On Modeling Cognition and Culture

Why cultural evolution does not require replication of representations

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ABSTRACT

Formal models of cultural evolution analyze how cognitive processes combine with social interaction to generate the distributions and dynamics of ‘representations.’ Recently, cognitive anthropologists have criticized such models. They make three points: mental representations are non-discrete, cultural transmission is highly inaccurate, and mental representations are not replicated, but rather are ‘reconstructed’ through an inferential process that is strongly affected by cognitive ‘attractors.’ They argue that it follows from these three claims that: 1) models that assume replication or replicators are inappropriate, 2) selective cultural learning cannot account for stable traditions, and 3) selective cultural learning cannot generate cumulative adaptation.

Here we use three formal models to show that even if the premises of this critique are correct, the deductions that have been drawn from them are false. In the first model, we assume continuously varying representations under the influence of weak selective transmission and strong attractors. We show that if the attractors are sufficiently strong relative to selective forces, the continuous representation model reduces to the standard discrete-trait replicator model, and the weak selective component determines the final equilibrium of the system. In the second model, we assume inaccurate replication and discrete traits. We show that very low fidelity replication of representations at the individual level does not preclude accurate replication at the population level, and therefore, accurate individual-level replication of representations is not necessary for either cultural inertia or cumulative cultural adaptation. In the third model, we derive plausible conditions for cumulative adaptive evolution, assuming continuous cultural representations, incomplete transmission and substantial inferential transformations.

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Introduction

Formal models of cultural evolution (e.g. Boyd & Richerson 1985) provide an important tool for understanding cultural phenomena. Culture is shaped by both psychological processes that determine how people think and feel, and social processes that determine how people interact, whether they succeed or fail, and if they live or die. Formal cultural evolutionary models can combine cognitive and affective processes with patterns of social interaction to generate explicit predictions about the distributions and dynamics of ‘representations’ — ideas, beliefs, schemas, or mental models. With such predictions, empirical data on behavior and beliefs from populations can be used to select among alternative models, or among the cognitive and social components of these models, in a manner that informs our understanding of cognition and connects laboratory experiments with real life (e.g. Henrich 2001).

Recently Sperber (1996), Atran (2001, 2002), and Boyer (1994, 1999) have argued that these “Neo-Darwinian” models1 are inappropriate for studying cultural evolution because they are “based on a serious distortion of the relevant facts” (Sperber 1996:118). They identify three interrelated problems with such approaches: First, cultural transmission processes are usually incomplete and imperfect, so, unlike genetic systems, accurate replication rarely occurs. Replication is the exception, rather than the rule. Second, unlike DNA replication, inferential processes “transform” these representations during their transmission and reconstruction. This suggests that mutation-like processes are much more important than selection-like processes in shaping cultural variation. Third, unlike genes, cultural representations are rarely discrete units, suggesting that the idea of a ‘replicator’ (or meme) makes little sense for most types of cultural representations.2

1These scholars use the term “Neo-Darwinian” for such models even when they are explicitly derived from disease epidemiology (Cavalli-Sforza & Feldman 1981:46-53), learning theory (see Boyd & Richerson 1985, Chapter 4), or cognitive psychology (Lumsden & Wilson 1981).

2While such characterizations apply to the informal theorizing of memeticists (e.g. Dawkins 1982, Dennett 1995 or Blackmore 1999), they are wide of the mark for much formal cultural theory. Boyd and Richerson (BR 1985:75) explain that there is no need to assume particulate “units” in order to build evolutionary models (BR 1985:70-75), and 19 of
Building on these three points, these authors have claimed: First, because representations are non-discrete, using replicators and replicator dynamics is entirely inappropriate for culture systems (Sperber 1996: 101-103; Atran 2002: Chapter 8, 2001). Second, because of the high levels of inaccuracy and the incompleteness, selective cultural transmission itself does not lead automatically to ‘cultural inertia’ — the existence of social groups in which individuals share a set of relatively stable representations (Sperber 1996: 118). From this, Sperber (1996: Chapter 5) further asserts that inferential transformations create ‘strong cognitive attractors’ that swamp any influence from the selective components of cultural evolutionary processes. Such attractors create powerful forces that overwhelm the weaker effects of selective processes, and, therefore it is argued, selective processes do not usually generate adaptive cultural evolution.

Whether social learning is low fidelity, non-discrete and strongly affected by cognitive transformations are empirical matters, although it is plausible that Atran and Sperber are correct for a wide range of cultural domains. However, the deductions they make from these assumptions are matters of logic, and here we use three mathematical models to

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the 38 models presented in their book are continuous (non-discrete) trait models that allow for an arbitrary amount of transmission error. Similarly, Cavalli-Sforza and Feldman (CSF 1981) devote one of their five chapters entirely to continuous trait models. These continuous models allow for substantial error and other forms of non-replication. BR also explicitly distinguish public representations from mental representations (though using different terminology) throughout their book, and repeatedly specify the inferential transformation between observed behavior and representation formed. For example, in describing one model, they write, “The offspring uses each model’s actual behavior [public representation] to estimate [make inferences about] his or her cultural variant [mental representation]” (1985:75-79). They also make explicit reference to much research in psychology on the nature of social learning. After a review of the psychological literature, especially Bandura’s work on social learning, they specify the following pathway: Modeled events → Attention Processes → Retention Processes → Motor Reproduction → Motivation Processes → Matching. Of the 396 references, 101 were by psychologists, or published in psychology journals. Chapters 4 and 5 discuss how cognitive structures — what Sperber (1996) would later call “attractors” — bias cultural change so that some outcomes are more likely than others, and even use some of the same examples as Boyer (1999).

