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Participation in evolution and sustainability


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Abstract:
The modern synthesis of genetics with evolution slanted our understanding of evolution and of ourselves by rejecting Darwin’s view of animals as participating in their own evolution. Defining evolution in terms of genetics, the modern synthesis indulges excessive individualism and distorted self-images as self-made. At the same time such gene-centered thought, evoking images of master molecules making us who we are, hollows out volition and so also moral concerns and political alternatives. Drawing on the geography of thought, we argue that stubbornly tacit preformationist biological thought reflects and anchors social processes that limit adaptability in reaching toward sustainable living. We appeal for leveraging sustainability efforts by affirming in theory and in the public square an open image of human nature that recognises the participation of our ancestors in becoming who we are, obliging people to make their history together. Achieving the collective self-regulation sustainability requires may depend on correcting slanted reasoning about ourselves.

Keywords: evolutionary biology, human geography, human nature, participation, sustainability
Such a view … poses a severe threat to humankind because it links the
authority of science with an imagery of the human condition that can
only trivialize and obfuscate its beneficiaries.
Sigmund Koch (1981, 266)

[A] perverse and mistaken idea of human nature … endangers our
existence.
Marshall Sahlins (2008, 112)

Introduction
In his appeal for a more open image of humanity Kenneth Bock recalled Circe
magically turning Ulysses’ men into animals, lifting from their shoulders the
more acute self-consciousness and moral challenges of being human. In a turn
of Homer’s original plot, Gelli (1549) required Ulysses to convince his men to
choose becoming human again, but his arguments ‘about the refinements of
human life and the superiority of human understanding or intellect carry little
weight against rejoinders concerning the dire results of people’s incontinent
indulgence in luxury and the absurdities into which they have been led by their
finely honed but tortuous reasoning’ (Bock 1994, 17).

This variation on a Western canon rouses perennial tensions at the
boundary of social theory and biology. What is human nature? How are humans
similar to and different from each other and other animals and what does this
tell us about ourselves? How has reasoning been twisted, toward what ends, in
framing and answering these questions? How do our answers reflect and affect our indulgences and prospects for achieving the collective self-regulation sustainability requires?

Taking up these questions and drawing on geographies of thought and boundary crossings, we argue that stubbornly tacit preformationist biological thought reflects and anchors social processes that limit adaptability in reaching toward sustainable living. These implications for sustainability underscore the importance of human geography’s engagement with contemporary life sciences (Castree 2009) and the urgent need for ‘conversation about our “species being”’ (Harvey 2000, 207).

**Participating in evolution**

With varying degrees of emphasis Darwin recognized behaviour, use and disuse, as a causal factor in evolution. In Darwin’s view (1964 [1859], 134-36), insects’ use or disuse of their wings on windswept islands, where they are ‘frequently blown to sea and perish’, were part of the natural selection of enlarged or reduced wing sizes in successful fliers and successful if ‘indolent’ walkers, respectively. The ostrich’s defensive use of its legs and disuse of its wings played some part in the selection of a large, flightless bird. For Darwin, insects and ostriches participated in their own evolution.

In *On the Origin of Species*, Darwin (1964 [1859], 134-137, 143, 206, 447, 479) invoked use and disuse in ‘long-continued’ time frames, not one generation to the next, and as ‘quite compatible with’, ‘aided by’, ‘aiding’ or
largely combined with, and sometimes overmastered by’ natural selection. And he consistently used the term “acquired” with reference to species, not individuals, across longer time frames, not one generation to the next, and often with explicit reference to selection processes. Yet Huxley (1960, 14, 20) and other prominent contributors to the modern synthesis considered Darwin’s references to use and disuse ‘Lamarckian errors’, chalked up to 19th century ignorance of genetics. They presented use and disuse as an alternative to ‘blind’ and ‘automatic’ natural selection, not as part of the selection process.¹ 

Evaluated only in terms of direct genetic inheritance, one generation to the next, of “acquired characteristics”, adaptations within the lives of animals were ‘rendered obsolete’ in evolutionary theory and ‘exposed … as sins against Occum’s razor’ (Lerner 1959, 173).

Why this skewed reading of Darwin that minimised behaviour’s role in evolution? A good start at answering this question would note Darwin’s invoking use and disuse both as sources of variation and as influencing selection outcomes. A full answer would also recognize that the theoretical refinements of the modern synthesis were not simply hard science prevailing over soft folk wisdom. One aspect of folk biology, essentialist preformationism (we are born with something inside us that makes us who we are), superseded another, use and disuse (manners maketh the man). Nature as transmitted, context independent, centralized genetic program causing development eclipsed nature as constructed, context dependent, causally diffuse, phenotypic product of development (Oyama 2000a). The flip side of 19th century ignorance of
genetics was 20th century indulgence in excessive causal attributions to internal, stable characters: genes.

