Losing the influence of a common ancestor

Analyzing the rate at which languages lose the influence of a common ancestor

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Analyzing the rate at which languages change can clarify whether similarities across languages are solely the result of cognitive biases or might be partially due to descent from a common ancestor. To demonstrate this approach, we use a simple model of language evolution to mathematically determine how long it should take for the distribution over languages to lose the influence of a common ancestor and converge to a form that is determined by constraints on language learning. We show that modeling language learning as Bayesian inference of $n$ binary parameters or $n$ binary constraints results in convergence in a number of generations that is on the order of $n \log n$. We relax some of the simplifying assumptions of this model to explore how different assumptions about language evolution affect predictions about the time to convergence; in general, convergence time increases as the model becomes more realistic. This allows us to characterize the assumptions about language learning (given the models that we consider) that are sufficient for convergence to have taken place on a timescale that is consistent with the origin of human languages. These results clearly identify the consequences of a set of simple models of language evolution and show how analysis of convergence rates provides a tool that can be used to explore questions about the relationship between accounts of language learning and the origins of similarities across languages.
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Introduction

Human languages share a surprising number of properties, ranging from high-level characteristics such as compositional mapping between sound and meaning to relatively low-level syntactic regularities (Comrie, 1981; Greenberg, 1963; Hawkins, 1988). One explanation for these universal properties is that they reflect constraints on human language learning, with the mechanisms by which we acquire language only allowing us to learn languages with these properties (Chomsky, 1965). These mechanisms may also make certain languages more learnable than others, resulting in linguistic trends that favor particular properties (Krupa, 1982; Tily, Frank, & Jaeger, 2011). However, similarities across languages could also be partially the result of descent from a common ancestor (Bengtson & Ruhlen, 1994; Greenberg, 2002). While the theory of a common origin is controversial (Picard, 1998), recent work on phonemic diversity across languages has lent support to this theory (Atkinson, 2011), and there is evidence that some linguistic patterns, such as trends in word order, can be explained by lineage patterns in language evolution and that these patterns allow one to glean information about a common ancestral language (Dunn, Greenhill, Levinson, & Gray, 2011; Gell-Mann & Ruhlen, 2011).

Given that both a common ancestor and learning biases may contribute to similarities across languages, we wish to investigate how quickly the influence of a common ancestor is lost in the process of language evolution. As languages evolve, learning biases will become the dominant cause of similarities across different languages, and the influence of the common ancestor will eventually disappear. Modeling the process of language change makes it possible to determine how different assumptions about language evolution affect the rate at which this influence will dissipate. We demonstrate how this approach can be used, providing bounds on how quickly the influence of
a common ancestor will disappear in a simple model of language evolution. This allows us to identify assumptions about language learning (within this simple model) that result in bounds that are consistent with the origin of human languages. We then investigate how relaxing the simplifications of this model affects the rate of language change, and find that the relaxations tend to slow the rate at which the influence of the common ancestor is lost. These results provide a clear picture of the implications of the simple models that we consider, but more importantly they illustrate a methodology that can be applied to determine the implications of more complex and realistic models of language learning and language evolution.

Language transmission is a process in which those who are currently learning a language do so based on the utterances of other members of the population. These other members were also once language learners. Building on recent formal models of language evolution (Kirby, 2001; Smith, Kirby, & Brighton, 2003; Kirby, Smith, & Brighton, 2004; Kirby, Dowman, & Griffiths, 2007; Griffiths & Kalish, 2007) that share this feature of current learners learning from previous learners, the models we consider are based on *iterated learning*. Iterated learning models assume that each generation of people learns language from utterances generated by the previous generation. By modeling how languages change over many generations of transmission, the iterated learning framework provides an opportunity to examine how constraints on learning influence the process of language transmission. We begin by analyzing an iterated learning model that makes strong simplifying assumptions, such as a lack of interaction between learners in the same generation, allowing us to obtain analytic results quantifying how quickly languages lose the influence of an ancestor. The language of the initial generation is modeled as known, and each future generation has a distribution over possible languages. Previous research using this model has shown that after some number of generations, the distribution over languages converges to an equilibrium that reflects the constraints that guide learning (Griffiths & Kalish, 2007). After convergence, the behavior of learners is independent of the language spoken by the first generation. Prior to convergence, similarities across languages may reflect common ancestry. Our key
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contribution is providing asymptotic bounds on the number of generations required for convergence, known as the convergence time, which we obtain by analyzing Markov chains associated with iterated learning.

