Evolved Open-Endedness in Cultural Evolution: A New Dimension in Open-Ended Evolution Research

Abstract  The goal of Artificial Life research, as articulated by Chris Langton, is “to contribute to theoretical biology by locating life-as-we-know-it within the larger picture of life-as-it-could-be.” The study and pursuit of open-ended evolution in artificial evolutionary systems exemplify this goal. However, open-ended evolution research is hampered by two fundamental issues: the struggle to replicate open-endedness in an artificial evolutionary system and our assumption that we only have one system (genetic evolution) from which to draw inspiration. We argue not only that cultural evolution should be seen as another real-world example of an open-ended evolutionary system but that the unique qualities seen in cultural evolution provide us with a new perspective from which we can assess the fundamental properties of, and ask new questions about, open-ended evolutionary systems, especially with regard to evolved open-endedness and transitions from bounded to unbounded evolution. Here we provide an overview of culture as an evolutionary system, highlight the interesting case of human cultural evolution as an open-ended evolutionary system, and contextualize cultural evolution by developing a new framework of (evolved) open-ended evolution. We go on to provide a set of new questions that can be asked once we consider cultural evolution within the framework of open-ended evolution and introduce new insights that we may be able to gain about evolved open-endedness as a result of asking these questions.

Keywords  Cultural evolution, open-ended evolution, evolved open-endedness, zone of latent solutions, cumulative culture

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1 Introduction

Genetic evolution appears to be open-ended. Taking advantage of environmental regularities, gene expression and regulation can generate a potentially infinite number of traits and trait variations. Such evolutionary open-endedness has been characterized by a constellation of overlapping features yet can generally be understood as the ability of an evolutionary system to produce a continuous stream of novel units (Taylor et al., 2016). For those trying to create and understand open-ended evolutionary systems, the goal is to understand the underlying principles and dynamics of evolutionary systems in general. Such understanding is based on knowledge of the best explored and understood open-ended evolutionary system: genetic evolution. But it also can, and should, draw on the development of artificial evolutionary systems that explore the principles of life-as-it-could-be (Langton, 1989). Such artificial evolutionary systems depart from the particular implementation and substrate features of Darwinian genetic evolution while still meeting the general requirements of an evolving system. The interaction between the two systems can be consilient. Darwinian genetic evolution provides a source of valuable ideas and inspiration as well as justification for the designs of artificial systems. Despite this positive interplay, having only one concrete instance of an open-ended system is a problem. Such sparse epistemological situations can limit abilities to discern alternate possibilities, detect generalizable features, and develop robust theories and models.

It is increasingly being recognized, however, that there is another evolutionary system from which one can find inspiration: cultural evolution (Bedau, 2013, 2019; Bedau et al., 2019; Borg & Powers, 2021; Marriott et al., 2018). Minimally characterized, culture is information transmitted through mechanisms of social learning (Boyd & Richerson, 1985; Cultural Evolution Society, 2021; Whiten et al., 2022). And while this minimal characterization leaves out many distinctive features of human and nonhuman cultural groups (for instance, that species differ in the types of information they can transmit) (Whiten et al., 2022), and leaves open precisely how “social learning” should be construed (Lewens, 2015), its abstractness makes it exceptionally useful for designing models of cultural change and describing general evolutionary dynamics. On this characterization, cultural evolution is the change in frequency, or, of special interest here, the form of cultural information over time, where these changes are at least in part influenced by social learning (Neadle et al., 2017). Although cultural evolution is often described as being analogous to genetic evolution (Cavalli-Sforza & Feldman, 1981), there are clear differences in the way culture is inherited: (a) Although genetic evolution typically relies on two (sometimes one) parents, there are potentially unlimited numbers of cultural “parents”; (b) although genetic transmission is almost exclusively vertical, from parent to child, cultural transmission can involve substantial amounts of horizontal or oblique transmission; and (c) although genetic changes generally occur between generations, cultural change generally occurs within generations (Mesoudi, 2011; Mesoudi et al., 2006). Though these features distinguish cultural from genetic change, these do not imply that cultural inheritance is in any sense less (or not) “evolutionary”—only that its dynamics frequently differ.

Over the past 40 years, there has been increasing recognition that culture and cultural evolution exist within nonhuman animal populations (most prominently in birds and mammals) (Whiten, 2019, 2021a, 2021b) and that culture not only exists as a result of genetic adaptation but also plays an important coevolutionary role in guiding genetic evolution (Uchiyama et al., 2021; Whitehead et al., 2019). This coevolutionary relationship between genes, culture, and the environment is sometimes known as triple inheritance (Laland et al., 2000). Nonetheless, though many animal species exhibit culture, human cultural evolution appears both quantitatively and qualitatively distinct. Several dividing lines between human and animal cultures have been proposed, but the most prominent of recent formulations holds that human culture is distinctive by virtue of its cumulative nature, with human culture accumulating modifications over time and with these modifications building on one another (Tomasello, 1999). However, as more observations of cultural evolution in other species have been
made, it has become increasingly apparent that cumulative cultural evolution is not unique to hu-
man culture (Mesoudi & Thornton, 2018). This raises the following question: What, if anything,
is unique about human cultural evolution?

We think issues about the distinctiveness of human culture and the nature of open-ended evolu-
tion are overlapping—and that explorations of the two will be mutually illuminating, with potential
downstream consequences for Artificial Life. Here we situate cultural evolution within a broader
framework of open-ended evolution and argue the following:

1. Culture is an evolving system, coevolving alongside genetic evolution.
2. Within cultural species is a range of “types” of cultural evolutionary patterns: cumulative
   and noncumulative, tall and wide, unbounded and bounded.
3. Recognizing these “types” of cultural evolution allows Artificial Life researchers to better
   understand evolutionary dynamics and provides new perspectives from which to explore
   open-ended evolution.
4. Only humans demonstrate open-ended cultural evolution, and human cultural evolution
   has transitioned from a bounded to an unbounded evolutionary system in recent
   evolutionary history, thus providing a second instance of “evolved open-endedness.”
5. Existing Artificial Life methods can be fruitfully applied to the study of cultural evolution.

To develop these points, we outline a number of core concepts from the wider study of cultural
evolution. We then analyze “open-ended evolution” and explore how such analyses might improve
our understanding of evolutionary dynamics and the emergence of evolved open-ended evolution-
ary systems. Table 1 provides definitions of key terms.