Focused on integrating genetics with evolution, the modern synthesis cast behaviour more as consequence than cause of evolutionary change. Intention in the lives of animals was dismissed partly on the coattails of arguments regarding superagents in evolution, God or a watchmaker with vision. Mayr (1961, 1504) rejected all purpose as ‘singularly inapplicable to evolutionary change … If an organism is well adapted, … this is not due to any purpose of its ancestors or of an outside agency, such as “Nature” or “God”.’ This conflation of ancestral intention with externally directed evolution eased biology’s robust rejection of purpose in theory, even as the practical language of biology remained thick with purpose.2 Hence, Haldane’s quip that ‘Teleology is like a mistress to a biologist: he cannot live without her but he’s unwilling to be seen with her in public’ (quoted in Hull 1982, 298).

Evolutionary biology continues to present us with genes inside and an environment outside, which “interact” to produce an organism and its behaviour. A recent summary of evolutionary thought presents behaviour as evolving ‘just as morphological characteristics do’. Though neuro-developmental processes afford behaviour far more plasticity than anatomy and render behaviour less dependent on genetic mutations or recombinations presenting natural selection with different options, behaviour and morphology are treated similarly, as objects of evolutionary study explained by ‘genetic change, influenced by environmental circumstances’ (Meagher and Futuyma
2001, 3, 26). The scientific claim of computing an organism from its DNA is considered naïve not in principle, but for lack ‘even at present’ of an ‘instruction manual’, which awaits ‘improving technologies’ by which ‘genes that underlie behavioural variation will become increasingly easy to identify’ (Hoekstra 2010, 638-47).

Behaviour’s role has been shoe-horned into the internal (e.g., gene flow, Duckworth 2009) or external (e.g., niche, Lewontin and Levins 2007) side of this dualism or both, as “gene-environment interactions”. Innovative animal behaviours are called ‘evolutionary strategies’ (MacArthur and Wilson 1967, 6) or ‘evolutionary inventions’ (Grant 1998, 313), dodging the specter of intention while using the language of intention.

This bashfulness about behaviour reflects a causal privileging of genes and environments over the competent participation of organisms in their own development and evolution. Biological thought has long relied on tacit appreciation of this competence while explicitly minimising or neglecting it. Oyama (2000b, 336) describes several ways reasoning has been bent to this end by the intuitive appeal of a ‘homunculoid gene’. Traits are called “innate” or “acquired” based on analysis of variance, which is not the same as analysis of organisms (Lewontin 1974). While interacting only with other intracellular molecules, genes are said to “interact” with culture, humans being products of this “interaction”, not semiautonomous participants mediating and to some extent shaping it. Both similarities and differences across species are interpreted as indicating innateness, incest avoidance considered innate in humans because
it is seen in other animals, language considered innate because it is not. To
Oyama’s list, we would add the common misapprehension that genetic causes
of disease reflect the degree to which genes specify health. In biology, agency
is more comfortably ascribed at the scale of genes than of organisms.³

This causal privileging has been challenged repeatedly over the years.
Gullick (1905, iv) observed that ‘members of the same species, exposed to the
same environment in isolated groups, will often arrive at different methods of
dealing with the environment, and so subject themselves to divergent forms of
selection’. Waddington (1960, 401) included the formulations ‘animal chooses’
and ‘modifies environmental niche’ as part of the evolutionary system. Ewer
(1960, 162) noted that biology’s ‘reaction against teleology…went too far’ in
concluding, from the inability of animals’ activities to be directly inherited, that
these activities are irrelevant to evolution. Piaget (1971, 81) embraced
biologists like Waddington for making evolution depend in part on
development, ‘and not only the inverse’.

Recognizing complexity, these scientists avoided the rhetorical trap Taylor
(1998) calls simpling – reducing complexity to a manageable size for
convenience, subsequent claims being inflated by confusing heuristic simplicity
with valid generalization, confusing reductionism as method with reductionism
as explanation, and convenience amnesia. Simpling is deceptively subtle when
overextending otherwise sound concepts, such as “reaction norm”, which refers
to phenotypic variation of identical genes across different individuals in
different environments. Though this concept reveals limits on genetic control of
phenotypes, it has been routinely invoked to assert genetic explanation of any given variation, favouring overgeneralization from the reasonable proposition of partial genetic and environmental regulation of phenotypes to the sufficiency of genes and environments to explain any particular phenotypic feature, including behaviour (Sarkar 1999). Turning a blind eye to the participation of whole organisms in their own development and evolution is not parsimonious; it is simpling, after which much progress in science is filling in gaps of its own making.

With genes, environments and chance in the driver seat, organisms are to be explained and not heard. Though zebra finch singing induces gene expression (Jarvis et al. 1998), Clayton et al.’s (2009) model integrating songbird genomes, brains and behaviour recognizes bidirectional causation between physiology and genome but not between behaviour and physiology. This omission reflects the cognitive activity of biologists, not the lives of songbirds.

