

Fitness and Individuality in Complex Life Cycles

Matthew D. Herron*†

Complex life cycles are common in the eukaryotic world, and they complicate the question of how to define individuality. Using a bottom-up, gene-centric approach, I consider the concept of fitness in the context of complex life cycles. I analyze the fitness effects of an allele (or a trait) on different biological units within a complex life history and how these effects drive evolutionary change within populations. Based on these effects, I attempt to construct a concept of fitness that accurately predicts evolutionary change in the context of complex life cycles.

1. Introduction. Nearly all eukaryotes alternate haploid and diploid generations. In many cases one or the other generation is reduced to one or a few cells; in animals, for example, the haploid phase is a single cell with limited or no expression of the genes it carries (the gametes). In land plants, either the haploid phase (ferns, seed plants) or the diploid phase (mosses, liverworts, hornworts) is reduced to varying degrees. At the opposite extreme, some taxa have haploid and diploid phases that are virtually indistinguishable (e.g., the ulvophyte green algae *Ulva* and *Cladophora*). Animals and *Ulva* represent the ends of a continuum between isomorphic (phases indistinguishable) and heteromorphic (phases morphologically distinct) alternation of generations. Intermediate examples exist with varying degrees of heteromorphy, ranging from near isomorphy (e.g., the brown alga *Ectocarpus*) to so different that the

*To contact the author, please write to: Division of Biological Sciences, University of Montana, 32 Campus Dr., Missoula, MT 59812; e-mail: xprinceps@gmail.com.

†I would like to thank Maureen O'Malley, Peter Godfrey-Smith, and James Griesemer for organizing the symposium. During this work, I was supported by a postdoctoral fellowship from the NASA Astrobiology Institute and by grants from the John Templeton Foundation (43285), the NASA Astrobiology Institute (Cooperative Agreement Notice 7), and the National Science Foundation (DEB-1457701).

Philosophy of Science, 83 (December 2016) pp. 828–834. 0031-8248/2016/8305-0016\$10.00
Copyright 2016 by the Philosophy of Science Association. All rights reserved.

two phases were originally described as different genera (e.g., the red alga *Porphyra*).

At its most basic, alternation of diploid and haploid generations involves fusion of haploid gametes to form a diploid zygote and meiosis of diploid cells to produce haploid propagules or gametes. In plants and multicellular algae, the haploid phase is generally called a gametophyte, and the diploid phase the sporophyte. Many variations on this theme exist with and without the growth of multicellular structures in one or both phases. The life cycles of various species include phases that range from complete isomorphism (i.e., haploid and diploid phases morphologically indistinguishable) through various degrees of heteromorphism to the extreme of one phase or the other being reduced to (or remaining) a single cell. Even within a ploidy level, distinct types may co-occur that differ in morphology and in their mode of reproduction. Heteromorphic life history stages may also be ecologically differentiated, occupying different niches and having little or no effect on each other's viability and reproduction. An additional complication is that the haploid and diploid phases do not always strictly alternate. Many life cycles include clonal reproduction of one or both phases (reviewed in Herron et al. 2013).

Alternation of generations and more complex life cycles complicate the question of which biological units should be considered individuals. The concept of biological individuality is intertwined with that of its close conceptual cousin, fitness; in fact, for some authors, defining individuality and defining fitness are the same problem. For example, fitness plays central roles in the individuality concepts of Godfrey-Smith (2009), Clarke (2010), Folse and Roughgarden (2010), Strassmann and Queller (2010), and Metz (2013).

Yet in most eukaryotic life cycles, it is not possible to predict evolutionary outcomes by considering fitness differences in just one phase. Even in so-called 'paradigm' individuals (Wilson 1999), doing so is only an approximation, as selection can take place among gametes. The view that effective selection among gametes does not occur is no longer tenable, as an increasing body of evidence shows that at least some genes are expressed by gametes (Erickson 1990; Hendriksen et al. 1995; Engel, Holmes-Davis, and McCormick 2005; Torgerson and Singh 2006; Borges et al. 2008; Gou et al. 2009). As we move away from the extremely reduced end of the heteromorphy spectrum, this approximation gets worse and worse, until not even the sign of evolutionary change can be reliably predicted by considering only one phase of the life cycle (as I will show).