Bounding the convergence time of iterated learning is a step towards understanding whether similarities across languages are solely caused by constraints on learning or whether a common ancestor might also be contributing to these similarities. To bound the number of generations required for iterated learning to converge, we need to make some assumptions about the algorithms and representations used by learners. Following previous analyses (Griffiths & Kalish, 2007; Kirby et al., 2007), we assume that learners update their beliefs about the plausibility of a set of linguistic hypotheses using Bayesian inference. We outline how this approach can be applied in a simple model inspired by the Principles and Parameters framework (Chomsky & Lasnik, 1993; Gibson & Wexler, 1994; Niyogi & Berwick, 1996). In this model, grammars are represented as vectors of binary parameter values. We show that iterated learning with a uniform prior reaches equilibrium after a number of generations that is on the order of $n \log n$, where $n$ is the number of parameters.

By using a Principles and Parameters framework, our analysis assumes a finite hypothesis space where all languages can be represented using the same parameters. The convergence analysis is a tool for determining whether the ancestor language has an influence on descendant languages beyond this common representation. If it does not, then we could use the observed distribution over one of these parameters (e.g., word order) to draw conclusions about cognitive biases. Conversely, if the common ancestor still has an effect, we cannot draw strong conclusions about human biases based on this distribution, as there is a danger of underestimating the amount of variation that needs to be accounted for (such as the number of parameters required to represent the space of possible languages). Pairing the Principles and Parameters iterated learning model with data concerning the estimated amount of time since the origin of human languages, the number of parameters, and the rate of language change, we find that under this model it is possible that a common ancestor language (or languages) could still have some influence on similarities across modern languages.
The initial model makes a number of simplifying assumptions about the process of language transmission. As previously mentioned, we address the key simplifications by determining how the predictions of the model change when these assumptions are relaxed. These relaxations include using another formal model of language acquisition, inspired by Optimality Theory (Prince & Smolensky, 2004); modifying the assumed probability of different languages; allowing the parts of a language to change at different rates; incorporating learning from multiple previous generations; and including differential transmission of languages based on communicative success. All of these variations either reproduce our original results, or slow down convergence, in some cases dramatically. Considering this wider range of models thus reinforces our original conclusion – that under simple accounts of language learning it remains possible that a common ancestor could play a role in explaining trends across modern languages. These more comprehensive analyses also reinforce our methodological goal of showing how analysis of convergence rates can contribute to debates about the causes of similarities across languages, illustrating the steps that can be taken towards determining the convergence rates of more realistic models of language evolution.

A Simple Model of Language Learning and Transmission

To illustrate how convergence rates can be determined for models of language evolution, we focus on a simple model of language learning and transmission. This model has two parts: how language transmission occurs, and the nature of language learning. This section presents these two parts of the model in turn.

Transmission by Iterated Learning

Iterated learning has been used to model many aspects of language evolution, providing a simple way to explore the effects of cultural transmission on the structure of languages (Kirby, 2001; Smith et al., 2003; Kirby et al., 2004, 2007; Griffiths & Kalish, 2007). The basic assumption behind the model – that each learner learns from somebody who was themselves a learner – captures a phenomenon we see in nature: Parents pass on language to their children, and these
children in turn pass on language to their own children. The sounds the children hear are the input, and the child produces language (creates output) by combining this input with whatever constraints guide learning.

Formally, we conceptualize iterated learning as follows (see Fig. 1). A first learner receives data, forms a hypothesis about the process that generated these data, and then produces output based on this hypothesis. A second learner receives the output of the first learner as data and produces a new output that is in turn provided as data to a third learner. This process may continue indefinitely, with the \( t^{th} \) learner receiving the output of the \( (t - 1)^{th} \) learner. The iterated learning models we use in our initial analyses make the simplifying assumptions that language transmission occurs in only one direction (previous generations do not change their hypotheses based on the data produced by future generations) and that each learner receives input from only one previous learner. Later in the paper we examine the effects of relaxing the assumption of learning from a single learner.

**Insert Figure 1 about here**

We can analyze the consequences of iterated learning if we make some commitments about the way that learning take place. Our models assume that learners acquire languages by applying Bayesian inference. This makes it possible to express the degree to which learners are predisposed to certain hypotheses about language through a probability distribution over hypotheses. This *prior* distribution gives a probability \( p(h) \) to each hypothesis \( h \) in a hypothesis space \( H \).\(^2\) These probabilities might, for example, tend to favor word forms with alternating consonant-vowel phonemes, or grammars that support constituent structure. These constraints on learning are combined with data via Bayesian inference. The *posterior* distribution over hypotheses given data \( d \) is given by Bayes’ rule,

\[
p(h|d) = \frac{p(d|h)p(h)}{\sum_{h' \in H} p(d|h')p(h')}
\]

(1)

where the *likelihood* \( p(d|h) \) indicates the probability of seeing \( d \) under hypothesis \( h \). We assume
that learners’ expectations about the distribution of the data given hypotheses are consistent with
the actual distribution (i.e. that the probability of the previous learner generating data \(d\) from
hypothesis \(h\) matches the likelihood function \(p(d|h)\)). Finally, we assume that learners choose a
hypothesis by sampling from the posterior distribution, although we consider other ways of
selecting hypotheses later in this paper.\(^3\)

