of developmental biases and constraints or phenotypic plasticity. More generally, as life history traits are high-level phenotypes that depend on the ensemble of morphological and physiological traits, the mechanistic analysis of life history evolution can consequently be regarded as an extension of the principal objective of evo-devo, namely to understand which developmental and genetic changes underlie phenotypic evolution.

Uncovering the mechanistic basis of life history variation is a non-trivial challenge. Life history traits were defined by evolutionary ecologists with the intent of reducing phenotypic complexity by focusing on a small number of traits that summarize the essential fitness components and by ignoring the underlying genetic, developmental, and physiological mechanisms that govern the expression of these traits. A given life history trait can thus be thought of as a functionally complex phenotype resulting from the integration of a suite of morphological, physiological, or behavioral phenotypes. At the level of the individual, their characteristics have therefore to be understood in terms of both the construction of multiple individual traits as well as their spatial and temporal integration into a higher-level phenotype. As such, life history traits are a priori composite, quantitative, polygenic traits whose expression is often highly contingent upon plasticity, pleiotropy, and epistasis. All these properties render the mechanistic analysis of life history traits extremely difficult in practice.

1.4 How can mechanistic insights contribute to understanding life history evolution?

Despite the inherent difficulties in studying the proximate basis of life histories, considerable progress has been made in our mechanistic understanding of life history evolution, with major contributions stemming from molecular genetic studies on experimental model organisms. Here we briefly discuss the importance of integrating such mechanistic information into organismal life history research; many more detailed examples can be found throughout the chapters in this book. For further reading on integrative approaches in life history biology we recommend the reviews by Houle (2001), Leroi (2001), Barnes and Partridge (2003), Harshman and Zera (2007), Chapter 5 in Van Straalen and Roelofs (2006), Roff (2007b), and Flatt and Schmidt (2009).
1.4.1 Why understanding mechanisms is important for answering evolutionary questions

While it is clear that knowledge of the proximate basis of life histories does not provide information about the ecological or evolutionary relevance of such mechanisms, it enables evolutionary biologists to address several fundamental questions about life history evolution, including, for example:

- What is the function of genes that are genetically variable in natural populations and that contribute to ecological adaptation?
- Are major candidate genes, as identified by molecular genetics, variable in natural populations?
- If so, do polymorphisms at these loci actually contribute to the evolution of life history traits in the wild?
- Are the genes that impact life history evolutionarily conserved or lineage-specific?
- What genetic and physiological mechanisms determine or modulate the expression of ecologically and evolutionarily important trade-offs?
- Are such trade-offs, as commonly assumed, resource based, or are they due to mechanisms independent of energy allocation?
- What are the mechanisms that mediate life history plasticity?

1.4.2 The molecular identity and function of genes that affect life history

Studies in molecular and developmental genetics inform us about the molecular identity and function of genes, including those that affect life history traits and other fitness components. The functionally best-understood genes that affect life history traits have been analyzed in model organisms such as Arabidopsis, Drosophila, or C. elegans. Information about the function of such genes is useful, for example, when evolutionary biologists want to investigate the consequences of allelic variation at such loci in natural populations. Although natural alleles might have much more subtle phenotypes than laboratory induced mutant alleles, detailed knowledge about gene function might help organismal biologists to understand whether and how particular genes contribute to ecologically relevant phenotypes and thus why selection acts on such loci. This does not mean that every gene with a major phenotypic effect on a fitness-related trait, as identified by molecular genetics, is in fact ecologically or evolutionarily relevant in natural populations; many such genes might not harbor standing genetic variation affecting life history phenotypes and might therefore not contribute to evolutionary change in the wild. Yet, it is also clear that loci that do contribute to phenotypic variation in fitness-related traits and thus to ecological adaptation in natural populations are a subset of all genes, including those that have been functionally studied by molecular geneticists (e.g., Stern 2000, Flatt 2004, Flatt and Schmidt 2009).

While developmental and molecular genetic approaches do inform us about the ecological or evolutionary significance of specific genes, they have proved powerful in identifying the molecular mechanisms that affect life history traits, for instance their endocrine regulation (Tatar et al. 2003, Fielenbach and Antebi 2008). Perhaps the best examples are genes known to affect adult survival and longevity in the nematode, fruit fly, and mouse; these have received particular attention, not only from biomedical researchers because of their potential implications for human gerontology (see Chapter 16), but also from evolutionary biologists because of their potential relevance for understanding the evolution of aging. During the past 20 years, numerous mutations that extend lifespan have been identified in diverse model organisms (e.g., Kenyon 2010; also see Chapter 14). Many of these mutations were found to affect a key metabolic pathway—the insulin/insulin-like growth factor signaling pathway—indicating that decreased effectiveness of insulin/IGF-like signaling causes lifespan extension, linked to correlated responses in reproduction, growth, and metabolism. These pivotal discoveries, many of which are discussed in this book, not only demonstrate the feasibility of molecular genetic analyses of complex life history traits such as lifespan, but also suggest that certain evolutionarily conserved signaling pathways are potential key regulators of major life history traits (also see Chapters 27 and 28). Many of these findings have also contributed to our understanding of life history...
trade-offs (see below and Chapters 11 and 13). The molecular genetic analysis of lifespan has thus rapidly become of great interest to many researchers studying life histories, and this interest is now paving the way for an integration of mechanistic and evolutionary approaches towards the understanding of life history variation (e.g., Partridge and Gems 2006, Flatt and Schmidt 2009).