Atran (2001) refers to attractors as “cognitive elicitors.” Boyer (1999) speaks of “triggered representations,” arguing that these channel cultural evolution onto “cognitive tracks.”
show that these deductions do not follow from their assumptions. In the first model, we assume continuously varying representations under the influence of weak selective transmission and strong attractors. We show that if the attractors are sufficiently strong relative to selective forces, the continuous representation model reduces to the standard discrete-trait replicator model, and the weak selective component determines the final equilibrium of the system. In the second model, we assume representations are discrete, but that replication is very inaccurate. We show that very low fidelity replication of representations at the individual level does not preclude accurate replication at the population level, and therefore, accurate individual-level replication of representations is not necessary for selective forces to generate either cultural inertia or cumulative cultural adaptation. In the third model, we assume continuous (non-discrete) cultural representations, incomplete transmission and substantial inferential transformations. Despite these assumptions, the model shows that adaptive cultural evolution is likely in empirically plausible conditions — predictions derived from this model have been tested elsewhere using ethnographic data (Henrich 2002).

**Model 1: Strong attractors give rise to discrete replicators**

Formal models of cultural evolution have made use of both continuous and discrete representations of cultural variation, depending on the situation. Sperber, Atran and Boyer reject the use of discrete-representation models. At the same time, they emphasize the role of intra-individual cognitive transformations of representations (‘strong attractors’), and argue that, by comparison, selective cultural processes have little or no effect on the epidemiology of representations (e.g. Sperber 1996: Chapter 5). In this section, we analyze a simple model that assumes continuous representations, strong attractors and weak selective forces. We show that this model reduces to discrete-representation replicator-dynamics in which the weak selective forces determine the ultimate outcome. The stronger the cognitive attractors are relative to the selective transmission forces, the better is the discrete-replicator approximation. Therefore, assuming that human cognition gives rise to more than one attractor per domain, it is inconsistent to simultaneously advance the idea of strong attractors
(or ‘cognitive tracks’ or ‘triggered representations) and to reject the use of discrete-representations in models.

Here we study the evolution of the distribution of mental representations in a population. We assume that each individual’s mental representation in this domain is a single real number, $x$, between zero and 1.\footnote{We restrict the $x$ to $[0, 1]$ only for simplicity of exposition. Representations that varied over other ranges, or multidimensional variable produce the same basic result.} Individuals holding different representations (i.e. different values of $x$) behave differently on-average, or in Sperber’s terminology, generate different “public representations.” During each time period, people in the population observe the behavior of another individual, infer the mental representation of the model from his or her behavior, and adopt this inference as their own representation.\footnote{For the moment we do not allow individuals to observe a sample models and make use of the pattern of observed behavior in the sample. We return to this question below.} Following Sperber, cognitive ‘attractors’ strongly bias the inferential process that underlies social learning — an individual whose model has a representation $x$, on average, infers a representation that is nearer to one of two attractors. In particular, we assume that the attractors are $x = 0$ and $x = 1$, and that an individual who observes a model with representation $x$ infers a representation $x + \Delta x$, where $\Delta x < 0$ for $x < m$ and $\Delta x > 0$ for $x > m$. We assume that $\Delta x$ has the form shown in Figure 1.

The following example illustrates how domain specific cognition might create multiple attractors. Suppose individuals at Attractor-0 ($x = 0$) perceive the Moon as a self-aware, conscious, entity with goals, emotions, and motivations — thus the Moon’s behavior can be understood using folk psychology (Leslie 1994). This means that the quantity $(1 - x)$ tells us the degree to which an individual attributes the Moon’s shape, color, position and movements to underlying goals, emotions or motivations, or to what degree the Moon’s goal-driven actions influence events and individuals, such as weather, tides, personal moods, werewolves, etc. In contrast, individuals with $x = 1$ (those at Attractor-1) see the Moon as simply a big rock, lacking goals, consciousness, and emotions. These individuals attribute the Moon’s color, shape and movement to the effects of non-agentic interactions with light, and other mindless bodies, governed by physical laws that operate throughout the Universe. Any effects the Moon
Figure 1. Graphical representation of the assumptions in Model 1. A learner who observes a model with representation, \( x \), infers a representation \( x + \Delta x \), where \( \Delta x = -\beta_0 x \) if \( x < m \) and \( \Delta x = \beta_1 (1 - x) \) if \( x > m \). This creates a strong force that shifts representations in the population toward attractors at 0 and 1. The probability that an individual is chosen as a model is \( 1 + s x \). This selective transmission process creates a weak force that increases the frequency of larger values of \( x \) in the population.

has on things such as tides, moods and calendars, are merely unintentional consequences of the moon’s mass and movement. Now, it is possible to imagine Moon-concepts that mix these poles (\( 1 > x > 0 \)). One could believe, for example, that the moon’s movement and shape are out of its control (governed by physical laws), while its color or hue expresses its mood, which in turn influences the weather. Or, perhaps the Moon’s color is 23% controlled by its emotions and 77% controlled by the laws of light refraction. One might also believe that on Tuesdays and Thursdays the Moon is a goal-oriented agent, on Mondays, Wednesdays and Fridays the Moon is a big rock, and on the weekends these two alternate minute by minute. Such beliefs might seem odd to us because they violate intuitive expectations, and would consequently be transformed by cognitive attractors. In contrast, \( x = 1 \) or 0 are “easier to think.” To capture this example in a formal model, the underlying mental representations, \( x \), would likely need to be a multi-dimensional vector, and the analysis represented below could be easily adapted to such representations — none
of the qualitative findings would change. However, to keep the presentation tractable, we will limit our analysis to one-dimensional $x$ variables.