Yet this model does recognize behaviour’s role in natural selection, otherwise ‘largely unacknowledged in current evolutionary theory’ (Duckworth 2009, 514). Duckworth helps fill this gap by clarifying that behaviour can influence natural selection if it occurs in a large enough subset of the population and persists across several generations. Capacities for learning and cultural transmission subserved by a ‘robustly epigenetic’ mammalian brain (Goodenough and Deacon 2003, 806) expand significantly the means by which behaviour changes can pass these thresholds to influence selection.
Neglect of organisms’ evolutionary effectiveness extends from academia to public lectures, museums and popular books, feeding back into our folk biology. In a recent lecture, Leakey (2010) referenced his personal commitments and described elephants coming to the aid of a different species attacked by lions, yet consideration of elephant or human volition in evolution was out of bounds. Asked about the current status of Waddington’s inclusion of animal choices in modeling evolution, Leakey was ‘dismissive’ and recited the scientistic creed that chance alone is sufficient, with natural laws, to account for evolution.

The Smithsonian Institution’s David H. Koch Hall of Human Origins celebrates human inventiveness while presenting it more as a product – of ‘environmental change’, ‘large, complex brains’ and an evolutionary process defined by genetics – than as a causal force in human evolution (Potts and Sloan 2010, 53, 77, 168). Only obliquely intimated is the possibility that adaptive cultural innovations shaped human evolution by their incidental effects on selection pressures and outcomes, for example cooked food changing selection trade-offs for larger brains and smaller guts (Wrangham 2009) or hunting weapons leveling male hierarchies and reducing sexual dimorphism (Boehm 1999).

In his popular book, *Evolution for Everyone*, Wilson (2007, 159) misrepresents Boehm (1999) as showing that ‘egalitarianism is not a cultural invention that began in ancient Greece, as many have supposed, but is part of our genetic endowment’. Implicitly equating ancient Greece with culture and
deeper prehistory with ‘genetic endowment’, Wilson minimises the role of
moral sanctioning in creating and maintaining egalitarian dispositions. Boehm
emphasises that humans invented egalitarian politics by intentional, indeed
vigilant, cultural practices, creating ‘a major social-structural divergence at the
level of the phenotype’ (Boehm and Flack 2010, 76). There is good reason to
expect modest genetic contribution to such complex behaviour, which would
anyway be as much consequence as cause, actions harnessing genetic variation,
sometimes toward conscious ends (Boehm 2008).

This theoretical bias against organisms as participants in their own
evolution has been shaped in part by biologists’ denial of their own
participation in understanding life.

‘The achievements which form the subject matter of
biology can be identified only by a kind of appraisal which
requires a higher degree of participation by the observer in his
subject matter than can be mediated by the tests of physics and
chemistry. The current ideal of “scientificality” which would
refuse such participation would indeed destroy biology but for
the wise neglect of consistency on the part of its supporters’
(Polanyi 1957, 482).5

Also biasing theory against recognizing participation in evolution have
been practical and conceptual difficulties in studying behaviour, genes
providing more convenient “handles” (Gannett 1999); confusing science’s
pragmatic exclusion of purpose with the conclusion that purpose is not real,
‘limit[ing] a problem by reason of a method of attack’ (Whitehead 1962, 15); confusing the historical priority of genes with their current domination of organisms (Gould 2001); a folk biology that favours homunculoid genes (Bateson and Mameli 2007); prestige processes within the culture of science (Burkhardt 1970; Fracchia and Lewontin 1999); and political-economic interests in controlling populations and sowing division with biologically essentialist thought (Kearns 2010; Nally 2011). These among other leanings elevated ‘one aspect of evolution to its very definition’ (Oyama 2000a, 12).

Crossing boundaries

Transitions across boundaries underscore the role of animal behaviour in evolution. Evolutionary changes commonly attributed to changing environments, including changes associated with colonisation and understood as “ecological release”, are often initiated and mediated by animals’ activities. Innovative foraging behaviours are major determinants of successful colonisation of novel environments (Sol et al. 2002; 2005a), granting some birds the power to stay, while others are obliged to migrate (Sol et al. 2005b). This is one way behaviour drives speciation and may explain why behavioural flexibility is associated with higher rates of speciation (Sol et al. 2005c).

Colonisation events reveal the participation of organisms in changing their environments and consequent changes in behaviour and morphology. Conspicuously but not uniquely on islands, whose significance here stems more from their boundaries than their isolation (Greenough 2006), these
biogeographical processes indicate that behaviour ‘may be a potent force in driving evolution in novel directions’, while sometimes actually reducing the likelihood of genetic change (Price et al. 2003, 1433).