In addition to functional studies of individual mutations, genome-wide gene expression analyses have also been widely used by both molecular and evolutionary biologists to investigate the proximate basis of life history variation (as is discussed in detail in Chapter 2). For example, genome-wide transcriptional profiling has been used to identify candidate genes involved in lifespan regulation (e.g. Murphy et al. 2003), or to describe gene expression patterns associated with particular life history stages, for example dauer larva formation in C. elegans (Wang and Kim 2003). Many of these studies illustrate the complex and manifold changes in gene expression associated with life history variation and further indicate that life history trade-offs might emerge through “conflicts over gene expression”, i.e., antagonistic pleiotropic effects of genes involved in multiple functions (Stearns and Magwene 2003, Bochdanovits and de Jong 2004). However, the functional interpretation of such data remains challenging because the precise causal connections between transcriptional changes and the resulting phenotypes are rarely known. Thus, while it is clear from these few examples that we have learned a great deal about the molecular genetic basis of life history traits, a current key challenge is to integrate such mechanistic insights into the evolutionary framework (also see Chapters 27 and 28). One obvious question for the evolutionary biologist is, for example, whether the candidate genes identified by molecular geneticists actually matter in natural populations.

1.4.3 Are candidate life history genes ecologically and evolutionarily relevant?

Mutational, transgenic, and genomic analyses in model organisms have been successful in identifying at least some of the key mechanisms that affect life history traits. However, while many of these mechanisms show a surprisingly high degree of conservation across widely divergent taxa, their relevance in shaping evolutionary life history variation in natural populations is not yet sufficiently clear. Determining whether and how such mechanisms evolve to generate natural life history variation represents a promising starting point for the integration of functional and evolutionary analysis of life histories. In most cases, however, such studies are limited to model organisms. Such an analysis requires testing of whether the genes involved in these candidate mechanisms show actual variation in natural populations and, as a more challenging step, to functionally demonstrate that this allelic variation impacts the life history trait in question.

Several studies suggest that genes identified through molecular and developmental genetic analyses indeed harbor natural allelic variation that contributes to population variation in life history traits, for example in Drosophila (e.g., Schmidt et al. 2000, Paaby and Schmidt 2008, Paaby et al. 2010; also see Chapter 18), or in Arabidopsis (e.g., Todesco et al. 2010; also see Chapter 9). Although the screening of natural polymorphisms in candidate life history genes only provides a first glimpse of the molecular basis of life history variation, such initial findings are encouraging since they indicate that developmental and molecular genetic studies indeed generate valuable candidate genes of interest for evolutionary biologists.

In contrast to the analysis of natural allelic variants at major candidate loci identified by molecular and developmental genetics, quantitative trait locus (QTL) mapping provides a less biased, yet technically challenging, approach to the characterization of the genetic basis of polygenic quantitative traits, including life history traits (Falconer and Mackay 1996). While classical QTL mapping approaches have been useful in determining the basic genetic architecture of life history traits (e.g., the number and effect size of the involved loci), they rarely achieve sufficient resolution to pinpoint individual candidate genes (see discussion in Roff 2007b and Mackay et al. 2009). However, recent technological advances, such as rapid and cost-effective genotyping methods and refined statistical and mapping methods, have increased the feasibility of high-resolution mapping, now allowing the identifica-
tion of candidate genes within QTL regions for organisms with well-annotated genomes, in some cases down to the level of single nucleotide polymorphisms (e.g., Mackay et al. 2009). High-resolution mapping through recombinant inbred lines and genome-wide association studies have already been successful in characterizing natural polymorphisms underlying genetic variation in complex developmental or life history traits in C. elegans (e.g., Kammenga et al. 2007, Palopoli et al. 2008), Drosophila (e.g., De Luca et al. 2003, Schmidt et al. 2008, also see Flatt and Schmidt 2009 for a recent review), and in Arabidopsis (e.g., Atwell et al. 2010, also see Chapter 9). Moreover, recent progress in genomic methods now allows the researcher to treat genome-wide expression patterns as complex quantitative traits (e.g., Rockman 2008).

