

Resurrecting Biological Essentialism*

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The article defends the doctrine that Linnaean taxa, including species, have essences that are, at least partly, underlying intrinsic, mostly genetic, properties. The consensus among philosophers of biology is that such essentialism is deeply wrong, indeed incompatible with Darwinism. I argue that biological generalizations about the morphology, physiology, and behavior of species require structural explanations that must advert to these essential properties. The objection that, according to current “species concepts,” species are relational is rejected. These concepts are primarily concerned with what it is for a kind to be a species and throw little light on the essentialist issue of what it is for an organism to be a member of a particular kind. Finally, the article argues that this essentialism can accommodate features of Darwinism associated with variation and change.

Essentialism about species is today a dead issue. (Sober [1980] 1992, 249)

Folk essentialism is both false and fundamentally inconsistent with the Darwinian view of species. (Griffiths 2002, 72)

1. Introduction. The idea that biological natural kinds, particularly a spe-

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‡ This article started with “Some Heretical Thoughts on Biological Essentialism,” an eight-page piece I wrote in 2003, on the basis of little reading, and sent to a number of experts for comment. This had two surprising consequences. First, the volume of response was astounding: initial responses together with follow-up discussions amounted to one hundred pages. Second, given the consensus, I expected the experts to identify deep flaws in these “heretical thoughts.” Yet this did not happen. I was corrected, informed, and guided on many matters and yet my basic argument for biological essentialism seemed to me to survive fairly intact. The experts I am indebted to for their heroic attempts to set me straight at that point are Peter Godfrey-Smith, Paul Griffiths, Stephen Schwartz, Stephen Stich, and particularly, Joseph LaPorte,

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cies like dogs, have intrinsic underlying natures is intuitively appealing. It has been shown to be widespread even among children (Keil 1989). It was endorsed by a great philosopher, Aristotle. Under the influence of the logical positivists, Popper (1950), Quine (1960), and others, it fell from philosophical favor in the twentieth century until revived by Saul Kripke (1980), Hilary Putnam (1975), and David Wiggins (1980). Many philosophers probably now take the view for granted. If so, they are right out of touch with biologists and, especially, philosophers of biology. For, the consensus among philosophers of biology, and a wide-spread view among biologists, is that this sort of “Aristotelian essentialism” is deeply wrong, reflecting “typological” thinking instead of the recommended “population” thinking (Sober [1980] 1992, 247–248). This essentialism is thought to arise from a naive and uninformed view of biology, indeed to be incompatible with Darwinism.¹ This view is nicely presented and argued for in a paper by Samir Okasha (2002). I shall take that as my main text. I shall defend intrinsic biological essentialism. I think that the children are right and the philosophers of biology, wrong.²

I start by saying something about essentialism in general and about the essentialism I shall defend in particular. A property *P* is an *essential property* of being an *F* iff anything is an *F* partly in virtue of having *P*. A property *P* is *the essence* of being an *F* iff anything is an *F* in virtue of having *P*. The essence of being *F* is the sum of its essential properties. Essences can be fully intrinsic; for example, the essence of being gold is having atomic number 79. Essences can be partly intrinsic and partly

Karen Neander, and Samir Okasha. The first version of the article was delivered at the University of Queensland in November 2005 and later versions have been delivered at many other universities. The article has benefited greatly from those events and also from the written comments of Matt Barker, Alberto Cordero, Michael Dickson, Marc Ereshefsky, Philip Kitcher, Joseph LaPorte, Mike Levin, Georges Rey, Iakovos Vasilioi, John Wilkins, and Rob Wilson. Finally, my thanks to Macquarie University for the position of Visiting Associate in October and November of 2005, during which the first version of the article was mostly written.

1. Michael Ruse places Kripke, Putnam, and Wiggins “somewhere to the right of Aristotle” and talks of them showing “an almost proud ignorance of the organic world” ([1987] 1992, note 358). John Dupré argues that the views of Putnam and Kripke are fatally divergent from “some actual biological facts and theories” (1981, 66).

2. This article was prompted by writing another one defending the thesis that the notion of rigidity we need for kind terms is one of *rigid application* not one of *rigid designation* (Devitt 2005). The view that natural kind terms are rigid applicators has the metaphysical consequence that a member of a natural kind is essentially a member. This sort of “individual essentialism” needs to be distinguished from the “kind essentialism” that is the concern of the present article.

extrinsic and relational;³ for example, the essence of being a pencil is partly being an instrument for writing, which an object has in virtue of its relation to human intentions, and partly having the sort of physical constitution that distinguishes it from a pen, which an object has intrinsically. Finally, essences can be fully relational and extrinsic; being Australian is probably an example because it seems that anything—Rupert Murdoch, Phar Lap (a horse), the Sydney Opera House, a bottle of Penfolds’ Grange, the expression “no worries mate,” and so on—can have the property provided it stands in the right relation to Australia.⁴

The doctrine I want to defend, which I shall call “Intrinsic Biological Essentialism,” abbreviated sometimes to “Essentialism,” is that Linnaean taxa have essences that are, at least partly, intrinsic underlying properties. This calls for some clarification and comment:

- By “Linnaean taxa” I mean kinds that are thought to fall under the biological categories in the Linnaean hierarchy: kingdoms, phyla, classes, orders, families, genera, species, and even subspecies (varieties).⁵ I do not mean kinds like those of predators or parasites. And I do not mean the categories themselves. Essentialism is a thesis about what it is for an organism to be, say, a dog not a cat, not about what it is for, say, dogs to be a species not a genus. (This distinction will loom large in Sections 5–9.) The focus of my discussion will be on species but, I emphasize, Essentialism covers kinds that fall under all the categories.
- I include the qualification “at least partly” because I shall not take issue with the consensus that a species is partly an historical entity.⁶

3. Biological essentialism is usually taken to be concerned *only* with what is intrinsic (e.g., Mayr [1963] 1992, 16; Sober 1993, 146; Wilson 1999, 188). This reflects the influence of Aristotle. I think it more helpful to define essentialism in a more general way so that issues come down to the *sort* of essence that a kind has.

4. Locke called an underlying intrinsic essence that is causally responsible for the observable properties of its kind a “real essence.” This is contrasted with a “nominal essence” that is picked out by reference-determining descriptions associated with a kind term. So, having atomic number 79 is the real essence of gold and the essence of being Australian, whatever it may be, is merely nominal. Kripke and Putnam showed that natural kind terms like ‘gold’ are not associated with reference-determining descriptions and so do not pick out nominal essences; they pick out real essences without describing them. This is not to say that a term *could not* pick out a nominal essence that is also real; indeed, ‘having atomic number 79’ is such a term (cf. Boyd 1999, 146).

5. I say “thought to fall” because I sympathize with the doubts of some about this hierarchy; see Ereshefsky 1999, 2001; Mishler 1999.

6. However, I say that the essences are “at least, partly” intrinsic rather than simply “partly” because I do wonder whether *all* species are, or should be, partly historical. Citing the possibility of regularly produced hybrids like the lizard *Cnemidophorus*

- In sexual organisms the intrinsic underlying properties in question are to be found among the properties of zygotes; in asexual ones, among those of propagules and the like.⁷ For most organisms the essential intrinsic properties are probably largely, although not entirely, genetic. Sometimes those properties may not be genetic at all but in “the architecture of chromosomes,” “developmental programs,” or whatever (Kitcher [1984] 2003, 123).⁸ For convenience, I shall often write as if the essential intrinsic properties were simply genetic but I emphasize that my Essentialism is not committed to this.
- Intrinsic Biological Essentialism would certainly be opposed by the consensus because of its commitment to *intrinsic* essences. But the consensus should not be opposed to biological essentialism in general because, as I am understanding essentialism, the consensus is that species have essences but these are *extrinsic or relational*. And Kim Sterelny and Paul Griffiths, in their excellent introduction to the philosophy of biology, *Sex and Death*, are explicitly not opposed to this sort of essentialism: “the essential properties that make a particular organism a platypus . . . are historical or relational” (1999, 186). Of course, the very term ‘essentialism’ has become so distasteful to biologists because of its association with Aristotelian metaphysics that a biologist would doubtless be reluctant to admit to any sort of essentialism. But the essentialism I have defined need not come with those Aristotelian trappings. Many philosophers would be similarly reluctant because the term ‘essentialism’ strikes them as quaintly old-fashioned, scholastic, even unscientific. But such reluctance would be a merely verbal matter. The issue of essentialism would remain even if the term ‘essentialism’ were dropped. It is the issue of in virtue of what an organism is a member of a certain Linnaean taxon; the issue of what makes an organism a member of

tesselatus, Philip Kitcher claims that “it is not necessary, and it may not even be true, that all species are historically connected” ([1984] 2003, 117).

7. What I would like is a term for asexual organisms that is like ‘zygote’ for sexual ones in referring to the beginning of an organism. John Wilkins informs me that there is no one term for this. Others he mentions include ‘bud’ and ‘gemmule’. He has also drawn my attention to other uses of ‘propagule’. Thus, consider the following definition: “In animals, the minimum number of individuals of a species capable of colonizing a new area. This may be fertilized eggs, a mated female, a single male and a single female, or a whole group of organisms depending upon the biological and behavioral requirements of the species. In plants, a propagule is whatever structure functions to reproduce the species: a seed, spore, stem or root cutting, etc.” (available at <http://www.radford.edu/~swoodwar/CLASSES/GEOG235/glossary.html>).

8. Webster and Goodwin (1996) promote the idea of “morphogenetic fields.”

that taxon; the issue of the very nature of the taxon. I stick with ‘essentialism’ because it is the term that philosophers of biology use for the doctrine that they want to reject and I want to promote. Those who are offended by the term should replace it with one of the other ways of characterizing the issue.