Price (2008) notes social transmission of innovation in some birds and innovates himself in emphasising behaviour and development alongside ecology and genetics. The Grants (2008) also feature behaviour more prominently as causes of speciation among Galapagos finches. Attention to boundary crossings contributed to this constructive redirection in thinking about speciation, which had ‘tended to overemphasise genetics at the expense of … behaviour’ (Owens 2008, 185).

Human colonisation also depends on behavioural flexibility. John Terrell (1986) described such flexibility in the settlement of Pacific islands, seen in tolerance for diversity of thought, intensive sharing that was by geographic necessity also quite extensive, and remarkable navigational skills. Without benefit of sextant and meeting the demanding requirements of dead reckoning, Polynesian wayfinders routinely navigated many miles of open sea. Such ingenuity in the service of perpetual sharing allowed humans to cope with the harsh biogeographic contingencies of small numbers on small islands, known to impose high rates of extinction.

Having changed their environment toward a more continental abundance and connectedness, ‘it demands no impossible leap of imagination to view all of Polynesia as just another Melanesian village’ (Terrell 1986, 261). Socially focused and socially mediated mobility (Warf 2008) is part of a broader
adaptability by which humans made geographic boundaries more permeable, deferring the usual biological constraints on island dominants while rendering islandness less distinctive in human geography (Cliff et al. 2000).

**Geography of biological thought**

The geography of thought casts light on evolutionary theory’s entanglement in social issues. Compared with East Asians, Westerners attend more to actors than context, attribute more causal significance to fixed internal characteristics of categorized objects, make more causal attributions even while seeing fewer factors as relevant to understanding the world (Nisbett 2003), and recognize individual agency more than group agency (Kashima et al. 2005; Haslam et al. 2006). Causal theories and focus of attention reinforcing each other (Plaks et al. 2001), Western habits of thought arrive comfortably to a view of social, political and economic processes as driven by internal, stable, biologically “endowed” dispositions of individuals. Hence, twentieth century biology in the West ‘witnessed a general trend to emphasize the constancy and causal efficacy of the genotype at the expense of the complexity of its interactions’ (Sarkar 2006, 80-81). Less inclined to individualism, reductionism and human-animal dualism, the East, particularly Japan, readily accepted evolution including attribution of intentions to animals and pragmatic anthropomorphising, to the advantage of Japanese primatology (de Waal 2001a).

‘Culture seeps into science unbidden’ (de Waal 2001b 46), the study of behaviour being particularly vulnerable to cultural bias. Compared with
Western science, Asian scientists have paid little attention to individual differences in stereotyping or prejudice (Fiske 2000). More consequential examples include the shadows power relations and nationalism cast over scientific conceptions of race (Blakey 1998; Templeton 1998) and prehistoric archeology (Richards 2003).

Essentialist individualism finds expression in both scientific and lay theories (Keller 2005; Levy et al. 2006), between which reciprocal influences played out historically as Darwinian evolution’s conceptual consistencies with capitalism brought it wider acceptance. Livingstone (2003) sketches this geography, regional differences in receptiveness to Darwinism varying with religious commitments, racial politics and economic interests. For example, Darwinism was embraced in New Zealand as a ready justification for routing the Maori. Russia, lacking a market based middle class, favoured cooperation and rejected Malthusian elements of Darwinian theory. Kropotkin and other Russian critics recognized that ‘contrary enthusiasms’ regarding Darwin ‘might record the parochiality of his different surroundings’ (Gould 1987, 17).

Geography has also been exploited in presenting simplified cartographies of human lineages, ‘akin to creation myths’, that conducted ideological power (Livingstone 2010, 205; Nash 2005). But evolutionary thought may reflect and influence social and political-economic processes more by its corollary myth of development as genetically given – bolstered by mapping genes and ‘putting the gene in its place’ (Hall 2003, 159; cf. Hinchliffe 2001) – than by its origin story.

Ever since Malthusian thought simplified the Irish Potato Famine as a self-inflicted consequence of population growth and discounted Irish resistance as ‘the disorderly conduct of a people so degraded by their self-inflicted poverty’ (Ross 1998, 44), mutual influences between biological and political-economic thought have naturalised excesses of inequality7 by indulging self-made conceptions of one’s own fortune and others’ misery. Pragmatic and socially influenced selective attention to presumed centralised causal agents in gene-centered biological thought to some extent reflects and exacerbates this politics.