The recent advent of refined QTL and genetical genomics approaches is emblematic for an integrative and novel research program, namely the use of natural genetic variation as a tool to understand the causal connection between genotype and phenotype. By explicitly taking evolutionary variation into account, this approach holds great promise for facilitating the detection of mechanistic features that are involved in phenotype construction. However, the identification of individual genes or nucleotide polymorphisms that contribute to quantitative trait variation remains a major challenge because of subtle phenotypic effects, complex genetic interactions, pleiotropy, and genotype-by-environment interactions (e.g., Weigel and Nordborg 2005, Mackay et al. 2009).

1.4.4 How do trade-offs work?

One central and recurring theme in this book is the mechanisms that underlie life history trade-offs (see the chapters in Part 6). Given the central importance of such trade-offs in life history evolution, uncovering their mechanistic basis is one of the most fundamental but unresolved problems in life history research (e.g., Stearns 2000, also see Chapters 27 and 28). Despite numerous and seemingly obvious trade-offs between life history traits in a wide range of taxa, most reported trade-off relationships basically describe no more than a statistically inferred negative correlation. The description of trade-offs by means of trait correlations or covariances is, however, insufficient for evaluating how genetic architecture influences evolutionary trajectories (e.g., see Chapter 2 and Roff 2007b). Specifically, it remains to be determined to what extent presumptive trade-offs are conclusively due to actual competition for limited resources or caused by alternative mechanisms, such as hormonal signaling independent of resource allocation (see Chapters 11, 13, 27, and 28). The very limited knowledge on the mechanistic underpinnings of trade-offs therefore represents a current key problem in our understanding of life history evolution (e.g., Stearns 2000, Flatt et al. 2005, Roff 2007b, Flatt and Schmidt 2009).

Recent progress in this area comes again from the molecular genetic analysis of lifespan. Several studies on the relationship between lifespan and reproduction in worms and flies have challenged the fundamental notion that reproduction exacts an energetic cost in terms of reduced survival (e.g., see Chapter 11, Leroi 2001, Barnes and Partridge 2003). Of particular relevance was the observation of a C. elegans insulin receptor mutant with extended lifespan (Kenyon et al. 1993). Although this mutant exhibited decreased fecundity—consistent with a resource-allocation trade-off where investment in longevity extension lowers investment in reproduction—detailed experimental analysis of this relationship indicates that decreased reproduction is not the causal agent in extending longevity (e.g., Kenyon et al. 1993, Leroi 2001). Therefore, reproductive versus somatic investment may not necessarily be coupled through resource competition but rather via independent underlying signaling processes (see Chapters 11, 13, and 24, and Hsin and Kenyon 1999, Flatt et al. 2008b). While these findings do not prove the absence of a cost of reproduction (Barnes and Partridge 2003, Flatt and Schmidt 2009), they underscore the difficulty of inferring resource-allocation trade-offs without a precise understanding of the proximate mechanisms involved. For example, a major technical challenge in demonstrating the resource basis of trade-offs is to experimentally track resource allocation to different organismal functions by detailed measurement of relevant parameters, such as nutrient ingestion and assimilation (see Chapter 24 and O’Brien et al. 2008).
Other valuable information on the mechanistic basis of life history trade-offs comes from research exploring the fitness consequences of organismal defensive mechanisms against pathogens, parasites, stresses, or toxins. For example, studies in both vertebrates and invertebrates indicate that elevated immune and other defense functions incur fitness costs in terms of reproduction and survival (see, for example, Chapters 2 and 23, Flatt et al. 2005, Harshman and Zera 2007). Similarly, the evolution of pesticide tolerance in insects often results in a fitness cost, which is generally supposed to stem from increased energy allocation to corresponding detoxification mechanisms. Remarkably, however, it turns out that such fitness costs can result from collateral metabolic costs rather than energetic costs due to the detoxification mechanism (Van Straalen and Hoffmann 2000).

Thus, while many observations support the existence and evolutionary relevance of life history trade-offs, their underlying causal mechanisms still remain rather poorly understood. Importantly, one of the central postulates of life history theory, namely that trade-offs are caused by competitive resource allocation, might not necessarily always hold. As discussed in many chapters throughout this book (e.g., Chapters 11, 13, 27, and 28), major efforts are currently under way to dissect the mechanistic basis of life history trade-offs.

1.5 Conclusions

Combining mechanistic and evolutionary analyses of life history variation is a fundamental yet ambitious aim in current biology. On the one hand, there are inherent biological and technical problems with studying complex quantitative phenotypes such as life history traits. On the other hand, there are cultural divides that necessitate a combination of diverse research approaches and concepts from both molecular and organismal biology. Despite these challenges, the chapters in this book illustrate that the successful integration of mechanisms into life history research is fully under way.