- There is some controversy over whether species are natural kinds or individuals. Michael Ghiselin ([1974] 1992) and David Hull ([1978] 1992) seem to see individualism as an antidote to essentialism.⁹ But, as Okasha points out, “the issues about essentialism . . . do not depend on which view of the ontological status of species we favour” (2002, 193–194; see also Kitcher [1989] 2003, 137–140).¹⁰ Thus, if a species is an individual rather than a kind, our essentialism issue for species becomes that of saying in virtue of what organisms are *parts of* a certain species, for example, the species *Canis familiaris*. And the consensus answer should be that it is entirely in virtue of the organisms’ historical or relational properties, entirely “because they are part of the genealogical nexus” (Hull [1978] 1992, 309), whereas my Essentialism’s answer would be that it is, at least partly, in virtue of the organisms’ intrinsic underlying properties.¹¹ Indeed the essentialism issue can be posed “nominalistically” in a way that is non-committal on the ontological status of species: In virtue of what is an organism, say, a *Canis familiaris*? And that is how I *do* pose the issue in Section 5. I mean to be neutral on the ontological issue but for convenience will mostly talk of species as if they were kinds.
- Essentialism is primarily concerned with the natures of the *actual* groups identified by the folk and biologists for explanatory purposes. This “descriptive” issue needs to be distinguished from the “normative” issue of the natures of the groups that we *should* identify for explanatory purposes. Clearly, we might not be doing what we should be doing. However, I shall write as if we are. If we are not, then my Essentialism should be taken to cover the groups that we should be identifying for explanatory purposes as well.

In Section 2, I give evidence that the consensus really is opposed to

9. “*Individualism* about species is an idea with close links to antiessentialism, both conceptually and historically” (Griffiths 1999, 211).

10. Richard Boyd goes so far as to say that the distinction between species being individuals or kinds “is almost just one of syntax” (1999, 164).

11. Also, we should note, if *Canis familiaris* is an individual, we can ask about its *individual* essence just as we can about that of any individual (note 2). And the consensus answer should be that its essence is its being constituted by organisms that share historical or relational properties whereas my Essentialism’s answer would be that those organisms must also share certain intrinsic underlying properties.

Intrinsic Biological Essentialism. In Section 3, I argue for the doctrine: explanations in biology demand that there be essential intrinsic underlying properties. I turn then to objections. In Section 4, I describe the standard relational views of species which, according to the consensus, make Essentialism untenable. In Section 5, I emphasize a distinction which is crucial to showing that the consensus is wrong about this. The distinction is between two questions:

1. What is it to be a member of any group that happens to be a species?
2. What is it for a group to be a species?

In Sections 6–9, I argue that the relational views of species are, primarily at least, answers to 2. Essentialism, in contrast, is an answer to 1. Indeed, these relational views can, mostly, be happily wedded to Essentialism. Even the influential phylogenetic-cladistic view can be wedded if it loses some implausible features. How has the consensus got it so wrong? My tentative diagnosis is that the error has arisen from conflating questions 1 and 2, a conflation encouraged by some mistaken thoughts about conspecificity. Finally, in Section 10, I accommodate some general features of Darwinianism, associated with variation and change, features that are thought to undermine Essentialism. The accommodation requires acceptance of some indeterminacy in what constitutes a biological kind. But we must all accept that, whatever our views of Essentialism.

If the arguments in Sections 4–10 are right, the stated objections to an essentialist doctrine like Intrinsic Biological Essentialism fail. Perhaps there are some unstated objections that would succeed. And perhaps these could provide the basis for showing that the arguments in Section 3 in favor of Essentialism are inadequate. Given the strength and longevity of the consensus in biology against intrinsic essentialism, it seems reasonable to predict this. Still, it remains to be seen whether it is so. At the very least I hope to show that the case for the consensus needs to be made a lot better than it has been.

2. Evidence of the Consensus. I have claimed that the consensus among philosophers of biology is that doctrines like Intrinsic Biological Essentialism are wrong. Among those philosophers, the claim hardly needs support because the consensus is so established. Still, among philosophers in general, the claim does need support because, influenced by Kripke and Putnam, many find the claim incredible and so think I must be struggling with a straw man. The epigraphs to this article, drawn from the works of Elliott Sober and Paul Griffiths, two leading philosophers of biology, are some evidence that I am not. Here is some more.

The consensus starts by denying that members of a species share a distinctive set of genetic properties. Thus, according to Okasha, “virtually

all philosophers of biology agree that . . . it simply is not true that the groups of organisms that working biologists treat as conspecific share a set of common morphological, physiological or genetic traits which set them off from other species” (2002, 196).

Clearly, if the members of a species do not share a distinctive set of genetic properties then those properties could not be essential properties of that species. Indeed, Okasha claims that “biologists and philosophers of biology typically regard essentialism about species as incompatible with modern Darwinian theory” (2002, 191). And John Dupré claims that “it is widely recognized that Darwin’s theory of evolution rendered untenable the classical essentialist conception of species” (1999, 3). Alex Rosenberg says: “The proponents of contemporary species definitions are all agreed that species have no essence” (1985, 203). Mohan Matthen claims that “species . . . are associated with no nonrelational real essence” (1998, 115). Sober expresses this consensus as follows: “biologists do not think that species are defined in terms of phenotypic or genetic similarities”; tigers are “not *defined* by a set of traits” (1993, 148). Sterelny and Griffiths put the point bluntly: “no intrinsic genotypic or phenotypic property is essential to being a member of a species” (1999, 186). Ghiselin puts it even more bluntly: “That John Doe has a particular set of genes is about as relevant to his being a specimen of *Homo sapiens* L. as it is to his working for the manufacturers of Brand X” ([1974] 1992, 283).

Finally, if the essence of a species is not in the least intrinsic then it must be entirely relational. I have already quoted Sterelny and Griffiths’s claim to this effect about the platypus. And they think that nearly everyone agrees with them: there is “close to a consensus in thinking that species are identified by their histories” (1999, 8). Their view is endorsed by Okasha (2002, 202). Sober declares that tigers are “*historical entities*” (1993, 148, his emphasis). “Two organisms are conspecific in virtue of their historical connection to each other, not in virtue of their similarity” (1993, 150). Marc Ereshefsky makes similar comments, speaking for “Darwin, the founders of the Modern Synthesis, and most cladists” (2001, 209). Finally, Hull claims: “If species are interpreted as historical entities, then particular organisms belong in a particular species because they are part of that genealogical nexus, not because they possess any essential traits. No species has an essence in this sense” ([1978] 1992, 313). Ruth Millikan says much the same (2000, 19).

The consensus is broad but some are not part of it. Thus David B. Kitts and David J. Kitts (1979) urge an intrinsic essentialism like mine. According to Richard Boyd (1999) and Robert Wilson (1999), species are “homeostatic cluster kinds” and I take it that they think that they have

at least partly intrinsic essences.¹² And Philip Kitcher has this to say: “I want to remain agnostic on the issue of whether any species taxon has a nontrivial essence” ([1984] 2003, 132, note 16).

I take the opposition to Intrinsic Biological Essentialism to be established. It is now time to argue for the doctrine.

3. An Argument for Intrinsic Biological Essentialism. I shall offer two reasons for believing Essentialism. The first is superficial but still, it seems to me, indicative of where the truth lies. Such essential properties seem to be part of what “genome projects” are discovering. The projects seem to be throwing light on the very nature of certain species. Thus the *New York Times* recently reported that researchers hope “to discover, from a three-way comparison of chimp, human and Neanderthal DNA, which genes have made humans human” (Wade 2006).¹³ Philosophers of biology disparage this common view (Sterelny and Griffiths 1999, 7; Okasha 2002, 197), but the view is certainly appealing.

The second reason is deep and shows why the view is appealing. We group organisms together under what seem, at least, to be the names of species or other taxa and make generalizations about the morphology, physiology, and behavior of the members of these groups: about what they look like, about what they eat, about where they live, about what they prey on and are prey to, about their signals, about their mating habits, and so on. These generalizations are the stuff of popular nature programs and are to be found throughout the writings of biologists and philosophers of biology. For example, we are told that ivy plants grow toward the sunlight (Sober 1993, 6); that polar bears have white fur; that Indian rhinoceri have one horn and Africa rhinoceri have two (1993, 21); that Hawaiian *Drosophila* “routinely form interspecific hybrids in the wild” (1993, 156); that the Australasian bittern is superbly camouflaged (Sterelny and Griffiths 1999, 32); that “Major Mitchell cockatoos occasionally hybridize with galahs” (1999, 189); that “Australian trees . . . are not just drought-proof; they are fireproof as well” (1999, 203); that “magnetotactic bacteria . . . come equipped with little compasses called

12. Hilary Kornblith favors the view that species are homeostatic cluster kinds, notes that the members of the cluster need not be intrinsic, but does not take a stand on whether any of them are (1993, 111, note 10). Griffiths writes approvingly of the homeostatic cluster view but argues that species have purely historical essences (1999, 217–222).

13. Consider also this recent news report in the *Scientific American* online: “‘DNA barcodes are giving us a direct signal of where species boundaries lie,’ says Paul Herbert, an evolutionary biologist at the University of Guelph in Ontario and a progenitor of the genetic bar code effort” (Biello 2007).

magnetosomes, which they use to navigate away from oxygen-rich surface water because oxygen is toxic to them” (1999, 209).

Generalizations of this kind demand an explanation. *Why are they so?* Why, for example, is there this difference between the Indian and African rhinos? Such questions could, of course, be seeking an explanation of the evolutionary history that *led to* the generalization being true. Set that aside for a moment. The questions could also be seeking an explanation of *what makes* the generalization true. Regardless of the history of its coming to be true, in virtue of what is it now true? What are the mechanisms? The truth of these generalizations cannot be brute facts about the world and so must be explained. Explanations will make some appeal to the environment,¹⁴ but they cannot appeal only to that. There has to be something about the very nature of the group—a group that appears to be a species or taxon of some other sort—that, given its environment, determines the truth of the generalization. That something is an intrinsic underlying, probably largely genetic, property that is part of the essence of the group. Indeed, what else could it be?¹⁵ Some intrinsic underlying property of each Indian rhino causes it, in its environment, to grow just one horn. A different such property of each African rhino causes it, in its environment, to grow two horns. The intrinsic difference explains the physiological difference. If we put together each intrinsic underlying property that similarly explains a similar generalization about a species, then we have the intrinsic part of its essence.¹⁶

The generalizations we have been discussing reflect the fact that it is *informative* to know that an organism is a member of a certain species or other taxon: these classifications are “information stores” (Sterelny and Griffiths 1999, 195). But being a member of a certain taxon is more than informative, it is *explanatory*. Matthen points out that “many biologists seem committed to the idea that something is striped *because* it is a tiger” (1998, 115). And so they should be: the fact that an individual organism is a tiger, an Indian rhino, an ivy plant, or whatever, explains a whole lot about its morphology, physiology, and behavior. At first sight, the explanation of the animal’s stripes may seem rather superficial, but it is not really. For, when biologists group organisms together under some name on the basis of observed similarities, they do so partly *on the assumption that those similarities are to be explained by some intrinsic underlying nature*

14. The role of the environment is very obvious with plants. Thus the height of corn in a field depends on the temperature, the soil, and so on.