The importance of attention is underscored by Stotz et al.’s (2004) unique empirical study of biologists’ cognitive activity finding neglect of contextual factors by molecular more than developmental biologists. Hoekstra’s (2010) informal survey found geneticists more optimistic than organismal biologists
about imminently “knowing it all”. Beyond its partial basis in epistemological
naivety and cognitive miserliness, what social processes surround this
exhuberance? Stotz et al. suggest that for biologists, the gene concept may
function like a stereotype. The point is not that biologists are ideologues, but
that science is fundamentally a social process. Social influences on, and
consequences of, the science of who we are and how we came to be tend to
remain tacit, naturalised and resistant to critical reflection.8 Scientists who
warn against social and ideological influences (e.g. Segerstrale 2000, 341) are
themselves influenced (Laland and Brown 2002, 97).9

‘Scientific progress is very much about improving the metaphors we use so
that our interventions in the world are more successful’ (Sayer 2000, 78).
Whatever their own understanding of the complexity of genes’ interactions with
other developmental resources, biologists should appreciate the social
implications – and consequences for their own thinking – of their metaphorical
references to genes as blueprints or programs making ‘a deterministic reading
of claims about the role of genes in development almost inevitable’ (Griffiths
2006, 192). Intended or not, the idea of evolution thus accommodates
historicism’s “inevitable” (Popper 1957; Beatty 2001), seen today more in
geographically distributed transnational capitalism than in national
totalitarianism. Both rely on shrinking volition and thereby also moral concerns
and political alternatives, succinctly expressed when Thatcher dusted off
Spencer’s ‘there is no alternative’.
Also accommodated is the tendency, mischievous because commonsensical, to draw lines among ourselves defining essentially different categories of people. Current expressions of biologically essentialist thought include framing wage suppression in terms of immigration by the alien “other” rather than fair wages for labor. Especially vulnerable are portrayals of the past, as when ethnic labels are applied to genetic lineages. While genes’ bookkeeping features provide useful tools for phylogeography and researchers themselves generally appreciate the complexity of the history they model, common sense uses of categorical terms to summarise findings can bias readers’ understanding toward simplified and divisive origin myths that exaggerate group stability, purity and distinctiveness. Freighted with social significance, genetic studies demand great interpretive vigilance (Terrell 2010).

Biological thought includes an understanding that we are made by our participation in relationships and communities in for example developmental systems theory (Oyama, Griffiths and Gray 2001), developmental psychobiology (Gottlieb 1992, 2003), the expanded evolutionary synthesis of Jablonka and Lamb (2007) and ecological developmental biology (Gilbert and Epel 2009). The developmental biological insight that ‘to be one is always to become with many’ (Haraway 2008, 4) is expressed in political-economic activity: in appreciating the broad benefits and responsibilities, for each individual, of public investments and citizenship; when political processes maintain healthy levels of equality by affirming the fundamental complementarity – not simple opposition – of autonomy and solidarity,
individual and collective, economic performance and social justice (Bauman 2008a; Clark and Clark 2009; Sayer 2011); and when community-based co-management of resources proves an effective alternative to wholesale privatization or state control, driven by these same slants of theory (Gutierrez et al. 2011; cf. Scott 1998).

By contrast, when US Treasury Secretary Paulson described escalating inequality as ‘simply an economic reality’ about which ‘it is neither fair nor useful to blame any political party’ (quoted in Bartels 2008, 296), markets are construed as pre-political facts of nature. This illusion relies on those strands of preformationist, essentialist biological thought (Keller 2005; Bastian and Haslam 2006) that ease the reduction of society to a population of atomized individuals, which together: buttress the individualistic basis of neoclassical economics and associated neoliberal politics; lower reputational costs of placing self-interest over the group; exacerbate tendencies to blame individuals when markets fail, leading to system under-correction (Jost et al. 2003); entrench fair market ideology and its unequal consequences (Blount 2000); expand the political subjectivity of possessive individualism; and divide rank and file with naturalised competition between individuals and their more or less praiseworthy genes (Clark 2007). Where this individualism by molecularisation prevails, a robust politics is less thinkable.

The relation between essentialist attributions to individual biology and tolerance for inequality is reflected in remarkable temporal coincidences within the US, now the most unequal among wealthy countries. Through most of the
mid-century “Great Compression” of US incomes (Goldin and Margo 1992) human genetics was ‘largely silent about those complex patterns such as insanity, criminality, or vagrantism that had so infatuated the eugenicists’ prior to this period of relative income equality (Sarkar 1998, 1-2). The post-1979 “Great Divergence” of US incomes (Krugman 2007) coincided with a return to immodest claims of genetic explanations of complex human behavioural patterns, ‘offered with a qualitatively new degree of precision or … the appearance of such precision’ (Sarkar 1998, 2). This shift followed Wilson’s (1994, 333) ‘exceptionally strong hereditarian position … at a time when nurture had seemingly won.’ Similar divergences characterize many other countries under the global wave of neoliberalisation (Harvey 2005), with far-reaching consequences and human costs (Sayer 2005; Wilkinson and Pickett 2009; Hedin et al. 2012).