2. The Model. I start by developing a single-locus model of a simple haploid-diploid life cycle. Diploid sporophytes produce haploid spores by meiosis, which develop into haploid gametophytes. Gametophytes produce gametes by mitosis, which fuse to produce diploid zygotes. Assuming Lotka–Volterra-like dynamics, we can calculate the dynamics of such a population as follows:

$$\begin{aligned}\frac{dH}{dt} &= r_D D \left(1 - \frac{H + \alpha_{D,H} D}{K_H} \right) - \varnothing_H H, \\ \frac{dD}{dt} &= r_H H \left(1 - \frac{D + \alpha_{H,D} H}{K_D} \right) - \varnothing_D D,\end{aligned}$$

where dH/dt and dD/dt describe the dynamics of haploid gametophytes and diploid sporophytes, respectively. Haploids produce diploids at an intrinsic per-capita rate r_H , diploids produce haploids at r_D , and the carrying capacities of haploids and diploids are K_H and K_D , respectively. The relative strength of competitive effects is represented by a matrix of competition coefficients, where $\alpha_{x,y}$ represents the competitive effect (reduction of growth rate) of type y on type x . The terms \varnothing_H and \varnothing_D represent constant per-capita death rates for haploids and diploids.

I introduce evolutionary dynamics by modeling two alleles at a single locus. I assume that genotype does not affect competitive ability or death rate. Gametes combine randomly, so diploids are produced in Hardy–Weinberg frequencies (A^2 , $2Aa$, and a^2):

$$\begin{aligned}\frac{dA}{dt} &= (2r_D AA + (r_D + hs)Aa) \left(1 - \frac{A+a+\alpha_{D,H}(AA+Aa+aa)}{K_H} \right) - \varnothing_H A, \\ \frac{da}{dt} &= (2(r_D + s)aa + (r_D + hs)Aa) \left(1 - \frac{A+a+\alpha_{D,H}(AA+Aa+aa)}{K_H} \right) - \varnothing_H a, \\ \frac{dAA}{dt} &= \frac{r_H^2 A^2}{r_H A + (r_H + u)a} \left(1 - \frac{AA+Aa+aa+\alpha_{H,D}(A+a)}{K_D} \right) - \varnothing_D AA, \\ \frac{dAa}{dt} &= \frac{2(r_H A(r_H + u)a)}{r_H A + (r_H + u)a} \left(1 - \frac{AA+Aa+aa+\alpha_{H,D}(A+a)}{K_D} \right) - \varnothing_D Aa, \\ \frac{daa}{dt} &= \frac{(r_H + u)^2 a^2}{r_H A + (r_H + u)a} \left(1 - \frac{AA+Aa+aa+\alpha_{H,D}(A+a)}{K_D} \right) - \varnothing_D aa.\end{aligned}$$

In this version, dA/dt and da/dt represent the dynamics of the wild-type and mutant haploids, while dAA/dt , dAa/dt , and daa/dt represent the dynamics of wild-type homozygotes, heterozygotes, and mutant homozygotes, respectively. The terms s and u represent the selective effects of the mutation, in terms of production of propagules, on the haploid and diploid phases, respectively. If $u = s$, the mutant has the same selective effect on the haploid (a) and homozygous diploid (aa). If u and s differ, the effect of the mutation differs between ploidy levels. The relationship between u and s reflects the type of pleiotropy: when u and s are both positive (or both negative), the mutation is beneficial (or harmful) regardless of ploidy level (positive pleiotropy); when the signs differ, pleiotropy is negative. The degree of dominance (h) deter-

mines the strength of selection on the heterozygote (Aa) relative to the homozygous mutant (aa).

The competitive coefficients across ploidy levels ($\alpha_{H,AA}$, etc.) roughly reflect the degree of isomorphy: when haploids and diploids are isomorphic and consuming the same resources, these terms will be near unity; for extremely heteromorphic species in which haploids don't compete with diploids, they will be near zero.

In this form, the system of equations approximates the dynamics of a new mutation in a haploid-diploid population and allows us to calculate the invasion fitness of such a mutation. The frequency of the mutant allele in such a population is

$$q = \frac{a + Aa + 2aa}{N},$$

where the total number of alleles in the population is $N = A + a + 2(AA + Aa + aa)$, and the change in frequency of the mutant allele is represented by

$$\frac{dq}{dt} = \frac{\left(\frac{da}{dt} + \frac{dAa}{dt} + 2\frac{d aa}{dt}\right)N - (a + Aa + 2aa)\frac{dN}{dt}}{N^2},$$

$$\frac{dq}{dt} = \frac{\left(\frac{da}{dt} + \frac{dAa}{dt} + 2\frac{d aa}{dt}\right)(A + a + 2AA + 2Aa + 2aa) - (a + Aa + 2aa)\left(\frac{dA}{dt} + \frac{da}{dt} + 2\frac{dAA}{dt} + 2\frac{dAa}{dt} + 2\frac{d aa}{dt}\right)}{(A + a + 2AA + 2Aa + 2aa)^2},$$

which will be positive whenever $\left(\frac{da}{dt} + \frac{dAa}{dt} + 2\frac{d aa}{dt}\right)N > (a + Aa + 2aa)\frac{dN}{dt}$.