15. The point is not, of course, that the explanation of any generalization, even any biological one, demands an intrinsic property, just that the explanation of a generalization of the kind illustrated demands one.

16. So this intrinsic part is a *real* essence, in Lockean terms; see note 4.

of the group. It seems to me clear that this is their practice, whatever they say about essentialism.¹⁷ So the apparently superficial explanation points to the deep fact that there is something intrinsic, probably unknown, partly in virtue of which the animal is a tiger and which causes it to be striped. That something is an essential intrinsic property. The sum of those properties, together perhaps with some historical ones, constitute the essence of a tiger. Sober rightly insists that the essence of a species must explain why its members are the way they are. It must be “a causal mechanism that acts on each member of the species, making it the kind of thing that it is” ([1980] 1992, 250). That is exactly what this (partly) intrinsic essence is.

I distinguished two sorts of explanation that might be sought in asking why members of a species have a certain property. In so doing I am following in the footsteps of Ernst Mayr (1961). He regards an explanation of the mechanisms within members of a species that make a generalization true (regardless of the history) as concerned with “proximate” causation and part of “functional biology.” In contrast an explanation of the evolutionary history that led to the mechanism being present in the members of a species is concerned with “ultimate” causation and is part of “evolutionary biology.” The use of “ultimate” to describe the latter explanation seems like a gratuitous put-down of the former. Kitcher’s even-handed description in response to Mayr’s is *prima facie* more appropriate: “there are indeed two kinds of biological investigation that can be carried out relatively independently of one another, neither of which has priority over the other” ([1984] 2003, 121). Adopting Kitcher’s terminology, I shall call the former sort of explanation “structural” and the latter “historical.” Structural explanations, as he says, seek to “explain the properties of organisms by means of underlying structures and mechanisms.” He gives a nice example ([1984] 2003, 121):

A biologist may be concerned to understand how, in a particular group of bivalve mollusks, the hinge always comes to a particular form. The explanation that is sought will describe the developmental process of hinge formation, tracing the final morphology to a sequence of tissue or cellular interactions, perhaps even identifying the stages in ontogeny at which different genes are expressed.

17. The following comment of Sterelny is interesting in this respect: “Some, perhaps most, evolutionary biologists take speciation to occur only when there have been intrinsic changes.” He finds this “puzzling for the view that species are historically defined entities is close to the consensus view in evolutionary biology.” He is inclined to blame the influence of the folk who, as we noted (Section 1), tend to be intrinsic essentialists (1999, 130). I think that the biologists and the folk are, deep down, tuned into the demands of explanation.

He goes on to claim that “explanations of this type abound in biology: think of the mechanical accounts of normal (and abnormal) meiosis, of respiration and digestion, of details of physiological functioning in all kinds of plants and animals” ([1984] 2003, 121). And my point is that, at bottom, structural explanations will advert to essential intrinsic, probably largely genetic, properties.¹⁸ It is because the bivalve mollusks have a certain intrinsic underlying nature that the hinge takes that form. That is the deep explanation.¹⁹

This discussion generates a number of questions. Here are two:

- “Surely any of the generalizations we have been discussing could have exceptions: a small mutation may lead to an organism that seems to be a member of a species and yet lacks the property attributed to the species by a generalization. So the generalizations do not seem to be lawlike. How does Intrinsic Biological Essentialism deal with that?”²⁰
- “It is of course the case that the truth of any such generalization must be explained by an intrinsic, probably largely genetic, property, but why does that property have to be an *essential* property of the kind in question?”²¹

I shall consider these questions in Section 10, along with others arising from Darwinian views of variation and change. I shall argue that Essentialism has an adequate answer to all these questions.

Sober claims that “evolutionary theory has removed *the need* for providing species with constituent definitions” and hence with intrinsic essences ([1980] 1992, 255). I suspect that this sort of focus on evolution, hence on historical rather than structural explanations, has misled biologists and philosophers of biology about essentialism.²² This having been said, I suspect that even historical explanations demand a partly intrinsic essence; that, for example, the explanation of how polar bears came to

18. Griffiths thinks not but his argument conflates structural explanations with historical explanations (1999, 210–211 and 219–221).

19. Hull unfavorably contrasts “classificationists” seeking “the unit of identification” with “phylogeneticists” seeking “the unit of evolution” ([1965] 1992, 204). I think that the classificationists should be seen as seeking units of *structural explanation*, a very worthwhile pursuit.

20. “Philosophers of biology have often noted that there seem to be no laws which apply to all and only members of a species taxon (Hull [1978] 1992; Rosenberg 1985)” (Okasha 2002, 209).

21. I am indebted to Peter Godfrey-Smith for raising this question.

22. See, e.g., Matthen (1998, 117–121), Griffiths (1999, 219–222), and Millikan (2000, 18–20).

be white will ultimately depend on essential intrinsic properties of polar bears and of their grizzly ancestors. But I shall not attempt to argue this.

I have presented a positive argument for Intrinsic Biological Essentialism. We might sum it up: *structural explanations in biology demand that kinds have essential intrinsic properties*. That is my *first* main point in defense of Essentialism. In the rest of the article I will develop the case for Essentialism in the course of responding to objections.

4. Relational Species Concepts. I start with what is alleged to be the central objection to Intrinsic Biological Essentialism: according to nearly all current “species concepts”—theories about the nature of species—species are *relational*. Okasha expresses this consensus as follows: “On all modern species concepts (except the phenetic), the property in virtue of which a particular organism belongs to one species rather than another is a relational rather than an intrinsic property of the organism” (2002, 201).

Despite the consensus that these species concepts make Essentialism untenable, the nature of biological species is, and always has been, an extremely controversial issue: “The species problem is one of the oldest controversies in natural history” (O’Hara 1993, 231); it is “one of the thorniest issues in theoretical biology” (Kitcher 2003, xii).²³ There are around two dozen species concepts and “at least seven well-accepted ones” (Ereshefsky 1998, 103). I shall follow Okasha in placing them in “four broad categories.” In this section I shall briefly describe these concepts. In the next, I shall draw a distinction which is crucial to showing, in Sections 6–9, that the consensus is wrong.

- *Phenetic* concepts. On this sort of view, organisms are grouped into species on the basis of overall similarity of phenotypic traits. This is thought by its proponents to have the advantage of being fully “operational.” Okasha says that phenetic concepts are “the least popular” (2002, 199) and this is hardly surprising because they arise from the “philosophical attitude . . . of empiricism” (Sokal and Crovello [1970] 1992, 29). “Phenetic taxonomists have often wanted to segregate taxonomy from theory” (Sterelny and Griffiths 1999, 196).²⁴ This category of species concept is the only one of the four that is *not* in the least historical and relational. I shall set it aside.

23. Although, interestingly enough, an issue that Darwin himself was skeptical about: he talks of “the vain search for the undiscovered and undiscoverable essence of the term species” ([1859] 2004, 381).

24. Sterelny and Griffith include under phenetic concepts those that define species in terms of *genetic* similarity (1999, 184). I clearly do not include these. I take the phenotype of an organism to be observable properties of it distinct from, but caused by, its genotype (along with the environment).

- *Biological Species* concepts (BSC). The most famous example of BSC is due to Mayr. He defined species as “groups of interbreeding natural populations that are reproductively isolated from other such groups” (Mayr 1969, 26). Sterelny and Griffiths remark that “If the received view has a received species concept” it is BSC (1999, 188).²⁵
- *Ecological Niche* concepts (ENC). According to ENC, a species occupies a certain ecological niche. “A species is a lineage . . . which occupies an adaptive zone minimally different from that of any other lineage in its range and evolves separately from all lineages outside its range” (van Valen [1976] 1992, 70). Okasha puts the view succinctly: species “exploit the same set of environmental resources and habitats” (2002, 200).²⁶
- *Phylogenetic-Cladistic* concepts (P-CC). On this view we “identify species in terms of evolutionary history . . . [with] particular chunks of the genealogical nexus. . . . Species come into existence when an existing lineage splits into two . . . and go extinct when the lineage divides, or when all members of the species die” (Okasha 2002, 200). Sterelny and Griffiths claim that “something like a consensus has emerged in favor of a *cladistic* conception of systematics” (1999, 194). Nonetheless, it has some surprising features, as we shall see (Section 9).

But perhaps the most important feature of the P-CC concept for the purposes of this article is that it is, as everyone agrees, *incomplete*. It needs to be supplemented by a theory of speciation, a theory that explains when a lineage *has* split in two. For this, as Okasha says, P-CC “will have to rely on a concept of one of the other types” (2002, 201).

5. A Crucial Distinction. It is alleged that, according to each of these species concepts, except the phenetic which we are setting aside, species are relational. These concepts are thought, therefore, to show that the nature of a species could not be partly intrinsic and hence that doctrines like Intrinsic Biological Essentialism are false. In assessing this thought it is very important to distinguish the question that Essentialism is supposed to answer from another which it isn't.

Let *F*'s be some group that has been classified for biological purposes under one of the taxa; for example, a group of poodles, dogs or *Canis*.

25. Popular as it is, BSC has been the subject of extensive criticism; see, e.g., Sokal and Crovello 1970; van Valen [1976] 1992; Cracraft [1983] 1992; Sober 1993, 155–156; Kitcher [1984] 2003, 118–120; [1989] 2003, 141–145; Mallet 1995; Dupré 1999; Sterelny and Griffiths 1999, 186–190. Mallet claims provocatively that the BSC concept “owes nothing either to genetics or to Darwinism” (1995, 295).

26. Ghiselin ([1987] 1992, 374–378) has some severe criticisms of ENC.

The question that Essentialism answers has many forms, as already indicated:

1. In virtue of what is an organism an *F*?
 - What makes an organism an *F*?
 - What is the nature of being *F*?
 - What is the essence of being *F*?

This is a question about the *properties of organisms*. When it concerns *F*'s that form a species, Mayr calls it the species "taxon" problem (1982, 253–254). So let us generalize this as "the taxon problem." It needs to be distinguished from the very different, "higher level," problem about the *properties of those properties*:

2. In virtue of what are *F*'s a subspecies, a species, a genus, etc.?
 - What makes a group of *F*'s a subspecies, a species, a genus, etc.?
 - What is the nature of being a subspecies, a species, a genus, etc.?
 - What is the essence of being a subspecies, species, genus, etc.?