2 Cultural Evolution

What is culture, and how does it evolve? As suggested, culture can be minimally defined as the
transmission of information—traits—through mechanisms of social learning (Boyd & Richerson,
1985). This minimal and abstract characterization of culture permits information and traits to be read
in an encompassing way, to include a wide variety of techniques, technologies, and behaviors. Ex-
amples of such traits include the extractive foraging techniques among chimpanzees (Sanz et al.,
2010) or methods for lighting a fire (MacDonald et al., 2021). Culture may also incorporate behav-
iors with communicative effects, such as warning calls (Griffin, 2004), birdsong, or language (Janik
& Slater, 2000). The definition also incorporates population-level conventions among conspecifics
for greeting and leave-taking (Baehren, 2022; Duranti, 1997) as well as normative behaviors, such as
styles of dress or decoration (Baehren, 2022; Richerson & Henrich, 2009). Again, the key is that the
acquisition of these behavioral traits or beliefs is and must be influenced by social learning—when
it is not, the traits are not cultural.

2.1 Does Culture Evolve?

An evolutionary process does not require a particular kind of physical instantiation or biological
substrate. Whereas familiar processes of biological evolution are grounded mainly in the ma-
ipulation and modification of genes, cultural evolution (and evolution more generally) is under
no such obligation. Consider Dennett’s (1996) conception of evolution as being both algorithm-
ic and substrate neutral. Evolution is algorithmic in the sense that if certain conditions are met,
a certain sort of outcome is necessarily produced (Dennett, 1996). Where there is reproduction
with variation under selection at a population level, the frequency of adaptive outcomes is increased
in the population over time. In cultural evolution, adaptive may refer to the cultural trait and the suc-
cess the trait has in spreading from mind to mind (Rosenberg, 2017), or it may refer to the effects
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Table 1. Definitions of key terms.

| Term                     | Definition                                                                 | Reference(s)                                             |
|--------------------------|---------------------------------------------------------------------------|----------------------------------------------------------|
| Culture                  | Information transmitted through mechanisms of social learning            | Boyd and Richerson (1985), Cultural Evolution Society (2021), Whiten et al. (2022) |
| Cultural evolution       | The change in frequency or the form of cultural traits over time, where these changes are at least in part influenced by social learning | Neadle et al. (2017)                                     |
| Open-ended evolution     | An evolutionary process that is capable of producing a continuous stream of new adaptive novel units, with no a priori limitations on the generation of such novelty | Gabora and Steel (2017), Taylor et al. (2016)             |
| Cumulative culture       | A process whereby a culturally transmitted trait accumulates modifications over time with a ratchet-like effect | Boyd and Richerson (1985), Tomasello (1999)              |
| Unbounded evolution      | A continuous demonstration of new adaptive novelty and/or the ongoing growth in trait diversity. Term used interchangeably with open-ended evolution, but often used to contrast with bounded evolution | Bedau et al. (1998), Channon (2006)                       |
| Evolved open-endedness   | Open-endedness as the outcome of an evolutionary process as opposed to an assumed pre-condition | Pattee and Sayama (2019)                                 |
| Wide evolution           | A characterization of the disparity of traits and traditions; increased through processes of recombination, innovation, or the exploration of previously underappreciated affordances | Buskell (2022), Derex (2022)                             |
| Tall evolution           | A characterization of the typical length (measured in relevant changes generated through cumulative evolution) of independent trait traditions |                                                        |

the trait has on its bearers’ (adaptive) behavior. Importantly, these adaptations may only emerge out of complex coevolutionary interactions between culture and biology (Henrich & McElreath, 2007).

Ultimately, the target of reproduction is the informational content carried by some vehicle—whether this vehicle is expressed behavior, an artifact, or the instructions of a written account (though it is of course the case that the vehicle can itself have “fitness”). Artificial Life has often equated such a characterization with the idea of a “meme” (Bedau, 2013; Bull et al., 2000; Bullinaria, 2010): a discrete, particulate unit of information that is copied intact between brains, analogous to the way that genes are copied between parents and offspring (Dawkins, 1976). Cultural evolution, however, does not require the process of reproduction and cultural inheritance to be understood in terms of strict copying. Though the literature on this point is vast, Rosenberg (2017) provides a clear summary of the arguments:

1. Replication in biology has not always involved high-fidelity replicators—the “major transitions in evolution” literature explains how evolution itself has gradually generated higher-fidelity transmission processes. Though the first replicating molecules were not DNA, nor did they have accurate copying mechanisms, increases in fidelity are evolutionary
achievements that have been and can be selected for over time (Maynard Smith & Szathmáry, 1995).

2. Even in genetic evolution, a single gene can rarely be equated with a single trait, the vast majority of biological traits result from complex interactions between the proteins expressed and regulated by many genes, so why should one demand in cultural evolution that a trait is the product of one discrete meme?

3. Many features of human institutions are adapted to preserve and proliferate cultural traits even under low individual copying fidelity. Variation is introduced in the form of the (re)combination of existing traits, innovation of new traits by individuals (which may involve rational thought), or copying error (loosely analogous to mutation in genetic evolution). Meanwhile, selection may occur in multiple ways. This includes biological selection, that is, the effect that cultural traits have on biological fitness (for instance, being led to believe that something is safe to eat when it is not).

If we accept that evolution is algorithmic (i.e., it follows a series of processes to produce a certain outcome: selection + reproduction + variation = evolution), it follows that we are not bound to particular features of biological processes (e.g., sexual reproduction), nor are we bound to a specific substrate (e.g., DNA). Though Dennett’s conception of cultural evolution has changed over the years (e.g., Dennett, 2017)—perhaps in response to critics (Uhlíř & Stella, 2012)—the fundamental insight we take from him still applies: The idea of an algorithmic process makes it all the more powerful, because the substrate neutrality it thereby possesses permits us to consider its application to just about anything (Dennett, 1996). This is, of course, true: That one can create an evolutionary process within a computer is evidence that the process itself need not be strictly biological but merely algorithmic (Lehman et al., 2020).

2.2 Interdependent Evolutionary Systems

Cultural evolution is deeply intertwined with biological evolution. Although these evolutionary processes and their products can generate complicated coevolutionary feedback loops, each evolutionary system can be understood, studied, and modeled separately (Boyd & Richerson, 1985; Mesoudi, 2011). For instance, as we suggest in more detail later, premodern hominin cultural evolution contributed to biological fitness in the form of ecological knowledge and technological production. Nonetheless, over time, cultural evolution has become increasingly unmoored from genetic fitness effects, producing a wide range of behavioral, social, and technological change (Henrich, 2015). The reason for both the intimacy and the relative independence of the two systems should be evident: The substrate of culture is biological—the brain.