3. Discussion. Among eukaryotic lineages, complex life cycles are the rule rather than an exception. Understanding how such lineages respond to selection is thus crucial for understanding adaptive evolution in eukaryotes. Models that consider selection in only one phase of the life cycle at best only approximate this process, and except when the dominance of one phase is extreme, predictions from models that consider only one phase are likely to be substantially wrong.

This model shows that the fitness effect of an allele cannot be accurately calculated without taking its influence on all life cycle phases into account. Furthermore, fitness is a function not only of allele frequency but also of the proportion of alleles represented in each phase of the life cycle. In cases in which the sign of selection differs between haploid and diploid phases, not only the magnitude of selection but also the direction depends on the composition of the population, both allele frequency and the proportion of those alleles in haploid versus diploid phases.

For concepts of individuality that include fitness or adaptation as an important component, the realization that calculating fitness requires consider-

ing the entire life cycle presents a challenging complication. The concept of fitness is useful to the extent that it predicts, at least probabilistically, the direction and magnitude of short-term evolutionary change. However, I have shown that such predictions will often be in error when only one phase of the life cycle is considered. Privileging one or the other phase as ‘the’ individual therefore neglects potentially important selective effects.

Previous authors have recognized that alternation of generations and more complex life cycles present problems for concepts of biological individuality, sometimes arriving at very different solutions. Godfrey-Smith (2009, 2012, 2014) considers individuals to be members of ‘Darwinian populations’, those groups that have the necessary ingredients for adaptive evolutionary change. One factor in the cohesiveness of such populations is the strength of intraspecific ecological interactions, which plays a role nearly identical to the α parameter in my model (and in Lotka–Volterra models of interspecific competition). At the extreme of heteromorphy, $\alpha = 0$, the haploid and diploid phases are ecologically distinct, such that the density of one life history phase has no competitive impact on the growth of the other. Nevertheless, a shared gene pool dictates that they do affect each other’s evolution, as selection in one phase will inevitably affect allele frequencies in the other.

In the view of Griffiths and Gray (1994), the life cycle itself is the individual. While I agree that the entire life cycle needs to be taken into account, it is not clear how to square this developmental view with the evolutionary view of individuals as subjects of selection. How, for example, does selection operate among life cycles, and do life cycles have the requisite conditions of heritable variation in fitness?

Much of the literature on biological individuality has focused on the question of which level (or levels) in the hierarchy of life comprises individuals, for example, cells versus multicellular organisms, ramets versus genets, multicellular organisms versus eusocial ‘superorganisms’. This remains an important question, but my focus here has been quite different, looking instead at a single hierarchical level, that of the multicellular organism (though most of the conclusions would be identical for facultatively sexual unicells).

If we can draw any conclusion from this, it is probably that approaches to biological individuality that privilege one life history stage at the expense of the other(s) are not viable unless we are willing to give up the idea that individuals are the bearers of fitness. First, such an approach may work acceptably for a particular case, but even then it is an approximation; accurately predicting the outcomes of selective processes will require considering fitness differences in both (or all) phases of the life cycle. Second, the choice of which phase to privilege is at least somewhat arbitrary, and the dominant phase will change depending on what taxon we are looking at: in animals and land plants, the diploid; in *Porphyra* and *Volvox*, the haploid. Finally, in cases of isomorphic alternation of generations (such as *Ulva*), haploid and diploid

phases are on equal footing, and there is no basis on which to choose which phase to privilege.

I have taken a gene-centric and adaptationist approach, neglecting the possible effects of stochastic processes such as genetic drift. It seems unlikely, however, that including such effects would change the qualitative outcomes. The model could be extended to capture stochastic effects as well as to consider quantitative traits, pleiotropic effects within a phase, and development. In addition, the model could be extended from the simple alternation of generations I have considered to include complications such as clonal propagation of one or both phases, reproduction of unicellular phases (as in the red alga *Porphyra*; Polne-Fuller and Gibor 1984), additional multicellular phases (such as parthenosporophytes in the green alga *Ulva*; Hoxmark and Nordby 1974; and the brown alga *Ectocarpus*; Charrier et al. 2008), and chromosome doubling during development (as in *Ulva*; Föyn 1958; Fjeld 1972).