When it concerns species, Mayr calls it the species "category" problem (1982, 253–254). So let us generalize this as "the category problem."

"What is it to be a poodle not a bulldog?" is an instance of the taxon problem 1, "What is it for poodles to be a subspecies not a species?" is an instance of the category problem 2. The distinction between the two problems may seem obvious and yet it is easily conflated by certain forms of words. In particular, consider the question, "What is a species?" or "What is the nature/definition of a species?" These questions are ambiguous. They could be asking what sort of a nature any group has that happens to be a species, an instance of the taxon problem 1. But they are more likely to be asking what is it for any group to be a species, an instance of the category problem 2.

The distinction between the two problems is absolutely crucial to this article. Which problem are the species concepts supposed to answer? According to Okasha, speaking for the consensus, they are at least supposed to answer problem 1 for species: they are supposed to show, as we noted, that "the property in virtue of which a particular organism belongs to one species rather than another is a relational rather than an intrinsic property of the organism" (2002, 201). Yet Ereshefsky, in his introduction to a collection that includes many classic papers on species concepts, says that "our concern is with a definition of the species category" rather than of the species taxon (1992a, xiv; see also Kitcher [1984], 120). And according to Sterelny and Griffiths (1999, 211) and Wilson (1999, 191–192), the species concepts provide answers to *both* taxon and category problems.

I shall argue that, on the contrary, the species concepts are primarily concerned with 2 and throw little light on 1.²⁷ Yet Intrinsic Biological Essentialism is concerned with 1. So, the central objection to Essentialism, based on the species concepts, fails. This is my *second* main point in defense of Essentialism.

In Section 6, I shall show how much the species concepts bear on the category problem 2. In Sections 7 and 9, I shall show how little they bear on the taxon problem 1.

6. Species Concepts and the Category Problem 2. The species concepts straightforwardly answer problem 2 for species: they tell us about “the species category” (Sterelny and Griffiths 1999, 184).²⁸ Thus the popular BSC tells us that a group is a species in virtue of being an interbreeding natural population that is reproductively isolated from other such groups. And the ENC tells us that a group is a species in virtue of being a lineage which occupies an adaptive zone minimally different from that of any other lineage in its range and evolving separately from all lineages outside its range. And both these answers do indeed entail that being a species is *relational*: a group is a species in virtue of its breeding or niche relations to other groups.

The story for the influential P-CC is a bit more complicated. As we have noted, the P-CC account of species adverts to the splitting of a lineage and so needs to be supplemented with a theory of splitting, a theory of speciation. Thus, according to P-CC, a group of organisms constitute one species at time t_1 and their descendents constitute two daughter species at t_2 . But what makes it the case that the descendents are members of the daughter species rather than the original species? An account of speciation will tell us.

When faced with the need to supplement P-CC it is customary to wave a hand toward other species concepts to provide the needed theory of speciation. “The biological species concept, perhaps supplemented by the ecological species concept or by something else, reemerges as an account of speciation” (Sterelny and Griffiths 1999, 192). Thus, if P-CC is supplemented by BSC it will take a lineage to split when it yields two groups each of which is interbreeding but reproductively isolated from the other.

27. Note that this is *not* the general claim that answers to 2 will throw little light on 1; indeed, see note 29. It is a claim that *the relational species concepts* throw little light on 1.

28. We are setting aside the phenetic concept but it is interesting to note that it is not obvious that this concept *does* answer 2. After all, subspecies like poodles and genera like *canis* could equally be identified by an overall similarity of phenotypic traits. What sort of similarity marks out species in particular?

And if it is supplemented by ENC, it will take a lineage to split when it yields two groups exploiting different sets of environmental resources and habitats. With some such supplement in mind, P-CC, just like the other species concepts, straightforwardly yields an answer to problem 2 for species. Thus, return to Okasha's statement of P-CC: "species come into existence when an existing lineage splits into two . . . and go extinct when the lineage divides, or when all members of the species die" (Okasha 2002, 200). Supplemented by a theory of speciation, this tells us what it is for a group of organisms to be a species rather than, say, a subspecies or a genus. And it tells us that this is a relational matter.

Not only do the species concepts straightforwardly yield answers to problem 2 for species, that seems to be what they are intended to do. Ever since Darwin, the species concepts have been tied closely to views of speciation and to distinguishing when two groups constitute subspecies of the one species and when they constitute two distinct species of a genus. They are concerned, for example, with whether the British red grouse (*Lagopus lagopus scoticus*) and its continental relative (*L. l. lagopus*) are separate species; and with whether the divergent forms of the bluegill sunfish (*Lepomis macrochirus*) constitute a single species. They are concerned with *what distinguishes species from other taxa*. And, we should note, they do not seem to be concerned with the taxon problem 1; with, for example, the nature of the British red grouse or the bluegill sunfish.²⁹

Consider also the earlier-mentioned controversy over species (Section 4). The arguments here are all over problem 2. Thus, alluding to the controversy, Ereshefsky says: "Biologists differ widely on how to define the species category . . . [on how] to provide the essential property of the species category—a property found in all and only species taxa" (1992a, xiv–xv). The controversy has led some (Mishler and Donoghue [1982] 1992; Kitcher [1984] 2003; Sterelny and Griffiths 1999, 194–201) to the view that "there is no unique factor common to all species" (Ereshefsky 1992a, xv). This pluralism, as Robert Wilson notes, is "about the species category" (1999, 192). Biologists are concerned with whether groups that we have picked out for biological purposes should be counted as a subspecies, a species, a genus, or whatever. And with whether various

29. I take these examples from Mallet 1995. This article starts with a nice discussion of the history of species concepts that makes their concern with 2 very apparent. Mallet himself urges a genotypic cluster definition of species as an answer to 2: "we see two species rather than one if there are two identifiable genotypic clusters. These clusters are recognized by a deficit of intermediates" (1995, 296). This view puts him right outside what the philosophers of biology consider the consensus. Yet, he claims, "many, perhaps most, systematists are currently using the genotypic (or morphological) cluster definition" (1995, 298). Whether or not Mallet's answer to 2 is correct, it implies an answer to 1 that clearly is as congenial as could be to Essentialism.

considerations do, and should, play a role in settling such issues. In response to these issues, and inspired by cartographic generalization, Robert O'Hara urges that we take up the perspective of "systematic generalization" and then "we will be better able, not to solve the species problem, but rather to get over it" (1993, 232) It is quite clear that what he thinks we will be able to get over is fussing about when to judge a group to be a species. And, once again, the issues are not over problem 1, not over what is it for an organism to be a member of a group that we have picked out for biological purposes, irrespective of whether that group is a subspecies, species, genus, or whatever. So the issues are largely orthogonal to Essentialism. Essentialism is concerned with the nature of a group whatever the category it falls under.

I turn now to consider the bearing of species concepts on the taxon problem 1. In Section 7, I will argue that BSC and ENC do not bear on this problem and so the consensus is wrong. In Section 8, I will try to diagnose the error. In Section 9, I will consider the bearing of P-CC.

7. BSC, ENC, and the Taxon Problem 1. One reason for thinking that species concepts are not intended to answer the taxon problem 1 for species is that they quite obviously have nothing to say in answer to this problem for taxa other than species.³⁰ Yet if they were answering 1 for species we would expect them to be like Essentialism in saying *something*, at least, in answer to 1 for the other taxa, in saying *something* about what it is to be a member of a particular genus, for example.

Despite the consensus, neither BSC nor ENC provides an answer to 1. Let *F*'s be a group of organisms that is a species according to BSC or ENC; for example, the group of tigers. What do BSC or ENC, as they stand, tell us about how to complete 'Some organism is an *F* in virtue of . . .'? Since *F*'s are a species, BSC or ENC obviously tell us that whatever the completion it must specify *some property or other* of an organism that is at least *compatible* with the organism being a member of *some group or other* that, briefly, interbreeds or occupies a niche. But, beyond that, *they tell us nothing at all!* They don't tell us what property makes an organism a member of the group of *F*'s *in particular*. Indeed, they don't even tell us *what sort* of property that must be. As we have just noted, BSC and ENC tell us what it is for *F*'s to be a species rather than, say, a subspecies or genus, but they are silent on what it is for *an organism to be an F*, say, a tiger rather than a lion. Analogously, an account of what it is for a group of objects to be tools rather than, say, pets or toys would not tell us what it is for an object to be a hammer rather than,

30. But the phenetic concept does answer the species taxon problem and could answer the taxon problem for other taxa; cf. note 28.

say, a saw. In brief, BSC and ENC, as they stand, say nothing about what *identifies* a particular species, hence nothing about what constitutes its essence. *The consensus view expressed by Okasha is simply false*: it is not the case that, according to BSC and ENC, “the property in virtue of which a particular organism belongs to one species rather than another is a relational rather than an intrinsic property of the organism” (2002, 201). These concepts, as they stand, say nothing about this matter and so need not be at odds with any doctrine that does. Indeed, they are not at odds with Intrinsic Biological Essentialism.

Essentialism’s answer to taxon problem 1 is that the group *F* is identified (partly at least) by certain intrinsic underlying properties: it is in virtue of having such properties that an organism is a tiger rather than a lion. This is compatible with the BSC view that what makes *F*’s a species rather than a subspecies or a genus is a matter of interbreeding, and with the ENC view that it is a matter of occupying a niche. Indeed, it is better than compatible: Essentialism *partly explains* why the members of a species have the characteristics which, according to BSC or ENC, make them a species: it is partly *because* those members have a certain essential intrinsic properties that, in the given environment, they interbreed and occupy a niche. Far from being undermined by these species concepts, Essentialism is complementary to them.

BSC and ENC do not *entail* a relational answer to 1 despite the consensus that they do. But perhaps we can see them as *associated with* such an answer. We can indeed find signs of two such answers in the literature. But neither could be a serious rival to Essentialism’s answer.

The first answer is the simple idea that what makes something an *F* is that its parents are *F*’s. This seems to be suggested by the following: “the reference of an individual to a species is determined by its parentage, not by any morphological attribute” (Griffiths 1974, 102). Consider also a statement by Ruse: “if we suppose that humans first appeared about a half million years ago, *Homo sapiens* is the name for the group that descended from the original organisms” (1987, 344).