Culture is bound to a biological substrate, but a substrate that is different from the classical understanding of genetic evolution in which traits are encoded (directly or indirectly) by genes. Gene expression may produce brains, and (some) brains may acquire culture, but one cannot skip the middle step and claim that genes produce culture. Although humans may be biologically prepared to acquire language (Fitch, 2011), they are not biologically determined to learn English, Farsi, or Korean. Clearly accessibility and exposure to certain kinds of inputs—the presence of English, Farsi, or Korean language cues—determine what language any given human ultimately produces. Or put another way, the acquisition, production, and transmission of language are largely influenced by social learning. Thus one cannot simply claim that the process of cultural evolution is independent from biology. Biological and cultural evolution are interdependent.

The idea that cultural species, and particularly cumulatively cultural species like *Homo sapiens*, have two interdependent systems of inheritance has been labeled *dual inheritance* (triple inheritance if the environment is also included; Laland et al., 2000). On this account, human offspring inherit a genotype from their parents through sexual reproduction, and they inherit a body of cultural information over the course of their postnatal lives via processes of social learning (Henrich
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& McElreath, 2007)—processes that themselves may be culturally evolved tools (Heyes, 2018). Just as one’s genotype has been dictated by a history of selection pressures acting on genetic variation, one’s cultural inheritance is similarly shaped by selective pressures and the variation introduced through the innovation, recombination, and error involved in social learning. Thus, in the same way that certain phenotypic features are adaptations—increasing the biological fitness of individuals—elements of culture may also be adaptations. Consider food taboos present in Fijian society that apply exclusively to pregnant women (Henrich & Henrich, 2010; McKerracher et al., 2016). Despite the causal opacity of the underlying process, these taboos protect women from miscarriage. Alternatively, consider the ritualized process of cassava production. Again, despite the causal opacity of the underlying process, populations have developed practices that remove toxic cyanogenic elements that would have long-term health consequences if regularly consumed (Banea et al., 1992; Bradbury & Denton, 2011; Cardoso et al., 2005; McKerracher et al., 2016). Of course, it can also be adaptive to acquire cultural elements idiosyncratic to local cultures. Regardless of whether the practice of female or male circumcision has biological benefits, within a circumcising culture, it can be adaptive to demonstrate commitment to the group by engaging in such a costly signal. This can ensure inclusion among and support by the group and prevent ostracism (Howard & Gibson, 2017; Sosis, 2004), thus enhancing reproductive outcomes.

Cultural organisms inherit not only genes and cultural information but also an environment, that is, a habitat that has been selected, modified, and partly created by their ancestors. All organisms change their habitats through their actions—of which spider webs, termite mounds, or human-made earthworks are just a few notable examples—with more or less transitory effects. Such organism-modified environments are evolutionarily relevant insofar as they modify selection pressures or transmission opportunities—what the evolutionary literature calls niche construction (Laland et al., 2000). Systematic and long-lasting modifications, such as beaver dam building or human agriculture, can have profound effects on both biological and cultural evolutionary processes of the species producing these modifications and on others in the habitat.

Although niche construction is not uniquely human, humans are distinctive in that most of their niche construction activities are cultural (e.g., making dams, fences, schools, roads, clothes). Over evolutionary time, the hominin lineage has created a cultural niche that has not only affected their biological and cultural evolution by creating new selection pressures but increasingly become crucial for their survival (Laland & O’Brien, 2011; Uchiyama et al., 2021). For example, the use of fire and cooking may have facilitated selection for larger brains alongside smaller guts and jaws. Lacking fire or cooking, hominins would have been poorly adapted to their environments (Aiello & Wheeler, 1995). The second inheritance system, culture, can thus indirectly affect the first, genes, through niche construction. Genes and culture have coevolved: Cultural activities, such as tool use and tool making, have generated selection pressures for social tolerance and cognitive skills like social learning, attention, working memory, and language, which in turn have opened up ever greater capacities for cultural innovation, social learning, and large-scale cooperation (Henrich, 2015), creating the biological and cultural conditions for the emergence of open-ended cultural evolution.

Cultural evolution is often faster than genetic evolution: A cultural variant can emerge and recombine quickly and repeatedly within the lifetime of its carrier and can die independently of the death of the individual (Boyd et al., 2013). Alongside the speed of cultural evolution, humans’ capacity for planning and foresight suggests that many human adaptations are cultural or have cultural origins (Uchiyama et al., 2021). Thus cultural evolution can not only produce solutions to (ecological) problems but also create new opportunities and niches that cultural evolution can exploit—an autocatalytic process, resulting in the emergence of open-ended, cumulative culture.

3 Open-Ended Cultural Evolution

As noted in the Introduction, open-ended evolution is an umbrella term for a constellation of features associated with evolutionary change. These include the ongoing generation of novelties, adaptations,
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and evolutionarily salient entities (Taylor et al., 2016). For simplicity, we hold that an evolutionary system can generate open-ended evolutionary change if it is able to produce a continuous stream of novel units (evolutionary individuals, traits) with no a priori limits to the generation of such novelties (Gabora & Steel, 2017; Taylor et al., 2016). As several commentators have noted (Bedau, 2019; Bedau et al., 2019; Pattee & Sayama, 2019; Tennie et al., 2018), human cultural evolution appears to be just such an open-ended evolutionary system.

More recently, cultural evolution researchers have used the term open-ended to describe what is unique about human culture (Tennie et al., 2018). This acknowledges that human culture frequently involves processes of cumulative cultural evolution—processes that generate traits (e.g., behaviors, beliefs) that build on previous traits, perhaps also making them more complex, efficient, and adaptive. But calling human culture open-ended is also meant to suggest that cultural solutions to problems do not need to be stuck at local optima but can break free and further improve, for instance, by harnessing new affordances (Arthur, 2009; Derex, 2022). Focusing on this putative “uniqueness” of human culture, researchers have identified important transitions, cognitive capacities, and patterns of cultural evolution as hominins have evolved over the past 8 million years.

In sections 3.1–3.3, we make distinctions between patterns of cultural evolutionary change: between cumulative and noncumulative cultural traditions, between “building up” tall traditions and “building out” wide repertoires of traditions, and between bounded and unbounded evolution. These patterns capture important differences in cultural evolutionary dynamics. Though these patterns are distinct, they likely overlap in many instances. In section 3.4, we turn to considering how these distinct kinds of evolutionary patterns help characterize and explain the evolution of open-ended cultural evolution in hominins.

In focusing on distinct kinds of evolutionary patterns, and tracing these patterns back to concrete changes in selection pressures, cognitive mechanisms, and social arrangements, the approach taken here differs from recent attempts at describing hallmarks of open-ended evolution (Taylor et al., 2016). Hallmarks are signals, such that if one were to encounter them, it would be good evidence that the evolutionary system is capable of open-ended evolution. By contrast, our approach distinguishes patterns that are associated with processes supporting cultural evolutionary change. These processes are critical to, but not necessarily sufficient for, open-ended evolution—and thus are poor candidates for a hallmark approach. Nonetheless, distinguishing these processes helps to identify those important for evolving open-endedness, as well as how the interaction between such processes may be important to the eventual emergence of a system supporting full-blown, open-ended evolution.