REFERENCES

- Borges, F., G. Gomes, R. Gardner, N. Moreno, S. McCormick, J. A. Feijó, and J. D. Becker. 2008. "Comparative Transcriptomics of *Arabidopsis* Sperm Cells." *Plant Physiology* 148:1168–81.
- Charrier, B., S. M. Coelho, A. Le Bail, T. Tonon, G. Michel, P. Potin, B. Kloareg, C. Boyen, A. F. Peters, and J. M. Cock. 2008. "Development and Physiology of the Brown Alga *Ectocarpus siliculosus*: Two Centuries of Research." *New Phytologist* 177:319–32.
- Clarke, E. 2010. "The Problem of Biological Individuality." *Biological Theory* 5:312–25.
- Engel, M. L., R. Holmes-Davis, and S. McCormick. 2005. "Green Sperm. Identification of Male Gamete Promoters in *Arabidopsis*." *Plant Physiology* 138:2124–33.
- Erickson, R. P. 1990. "Post-meiotic Gene Expression." *Trends in Genetics* 6:264–68.
- Fjeld, A. 1972. "Genetic Control of Cellular Differentiation in *Ulva mutabilis*. Gene Effects in Early Development." *Developmental Biology* 28:326–43.
- Folse, H. J., and J. Roughgarden. 2010. "What Is an Individual Organism? A Multilevel Selection Perspective." *Quarterly Review of Biology* 85:447–72.
- Föyn, B. 1958. "Über die Sexualität und den Generationswechsel von *Ulva mutabilis* (N.S.)." *Archiv für Protistenkunde*. 102:473–80.
- Godfrey-Smith, P. 2009. *Darwinian Populations and Natural Selection*. Oxford: Oxford University Press.
- . 2012. "Darwinian Individuals." In *From Groups to Individuals: Perspectives on Biological Associations and Emerging Individuality*, ed. F. Bouchard and P. Huneman, 1–25. Cambridge, MA: MIT Press.
- . 2014. "Individuality and Life Cycles." In *Individuals across the Sciences*, ed. T. Pradeu and A. Guay, 1–20. Oxford: Oxford University Press.
- Gou, X., T. Yuan, X. Wei, and S. D. Russell. 2009. "Gene Expression in the Dimorphic Sperm Cells of *Plumbago zeylanica*: Transcript Profiling, Diversity, and Relationship to Cell Type." *Plant Journal* 60:33–47.
- Griffiths, P. E., and R. D. Gray. 1994. "Developmental Systems and Evolutionary Explanation." *Journal of Philosophy* 91:277–304.
- Hendriksen, P. J., J. W. Hoogerbrugge, A. P. Themmen, M. H. Koken, J. H. Hoeijmakers, B. A. Oostra, T. van der Lende, and J. A. Grootegeod. 1995. "Postmeiotic Transcription of X and Y Chromosomal Genes during Spermatogenesis in the Mouse." *Developmental Biology* 170:730–33.
- Herron, M. D., A. Rashidi, D. E. Shelton, and W. W. Driscoll. 2013. "Cellular Differentiation and Individuality in the 'Minor' Multicellular Taxa." *Biological Reviews of the Cambridge Philosophical Society* 88:844–61.

- Hoxmark, R. C., and Ø. Nordby. 1974. "Haploid Meiosis as a Regular Phenomenon in the Life Cycle of *Ulva mutabilis*." *Hereditas* 76:239–50.
- Metz, J. A. J. 2013. "On the Concept of Individual in Ecology and Evolution." *Journal of Mathematical Biology* 66:635–47.
- Polne-Fuller, M., and A. Gibor. 1984. "Developmental Studies in *Porphyra*. I. Blade Differentiation in *Porphyra perforata* as Expressed by Morphology, Enzymatic Digestion, and Protoplast Regeneration." *Journal of Phycology* 20:609–16.
- Strassmann, J. E., and D. C. Queller. 2010. "The Social Organism: Congresses, Parties, and Committees." *Evolution* 64:605–16.
- Torgerson, D. G., and R. S. Singh. 2006. "Enhanced Adaptive Evolution of Sperm-Expressed Genes on the Mammalian X Chromosome." *Heredity* 96:39–44.
- Wilson, J. 1999. *Biological Individuality: The Identity and Persistence of Living Entities*. Cambridge: Cambridge University Press.