Hull quotes the Griffiths passage approvingly ([1978] 1992, 305). He goes on to consider whether a human-like organism made by a scientist would *be* a human and this leads him to modify the simple idea: what makes something human is “being born of human beings *and/or mating with human beings*” ([1978] 1992, 306, emphasis added). A difficulty with the simple idea is that it rules out speciation: all organisms will be conspecific with their ancestors, however distant. Set that aside until later (Section 10). The idea is open to an obvious objection: it is not really an answer to 1. It tells us that an organism is an *F* if its parents are *F*’s. But what is it for *them* to be *F*’s? The idea does not solve our problem, it simply moves it back a generation. And Hull’s modification of the simple

idea is open to a similar objection. It tells us that an organism is an *F* if it mates with *F*'s. But what makes the organisms it mates with *F*'s?

The second relational answer might be considered an elaboration of the first. It gets its inspiration from the typical naming practices of biologists: "Biologists coin new species terms by providing a *sample*, called a 'type specimen'" (LaPorte 2004, 5). Could we then identify a species by referring to its type specimen? So, what makes this interbreeding or niche-occupying group *F*'s is that it contains a certain type specimen. This idea seems to be suggested by some other remarks of Hull: "The taxonomist . . . selects a specimen, any specimen, and names it. . . . A taxon has the name it has *in virtue of* the naming ceremony, not in virtue of any trait or traits it might have" ([1978] 1992, 308). He goes on to write that "any organism related to [the type specimen] in the appropriate ways belongs to its species, regardless of how aberrant the type specimen might turn out to be or how dissimilar other organisms may be" ([1978] 1992, 311–312). And consider this (entertaining) proposal for using BSC for taxonomic definitions: "Specify some individual, say Brigham Young, as your reference point, and then members of the same taxon are potential and actual interbreeders" (Ruse [1987] 1992, 344).³¹

Combining this idea with BSC or ENC seems to suggest that what makes an organism a lion is that it is part of an interbreeding or niche-occupying group that contains a certain historically identified type specimen, say Leo; and what makes this other organism a tiger is that it is part of another interbreeding or niche-occupying group that contains a certain other historically identified type specimen, say Benji.

This answer is transparently inadequate. Relating an organism to a type specimen may sometimes be a convenient way to *tell* what species the organism belongs to—for example, if the type specimen is held in some museum—but it should not be taken seriously as an account of what *constitutes* being a member of a species. The answer may be epistemically useful but it is *metaphysically hopeless*.³² Why? Briefly, because being an *F*—for example, being a lion or being a tiger—is an *explanatory* property, as we noted in discussing the generalizations in Section 3.

The hopeless answer itself immediately generates the demand for an explanation that it cannot possibly fulfill: *Why can lions interbreed with Leo and not Benji?* It is clearly no help to be told that that is what it is to be a lion. And this failure is just the tip of the iceberg. Consider the

31. Similarly, Matthen takes an organism to be a member of a particular species in virtue of belonging to the same extended reproductive community "as the originally ostended individual" (1998, 120).

32. And for that reason it is charitable to construe any remark that seems to suggest this answer as making only an epistemic point.

following questions (construed structurally not historically): Why do tigers have stripes? It is no help to be told that it is because they can interbreed with Benji. That does not tell us why any tiger, including Benji, has stripes. Why do polar bears have poor eyesight? Once again, their relation to some Ur-bear gives no explanation just as the relation of pieces of gold to the stuff in Fort Knox gives no explanation of why they are malleable. And so on through indefinitely many structural questions about the morphology, physiology, and behavior of species.

All these questions concern facts about species that could not be brute: the facts have to be explained. The suggested answers to 1 cannot provide adequate explanations. The moral of this discussion is that any adequate explanation cannot appeal only to relational properties of members of the species in question because those relations cannot bear the explanatory burden. An adequate explanation must appeal to intrinsic properties of the organisms. It is something about the intrinsic natures of lions, tigers, polar bears, and so on that provides the explanation (along with some environmental factors).

It is worth noting that the main point of the argument is not restricted to biology. Suppose we ask: Why do paperweights make good weapons? The answer is not that they are the same tool as a certain specimen paperweight kept in some museum. The answer is that the nature of paperweights makes them suitably heavy and easily grasped. And if they had a different nature, say that of buttons, then they would not make good weapons.³³

So, contrary to consensus opinion, BSC and ENC do not give relational accounts of species identity. Indeed they do not give *any* account of species identity. Furthermore, they cannot be happily wedded to a relational account because such accounts are explanatorily hopeless. They can, however, be happily wedded to Intrinsic Biological Essentialism because intrinsic underlying properties can bear the explanatory burden.

We are left with a puzzle. BSC and ENC do not give a relational answer to the taxon problem 1 for species and yet the consensus is that they do. What has gone wrong? How can we diagnose the error?

8. The Conspicuity Route to Error about the Taxon Problem 1. The obvious answer to the diagnostic question is that the error has come from somehow conflating the problems 1 and 2 that Mayr distinguished.³⁴ BSC

33. Despite this, I am not making any claim about explanation in general, just one about what is required for explanations of these phenomena in biology and similar ones elsewhere.

34. Mayr's distinction is established but it is often overlooked; see Dupré 1981; Stanford 1995 (on which see Devitt 2008); Griffiths 1999; Sterelny 1999.

and ENC do indeed offer relational accounts of what it is to be a species. But that is a very different matter from offering a relational account of what it is to be a member of a group that is a species.

Sterelny and Griffiths are interesting on this score. They nicely distinguish the taxon problem 1 from the category problem 2 in the “Further Reading” that concludes a chapter discussing the species concepts (1999, 211). Yet, surprisingly, the preceding discussion itself does *not* distinguish the problems (1999, 184–194). Indeed, they themselves draw attention to this conflation! Their defense is that “an answer to the taxon problem should solve the category problem, and vice versa” (1999, 211).

This raises our puzzle in an acute form. *Why* would anyone think that an answer to the one problem would answer the other? The discussions in Sterelny and Griffiths, and in Okasha (2002), suggest that the route from a category answer to a taxon answer may be via an answer to the *consppecificity* problem, the problem of saying in virtue of what organisms are in the *same* species. This route is most explicit in Wilson’s discussion (1999). Wilson takes what are, in effect, the BSC and P-CC answers to the species category problem to imply answers to the taxon problem: they “imply that the properties determining species membership for a given organism are not intrinsic properties of the organism at all, but depend on the relations the organism bears to other organisms” (1999, 192). How so? Because they imply that “consppecificity is not determined by shared intrinsic properties, but by organisms’ standing in certain relations to one another” (1999, 193). So the idea is that (a) BSC and P-CC imply a relational nonintrinsic answer to the conspecificity problem, and (b) this implies a relational nonintrinsic answer to the taxon problem. And the problem with the idea is that, although (a) is clearly tempting, it is false. However, (b) is true. I shall start with it.

The first thing to note in considering (b) is that an answer to the conspecificity problem does not alone provide an answer to the taxon problem. A conspecificity answer tells us what it is for two organisms to be members of the same species and hence what makes Leo not conspecific with Benji. A taxon answer tells us what it is to be a member of a particular species and hence what makes Leo a lion and Benji a tiger. These are two very different matters. An answer to the conspecificity problem does not answer the taxon problem because it does not determine *which* species conspecific organisms are members of; it does not determine species *identity*; it does not tell us that these conspecifics are lions, those, tigers. So, even if a species concept did answer the conspecificity problem, as (a) claims it does, more would still need to be done to answer the taxon problem. Still, the two problems are related in a way that sustains (b).

Suppose that the answer to the taxon problem is that an organism is a member of species *F* in virtue of being *Q*; and the answer to the con-

specificity problem is that two organisms are conspecific in virtue of being *R*-related. Now, necessarily, if two organisms are both *Q* then they are both *F*'s and hence conspecific. So the fact that they are both *Q* must *determine*, in a very strong way, that they are *R*-related and hence conspecific: *R* must *be* the relation of sharing a property of the *Q* sort, whatever that sort may be. Now suppose, as Intrinsic Biological Essentialism does, that *Q* is a partly intrinsic property of *F*'s then *R* must be partly the relation of sharing that sort of intrinsic property. So if a species concept did imply a nonintrinsic answer to the conspecificity problem, as (a) claims it does, that would indeed count against Essentialism, as (b) claims.

Turn now to (a) and consider BSC. I have allowed that (a) is tempting. Yet BSC, as it stands, *says* nothing at all about conspecificity. So why is (a) tempting? Because, as Wilson notes, citing Mayr, BSC *implies* something about conspecificity: it implies that "a given individual organism is conspecific with organisms with which it can interbreed" (1999, 192–193). And it is easy to think that this amounts to (a). But it doesn't, because BSC does *not* imply that organisms are conspecific *in virtue of* interbreeding. The crucial error is to suppose that it does imply this, to suppose, quoting Wilson again, that BSC implies that "conspecificity is . . . determined by . . . organisms' standing in certain relations to one another" (1999, 193, emphasis added).³⁵ The point is a bit subtle and so I shall provide more details.

BSC, as it stands, straightforwardly provides an answer to the category problem. Now that answer implies that conspecific organisms are members of a group that is, as a matter of fact, an interbreeding (and reproductively isolated) group. For, to be conspecific is, by definition, to be members of a group that is a species and, according to BSC's category answer, what makes a group a species is being an interbreeding group. The category answer tells us that conspecific organisms are members of an interbreeding group but it does not tell us *in virtue of what* they are members of that group. Indeed, BSC's category answer is compatible with Intrinsic Biological Essentialism's answer to the conspecificity problem: it is compatible with the view that organisms are conspecific in virtue of sharing a certain intrinsic underlying property and, perhaps, a history. The compatibility is easy to see. Essentialism is motivated by the need to *explain* the observable properties of a group of organisms (Section 3). These properties include, of course, the property of interbreeding. So, according to Essentialism, it is *because* the members of a species share the intrinsic underlying properties necessary to make them conspecific that, in the given environ-

35. Matthen provides another clear example of the error (1998, 117–121).

ment, those members interbreed and hence have the property than makes them a species according to BSC.

Now we could, of course, *supplement* BSC as it stands with a relational answer to the conspecificity problem: organisms are conspecific *in virtue of* being able to interbreed. But, first, this supplement is not entailed by the BSC answer to the category problem and gets no support from the considerations that motivate that answer. Without further argument, the supplement is gratuitous. And, second, the supplement is a very bad answer to the conspecificity problem. It is a very bad answer *because*, as (b) shows, it is at odds with Intrinsic Biological Essentialism. It is shown by (b) that a relational conspecificity answer entails a relational taxon answer and our earlier argument shows that relational taxon answers are explanatorily hopeless. The supplement is not something that BSC should be saddled with.