A second example illustrates how an attractor might be represented by a single quantitative dimension. Young and Burke (1998) have shown that an overwhelming majority of the share cropping contracts in Illinois are one of two types: In some communities the farmers and the landowners split the crop yield, while in other communities the landowners receive two shares for every one received by the farmer. Interestingly, despite wide variation in land quality within communities, there is little variation in share cropping contracts within communities. Let us suppose that we were studying a population of farmers in this region, and that $x_i$ is individual $i$’s mental representation of the fair or appropriate share for the landowner in a share-cropping contract. We scale these representations so that Attractor-0 ($x = 0$) is a $1:1$ share, while Attractor 1 ($x = 1$) represents a $2:1$ share. Cognitively, such attractors might arise because integer ratios involving numbers that humans intuitively understand are easier to represent and remember than other ratios (e.g., $1.38:3.13$). When a person is observed to use other contracts, he or she is often thought to be using (and thinking about) one of the two focal contracts.

We also assume that individuals are selective in picking their cultural models — that is, in deciding on whom to focus their inferential processes. In particular, we assume individuals focus on people with higher values of $x$. Individuals with higher values of $x$ might have more success, interact more, or be more salient for other reasons. This assumption about human psychology is well founded: A substantial amount of evidence from both laboratory and field research demonstrates that humans pay particular attention to, preferentially interact with, and tend to imitate successful/prestigious individuals (Henrich & Gil-White 2001 summarize this evidence and lay an evolutionary foundation). Using our above example, it could be that individuals who believe in a $2:1$ contract have more success, on average, than individuals who have other mental representations, and that their relative success will make them preferred models. We let $w_i$ be the likelihood that individual $i$ is selected as a model, and assume that there is a positive correlation between $x_i$ and $w_i$.

We analyze this model using the Price Equation (Price 1970). This equation (1) says that the mean change in the mean of any property of
a population of things, genes, mental representations, or the hydrogen atoms in the Andromeda galaxy, can be decomposed into two parts: the extent to which the properties of things covary with the effect of selection, and the rate at which the things themselves change with time. There are no assumptions about replication fidelity, discreteness or underlying distributions. This form is particularly useful here because it partitions the change in a population of mental representations into the effect of selecting particular cultural models (“selective transmission”) from the effect of inferential processes (“incomplete inference”).

\[
\bar{w} \Delta \bar{x} = \text{Cov}(w_i, x_i) + E(w_i \Delta x_i)
\]

\( \Delta \bar{x} \) is the change in the average value of \( \bar{x} \) in the population per time step, \( \Delta x_i \) captures the effect of the cognitive attractors, and \( \bar{w} \) is the average replicability of the \( x_i \) values carried in the minds of the \( n \) individuals in the group.

Another useful property of the Price equation is that it makes it easy to deal with populations that are structured into groups (Price 1972). Here, we divide the population into two groups, one for each domain of attraction, and express the dynamics for the change in the average value of \( \bar{x} \) in each domain of attraction. Group 0 is composed of individuals whose mental representations are less than \( m \), and group 1 contains those individuals whose mental representations are greater than \( m \). Let \( \bar{x}_j \) and \( \bar{w}_j \) be the mean value of \( x \) and the mean replication rate for group \( j \), and let \( x_{ji} \) and \( w_{ji} \) be the mental representations of the \( i \)th individual in group \( j \). Then we can rewrite (1) as follows:

\[
\bar{w} \Delta \bar{x} = \text{Cov}(\bar{w}_j, \bar{x}_j) + E(\bar{w}_j \Delta \bar{x}_j)
\]

That is, the change in the mean value over the whole population can be decomposed into the covariance between group means and group mean replication rates, and the average change within each group. But, notice that the average change within each group has the same form as the left hand side of the Price equation. Thus, we can apply the Price equation

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\(^{6}\text{In the form expressed here we have assumed that the x’s proportional representation from one time step to the next is a function of its relative replicability (or fitness) and its current representation in the population.}\)
again, this time to the dynamics within each group — substituting the expressions given in (3) into the expectation term in (2).

\[ w_j \Delta \bar{x}_j = \text{Cov}(w_{ji}, x_{ji}) + E(w_{ji} \Delta x_{ji}) \]  

(3)

The covariance term gives the change within the groups due to the psychology of model selection, and the second term gives the change within each group due to inferential transformation by attractors.