3.1 Cumulative Versus Noncumulative

A key distinction drawn by cultural evolution researchers is that between cumulative and noncumulative culture. As many researchers see it, cumulative culture is central to explaining how human beings could have developed the sophisticated technical toolkits that allowed them to survive and thrive across varying—and sometimes extreme—ecologies (Grove, 2011; Henrich, 2015; Potts, 2013; Richerson & Boyd, 2005). On the basis of extensive human and nonhuman experiments, and a number of computational and mathematical models, Mesoudi and Thornton (2018) have suggested “core” criteria that cultural evolutionary processes would have to satisfy to be classified as cumulative: (a) a change in behavior, followed by (b) transfer of the modified or novel trait via social learning, where (c) the learned trait results in an “improvement” in performance/fitness (cultural or genetic), with (d) the previous steps repeated in a manner that results in (sequential) modification and improvement over time. However, we follow recent work in denying that “improvement” over time is a necessary feature of cumulative cultural evolution and instead favor a minimal formulation that sheds this requirement (Buskell & Tennie, in press).

On this minimal formulation, cumulative culture is simply the modification to, and retention of, socially transmitted cultural traits (Buskell & Tennie, in press). What we have called processes of cumulative culture in the preceding discussion are whatever cognitive and social capacities are
sufficient to bring about trait modification and retention over time. But these processes generate patterns in the evolutionary record. Because cumulative culture involves retained modifications, they have histories—and can be considered “traditions.” The histories of such traditions can, at least in principle, be reconstructed as sequences of step-by-step changes (akin to what Calcott, 2009, calls “lineage explanations”). This minimal formulation better aligns cumulative culture with evolutionary theory such that cumulative changes can generate not only adaptive traditions but also neutral and maladaptive ones (Buskell & Tennie, in press).

Contrasting with cumulative culture is noncumulative cultural evolution. The latter is a process of cultural change that does not retain modifications for one reason or another. This might be because there is no retention of past behavior, no introduction of modifications, or no social learning sophisticated enough to pick up on relevant modifications. These situations might occur if individuals can only innovate new traits, can cycle through a set of traits, or do not learn from one another. In these cases, histories of modifications will be nonexistent, uninformative, or based in noncultural inheritance systems.

3.2 Tall Versus Wide Evolution

Recent work has built on analyses of cumulative culture to distinguish further cultural evolutionary patterns that had been unhelpfully lumped together. This work distinguishes between patterns involving an increasing stock of cultural traditions (cultural disparity) and important aspects of cumulative cultural traditions (e.g., increases in adaptiveness, efficacy, or complexity) (Buskell, 2018, 2022). This and other work (Dean et al., 2014; Tennie et al., 2009) points to a helpful distinction between cultural evolutionary patterns: between “building upon” traditions and “building out” to generate new traditions—or just tall versus wide evolution.

Figure 1 provides a visual example of both tall and wide evolution, with tall evolution displaying a series of path-dependent adaptations within a single tradition. Each step in the sequence could only have occurred if the previous evolutionary steps had already arisen. While tall traditions need not be path-dependent—for instance, if evolution is highly constrained—it is a common assumption that evolutionary change is so, and we emphasize path dependency here. Wide evolution, by contrast, is about the novel instancing of new traits. Paradigmatically, this involves the innovation of completely new traditions that need not follow any a priori sequence. Of course, some new traditions may arise only through path-dependent cumulative evolution and recombination—but we put those instances to the side in this illustration. Thus, in Figure 1, one could rearrange the wide axis (because new traditions need not appear in any sequence) but not the tall (because each step is strongly determined by the prior step).

By way of example, let us consider some kind of adaptive problem that may have multiple starting points that are either equiprobable (equally likely to occur in the same environment) or equally efficient at solving the problem but the product of different affordances due to different environments. This might include capturing fish or preserving meat, or it could include producing housing or clothing, refining ore into more valuable products, or skinning cats. The specifics matter less than the principle being illustrated. Along the x axis, we have multiple starting points. Let us consider the fishing example. One equiprobable starting point may be to wait in the shallows and bash a fish with a rock or a stick as it swims by. Another example may be to wait at a certain point on the beach that, at low tide, forms a natural pool from which fish cannot escape. Another yet may involve poisoning the water with certain plant foliage. It can be true that these starting points are (a) all equally likely due to the affordances of the environment or (b) all arrived at by different groups who live in different environments with different affordances. Whether either is true in any given situation is less important than accepting that these are (some of) the starting points for acquiring fish.

Tall evolution may involve the rock culture innovating upon the basic rock-bashing behavior, perhaps first by throwing the rock, then by tying a fiber to the rock before throwing it (so as to recover the rock more quickly through a pulling motion), and then by using multiple rock–fiber devices to expand the range of striking. Later innovations might eschew the bashing or throwing
Figure 1. Our conception of tall and wide evolution. Full details are available in the appendix. Each box represents some kind of technology or cultural practice. Each pixel within each box represents a piece of discrete (but arbitrary) information. The eight squares in row 1 (the bottom row) were generated by asking each of the pixels to become black or white at a probability of 0.5. Thus all initial configurations of aggregate information are equiprobable. Thereafter one of eight arbitrary rules was applied over 10 iterations. These rules were not grounded but represent changes of “information” within the aggregate or that introduces structure (such as symmetry) in the aggregate. As can be seen, as the aggregate information cumulatively changes over time, it becomes more complex and structured and increasingly dissimilar from other traditions. Each column is independent of all other columns, and “movement” along the wide axis is not possible without violating the cumulative principle of tall evolution.

motion for connecting the fibers together to make a rake or net. Further innovating might then improve the netting technology or the casting technique, and so on.

Meanwhile, the stick culture may innovate upon the bashing motion by innovating a sharp point—now preferring to pierce rather than to bash. Later innovations might make spears much longer than would ever be practical for bashing, so as to stand farther away from the fish without
scaring them. Then, perhaps, innovations might lead to a stone tip for the spear and, later still, a spear-throwing device like an atlatl or woomera to bring down larger prey, and so on.

It may be the case in the first instance that stick bashing and rock bashing are equally (in)efficient and that—assuming an abundance of rocks and sticks—one individual or one culture may switch between techniques with little cost. However, once groups begin to innovate upon their starting point, horizontal movement comes with greater cost and relies on different principles. A raking technique does not beget a spear thrower, and vice versa. After tall evolution has progressed beyond a certain point, horizontal movement cannot be integrated or combined with the existing “advanced” approach, and switching comes at a greater cost to the individual or culture.