Although (a) is certainly tempting it remains puzzling that people would give in to the temptation: it remains puzzling that they would not have doubts about this conspecificity route to anti-Essentialism. For, although the answer to the conspecificity problem that BSC is wrongly thought to give does entail that the answer to the taxon problem must be relational and nonintrinsic it does not *give* such an answer. And as soon as one tries to give one, it should become apparent how explanatorily inadequate a relational answer must be. At least, this should be apparent if one keeps in mind the needs of structural explanations. So, it remains puzzling that a person who starts with the idea that BSC implies a relational conspecificity answer would not be led to contemplate one of two responses: abandoning BSC *simply because* of that implication; or, abandoning the idea that BSC has the implication.³⁶

Our discussion of the puzzling conflation of the category problem with the taxon problem has focused on BSC. Yet the discussion applies just as much to ENC.

This concludes our discussion of how little bearing BSC and ENC have on the taxon problem 1. We turn now to consider the bearing of P-CC on that problem

9. P-CC and the Taxon Problem 1. At first sight P-CC, unlike BSC and ENC, may seem to provide an answer to the taxon problem 1 for species. Let *F*'s be a group of organisms that is a species according to P-CC. Whereas BSC and ENC told us little about how to complete 'Something

36. Could the lack of doubt arise from conflating the false view that organisms are conspecific in virtue of being able to interbreed with the true view (assuming BSC) that they are conspecific in virtue of that in virtue of which they are able to interbreed? (Thanks to Michael Dickson.)

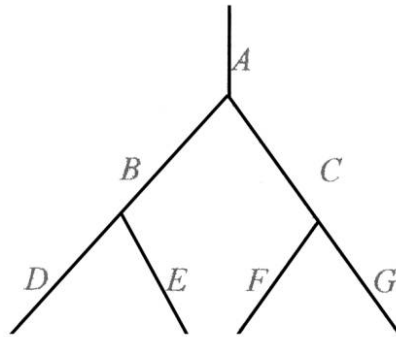


Figure 1.

is an *F* in virtue of . . . ,’ P-CC may seem to tell us a lot. The idea would be that since *F*’s are a species, an organism is an *F* in virtue of being a member of a group with a particular “evolutionary history;” the species is “this particular chunk of the genealogical nexus.” What makes this organism a lion not a tiger is that it is a member of a species having this particular place in “the branching tree-of-life” (Okasha 2002, 200). But this idea amounts to identifying a species simply by its relation to other species, to the species that preceded and succeeded it in its lineage: “You and I are members of *Homo sapiens*, therefore, because we both belong to the segment of the genealogical nexus which originated in Africa some 300,000 [years] ago (on current estimates), and which has not budded off any daughter species since that point” (2002, 200–201). But this answer to 1 is inadequate in just the same sort of way as our earlier second answer in discussing BSC and ENC: its relational identification of a species is explanatorily hopeless. To see this, it helps if we drop the actual names of species (like “*Homo sapiens*”), which might provide some illicit information, and replace them with schematic names using which we can capture the relational information that is all we are entitled to on this P-CC view. Thus suppose that species *A* splits into species *B* and *C*, then *B* splits into *D* and *E* and *C* splits into *F* and *G*. This is represented in the “tree-of-life” as depicted in Figure 1.

What does the P-CC view tell us about the nature of *B*? That *B* is descended from *A* and that *B* is distinct from *C*. And that’s what P-CC tells us that about the nature of *C* too. So that clearly does not distinguish *B* from *C*. No more does it distinguish *D* and *E* to know that each is descended from *B*, and *F* and *G* to know that each is descended from *C*. Furthermore, since P-CC does not distinguish *B* from *C* it does nothing to distinguish *D* and *E*, descended from *B*, from *F* and *G*, descended from

C. Suppose *D* were lions and *G*, tigers. Relational facts of the sort captured in this representation, which are the only facts that P-CC allows to constitute the natures of species, would do nothing to distinguish lions from tigers, hence nothing to explain the morphological, physiological, and behavioral differences between them. Suppose that we want to explain why *C* has poor eyesight, all we could appeal to on this view is its relation to *A*, *F*, and *G*. This is no more helpful in explaining the poor eyesight of *C* than was the relation of Polar Bears to some Ur-bear in explaining theirs. The structural explanations we need must appeal to the intrinsic properties of a species.

In sum, if P-CC is taken to give an answer to the taxon problem 1 for species, its answer is a very poor one. But it would be more charitable to suppose that it does not really intend to give an answer. Indeed, why would anyone think that it does? We have already noted a possible explanation in Section 8: the route from a category answer to a taxon answer may be via an answer to the *consppecificity* problem, the problem of saying in virtue of what organisms are conspecific.

We have agreed with Wilson that a relational answer to the conspecificity problem implies a relational answer to the taxon problem; that is (b) in Section 8. Wilson also thinks that P-CC, like BSC, implies such an answer to the conspecificity problem: P-CC implies that “consppecificity is determined by a shared pattern of ancestry and descent” or by something similar (1999, 193); that is (a) in Section 8. Now P-CC does indeed seem to give such a relational answer to the conspecificity problem, just as it seems to give a relational answer to the taxon problem. But that conspecificity answer would be bad because that taxon answer would be explanatorily hopeless, as we have seen. So, if P-CC really did involve such an answer it should be abandoned. But it is more charitable to suppose that, despite appearances, P-CC does not really propose an answer to the conspecificity problem.

In Section 8 I argued that the account given by Intrinsic Biological Essentialism of what it is for an organism to be a member of a particular species—an answer to 1—can be wedded happily to the accounts that BSC and ENC give of what it is for a group to be a species—answers to 2. Can it also be wedded happily to the influential P-CC? No. Adjustments would have to be made.

There is no problem wedding Essentialism to P-CC’s view that species are historical entities because Essentialism is not committed to a fully intrinsic essence. The wedding would yield the view that a species is constituted partly by intrinsic, probably genetic, properties and partly by a particular history: an organism is a member of a certain species *F* in virtue of having a certain intrinsic properties and being part of a particular chunk of the geneological nexus. Those intrinsic properties are the ones

that, together with environmental properties, explain the morphological, physiological, and behavioral properties of members of *F*. The chunk of the genealogical nexus in question should be the one that plays a role in explaining the evolution of the species.

But this wedding of Intrinsic Biological Essentialism to P-CC could not include two rather surprising features of P-CC.

- P-CC does not allow anagenesis, the forming of a new species without any split in the old. No matter how dramatically a lineage changes it will not form a new species unless it splits (Hennig 1966; cf. Simpson 1945). So if *Homo sapiens* had evolved from protists without any splits, all the organisms in this lineage would be in the same species. Kitcher aptly notes that “this strikes many people as counterintuitive (even insane)” ([1989] 2003, 151). Essentialism could not go along with this rejection of anagenesis (assuming, as we obviously do, that some groups covered by Essentialism *are* species).³⁷ For, as Sterelny and Griffiths point out, the rejection of anagenesis has the consequence that “there is no upper limit to the amount of evolutionary change that can take place within one species” (1999, 7). So there would be no limit to genetic variation in a species and hence its essence could not consist of genetic properties. But the rejection of anagenesis is deeply implausible. Why? Because a doctrine that, in some possible world, places *homo sapiens* and protists in the same species seems to have lost track of the explanatory significance of an organism being a *homo sapiens* or a protist.
- Suppose that a species *A* splits off a side branch that forms a daughter species *B* but the population otherwise remains unchanged; that is to say, apart from the members of *B*, the descendants of the members of *A* do not differ phenotypically or genetically from their ancestors. So, had there not been the split forming *B*, those descendants would have all been members of *A*, on anyone’s view of species. Yet P-CC has the consequence that because *B* did come into existence those descendants form a new species *C*. Essentialism cannot accept this. According to Essentialism, if a population remains unchanged then its members must be conspecifics. But the P-CC view is another implausible one, and for a closely related reason. If a population remains unchanged then its members should all be grouped together in explaining their common morphological, physiological, and be-

37. It is an interesting empirical question whether there are many plausible actual cases of anagenesis.

havioral properties. That's what the explanatory role of species seems to require.³⁸

In sum, Essentialism could be wedded to P-CC at the cost of dropping P-CC's two surprising, and implausible, features.

This concludes the discussion of the bearing of species concepts on the taxon problem 1 for species. Despite the consensus, BSC and ENC do not give a relational answer to that problem and if P-CC is taken to do so, its answer is a very poor one. My tentative diagnosis of the error in the consensus is that it arises from a conflation of the taxon problem with the category problem, a conflation encouraged by some mistaken thoughts about conspecificity. Perhaps also a focus on evolution has taken attention away from the needs of structural rather than historical explanations.

10. Variation and Change. The consensus among philosophers of biology is that doctrines like Intrinsic Biological Essentialism are at odds with Darwinian evolutionary theory (Sober [1980] 1992; Griffiths 2002). We have been discussing what is thought to be the most important problem for such doctrines: contemporary historical views of species. We must now consider a number of other alleged problems centering on variation and change.

Variation A. Sober claims that “no genotypic characteristic can be postulated as a species essence; the genetic variability found in sexual populations is prodigious” ([1980] 1992, 272). Others write in the same vein. Thus Wilson rejects genetic essentialism because “the inherent biological variability or *heterogeneity* of species with respect to both morphology and genetic composition is, after all, a cornerstone of the idea of evolution by natural selection” (1999, 190). And Okasha claims that

Intra-specific genetic variation is extremely wide—meiosis, genetic recombination and random mutation together ensure an almost unlimited variety in the range of possible genotypes that the members of a sexually reproducing species can exemplify. It simply is not true that there is some common genetic property which all members of a given species share and which all members of other species lack. (2002, 196).

38. Sterelny and Griffiths claim that, according to P-CC, the levels of the traditional Linnaean hierarchy above the base level of species “make little sense” (1999, 201). If this were taken as a view of the *taxa* then Essentialism should not go along with it. But P-CC does not support such a view of the *taxa*. The claim should be taken rather as a view of the *categories*, in which case it is quite compatible with Essentialism (Devitt 2008).