Other political-economic processes played out during both transitions, most notably the New Deal and World War II at the beginning of the “Great Compression”. Its ending around 1980 came with pervasive and intensifying neoliberalisation, increasing frequency of financial crises, precipitous increase in global capital mobility, vast transfers and concentrations of wealth, income and power, and egalitarianism being increasingly identified with markets and individual property rights (Harvey 2010). Science’s relevance and participation are evident when for example US Republicans, appreciating the coalitional implications of essentialist thought, asked *The Bell Curve* co-author Charles Murray to help orient new congressional representatives (Beder 1997).
However it happens, trends in biological thought fit too easily into the nexus of the currently ascendant ideology of privatization (Bauman 2008b), legitimising by naturalising growing inequalities.

Though criticism of biological thought turned our focus on the geography of thought, this review converges with geographies of emotion and affect (Bondi 2005; Thrift 2005; Barnett 2008; Pile 2010) in underscoring the importance of participation, both in the scientific process and as a scientific construct (Demeritt 2008; kinpaisby 2008). The geography of emotional responses to the suite of social consequences of essentialist and deterministic lay thought would complement our analysis, much as inclusion of social psychological processes in this analysis supplements Thrift’s (2004) review. Though further attention to emotion is beyond the scope of this article, we caution against using spatial metaphors of emotions as “underlying” social processes (Oyama 2000a; Moore 2008) and overextending non-intentional modes of relating in non-representational theory and other relational ontologies.

**Participation in Sustainability**

The meanings of sustainability are deeply contested (Davison 2001). Its sloganeering rests on shaky ground (Worster 1993). Heavily exploited for marketing commodities and places, sustainability has become a discursive resource for enhancing profits and legitimising various forms of accumulation by dispossession – unsustainable from just about any social-ecological perspective. Green-washing is a reminder that the power of representation does
not necessarily correspond with its accuracy. Yet the Holocene extinction
(Eldridge 1998) associated with habitat destruction, massive production and
diffusion of toxins (Colborn et al. 1997) and the more publicised changes in
climate make it clear that neglecting sustainability is a sure pathway to ‘self-
organized extinction’ (Gowdy 2007, 27). Sustainability is ‘the art of keeping
the future navigable’ (Hägerstrand 2009, 187 our translation). There are many
alternative pathways within the unclear yet ultimate limits of sustainability
(Schellnhuber 1999).

Biological thought affects sustainability by informing how we farm, fish,
make sense of ourselves, and regard each other. With life forms given in
genetic packets and markets comprised of selfish, competing individuals the
natural, pre-political order, large-scale crop monoculture makes good sense.
Having lent efficiency to the accumulation of capital through labor intensive,
extractive, colonial plantations, monoculture has become synonymous with
genetically engineered or otherwise uniform crops in pursuit of the “common
good” of economic efficiency. Small, complex farms that minimize and
internalize ecological costs, protect genetic diversity, and sustain agrarian
communities make better sense when life is understood as constructed by
diffuse and reciprocal causation within locally adapted communities (Netting
1993). Much of the efficiency achieved on these farms remains off the books in
an economy that does not value social and ecological gains and losses. From
the vantage point of such farms, the expanding privatization of agricultural
biotechnology – what Nally (2011, 46) calls ‘accumulation by
molecularisation’ – is readily seen as meeting near term private interests more than long term public interests (McAfee 2003). The merits of sufficiency (Princen 2005) and the paradox of efficiency (Polimeni et al. 2007) are more acutely appreciated.

In 1883, Thomas Huxley argued that ‘Any tendency to overfishing will meet with its natural check in the diminution of the supply, … this check will always come into operation long before anything like permanent exhaustion has occurred’ (Roberts 2007, 272). Relying heavily on this enticing notion that nature regulates itself, including human activity by way of markets, the Royal Commission of Inquiry on trawling concluded that regulation was not necessary. Garstang (1900) urged greater caution, based on both better data and appreciation of the complexities and contingencies of development. Though Huxley’s optimistic projection did not hold up, his argument carried the day and much of the twentieth century as well. While specifics of this argument have fallen out of favour, the broader mind-set persists, complicating efforts to develop effective governance structures to manage global resources (Dietz et al. 2003) by rationalizing the commodification of nature under idealized market efficiency narratives (McAfee and Shapiro 2010).

Beyond feeding and provisioning ourselves, biological thought affects prospects for sustainability because the sense we make of ourselves shapes who we become, including our capacities for learning, cooperation and self-regulation. “Knowing” that intelligence is fixed inhibits learning (Blackwell et al. 2007). “Knowing” that personality attributes are inherited impels hasty
negative judgments about others, foreclosing opportunities for constructive encounter (Dweck 2000). “Knowing” that free will is illusory engenders cheating (Vohs and Schooler 2008) and aggression (Baumeister et al. 2009). And “knowing” that humans are economically selfish by nature favours policies that crowd out reciprocity and trust, inducing self-interested behaviour (Bowles 2008).