Another case study is the tool use of chimpanzees. Chimpanzees are capable of spontaneously innovating tools given available resources, such as using blades of grass for termite fishing, sticks for obtaining out-of-reach objects, or branches for scooping algae out of water (Bandini & Tennie, 2017; Boesch & Boesch, 1990; Sanz et al., 2010). Each and all of these innovations can exist within a population of individuals, but the existence of one need not depend on the existence of any other. Theoretically, any of these innovations can be selected for and spread within the population independently of the others. This is wide evolution. Nonetheless, modifications could be added to these innovations—introducing an anvil prop to nut cracking, chewing and stripping the grass to produce ant-catching bristles—that put them on the vertical road to becoming a tall cultural evolutionary tradition.

This example also points to an important corollary of the distinction between tall and wide evolution. The capacities underlying each plausibly come apart. This seems clear when one looks at hominin evolution, where early capacities for social learning led to wide knowledge bases of disparate ecological traditions prior to building up any particular tradition into more complex forms (Buskell & Tennie, in press; Sterelny, 2021) (more on this later).

More generally, we want to resist identifying tall or wide evolution patterns as hallmarks of open-ended evolution. It is an open question of how tall (or short), wide (or narrow), evolutionary patterns relate to open-ended evolution, as well as the transition to open-ended evolution. As examples herein suggest, capacities that support tall and wide evolutionary patterns likely existed well before ecological and evolutionary circumstances permitted their expression. And indeed, open-endedness most likely emerged from the gradual accumulation of new traditions; their elaboration into tall, path-dependent traditions; and their recombination and exaptation into bushy, wide, and novel traditions—we can see this visually in the patent record genealogies produced by Bedau (2013, 2019), with both the gradual accumulation of new patent traditions and long sequences of traditions building up being easy to identify. There’s no reason to take either tall or wide evolution as a hallmark of open-ended evolution; ultimately, they just describe the patterns of change that underpin the emergence of open-ended evolutionary process. We suspect that both are necessary for open-ended evolution to emerge, but only further empirical analysis of the patterns of change found in open-ended evolutionary systems will allow us to ascertain whether common patterns exist or whether a multitude of patterns can ultimately underpin open-endedness. We think it unsurprising that capacities underlying both tall and wide evolution should be needed. Both formal modeling (Enquist et al., 2010; Kolodny et al., 2015; Winters, 2020) and cultural evolutionary theory (Buskell et al., 2019; Charbonneau, 2016; Richerson & Boyd, 2005) emphasize the role of cultural recombination as a potent force in generating new innovations: This occurs when distinct cultural traditions (or their constituent elements) are combined, and potentially exapted (Mesoudi & Thornton, 2018), to generate new traits. We expand on this line of thinking later and go on to ask whether these variations in the progression of evolution (tall, wide, recombinative, exapted) are detectable within the “AI life test” introduced by Bedau et al. (1998; see also Channon, 2001, 2003, 2006).

### 3.3 Unbounded Versus Bounded Evolution

A conceptually distinct and contrasting set of evolutionary patterns is that between bounded and unbounded evolution. Bounded evolution occurs when abilities for transmission, retention, or the
production of modifications are limited or absent. This leads to evolutionary exploration of a parochial, bounded space of traits. Unbounded evolution, by contrast, occurs when the abilities for transmission, retention, or the production of modifications are present and when the environment facilitates evolutionary exploration. This might occur, for instance, when the environment is rich in natural resources that can be exploited in technological production (Derex, 2022).

To get a grasp on this distinction, it is useful to look at a domain in cultural evolutionary research where issues of boundedness or unboundedness arise. A good example is work on the zone of latent solutions (ZLS) theory (Tennie et al., 2009), which analyzes the cultural and putative cumulative cultural traditions of nonhuman animals—putative because although several species have capacities for social learning, they appear to have minimal capacities for building on previous traits. Speaking generally, the ZLS theory suggests that the cultural capacities of nonhuman animal species are “bounded,” limited by a possible range of features. Explanations for why this might be the case have centered mainly on the great apes (hereinafter “apes”), but developing work suggests that similar explanations may hold true with other animals, such as some birds and whales (Aplin, 2019; Perry, 2011; van Schaik et al., 2003; Whitehead & Rendell, 2015; Whiten et al., 1999).

According to the ZLS theory, many putative instances of ape (and perhaps other animals’) cumulative culture are not, in fact, instances of cumulative culture. The ZLS theory argues that apes lack (or have minimal, or rarely expressed) capacities for transmitting and retaining trait modifications. What appears to be cumulative culture is instead likely to be socially influenced reinnovation. When apes reinnovate, they draw on a baseline repertoire of behaviors—behaviors that any able-bodied ape would be able to express—to individually strike upon the trait of interest. Though this reinnovation may be socially facilitated, in the sense that other apes may draw attention to relevant or highly salient environments or objects, the trait is developed by each learner anew.

The basic idea of the ZLS is that this baseline repertoire—and the artful combinations thereof—largely sets the bounds of possible cultural evolution (together, perhaps, with other cognitive features). Absent of more sophisticated forms of social learning, apes are unable to add novel traits or to build cumulative traditions that progress beyond the boundary of “latent solutions.” Apes, but not humans, do not seem to copy—or transmit—traits beyond their ZLS (be it in the technical [Tennie et al., 2009] or social domain [Clay & Tennie, 2017]). As said earlier, the appearance of cumulative culture can largely be accounted for by socially facilitated reinnovation (Tennie et al., 2020).

What might explain the transition between bounded ape culture and unbounded human culture? Though a full catalog of important underlying processes has not yet been completed, a key capacity seems to be abilities for copying “know-how,” that is, capacities for attending to, perhaps understanding, and copying/reconstructing the elements and interrelationships of any particular behavior (including the making of artifacts and of artifact structures themselves). Other relevant capacities, at least for modern humans, plausibly include language and special types of teaching (especially those that can transmit know-how).

ZLS research thus helps the current project in two ways. First, it helps to sharpen the notion of cultural evolutionary boundedness. Boundedness involves a limited exploration of cultural evolutionary space, due to minimal, lacking, or rarely expressed capacities for transmission, retention, or the production of modifications. Second, it helps to illuminate the devilish empirical issues involved in understanding the transition from boundedness to unboundedness. Focusing on the tall, wide, and unbounded cultural evolution of humans alone may not be helpful for understanding this transition (Buskell & Tennie, in press), but a combined focus that also includes understanding the patterns of change in evolutionary systems that ultimately fail to break away from boundedness may.