Now we make use of the specific form of the inferential transformation shown in Figure 1. In group 0, \( \Delta x_{0i} = -\beta_0 x_{0i} \), and in group 1, \( \Delta x_{1i} = \beta_1 (1 - x_{1i}) \). Further, assume that an individual's likelihood of being selected as a model depends linearly on their value of \( x_{ji} \) so that \( w_{ji} = 1 + sx_{ji} \). With these assumptions

\[ \tilde{w}_0 \Delta \bar{x}_0 = -\beta_0 \bar{x}_0 + s(\text{Var}(x_{0i}) - \beta_0 E(x_{0i}^2)) \approx -\beta_0 \bar{x}_0 \]  

(4)

and

\[ \tilde{w}_1 \Delta \bar{x}_1 = \beta_1 (1 - \bar{x}_1) + s[\text{Var}(x_{1i}) - \beta_1 (\bar{x}_{1i} - E(x_{1i}^2))] \approx \beta_1 (1 - \bar{x}_1) \]  

(5)

If intra-psychic transformations are strong relative to the selective forces (\( \beta_1, \beta_0 \gg s \)), then the first terms on the right-hand sides of (4) and (5) will dominate the dynamics and determine the equilibrium values of the mean values of \( x \) in each group.

We can now write down the approximate epidemiological dynamics for the entire population. First, let \( p \) be the fraction of individuals in group 1 and \( 1 - p \) the fraction of the population in group 0. Then, substituting (4) and (5) into (2) yields:

\[ \tilde{w} \Delta \bar{x} \approx \left[ p(\tilde{w}_1 \bar{x}_1) + (1 - p)(\tilde{w}_0 \bar{x}_0) - \bar{x} \tilde{w} \right] \]

Covariance Term from (2)

\[ + \left[ p\beta_1 (1 - \bar{x}_1) - (1 - p)\beta_0 \bar{x}_0 \right] \]

Expectation Term from (2)

(6)

Because the \( \beta \)'s are so much larger than \( s \), the dynamics of the system will be initially governed by the attractors: \( \bar{x}_0 \) will rapidly evolve toward zero, \( \bar{x}_1 \) toward one, and thus \( \bar{x} \approx p \). Substituting these values into (6) gives the following equation for the approximate dynamics after this initial period:

\[ \Delta \bar{x} \approx \Delta p \approx \frac{p(\tilde{w}_1 - \tilde{w})}{\tilde{w}} \]  

(7)
Equation (7) is the standard formulation for discrete-trait replicator dynamics. With this formulation, we see that the longer-term dynamics and the final equilibrium, $\bar{x} = 1$, are determined by selective cultural transmission and approximated with standard replicator dynamics. The stronger cognitive attractors are relative to selective cultural transmission, the better the discrete replicator approximation.

These findings are very general. Elsewhere, we (in prep) show this finding applies to more than two attractors, for a wide range different intra-psycho transformations, and to representations in any number of dimensions. There we also discuss the effect of more complex forms of selective cultural transmission.

Numerical simulations of the model indicate that the approximate analysis derived above is quite accurate. In our simulation, we assumed an initially uniform distribution of $x_i$ in a finite population. For parameters, $n = 200$, $b_0 = b_1 = 0.50$, $s = 0.05$ and $m = 0.60$, Figure 2 shows the dynamics of the average value of $x$ in group 0 ($\bar{x}_0$), the average value of $x$ in group 1 ($\bar{x}_1$), the mean of $x$ (overall: $\bar{x}$) for one run of the simulation, the mean value of $x$ for 10 simulations, and the predictions of replicator dynamics. As assumed above, $\bar{x}_0$ goes rapidly to 0 and $\bar{x}_1$ goes rapidly to 1. The trajectory of $\bar{x}$ from one run of the simulation shows the random

Figure 2. Results from simulating model described in text. The overall evolution of the population is very well approximated by a discrete model in which only weak selective forces are present.
effects of drift produced by finite populations, but still clearly tracks the replicator approximation. Across 10 simulations, the effects of drift are averaged out and replicator dynamics tracks $\bar{x}$ quite closely. A series of simulations with a variety of parameter combinations confirms the findings derived above. In addition, it shows how cultural drift is affected by $s$, $n$ and $m$. As $n$ decreases, the force cultural drift increases, making it more likely for $x$ to drift into Domain 0. If $s$ gets too small, drift is also more likely to drive $x$ to zero. As $m$ gets large, $x$ values are more likely to drift into Domain 0 and be driven to zero.

**Model 2: Cultural inertia and cumulative cultural adaptation can occur even when inference is inaccurate**

In this second model, we explore the effect of inaccurate replication. We assume that cognitive processes generate strong attractors, but that inferences, based on the available public representations, are highly inaccurate. We use discrete-representations to show that, even when transmission fidelity is very low, cultural transmission can still create cultural inertia and adaptive cultural evolution.

Much of the confusion about whether selective cultural transmission can produce cultural inertia, shared traditions, and adaptive evolution arises from a failure to appreciate the differences between cultural and genetic evolution. In genetic evolution both evolutionary inertia and the possibility of gradual cumulative cultural change have the same cause: accurate, unbiased genetic replication. In sexual species like humans, each individual has two parents and this has important consequences for the distribution of genotypes in populations. However, except for extremely rare mutations, each gene is a faithful copy of a single gene carried by a member of the previous generation — genetic replication is very accurate. Moreover, each gene that an individual carries, again with a few exceptions, is equally likely to be included in the individuals’ gametes — thus genetic reproduction is unbiased. These properties of genetic reproduction result in Mendel’s laws for the distribution of genotypes.