Surely, one thinks, this must be an exaggeration. Surely there are genetic properties that humans share and that say chimpanzees, let alone worms, lack. Indeed, even Mayr, no friend of essentialism, talks of “the historically evolved genetic program shared by all members of the species” ([1963] 1992, 17).³⁹ And Okasha continues: “This is not to deny, of course, that there are important genetic similarities between members of a single species . . . species taxa are distinguished by clusters of covarying [chromosomal and genetic] traits, not by shared essences” (2002, 197). Great! So the clusters *are* the essences!! On the strength of these remarks, it seems as if the consensus should be simply that the crude idea that there is, say, “a tiger gene” is wrong. But to reject that crudity is not to reject the idea that a certain cluster or pattern of underlying, largely genetic, properties is common and peculiar to tigers.⁴⁰ So my *third* main point in defense of Intrinsic Biological Essentialism is: *an intrinsic essence does not have to be “neat and tidy.”* And, because the intrinsic essence is identified by its causal work, we need not be concerned that the identification will be *ad hoc*: the essence of the Indian rhino is the underlying property that does, as a matter of fact, explain its single horn and other phenotypical features.

Variation B. Okasha emphasizes the importance of variation to natural selection: “Darwinianism leads us to expect variation with respect to organismic traits, morphological, physiological, behavioural and genetic. For genetically-based phenotypic variation is essential to the operation of natural selection” (2002, 197). Sober thinks that this variation clashes with essentialism’s commitment to the Aristotelian “Natural State Model”: essentialism takes the variation to be the result of “interfering forces” taking an organism away from its “natural state” ([1980] 1992, 257–259); to be “the result of imperfect manifestations of the idea implicit in each species” (Mayr [1963] 1992, 16); to be “deviation” from an “ideal” (Griffiths 2002, 78–79). This contrasts with the Darwinian view: “Individual differences are not *the effects* of interfering forces confounding the expression of a prototype; rather they are *the causes* of events that are absolutely central to the history of evolution” (Sober [1980] 1992, 264). Furthermore, “the Natural State Model presupposes that there is some

39. Kitcher ([1984] 2003, 132–133, note 27) refers to other similar suggestions in the literature. And note this claim by Hebert, as reported in the earlier-cited item from the *Scientific American* online (note 13): “We have very low levels of variation within a species and this deep divergence between species” (Biello 2007).

40. The evidence seems to point to genes that switch other genes on and off—for example, Hox genes—being particularly important to the nature of a biological kind (Carroll 2005).

phenotype which is the natural one *which is independent of a choice of environment*" (Sober [1980] 1992, 268).

Essentialism need not go along with the teleological thinking of the Aristotelian Model and Intrinsic Biological Essentialism does not.⁴¹ That doctrine can and should accept the Darwinian view of variation: variation within a species is indeed to be expected; species are indeed, as Griffiths says, "pools of variation" (2002, 78). Essentialism is committed simply to the view that in the pool of variation among the members of a species there are shared intrinsic, probably largely genetic, properties. And Essentialism rejects the idea that it is "not natural" for a corn plant of a particular genotype to wither and die, owing to the absence of trace elements in the soil (cf. Sober [1980] 1992, 268).

Gradual Change. Hull puts the problem for essentialism thus: "according to evolutionary theory, species develop gradually, changing one into another. If species evolved so gradually, they cannot be delimited by means of a single property or set of properties" ([1965] 1992, 203; see also Ruse [1987] 1992, 347). According to Griffiths, essentialism "is precisely the 'typological' perspective on species that Darwin had to displace in order to establish the gradual transformation of one species into another" (2002, 77; see also Caplan 1980, 73).⁴² But Darwin didn't have to. Suppose that *S1* and *S2* are distinct species, on everyone's view of species, and that *S2* evolved from *S1* by natural selection. Essentialism requires that there be an intrinsic essence *G1* for *S1* and *G2* for *S2*. *G1* and *G2* will be different but will have a lot in common. This picture is quite compatible with the Darwinian view that the evolution of *S2* is a gradual process of natural selection operating on genetic variation among the members of *S1*. Indeed, gradual change is obviously compatible with having essential intrinsic properties: rivers, mountains, continents, planets, and so on, are all the result of gradual change and yet all have partly intrinsic natures.

Still, there may seem to be a worry, nicely expressed by Sober:

Evolution is a gradual process. If species *A* gradually evolves into species *B*, where in this lineage should one draw the line that marks where *A* ends and *B* begins? Any line will be arbitrary. Essentialism, it is alleged, requires precise and nonarbitrary boundaries between natural kinds. (1993, 147)

41. Nor need it go along with Hull's "three essentialist tenets of typology" ([1965] 1992, 201).

42. Similarly Ereshefsky, writing about the essentialism of Lyell and Lamarck, claims that "their conception of species as evolving entities conflicts with this essentialist requirement" (1992a, xv).

This raises three issues: indeterminacy (or vagueness), arbitrariness, and “worldmaking.”

Indeterminacy. Ereshefsky, paraphrasing Hull ([1965] 1992), starkly puts the problem that indeterminacy is alleged to pose: “The boundaries of species are vague . . . there is no genetic or phenotypic trait that marks the boundary from one species to the next. Therefore no trait is essential for membership within a species” (1992b, 188–189). But this is a mistake: Essentialism does *not* require sharp boundaries between species. On the *Essentialist* picture, the evolution of *S2* from *S1* will involve a gradual process of moving from organisms that determinately have *G1* to organisms that determinately have *G2* via a whole lot of organisms that do not determinately have either. There is no fact of the matter about where precisely the line should be drawn between what constitutes *G1* and what constitutes *G2*, hence no fact of the matter about where precisely to draw the line between being a member of *S1* and being a member of *S2*. Essences are a bit indeterminate.

There are two reasons not to be worried by this. First, indeterminacy is everywhere. It is indeterminate whether a certain *x* is a mountain, or a certain *y*, a planet,⁴³ but this does not show that there is no essence to being a mountain or a planet. Mount Everest has the somewhat indeterminate essence of being a mountain and yet is determinately a mountain; Mars has the somewhat indeterminate essence of being a planet and yet is determinately a planet. Second, there is just the same level of indeterminacy about species *whatever one’s (Darwinian) view of them and of essentialism*, as indeed the passage from Sober indicates. For, *everyone agrees* that there comes a point where two organisms that have some common ancestor are nonetheless of different species. Yet there is no determinate matter of fact about precisely where that point is. And it is very easy to spot the root of the problem. We are tempted to say that an offspring and its parent are conspecific whatever the mutation (as Okasha indicates; see 2002, 197). But if we do say this, it is obvious that *all* organisms will (probably) come out conspecific. Biology faces a classic *sorites* problem. *The indeterminacy that biology must learn to live with is no special problem for Intrinsic Biological Essentialism.*⁴⁴ That is my *fourth* main point in the defense of the doctrine.

We have been talking about the issue of indeterminacy in the world.

43. The recent debate by the International Astronomical Union shows that Pluto is a good example of this indeterminacy.

44. “Essentialism is in principle consistent with *vague essences*” (Sober [1980] 1992, 253). Sober also draws attention to the fact that Aristotle was aware of “line-drawing problems” ([1980] 1992, 252–253).

This is likely to raise a worry about arbitrariness. There may indeed be some arbitrariness in the way we talk about the world. But that is not to say that there is arbitrariness in the world.

Arbitrariness. Biologists choose, for various explanatory purposes, to introduce names for certain groups of organisms thought to be species. There could be some arbitrariness about what groups to choose. Let us start with the worst case. Suppose that we allow for anagenesis, as my Essentialism must (Section 9): a new species can be formed without any split in a lineage. Suppose next that evolution were not only gradual but also *steady*: the morphological, physiological, behavioral and genetic properties of organisms in the lineage change at a steady rate. Essentialism alone does not rule this out. How then would we choose where to draw our (indeterminate) lines in naming the species of this lineage? Clearly, there would be a deal of arbitrariness about this choice.⁴⁵ But we should not exaggerate how much. Our explanatory purposes in introducing a name for a species demand that we draw the lines around a group that is small enough to share a whole lot of important properties and large enough to yield broad generalizations. That is what is required for structural explanations. And, as G. G. Simpson points out, “such arbitrary subdivision does not necessarily produce taxa that are either ‘unreal’ or ‘unnatural’” (1961, 60–61). Furthermore, *wherever we draw the lines* in naming a group “*F*,” it is still the case that the intrinsic essence of being *F*, together with the environment, explains the morphological, physiological, and behavioral properties typical of *F*’s.

Turn next to the best case. This is the situation if the hypothesis of *punctuated equilibrium* (Eldredge and Gould 1972) is right. On this hypothesis, evolution is far from steady. Species do not change much over most of their existence and then, in a relatively short period of time, either go extinct or evolve into daughter species. So, on my *Essentialist* picture, the need to explain the morphological, physiological and behavioral properties of organisms would dictate that organisms in the period of stasis formed a species that should be named. We would draw our (indeterminate) lines in the period of rapid change. Our choice would hardly be arbitrary at all.

The important point for our purposes is that, wherever the truth lies between the worst and the best cases, arbitrariness poses no threat to Essentialism. The groups we name will still have partly intrinsic essences.

45. “The idea then is that if phenotypic change does not proceed by large jumps (*saltations*), then species are not objectively identifiable over time” (Sterelny and Griffiths 1999, 180). They do not endorse this idea.

Indeed, arbitrariness is really a problem for the species *category* rather than the taxa.

“*Worldmaking.*” We *do* have a choice about what groups of organisms to name “*F*.” It is sadly common to confuse this with a choice we certainly *do not* have: the choice about which things *are F*. This is, in effect, the distinction between *making theories* and *making worlds*, a distinction the importance of which can hardly be exaggerated.⁴⁶ We name a group of organisms “*F*” for explanatory purposes and hence, even at worst, the choice of what group to name is mostly not arbitrary. But, *however arbitrary it is*, indeed even if it was *totally* arbitrary, we would not thereby make those organisms *F*. When biologists chose to apply the name ‘*Drosophila melanogaster*’ to a vast number of insects, they did not thereby *make* those insects *Drosophila melanogaster*. They always were *Drosophila melanogaster* and would have been even if there had been no biologists around to call them anything.⁴⁷ It is common to talk as if, in doing science, we impose our concepts to “divide up reality.” But this is not literally so: we choose our concepts in an attempt to discover the causally significant features of a nature that is already “divided up.”⁴⁸

Monsters. This discussion provides the wherewithal to deal with “monsters,” offspring that differ greatly from their parents. Monsters are thought to refute Essentialism because they lack what might plausibly be proposed as the intrinsic essences of their parents’ species. For example, Okasha claims that “if a member of the species produced an offspring which lacked one of the [essential] characteristics, say because of a mutation, it would be very likely to be classed as conspecific with its parents” (2002, 197). Sterelny and Griffiths put the point more firmly: “No intrinsic genotypic or phenotypic property is essential to being a member of a species. . . . People born with the wrong number of chromosomes, eyes,

46. I have argued this at length elsewhere (Devitt 1997, particularly Chapter 13). Overlooking the distinction seems to rest on something like a use/mention confusion.