### 3.4 Evolved Open-Endedness in Action

According to Pattee & Sayama (2019), “conditions for increased open-endedness must have been gradually acquired in the course of evolution” (p. 5). In justifying this claim, Pattee & Sayama point not only to concepts from the foundations of the modern synthesis (Haldane, 1932/1990) and
other more recent attempts to frame evolution as a progression of steps toward increased evolvability (Maynard Smith & Szathmáry, 1995; Szathmáry, 2015; Wagner & Altenberg, 1996; Wilson, 1997) but also to numerous examples of evolved mechanisms that have “significantly facilitated the open-endedness in the evolution of life” (Pattee & Sayama, 2019, p. 6). Notable among these examples are (a) the evolution of symbolic languages used by humans, which are noted as being “evolved from simpler, less open-ended languages” (Pattee & Sayama, 2019, p. 6); (b) the formation of cooperative groups of increasing scale and complexity (colonies → societies), with higher levels of organizational and institutional formation requiring the evolution of new mechanisms not previously seen in lower-level organizational entities; and (c) the evolution of new information-processing abilities, sensory modalities, and the brain, all providing organisms with new possibilities to explore and exploit. From these examples, it is clear that Pattee and Sayama consider what we describe as the evolution of culture (e.g., languages and social institutions) and the biological mechanisms that support culture (e.g., the brain and culture supporting sensory modalities) as clear examples of evolved open-endedness. Therefore we believe that in human cultural evolution (including dual and triple inheritance), we have a real (and recent) example of evolved open-endedness in action. In what follows, we outline the case for human cultural evolution as an instance of evolved open-endedness in action.

Within cultural species more broadly, we can differentiate between different types of cultural evolution: bounded noncumulative, bounded cumulative, unbounded noncumulative, and unbounded cumulative. Whereas cumulative culture may or may not be uniquely human (Mesoudi & Thornton, 2018), unbounded cumulative culture plausibly is. Indeed, human cultural evolution appears to be the only instance of unbounded cumulative cultural evolution.

Evidence suggests that the transition toward unbounded cumulative cultural evolution has taken place over the last few hundred thousand years with the origin and evolution of Homo sapiens (Stringer, 2016; Stringer & Galway-Witham, 2017) or even over a few million years with the advent of stone tool use in early Homo species (Lewis & Harmand, 2016). We thus have, in both archaeological remains and in our genes, the record of this transition into open-ended cultural evolution. Exploring this transition is valuable, for it offers a compelling insight into the problems, solutions, processes, and complex evolutionary dynamics that can jointly explain the emergence of a new open-ended evolutionary system. Though this is a particular instance, we suspect that the concepts, tools, and ideas can be generalized.

This is not to say that explaining the transition from primate ancestors to fully fledged cultural hominins is easy—it is anything but. Contemporary narratives point to a number of important changes that might have facilitated the evolution of a robust, quasi-independent system for cultural inheritance. These include changes in morphology (the bipedal stance, decreased gut size, larger crania), life history and population structure (social affiliation, intergenerational care, long developmental periods, extended family groups and social institutions), and cognitive attributes and machinery (greater executive control, social tolerance and attentiveness) (Aiello & Wheeler, 1995; Antón et al., 2014; Grove, 2017; Kaplan et al., 2000; Klein, 2008; Ostrom, 1990; Powers & Lehmann, 2013; Powers et al., 2016; Sterelny, 2012, 2021).

Just as important were cultural evolutionary feedback loops where early culture could facilitate selection for more and more effective social learning. Premodern hominin culture, for instance, generated an information environment seeded with cues as to how one should live. This includes “scaffolded” learning environments, where juveniles can learn in a relatively safe and low-cost manner by interacting with the products of adult cooperation. These low-cost and safe learning environments could be increasingly supplemented with real-world experience, perhaps teaching and, as learners developed, experimentation. Selection to improve capacities to navigate and explore this informational domain would in turn lead to greater informational structure in the world—and thus to further selection. This general story is one of humans as “evolved apprentices” (Sterelny, 2012).

The story of how hominins escaped the “boundedness” of their primate relatives exploits this evolutionary feedback loop, increasing capacities for both tall and wide culture and abilities to
recognize “task-independent” properties of artifacts and behaviors that could be transferred and combined with other behaviors to generate new kinds of cultural traditions. These cognitive and cultural capacities could open up new evolutionary domains by exploiting novel affordances (Arthur, 2009; Derex, 2022). As a result, human technologies capture and put to use a collection of phenomena: For example, a car exploits not only the phenomenon that rolling objects produce much less friction than sliding ones (resulting in the use of wheels) but also the phenomenon that chemical substances (say, diesel) produce energy when burned (Arthur, 2009). This discovery and exploitation of new solutions to old problems allow a potentially unbounded form of cumulative culture. As noted, we see evidence for the opening up of new evolutionary search spaces and the exploitation of new solutions in numerous domains within patent records (Bedau, 2013, 2019; Bedau et al., 2019).

Equally important is the way that human groups can support the increasing specialization of skills and knowledge, the circulation of knowledge, and participation in collective endeavors—pitching in on large or temporally distributed projects that could never be completed by a single agent in their own lifetime. These social features in turn could contribute to the changes in cognition, life history, and information dynamics discussed earlier. This is part of what some have called—with various slight differences—the cultural intelligence hypothesis (Herrmann et al., 2007; Muthukrishna et al., 2018; van Schaik & Burkart, 2011).

As this makes clear, the transition between a limited type of social learning and the more complex and open-ended form currently enjoyed by humans is a complex story. Despite this complexity, researchers in archaeology, comparative psychology, paleoanthropology, psychology, philosophy, and many other fields have been able to make progress on disentangling distinct causal pathways and to show how these can be put together again to explain the evolution of a distinct system of open-ended evolution: human cultural evolution (Boyd & Richerson, 1985; Tomasello, 1999).

4 Cultural Evolution, Open-Ended Evolution, and Artificial Life

Culture and cultural evolution have a long tradition in Artificial Life, appearing among both the grand challenges (C. Taylor & Jefferson, 1993) and open problems (Bedau et al., 2000) of the field and spawning a regular workshop series at the Artificial Life conference (Marriott et al., 2018). It is therefore curious that, until recently, open-ended cultural evolution has received relatively little attention as a possible avenue for fruitful research (see Bedau et al., 2019).