To see how accurate, unbiased replication gives rise to genetic inertia and makes cumulative adaptive change possible, consider a simple genetic system with only two alleles (or genes) at a single locus segregating in a large population. We label them $A$ and $a$. Let $q$ be the frequency of $A$
before genetic transmission, and \( q' \) the frequency after genetic transmission. Finally, suppose that each allele mutates to the other with probability \( u \). This means that,

\[
q' = (1 - 2u)q + u
\]

A plot of \( q' \) as a function of \( q \) is a straight line with a slope slightly smaller than one as shown in Figure 3. If the slope were exactly one (i.e., no mutation), then transmission would lead to no change in the population \( (q' = q) \) across the generations. A population that had 80% \( A \) alleles before transmission would have 80% after transmission, and a population that had 20% before transmission would have 20% afterward. However, any amount of mutation tends to reduce the frequency of the more common allele, and thus if no other evolutionary processes affect the population it will move toward a state in which both alleles have the same frequency. Because mutations are very rare \( (u \approx 10^{-6}) \), this will occur very, very slowly. Thus, in the absence of other forces, populations will change very slowly, and this explains much genetic inertia. Similarly, as long as natural selection is stronger than mutation, it can generate cumulative adaptation.

By analogy from the process just described, Sperber (1996: 103-118) and Atran (2001) argue that because cultural replication is highly inaccurate, the social processes of cultural transmission cannot give rise to cultural inertia or cumulative cultural adaptation. Unlike genes, mental representations are not replicated during cultural transmission. Instead, mental representations give rise to behaviors (or “public representations”) that are observed by others, who must then infer the underlying mental representations that gave rise to the behavior. Because individuals differ and public representations provide incomplete information, this inferential process is, Sperber, Boyer and Atran assert, highly inaccurate.

If they are correct, it does follow that cultural transmission cannot give rise to cultural inertia for the same reasons as genetic transmission. To see this, suppose there are only two possible mental representations in some domain, labeled \( A \) and \( B \). Each generates different but overlapping distributions of public representations. When cultural learning occurs, naïve individuals, perhaps children, observe a sample of individuals from these distributions, make inferences, and then adopt their own mental representation. Following Sperber, Atran and Boyer, we suppose this
process is very inaccurate; it is governed by equation (8) but now $u = 0.2$ (5 orders of magnitude larger than the rate genetic mutation). To get a feeling for the magnitude of $u$, remember that $u = 0.5$ means that there is no transmission at all — naïve individuals adopt mental representations at random, independent of who they observe. Thus, we are assuming that the acquisition of mental representations is 40% error and 60% transmission. As is shown in Figure 4, very high error rates lead to very different dynamics than accurate replication. Under such high error rates, populations very rapidly converge to a random distribution of mental representations. There is no inertia, and extremely strong selective forces would be required to generate cumulative adaptation.

However, this logic carries the genetic analogy too far. Even if cultural transmission is inaccurate, it does not follow there can be no cultural inertia or cumulative evolution of adaptations. Any transmission process that leads to accurate replication at the level of the population will lead to cultural inertia and allow cumulative adaptation. Put another way, any transmission process that produces a plot like that shown in Figure 3 will do the job. The mistake is to assume that the only process that can give rise
Figure 4. The frequency of representations after reproduction, \( q' \), as a function of the frequency before reproduction \( q \) when the error rate, \( u \), is high. As before, errors tend to drive representation frequencies toward 0.5; but, now this occurs very rapidly because the error rate is large.

Accurate replication at the level of the population is accurate replication at the level of individuals.

Here we show that a tendency to adopt the more common representation can lead to accurate replication at the population level even when individual inferences are highly inaccurate. Assume that there are two possible mental representations, \( A \) and \( B \), and that a fraction \( q \) of the experienced population has \( A \). Every naïve individual observes the public representations of \( n \) models and makes inferences about the mental representations that gave rise to these public representations. In each case, this inferential process is subject to frequent errors. The probability of inferring the correct mental representation is \( 1 - u \) and the incorrect representation is \( u \). Thus, the probability, \( p \), that a social learner infers that any one of the models has mental representation \( A \) is:

\[
p = q(1 - u) + (1 - q)u
\]  
(9)

Finally, suppose that individuals adopt, as their own, the mental representation that they believe is most common among their models. This means that the probability that a naïve individual acquires \( A \), and therefore, the frequency of \( A \) after transmission, \( q' \), is:

\[
q' = \sum_{i > n/2}^{n} \frac{n!}{i!(n-i)!} p^i (1 - p)^{n-i}
\]  
(10)
Because individuals tend to acquire the more common mental representation, this model is one way to represent a “conformist” bias in social learning.

The conformist bias has well-established theoretical and empirical foundations. Theoretically, evolutionary modeling of the cognitive capacity for conformist transmission suggests that genes leading to a conformist bias in social learning will be supported in virtually any environment that also favors social learning (Henrich & Boyd 1998). Empirically, psychological evidence for conformity from pre-1984 studies is summarized in Boyd and Richerson (1985). Much of recent evidence can be found in Baron et al. (1996), Insko et al. (1985), Smith and Bell (1994), Bond and Smith (1996), and Campbell and Fairey (1989). These studies analyze everything from economic decisions and strategy choice in strategic games to perceptual tasks and food choices. Moreover, by analyzing the dynamics of transmission, Henrich (2001) provides evidence of conformist transmission from field data on the diffusion of innovations.