So also, “knowing” that political dispositions are a genetic endowment sells short political prospects. And “knowing” that metabolism is natural while intention remains a supernatural specter (Mayr 1961, 1982) hedges responsibility for our extended metabolism – energy consumption – compromising our capacity to regulate our own inventions. Deterministic formulations of ourselves, whether environmental (Radcliffe et al. 2010) or genetic, can become self-imposed constraints, just as interpretations of history are themselves forces of history (Plumb 1969; Fairclough et al. 2004).

For example, gene-centered thought favours the conclusion that ‘evolutionary analyses are unlikely to provide any cures for our environmental problems and … often only help us to see why it is difficult to achieve our goals’ (Penn and Mysterud 2007, 8). Against this, understanding human nature as constructed by context-dependent, distributed and reciprocal causation in communities of participants opens up alternatives by moderating the essentialist thought that tends to reflect existing power relations and coalitional intuitions (Boyer 2001), justifying by naturalising socially generated inequalities. Though the relationship is complex (Baland et al. 2007), inequality is associated with
wasteful behavior and weaker environmental policies because of: greater
discounting of the future by both poor and rich, for different reasons (Boyce
2002); easier externalizing of ecological costs by the more powerful through
ecologically unequal exchange; reduced trust degrading cooperation in
managing resources sustainably; intensified competition to consume without
corresponding increases in well-being; and motivating indiscriminate economic
growth to “lift the poor”, accommodating instead of confronting excessive
inequality (Wilkinson and Pickett 2009). These ecological consequences of
inequality would be mitigated by broader appreciation of developmental
systems in understanding the “other”, to the extent this advantages political
alternatives that ‘institutionalize the commonality of fate’ over those
‘expressing and promoting the diversity of fate’ (Bauman 1994, 24).

The deeper our appreciation of the essential openness of human nature, the
better we face each “other” and, in turn, the better we are able to discern our
shared fates and cooperate in exercising our common right to and
responsibilities toward place. The right to place12 is a modern expression of
human sociality with deep roots in the vigilant egalitarian cultural practices of
our ancestors. Geo-history is agonistic (Mouffe 2005), consisting of ‘struggles
for power over the entry of entities and events into space and time’
(Hägerstrand 1986, 43 our translation). How such struggles contingently
play out facilitates or hinders sustainability by shaping peoples’ connections to
the land and their place.
While history and contingency are recognized in evolutionary theory alongside genes and environments, disproportionate attention is given to chance or seemingly random but physically determined events such as asteroid strikes (Briggs and Fortey 2005). However prudent scientific wariness of intention, choices forced on or seized by organisms are a distinct aspect of history, within the ineluctable indeterminacy that also shapes the course of evolution. Volitional activity is a force of nature, however conditioned on conservative genes, varied environments and untold physiological and psychosocial developmental processes. Recognizing animals’ choices – (mis)informed, impulsive, moral, or otherwise – among the contingencies that shape evolutionary change includes human activity more fully within evolutionary thought, exposing conceptual and moral blind spots that bear on sustainability.

From this viewpoint purpose has a place in evolution, only by way of the life histories of organisms. With sighted animals as participants, natural selection is not completely blind, consequences of which are ‘grandiose only in the aggregate, and in retrospect; up close and confined in time, they are ordinary’ (Margulis and Sagan 1995, 224). Within human biological development and evolution there is a margin of reach which perspective may render either trifling or abundant.

Keeping the future navigable demands reaching with the best of our egalitarian heritage and ideals, today’s economic fairness being a foundation for tomorrow’s intergenerational environmental justice (Wallerstein 2007). Some
innovations that succeeded by the standards of a growth economy may be counterproductive by the standards of fulfillment within a sustainable economy. Given the possible down-sides of this transition, success depends crucially on social dimensions of innovation (Kallis 2011). A more accurately open image of human nature in evolutionary and folk biologies, in scientific and lay theories, would itself be a social innovation helping us attend to alternatives, open up sites of transformative space (Langley and Mellor 2002) and develop the social capital to confront the constraints and uncertainties, and identify the opportunities, of transitioning toward sustainable living (Harvey 2000; Pretty 2003).

Conclusion

Sailing the Mediterranean, Ulysses would have seen ‘abundant monk seal, loggerhead turtles and porpoises’ (Roberts 2007, 375). Not today. The ‘juggernaut, improvement’ has rolled over habitation (Polanyi 2001, 191), fraying the weave of life. Meanwhile, the modern synthesis of genetics with evolution has been more committed to simpling man into causal theories than stretching theories to include the purposeful behaviour of man and other animals. Human inventiveness, regarded by the modern synthesis as the evolutionary consequence of climate variability (Potts 1996), has begun causing climate variability.

To take full account of insects, ostriches or humanity, biology must recognize the participation of animals in their own development and evolution.
More is at stake than a robust theory of evolution. Because sustainability is as much about managing ourselves and each other as managing resources, the sense we make of ourselves can limit or leverage sustainability achievements. Biology’s neglect of consistency regarding intention and participation, drolly conceded by Haldane and seeming wise when Polanyi diagnosed it over a half century ago, is no longer wise.