In the previous sections of this article, we have outlined many of the arguments and factors that we feel place cultural evolution firmly within the domain of open-ended evolution research. However, we also note a curious parallel between the work already taking place within the Artificial Life open-ended research community and the broader study of culture as an evolving system. A particular example of this can be seen in the work of Taylor (2019), where three classes of novelty, all capable of generating open-ended evolution, are introduced: (a) exploratory novelty, whereby existing traits are recombined to produce novel adaptations; (b) expansive novelty, resulting from the discovery and exploitation of new affordances; and (c) transformative novelty, resulting from the discovery of new state spaces, possibly via the exaptation of current traits. Within the cultural evolution literature, we can see clear parallels with each of these classes: exploratory novelty can be seen as a restricted process of cultural variation and accumulated modification within one domain or affordance (described as Type I cumulative cultural evolution by Derex [2022]); expansive novelty can be interpreted as an exploration of new affordances, expanding cultural evolution into new domains (described as Type II cumulative cultural evolution by Derex); and transformative novelty can be viewed as movement into an n-dimensional state-space through the recombination and exaptation of existing cultural traits, enabling the creation and exploitation of new cultural and ecological niches. Examples of cultural exaptation abound in numerous domains, for example, in technology (Bedau, 2019; Bedau et al., 2019; Boyd et al., 2013) and in pharmaceuticals (Andriani et al., 2015).
It is evident that open-ended evolution research in Artificial Life and cultural evolution research have been speaking about very similar things; the types of novelty discussed by Taylor (2019) and core aspects of cumulative cultural evolution outlined by Derex (2022) and Mesoudi and Thornton (2018) demonstrate such similarities. It should therefore be uncontroversial to suggest an open-ended evolutionary synthesis that combines genetic evolution, cultural evolution, and artificial evolution within a single theoretical framework. Combined with the exploratory work on open-ended technological innovation of Bedau et al. (2019) and Bedau (2019), the inclusion of social and cultural transitions emerging from earlier biological transitions within the major transitions framework (Calcott & Sterelny, 2011; Maynard Smith & Szathmáry, 1995; Szathmáry, 2015), and the clear articulation of evidence for both biological and cultural mechanisms for facilitating evolved open-endedness (Pattee & Sayama, 2019), we see a strong argument for including cultural evolution within the broader framework of open-ended evolution.

In the following sections, we argue that the transition from bounded to unbounded evolution that is evident within the recent hominin evolutionary history shines an important light on how evolved open-endedness might be achieved. We go on to consider tall and wide evolution within the context of the Bedau et al. (1998) “ALife test” and provide some initial thoughts on how this test could be expanded to detect tall and wide patterns in order to better delineate between the mechanisms driving (and halting) artificial evolutionary systems. Finally, we introduce a raft of new questions that including cultural evolution under the framework of evolved open-endedness allows us to ask.

4.1 Transitions From Bounded to Unbounded Evolution

As we saw in section 2, it is common to operationalize culture in informational terms: Culture is information, embedded (or carried) by heterogeneous vehicles, that can be transmitted between agents (Richerson & Boyd, 2005). On this understanding, one thread tying together the evolutionary history of hominin populations is an increase in and improvement of culturally transmitted information (Boyd & Richerson, 1985). This general observation has led some researchers to claim that culture represents a “major transition” in the sense of Maynard Smith and Szathmáry (1995) and Szathmáry (2015), building off the idea that such transitions involve changes in the quality and reliability of information transfer. For instance, Waring and Wood (2021) have argued that human cultural groups are a new kind of evolutionary individual, suggesting that cultural selection pressures now vastly outweigh biological selection pressures in determining the course of human diversification and change.

Waring and Wood’s (2021) arguments interpret the major transitions framework in a particularly strong way. This takes transitions to involve stabilizing a new evolutionary individual, here a cultural group (McShea & Simpson, 2011). But one need not understand the framework in this “unified” way (Michod, 1999). Instead, transitions may involve modifications of the “core elements of the evolutionary process itself” (Calcott & Sterelny, 2011, p. 4), irrespective of introducing a new level or kind of selection process (Godfrey-Smith, 2009). Thus, even if one is skeptical about cultural group selection (see, e.g., Chellappoo, 2022), one can usefully understand the introduction and refinement of cultural evolution using the ideas and machinery of the major transitions literature (Calcott & Sterelny, 2011; Maynard Smith & Szathmáry, 1995; Szathmáry, 2015).

We conceive “open-endedness” through this more expansive understanding. It characterizes an increase of informational content that can be (or is) transmitted in a given domain, potentially reflecting coordinated or piecemeal changes to the rate, quantity, or kind of variation that can be generated. In so doing, we elaborate and expand some ideas found in Pattee and Sayama (2019), as articulated in Borg and Powers (2021, p. 2): “Over time both biological adaptations that enable more complex and open-ended social and cultural behaviors (bigger brains, opposable thumbs, changes in the shape of the larynx, …), and cultural adaptations that open up access to new domains of knowledge (symbolic language, the scientific method, music and art, complex social institutions, …) have been selected for in a clear demonstration of selection in favour of open-endedness, with this same selection pressure being seemingly absent in our closest genetic relatives”.

[430]
4.2 Cultural Evolution and the ALife Test for Open-Endedness

Determining whether an evolutionary system exhibits unbounded evolutionary dynamics is still arguably the primary concern of open-ended evolution research. Without the ability to judge whether a system is open-ended, how can open-endedness be understood to any useful degree? Despite a general lack of use, we are of the opinion that the classification system of long-term evolutionary dynamics devised by Bedau et al. (1998) (sometimes known as the ALife test for open-endedness) provides us with the best method for determining whether an evolutionary system exhibits unbounded evolutionary dynamics. However, we believe that some of the key features of cultural evolution—wide versus tall evolution, transition from bounded to unbounded evolution, and evolved open-endedness—may necessitate some refinement of the ALife test.

The three primary measures of evolutionary activity Bedau et al. (1998) described are (a) the diversity of traits within the system at any given time, (b) the amount of “new evolutionary activity” observed in the system over time (i.e., the creation and maintenance of new adaptive traits), and (c) the mean cumulative activity of traits (i.e., the number of traits observed to date divided by the current diversity of traits in the system). For a system to exhibit unbounded evolutionary dynamics, it would need to always demonstrate positive new evolutionary activity (i.e., new traits are being created and maintained), alongside either unbounded diversity (as time progresses, the number of traits maintained in the system continues to grow) and/or unbounded mean cumulative activity.

What these measures of evolutionary activity do not take into account is whether the new activity is a result of cumulative evolutionary processes, noncumulative evolutionary processes, or recombinative processes. These distinctions matter because they can begin to shed light on how a system has progressed toward, and ultimately achieved, open-endedness. For instance, would we expect to see a “building out” of wide adaptations (as seems to be the case in hominin cultural evolution) before the emergence of tall accumulated modifications, ultimately leading to the combination of traits from disparate evolutionary lineages, forming recombinative adaptations (wide evolution providing the raw material for exploratory and expansive evolution as per Taylor, 2019)? Or are there numerous different pathways to open-endedness that can only be understood by breaking down the nature of the evolutionary patterns of change, adaptive processes, substrates, and mechanisms underpinning these evolutionary systems?