A conformist bias at the individual level leads to more accurate replication at the population level. Figure 5a plots $q'$ as a function of $q$ for $n = 10$ and $u = 0.2$. This figure shows that social learning processes come much closer to accurately replicating the population frequency of the mental representations, even though the transmission process is error prone at the individual level. The maximum error rate at the population level (at $q = 0, 1$) is approximately 0.02, one tenth of the individual rate. The reason for this is simple: errors have a bigger effect on populations in which one mental representation is common compared to populations in which both mental representations have similar frequencies. As a result, the conformist bias in transmission effectively corrects for the effect of errors. Furthermore, even higher error rates can be compensated for by larger samples of cultural parents: Figure 5b plots the effect of transmission when $u = 0.3$ and $n = 20$. Recall that $u = 0.30$ means errors occur 60% of time.

This conformist learning mechanism can lead cultural inertia because it tends to increases the frequency of common mental representations, and reduces the frequency of rare ones (Figure 6a). Observe that the conformist curves both closely follows the $q' = q$ line, and creates two stable equilibria near the points 0,0 and 1,1. These equilibria occur when the effect of
errors and the force of conformist transmission are balanced. Two initially different populations can remain different even in the same environment. Cultural inertia is created by conformist transmission even when error rates are high.

Conformist social learning also allows selective forces to generate more accurate adaptations to the environment. Suppose that individuals with mental representation $A$ are twice as successful as individuals with representation $B$ on average, and that the probability that individuals are chosen to be models is proportional to the ratio of an individual’s success to the average success in the population (‘prestige-biased transmission’). Figure 6b shows that when we combine prestige bias, conformist bias and high error rates the more successful representation readily spreads.

The combination of high error rates and a conformist bias does not result in the same kind of “frictionless” adaptation as genetic replication. Highly accurate, unbiased, genetic replication allows minute selective forces to generate and preserve adaptations over millions of years. Error prone cultural replication, even when “corrected” by a conformist bias, imposes modest, but still significant forces on the cultural composition of the population. This means that only selective forces of similar magnitude will lead to cumulative adaptation. We do not think this is a problem because the selective forces acting on cultural variation are probably much stronger.
Figure 6. (a) The combination of conformist bias and very high error rate ($u = 0.3$) acts to preserve variation between groups because common representations tend to increase in frequency. (b) When more successful individuals are more likely to be imitated, the more successful variant spreads.

than those that shape genetic variation because they work on shorter time scales, and are often driven by psychological not demographic events — e.g. the diffusion of innovations literature shows time scales of decades, not millennia (Rogers 1995).

**Model 3: Adaptive evolution in a model without accurate replication or discrete representations**

Evolutionary theory predicts, and both field and laboratory data confirm, that individuals have a psychological propensity to copy particularly skillful, successful and prestigious individuals. We call this *prestige-biased transmission*; Henrich and Gil-White (2001) lay out the theory and summarize the empirical data. This tendency creates a selective force that can, under the right circumstances, generate cumulative adaptation. It is easy to see how this would work if cultural replication were accurate. Suppose populations vary in their ability to perform some important task, say prey tracking or arrow manufacturing techniques. Further, suppose that in choosing models, people preferentially imitate the techniques used by the best trackers in the group. If the mental representations that underlie the success of the best trackers can be accurately copied, the mean tracking success of the population will increase through time (Boyd & Richerson 1985, Ch. 8) because good trackers will leave more cultural ‘descendants.’ However,
Sperber (1996) and Atran (2001) have argued that, because cultural representations are not discrete, and cultural replication is inaccurate, selective forces, such as prestige-biased transmission, are unlikely to be important (Note, Atran et al. (in press) now modifies that position).

Here we show that prestige biased transmission can lead to cumulative adaptation, even when cultural transmission is inaccurate and representations are not discrete. Consider a population of \( N \) individuals who are numbered \( i = 1, \ldots, N \). Individual \( i \) has a \( z \)-value \( (z_i) \). This value measures the individual’s skill in some domain like canoe making, arrow manufacture or medicinal plant selection. Later we will show that \( z \) may measure either an individual’s mental representation (e.g. how long they think arrows should be), or some phenotypic measure that aggregates many skills or measures success (like lifetime tapir kills). However, for now we assume that it is a mental representation. Each individual is also characterized by a variable \( f \) that specifies the relative likelihood that an individual will be chosen as a cultural model. If people’s social learning attention is drawn to skilled hunters (and larger values of \( z \) lead to hunting skill), then \( f \) and \( z \) will have a positive partial regression coefficient (other things can affect \( f \) as well). The pool of individuals could represent the same group of individuals (same farmers) from one year to the next, or it could be different generations or cohorts.

Once again, we use the Price Equation to study the combined effects of selective transmission and inaccurate inference on continuous representations.

\[
\Delta \tilde{z} = \underbrace{\text{Selective Transmission}}_{\text{Cov}(f_i, z_i)} + \underbrace{\text{Incomplete Inference}}_{E(f_i \Delta z_i)}
\]  

(11)

\( \Delta \tilde{z} \) is the change in the average value of \( z_i \) per time step. If \( \Delta \tilde{z} \) is positive, then adaptive evolution is taking place because, for example, people are becoming better trackers or arrow makers. \( \text{Cov}(f_i, z_i) \) is the covariation between \( f_i \) and \( z_i \), and gives the effect of selective cultural forces on \( \Delta \tilde{z} \). In this case, it captures our psychological tendency to copy successful/skillful people — here we will assume that all learners attempt to copy the most skilled individual. \( E(f_i \Delta z_i) \) is the replicability-weighted average effect of all the individual intra-psychic transformations on cultural representations.