47. Kyle Stanford (1995) has a different view; see Ereshefsky 1998 and Devitt 2008 for criticisms.

48. It is easy even for staunch realists to slip into loose ways of talking that suggest worldmaking. Thus Kornblith says that when we “group objects together under a single heading on the basis of a number of easily observable characteristics . . . we thereby create a nominal kind” (1993, 41). But we don’t! We create a *concept* that picks out a kind that may or may not be “real” in Locke’s terms (note 4 above) but which has its members independently of our creation. And Boyd, talking of kinds with nominal essences, says that their “boundaries” are “purely matters of convention” (1999, 142). But they aren’t! Our naming a kind picked out by a certain set of descriptions is conventional but the boundary of the kind thus picked out is not.

or arms are still human beings. So the essential properties that make a particular organism a platypus, for example, are historical or relational” (1999, 186). Now Okasha is surely right that we would very likely classify any offspring as conspecific with its parents. But the *sorites* problem shows that we cannot always be *right* to do so, whatever we think of Essentialism: as Hull says, “Obviously . . . there must have been instances in which non-horses (or borderline horses) gave rise to horses” ([1978] 1992, 306). So what should the *Essentialist* say about monsters? One of two things: (i) If the mutations are gross enough, we should indeed say that the offspring is not of the same species as its parents. And that surely is what we would say, as monster movies sometimes illustrate. I doubt that we would even hesitate to say it of embryos that are so monstrous that they would not grow into viable organisms and are spontaneously aborted. (ii) In other circumstances we should say that the status of the offspring is indeterminate. Return to our schematic example of the evolution of *S2* from *S1*. At the beginning of that process, there were organisms that determinately had *G1* and so were determinately members of *S1*, and at the end, there were organisms that determinately had *G2* and so were determinately members of *S2*. But in between there were organisms that did not determinately have *G1* or determinately have *G2* and so were not determinately members of *S1* or determinately members of *S2*. All we can say is that the further an organism gets from determinately having *G1* to determinately having *G2*, the further it gets from being determinately a member of *S1*. This is vague of course, but that’s the way a lot of the world is, not just living things. Monsters are no special problem for Essentialism.⁴⁹

Laws. We are now in the position to respond to the questions raised in Section 3 about my treatment of biological generalizations. Question 1 was: “Surely any such generalization could have exceptions: a small mutation may lead to an organism that seems to be a member of a species

49. Philip Kitcher, in commenting on an early version of this article, claimed that “knockout” mutants produced by modifying “normal” zygotes, show that my treatment of monsters is too quick. It seems to me that my discussion accommodates these knockouts well enough: a minor mutant of a *Drosophila melanogaster* may still count as a *Drosophila melanogaster* because it has the essential intrinsic property that explains the characteristics it shares with “normal” *Drosophila melanogaster*; a gross mutant would not count as a *Drosophila melanogaster* because it does not have that property; the status of other mutants, doubtless most of the mutants, is simply indeterminate. We can learn about *Drosophila melanogaster* from these mutants, as we did, even if they themselves are not determinately *Drosophila melanogaster*. There is plenty of room for subtlety here. And if I am right in my arguments, something along the lines of my proposal has no viable alternative.

and yet lacks the property attributed to the species by a generalization. So the generalizations do not seem to be lawlike. How does Intrinsic Biological Essentialism deal with that?" Essentialism surely does demand that these generalizations be lawlike rather than accidental. In a group of animals, it does not just *happen* to be the case that the members of a certain subgroup have one horn and the members of another, two. It is *because* the first subgroup are Indian rhinos and the latter, African rhinos; it is part of their very natures to have (in their environments) one horn and two horns respectively. But how can the generalizations be lawlike if there could be exceptions?⁵⁰

There are several things we might say in answer. First it is common, perhaps even the rule, for laws in the special sciences to have exceptions: they hold only *ceteris paribus*. So why should this be a problem for biology in particular? Indeed, if Nancy Cartwright (1983) is right the situation is not much different in physics. Second, *statistical* generalizations can be lawlike. Thus the claim that, say, 90% of *F*'s are *P* can be lawlike: it can sustain the subjunctive conditional that if something were an *F* it would very likely be *P*.⁵¹ Finally, we can say that universal biological generalizations are indeed lawlike but that there is some indeterminacy about precisely which organisms they would cover. 'All *F*'s are *P*' may be a law in that anything that would be determinately *F* would be *P* but there might be some organisms that would not be determinately *F* or determinately not *F* and so there would be no determinate matter of fact about whether the law covered them. Note that this is not primarily an epistemological problem of *telling* what organisms the generalizations cover: it is primarily a metaphysical problem. Of course, even where there is a determinate matter of fact that generalizations cover certain organisms there can still be a problem discovering this; thus, many black birds in Australia were determinately swans at a time when biologists believed that all swans were white.

Essentialism? Question 2 was: "It is of course the case that the truth of any such generalization must be explained by an intrinsic, probably largely genetic, property, but why does that property have to be an *essential*

50. Note that exceptions that arise from varying the environment are not a problem. Indeed, typical generalizations about an organism are implicitly restricted to its "normal" environment.

51. Griffiths points out that "the generalizations of the special sciences often fail to live up to the ideal of a universal exceptionless law of nature. . . . Nevertheless . . . they have "*counterfactual force*" (Griffiths 1999, 216). Referring to history, social sciences, geology and meteorology, Boyd notes that "causally sustained regularities . . . need not be eternal, exceptionless, or spatiotemporally universal" (1999, 152).

property of the kind in question?" Suppose that the generalization is 'All F 's are P ' and that the explanatory intrinsic property is G . So it is agreed that, ultimately, it is because F 's have G that they have P . The question asks why we must take G to be an essential property of F 's. My answer rests on the just-argued claim that the generalization is lawlike. So, anything that *would be* determinately an F would be P (in the appropriate environment). But now, in virtue of what is that the case? The answer is that anything that would be an F would have G . Indeed what other answer could we seriously entertain given that having G explains why all actual F 's are P ? We have now answered question 2. For, if anything that would be an F would have G then having G is essential to being an F : that is what it is to be an essential property.

11. Conclusion. I have proposed the doctrine, Intrinsic Biological Essentialism: Linnaean taxa have essences that are, at least partly, underlying, probably largely genetic, intrinsic properties. The consensus in biology and philosophy of biology is that any such essentialism is deeply mistaken. In Section 2, I set out evidence that this is indeed the consensus.

In Section 3, I presented my central argument for Essentialism: the ubiquitous generalizations of biology need structural explanations that rest on essential intrinsic underlying properties of kinds. That was my *first* main point in defense of Essentialism.

In Section 4, I described current species concepts. The consensus view is that these make doctrines like Essentialism untenable because, according to these concepts, species are *relational*. In Section 5, I emphasized a distinction that is crucial to my defense of Essentialism from this consensus view. It is the distinction between two problems, a taxon problem 1 and a category problem 2:

1. What is the essence of being F (where F 's are a group under one of the biological taxa)?
2. What is the essence of being a subspecies, species, genus, etc.?

This distinction yields two ways to understand the question "What is a species?" The question could be asking about the nature of any group that happens to be a species or it could be asking about what it is to be a species. My *second* main point in defense of Essentialism, argued for in Sections 6, 7, and 9, was that the (partly) relational species concepts are primarily concerned with 2 whereas Essentialism is concerned with 1.

In Section 6, I argued that, not only do the species concepts straightforwardly yield answers to question 2 for species, that seems to be what they are intended to do. In Section 7, I argued that, contrary to the consensus, the biological species concept and the ecological niche concept do not answer 1 nor are they even associated with relational answers that

are close to being explanatorily adequate. Indeed they can both be happily wedded to Essentialism's (partly) nonrelational answer. Where has the consensus gone wrong? My tentative diagnosis in Section 8 was that the error arises from a conflation of the taxon problem with the category problem, a conflation encouraged by some mistaken thoughts about conspecificity.

In Section 9, I considered the influential phylogenetic-cladistic concept (P-CC). P-CC might be taken to give a relational answer to 1 as well as 2 but, if it were, its answer would be explanatorily inadequate. Can Essentialism's answer to 1 be wedded to P-CC's answer to 2? Essentialism can easily accommodate P-CC's view that species are historical entities. However it cannot accommodate two of P-CC's features: its rejection of anagenesis and its view that a species must go extinct when it has a daughter. But those features seem unwelcome anyway.

Finally, in Section 10, I argued that some general features of Darwinism do not undermine Essentialism. Variation within a species can be seen to be compatible with Essentialism once one realizes that an intrinsic essence does not have to be "neat and tidy"—my *third* main point in defense of Essentialism—and that Essentialism is not wedded to the Aristotelian "Natural State Model." Essentialism can accept the gradual change of one species into another. Still, there are some concerns raised by the lack of sharp boundaries between species. First, Essentialism must accept a certain indeterminacy about species. But this is no worry because this indeterminacy has to be accepted whatever one's (Darwinian) view of species and of essentialism; biology faces a *sorites* problem. That was my *fourth* main point in defense of Essentialism. Next, Essentialism is compatible with there being a certain amount of arbitrariness in choosing which groups of organisms to name as species. But this choice, however arbitrary, must not be confused with a choice we do not have: the choice to make things member of a group we have named. The fact of indeterminacy enables Essentialism to deal with the problem of monsters—organisms lacking what might plausibly be proposed as the intrinsic essences of their parent's species—and to maintain the lawlike status of biological generalizations despite apparent exceptions.

I have dealt with the objections to Intrinsic Biological Essentialism that I have found in the literature. Perhaps there are other objections that would be more effective. Perhaps it can be shown that my argument in favor of Essentialism—the argument from explanation—is inadequate. Given the strength and longevity of the consensus against such an essentialist doctrine, it seems reasonable to predict this. Still, it remains to be seen whether it is so. At the very least I hope to have shown that the case for the consensus needs to be made a great deal better than it has been.

If the arguments of this article are good, the consensus relational view

about particular species is quite mistaken. And many claims that biologists make day in and day out about the living world require species to have natures that they do not have according to this consensus.

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