4.3 New Questions in Open-Endedness

Once we consider the implications and nature of cultural evolution from an open-ended evolution perspective, we can begin to ask new and important questions about evolved open-endedness, human cultural evolution, and the underpinning dynamics of all evolutionary systems. These questions include, but are not limited to, the following:

- Do the mechanisms underpinning cultural evolution more easily lead to open-endedness than those underpinning genetic evolution, or vice versa?

- What happens when a bounded aspect of an evolutionary system (e.g., animal cultural evolution) comes up against an unbounded aspect of the same evolutionary system (e.g., human open-ended cultural evolution)? Is there a sudden pressure for evolved open-endedness to emerge among species that have so far exhibited only bounded cultural evolution? And does the emergence of open-endedness always lead to the extinction of its bounded counterpart?

- Are there any bounded aspects of human cultural evolution? And could there also be bounded aspects of genetic evolution?

- Does an evolutionary system need to be cumulative to be open-ended, or is it possible to have noncumulative open-ended evolution? Note that if major transitions are one of the primary behavioral hallmarks of an open-ended evolutionary system (Taylor et al., 2016), and major transitions build up incrementally from one another (each transition is dependent
on subsequent levels), this would imply that open-ended evolution must result from a cumulative evolutionary process. But is it possible to generate open-ended evolution without cumulative major transitions, and could major transitions be the result of numerous independent innovations?

- Are cumulative evolutionary systems always open-ended? The numerous cases outlined in Mesoudi and Thornton (2018) would suggest not, nor do the criteria for cumulative cultural evolution necessitate an open-ended system (or logically lead to the conclusion that open-ended evolution is an unavoidable end point).

- What features of cultural evolution are common to all evolutionary systems capable of generating the open-ended evolution of novelty?

- Is an open-ended evolutionary synthesis that accommodates cultural evolution alongside genetic and artificial evolution viable and/or desirable?

- Is niche construction necessary for open-ended evolution? And are the autocatalytic processes resulting from the interplay between numerous interdependent evolutionary systems necessary for open-endedness?

5 Conclusion

In this article, we set out to outline culture as an evolutionary system and argue for its inclusion within the broader framework of evolved open-endedness. To make these arguments, we provided numerous examples of the unique aspects of cultural evolution that highlight important contrasts with biological evolution, but we also maintain a direct link between the core algorithmic features of biological and cultural evolution. We went on to discuss the key features and dynamics of cultural evolution, including tall, wide, cumulative, and noncumulative evolution; transitions from bounded to unbounded evolution; dual and triple inheritance; evolved open-endedness; major transitions; and the ZLS theory. Each of these features provides new insights into the nature of another model evolutionary system.

Going forward, we believe that two lines of enquiry are necessary to fully develop cultural evolution as an integral part of open-ended evolution research. First, following on from the work of Bedau et al. (2019), we believe that applying the ALife test to the vast number of available cultural evolution data sets, across numerous species, would be informative for both the open evolution community and the cultural evolution community. Second, including mechanisms of cultural transmission and the unique features of cultural evolution within artificial evolutionary models aimed at addressing the question of open-endedness to enable the exploration of cultural evolutionary dynamics—this may involve modeling culture as an independent system or including culture alongside genetic (and environmental) inheritance. To enable these two lines of enquiry, we believe that some work on refining the ALife test is necessary, as are the development of tall–wide–recombinative evolutionary theory and more interdisciplinary dialogue between the fields of cultural evolution and Artificial Life.

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Appendix: Elaborated Explanation of Figure 1

Following is an elaborated explanation of the logic behind the example found in Figure 1. Links to reproducible code can also be found herein.

In the example presented in Figure 1, each square is constituted by 441 pixels, where coordinates (0, 0) are located in the center and each axis is extending from −10 to +10. The world wraps (i.e., coordinate (−10, −10) is adjacent to (−9, −10); (−9, −9); (−10, −9); as well as (10, −10); (−10, 10); (10, 10); (−9, 10); (10, −9)). Each pixel represents a state of information, and though the colors have no grounded value, they indicate different states of information. In total, there are five states (black, white, yellow, blue, and pink). The aggregate of pixels within each square represents some kind of technology or cultural practice held by a distinct hypothetical population. In row 1 (bottom row), each pixel is turned white or black with a 0.5 probability. Thus each of the eight squares represents an equiprobable configuration of information. By this logic, each of these squares is solving the same kind of problem.

Over time, the hypothetical population alters its technology or practice, which is subsequently altered again and again in a cumulative fashion. In each case, the overall technology or practice becomes both more complex (an increasing diversity of informational [colored] states of pixels) and more structured (certain axis of symmetry). Like the colors, the axes of symmetry do not represent anything grounded but simply demonstrate a certain kind of orderliness and routinization of the technology or practice. In some cases, pixels are lost (rule golf and hotel), and the “shape” of the aggregate information changes. Again, this is not grounded but may represent accidental loss of information or a deliberate “pruning” of redundancy. In any event, the changes applied to information in each square are cumulative, influencing each subsequent iteration, beginning with the equiprobable starting point.

Eight rules are applied randomly over each iteration along the tall axis:

**Rule Alpha:** A random subset (of between 0 and 49) pixels are asked to turn yellow, blue, or pink.

**Rule Bravo:** All pixels with (negative) x coordinates (< 0) are to assume the informational states of their corresponding (positive) x coordinates (> 0). In effect, this creates an axis of symmetry along the vertical axis.

**Rule Charlie:** Rule Charlie is the inverse of Rule Bravo.

**Rule Delta:** All pixels with (positive) y coordinates (> 0) are to assume the informational states of their corresponding (negative) y coordinates (< 0). In effect, this creates an axis of symmetry along the horizontal axis.

**Rule Echo:** Rule Echo is the inverse of Rule Delta.

**Rule Foxtrot:** A random subset (of between 0 and 19) pixels is identified. The eight neighbors of this pixel become a single color (yellow, blue, or pink).

**Rule Golf:** Black pixels with max (top) and min (bottom) coordinates disappear. Colored pixels remain. (This is apparent in columns 1, 2, and 8 of Figure 1.)

**Rule Hotel:** Rule Hotel is the inverse of Rule Golf (applied to the left or right edge, rather than to the top or bottom).

Reproducable code can be found at https://doi.org/10.5281/zenodo.6948341

Code is written in netlogo, which is freely available at https://ccl.northwestern.edu/netlogo/