To capture the idea that inferential processes are incomplete and inaccurate, we assume that the inferential processes that underpin social
learning are inaccurate in two senses: first, they are noisy, so that copiers never accurately replicate the $z$-value of their model, and second, they are biased so that the behavior acquired by copiers is, on average, less skilled than the behavior of their model. More formally (as illustrated in Figure 7), individuals who attempt to copy a model with $z$-value, $z_i$, end up with a $z$-value drawn from a Gumbel probability distribution\(^7\) with mode $z_i - \alpha$ and dispersion $\beta$. Typically, copiers construct representations that are on average worse than their model’s $z$-values by an amount $\alpha$, but occasionally — through lucky guesses or errors — individuals construct representations that yield $z$-values higher than their model. The probability of that occurring for an individual is the area under the distribution to the right of the dashed line (the model’s $z$-value). It’s also worth noting that the probability of an exact copy is zero, and the probability of two people arriving at exactly the same copy is zero — no replicas in this model.

With these assumptions, equation (11) becomes:

$$\Delta \bar{z} = -\alpha + \beta(\varepsilon + \ln(N))$$

(12)

The first term represents the effect of systematic errors and is always negative — it operates against adaptive evolution. In the second term, $\ln(N)$ is the natural logarithm of $N$, the number of social learners in the population. The parameter $\varepsilon$ is the Euler-gamma constant, which equals 0.577. The parameters $\beta$, $\ln(N)$ and $\varepsilon$ are all always positive, so the second term is always positive — and favors adaptive evolution. This means that adaptive cultural evolution depends on the relative sizes of the two terms. Interestingly, the two components of inference, systematic bias (measured by $\alpha$) and random noise (measured by $\beta$), have opposite effects on adaptive evolution. Systematic bias operates against adaptive evolution, while noise — the tendency of individuals to make different inferences from observing the operation of the same underlying representation — favors adaptive evolution. The more individuals tend to make different inferences, the faster cultural evolution goes (or the more likely it is to be adaptive). Similarly, the larger the population of social learners, $N$, the faster adaptive evolution proceeds (or the more likely selective forces will favor adaptive processes).

\(^7\)The specific form of this distribution does not qualitatively impact our results (see Henrich 2002 for more information).
Figure 7. A graphical representation of the assumptions of Model 3. Learners observe a model with a behavior $z_i$, given by the dotted line. They infer a value drawn from the probability density function. Their most likely value is $\alpha$ less than the models value, and since larger values of $z$ are better, this represents systematic error. However, there is some probability that individuals acquire larger values (this is the area under the curves to the right of the dotted vertical line).

Setting $\Delta \xi > 0$ and solving (12) yields the conditions under which selective cultural transmission will drive adaptive cultural evolution:

$$N^* > e^{\frac{a}{\beta - \epsilon}}$$

(13)

$N^*$ is the critical number of social learners necessary to produce cumulative adaptive cultural evolution for a specified set of inferential processes ($\alpha, \beta$). Figure 8 plots this inequality and graphically shows the conditions under which selective transmission will drive adaptive evolution.

This result provides several insights about cultural adaptation in a noisy environment. First, cumulative adaptive evolution is facilitated by both a large pool of social learners and an inferential tendency to make different “mistakes” (large $\beta$, a tendency for individuals to form quite different representations). This finding contradicts Sperber (1996) and Atran (2001), who have argued that because cultural transmission leads to different representations in each individual’s mind, cumulative cultural evolution is not possible. Second, no matter how poor individuals are at imitating,
there are some values of $\beta$ and $N$ that generate cumulative evolution — elsewhere Henrich (2002) has shown how a sudden drop in population size ($N$) initiated a process of technological devolution in Tasmania over the last 8,000 years. So, both the social environment and the nature of cognition influence the conditions for cumulative adaptation. Looking only at the inferential processes and the conditions for adaptation, what matters is not $\alpha$, but $\alpha/\beta$. So, if the typical inaccuracy of copies doubles ($\alpha$), while the variation in different representations triples, cumulative adaptation will still occur. However, the model also illustrates that in particular cultural domains — those with large $\alpha$ and small $\beta$ (e.g., some religious beliefs) — we may not expect to observe cumulative adaptation. Finally, having incorporated Sperber, Atran and Boyer’s claims about the nature of culture into this simple model, that bears no resemblance to genetic models (other than both being about evolutionary processes), we derived insights that are not obvious from verbal reasoning alone.