Mayr’s (1959, 13) claim that ‘the very survival of man on this globe may depend on a correct understanding of the evolutionary forces and their application to man’ must be understood in light of Waddington’s (1960; cf. Lewontin 2000) view of man and other animals as both objects, to which evolutionary forces apply, and subjects, themselves evolutionary forces. Recognising our ancestors and ourselves as participants in evolution is vital to sustainability aspirations, as it animates healthful consideration of the responsibilities that come with our deft inventiveness. Theory that hollows out volition diminishes responsibility.

Awkwardly appended declarations of responsibility are poor substitutes for baking it into the theoretical cake. Simpson’s (1960; 973) is one of many such declarations in which it is unclear how people’s ‘sense of responsibility’ ensues from an evolutionary process in which ‘the mechanism of orientation, the non-random element’ is ‘blind’ natural selection. Vermeij (2010, 250) emphasizes the continuity of animal and human intentionality before inexplicably negating its evolutionary relevance, vitiating responsibility, by asserting that the ‘details of time, place and player’ of selection ‘reside fully in the realm of inscrutable
chance.’ Having at once trivialized participation and belittled our ancestors, Vermeij’s hopes for mankind emphasize philanthropy by naturally occurring superrich, not social and economic fairness and the broader participation in solutions this entails. Some biologists have wanted to own the bicycle of intention and responsibility without riding it (cf. Gould 2002, 256).

Bearing hallmark features of a major evolutionary transition (Wilson and Wilson 2007), human eusociality – good sociality – was achieved by the conscious participation of our ancestors in recognizing shared fates and cultivating responsibility to each other, in societies that cannot be coherently dissolved into populations of atomized individuals (Fracchia and Lewontin 1999). Having played some part in this transition, between-group conflict need not become a self-fulfilling, theoretically blinkered expectation distracting us from seeing clearly both our increasingly shared ecological fates and our capacities for cooperation between groups in achieving sustainability goals.

Recent work of biological and social scientists reveals misplaced concreteness (Whitehead 1978) in thinking about human nature as an unfounded constraint on human adaptability. Among other ways scientists stand to help manage anthropogenic global changes (Travis and Futuyma 1993; ICSU 2010), we appeal for affirming in theory and in the public square an image of human nature ‘that presents people as able to make their history … [and] as obliged to do so’ (Bock 1994, 116). Achieving the collective self-regulation sustainability requires may depend on correcting slanted reasoning about ourselves.
Notes

1 Huxley (1942) coined the term “modern synthesis” to describe the synthesis of evolutionary theory with other areas of biology, prominently including the young science of genetics while marginalizing developmental biology. Simpson (1960) and Mayr (1982) expressed similar views on Darwin’s position, consigning behavior, except as it is controlled by genetically given “programs”, to the realms of superstition and decisively refuted “soft inheritance”.

2 This conflation is echoed in EO Wilson’s (2005, 30) conclusion that evolution, lacking external guidance, is therefore ‘blind’, and in DS Wilson’s (2007, 92) pejorative ‘secular creationists’, describing social scientists who reject facile overextensions of “genetic evolution” as unfounded and socially consequential. Wilson’s epithet confuses recognition of emergent processes in development with belief in externally directed evolution.

3 The brain has been similarly privileged vis-à-vis behaviour (Bakker 1984).

4 See Plotkin (1988, 2, 143) on the occasional nods to behaviour.

5 Sadness evoked by this “scientificality” toward life registers an intuitive grasp of its failings. On the silencing of such emotion, see Anderson and Smith (2001).

6 Because of the advantages of plasticity, genetic assimilation of behavioral adaptations may have more costs than benefits (cf. Mayr 1963, 612).

7 A secular continuation of inherited power in monarchies based on divine essentialism.
The nature of these social consequences is ‘not strongly influenced by the efforts of the core scientists. They merely catalyze what is already happening …’ (Strand 2000, 453).

We make no exceptions for ourselves and warn instead against indifference toward or claims of immunity from such influences.

Clark places behaviour in a causal evolutionary role, but his attributions of “genetic” class differences rest on poor biological modeling – application to a non-isolated subpopulation within a short time frame.

Garstang (1922) later argued that modifications in ontogeny can influence the course of evolution, placing him in the company of Gullick, Waddington, Ewar and Piaget, noted above.

With right to place we refer to the right to the city (Harvey 2003; Mitchell 2003; Purcell 2008) in its broadest sense of citizenship, beyond the city limits (cf. Harvey 2000, 248-252).

Orians (2007) commends Monod’s (1971, 113) assertion of chance’s universal and exclusive role in evolutionary innovation. Darwinian in neither substance nor style, Monod’s pronouncement engenders more than it explains the resistance to evolutionary thought he dismisses as ‘instinctive’.
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