Sperber, Atran and Boyer have argued that evolutionary models cannot be applied to phenotypes (or public representations) alone; they are applicable only if there are identified ‘units’ of transmission. This is not the case, and suggests a failure to apprehend the more general nature of evolutionary processes (Boyd and Richerson 2000). Neither genetic, nor cultural evolutionary, models require ‘units of transmission’

![Figure 8.](image-url)
— genetic evolutionary models were providing insights into the nature of evolution long before anyone knew anything about DNA discreteness or replication, and genetic models of quantitative characters, like height, still ignore these “units.” The most general form of the Price Equation, for example, is exact for anything one can measure about the constituent parts of any evolutionary system. This feature allows us to study the evolution of, and interrelation between, phenotypically expressed behavior (“public representations”) and the epidemiology of the underlying mental representations. Suppose now, instead of a mental representation of a skill, \( z \) measures success in some domain such as hunting (perhaps quantified in lifetime tapir kills), combat (in ‘heads-taken’), canoe making, or farming (in sacks harvested per hectare of wheat sown). Equation (11) would still govern the evolution of \( z \). However, if we wanted to get at the underlying mental representations that produce particular values of \( z \), we would have to specify how each mental representation contributes to individuals’ behavioral expressions (to their success). For illustrative purposes, suppose \( z \) is hunting returns (a phenotypic measure) and \( y \) and \( \phi \) are mental representations related to prey pursuit time and arrow length — presumably there are many more relevant representations for hunting. Using a linear regression equation, we can express the causal relationship of between mental representations \( y \) and \( \phi \) on success, \( z \), as follows.\(^8\)

\[
z_i = \mu + \lambda_1 y_i + \lambda_2 \phi_i + \epsilon_i
\]

The \( \lambda \)'s give the relative contribution of an individual’s \( y \) and \( \phi \) mental representations to the observed success, \( z_i \). \( \epsilon_i \) gives uncorrelated random error, and \( \mu \) specifies the constant term. With this, we can express the change in the average value of representation \( y \) as:

\[
\Delta \bar{y} = \text{Selectiv Transmisson} \times \text{Incomplete Inference}
\]

But, because an individual’s likelihood of being selected as a cultural model depends on \( f \), and \( f \) depends on individuals observing \( z \) (e.g., hunting

\(^8\)In general, we can do this for any number of mental representations, and study the interaction of different mental representations.
returns), then this can be rewritten as,

$$
\Delta \tilde{y} = \lambda_1 \rho \text{Var}(y_i) + E(f_i \Delta y_i)
$$

(16)

where $\rho$ is the partial regression coefficient on $f$ on $z$. The introduction of $\lambda_1$ shows us that effective transmission of $y$ depends on how much it affects $z$. The $\Delta y_i$ term would have the same form as above; it depends on how difficult it was to infer the underlying $y_i$ by observing the models’ behavior. The reminder of the derivation proceeds as above.

**Conclusion**

Formal models help us develop a more complete of understanding of cognition and culture. Our minds are not well equipped to understand the population-level outcomes that arise from numerous minor interactions, weak cognitive biases, random errors, migration rates and micro-level decisions. The history of population ecology, epidemiology and evolutionary biology over the last 40 years provides strong evidence that simple mathematical models provide powerful tools for understanding such problems.

The simple models described in this paper yield several findings that challenge the untutored intuitions of many:

1. Strong attractors generate replicator dynamics, such that weak selective forces determine the final state of the system. The stronger the attractors, the better the assumption of discrete-trait replicator dynamics.

2. Conformist transmission can compensate for even very high error rates in social inference in a manner that violates the intuitions imported from genetics. Prestige-biased transmission combines with conformist transmission to spread adaptive representations; together they predict the conditions for inertia and change.

3. Adaptive cultural evolution can occur even when representations are continuous and inferences are biased against adaptation. Poor inferences can be compensated for by larger pools of social learners, or greater amount of random error (i.e., by the fact that people don’t all make the same inferential mistakes).

4. Discrete units of transmission are not necessary for adaptive evolution, Dawkins (1976, 1982) claims notwithstanding.
The crux of Sperber, Atran and Boyer’s position is that the transmission of culture requires domain specific cognitive mechanisms, and that therefore that population dynamic models of culture proceed from untenable assumptions. We accept that social learning, like all other forms of learning, requires innate expectations about objects in the environment and the nature of relationships among them. How these innate structures shape the human mind is obviously of great importance for understanding human culture. The mistake is to see these ideas as incompatible with making population dynamic models of cultural change. It will never be enough to focus on the mind and ignore the interactions between different minds. To keep track of such interactions some kind of population dynamic models will be necessary. What is needed is both more effort by coevolutionary theorists to incorporate rich cognition into formal models of social learning, and more effort by cognitive scientists to consider how innate cognitive structure interacts with social processes and the cognition of social learning to influence the epidemiology of representations and its associated behavioral products.

REFERENCES

ATRAN, S.


ATRAN, S., MEDIN, D., ROSS, N., LYNCH, E., VAPNARSKY, V., UCAN EK’, E., COLEY, J., TIMURA, C. & BARAN, M.


BARON, R., VANDELLO, J. & BRUNSMAN, B.


BLACKMORE, S.


BOND, R. & SMITH, P.B.


BOYD, R. & RICHERSON, P.J.


**BOYER, P.**

**CAMPBELL, J.D. & FAIREY, P.J.**

**CAVAILLI-SFORZA, L.L. & FELDMAN, M.**

**DAWKINS, R.**

**DENNETT, D.**

**HENRICH, J. & BOYD, R.**

**HENRICH, J. & GIL-WHITE, F.**

**HENRICH, J.**

**INSKO, C.A., SMITH, R.H., ALICKE, M.D., WADE, J. & TAYLOR, S.**

**LESLIE, A.M.**

**LUMSDEN, C. & WILSON, E.O.**

**PRICE, G.R.**
ROGERS, E.M.


SMITH, J.M. & BELL, P.A.


SPERBER, D.


YOUNG, H.P. & BURKE, M.A.