The analyses we present in this paper are based on the observation that iterated learning
defines a Markov chain. A Markov chain is a sequence of random variables \(X_t\) such that each \(X_t\) is
independent of all preceding variables when conditioned on the immediately preceding variable,
\(X_{t-1}\). The Markov chain is characterized by the transition probabilities, \(p(x_t|x_{t-1})\). There are
several ways of reducing iterated learning to a Markov chain (Griffiths & Kalish, 2007). We will
focus on the Markov chain on hypotheses, where the sequence of random variables corresponds to
the hypotheses selected at each generation. The transition probabilities for this Markov chain are
obtained by summing over the data from the previous time step \(d_{t-1}\), with
\[
p(h_t|h_{t-1}) = \sum_{d_{t-1}} p(h_t|d_{t-1}) p(d_{t-1}|h_{t-1})\] (see Fig. 1).

Identifying iterated learning as a Markov chain allows us to draw on mathematical results
concerning the convergence of Markov chains. In particular, Markov chains can converge to a
stationary distribution, meaning that after some number of generations \(t\), the marginal probability
that a variable \(X_t\) takes value \(x_t\) becomes fixed and independent of the value of the first variable in
the chain (Norris, 1997). Intuitively, the stationary distribution is a distribution over states in which
the probability of each state is not affected by further iterations of the Markov chain; in our case,
the probability that a learner learns a specific language at time \(t\) is equal to the probability of any
future learner learning that language. The stationary distribution is thus an equilibrium that iterated
learning will eventually reach, regardless of the hypothesis of the first learner, provided simple
technical conditions are satisfied (for details, see Griffiths & Kalish, 2007).

Previous work has shown that the stationary distribution of the Markov chain defined by
Bayesian learners sampling from the posterior is the learners’ prior distribution over hypotheses,
Losing the influence of a common ancestor (Griffiths & Kalish, 2007). This result illustrates how constraints on learning can influence the languages that people come to speak, indicating that iterated learning will converge to an equilibrium that is determined by these constraints and independent of the language spoken by the first learner in the chain. However, characterizing the stationary distribution of iterated learning still leaves open the question of whether enough generations of learning have occurred for convergence to this distribution to have taken place in human languages. Previous work has identified factors influencing the rate of convergence in very simple settings (Griffiths & Kalish, 2007). Our contribution is to provide analytic upper bounds on the convergence time of iterated learning with simple representations of the structure of a language that are consistent with linguistic theories.

Bayesian Learning of Linguistic Parameters

Defining a Bayesian model of language learning requires choosing a representation of the structure of a language. In this section, we outline a Bayesian model of language learning inspired by the Principles and Parameters theory (Chomsky & Lasnik, 1993), a proposal for how to characterize a Universal Grammar that captured all learnable languages. Under this theory, all languages are assumed to be representable using only a small number of parameterized principles, with the values of the parameters being set through exposure to language. This places strong constraints on the space of possible languages, downplaying the role of learning in language acquisition. These strong constraints provide one reason to use this as a starting point for our investigation of the rate of convergence of iterated learning: we might expect that reducing the space of possible languages to a finite discrete set would mean that convergence occurs quickly, giving us a bound on convergence for richer representations of the space of possible languages. Bayesian models of language learning consistent with other linguistic theories are also possible, and we consider the effect of adopting different theoretical approaches later in the paper.

The Principles and Parameters theory postulates that all languages follow a finite set of
principles, with specific languages defined by setting the values of a finite set of parameters (Chomsky & Lasnik, 1993). For example, one parameter might encode the head directionality of the language (with the values indicating left- or right-headedness), while another might encode whether covert subjects are permitted. We will assume that parameters are binary, as in previous models of language acquisition based on Principles and Parameters (Gibson & Wexler, 1994; Niyogi & Berwick, 1996). Learning a language is learning the settings for these parameters. In reality, learning is not an instantaneous process. Learners are presented with a series of examples from the target language and may change their parameters after each example. The exact model of learning varies based on assumptions about the learners’ behavior (e.g., Gibson & Wexler, 1994; Niyogi & Berwick, 1996). We do not model this fine-grained process, but rather lump acquisition into one computation, wherein a single hypothesis $h$ is selected on the basis of a single data representation $d$.

To specify a Bayesian learner for this setting, we define a hypothesis space $H$, a data representation space $D$, a prior distribution $p(h)$, and a likelihood $p(d|h)$. Assuming a set of $n$ binary parameters, our hypothesis space is composed of all binary vectors of length $n$: $H = \{0, 1\}^n$. We represent the data space as strings in $\{0, 1, ?\}^n$, where 0 and 1 indicate the values of parameters that are fully determined by the evidence and question marks indicate underdetermined parameters. For now, we assume a uniform prior, with $p(h) = 1/2^n$ for all $h \in H$. To define the likelihood, we assume the data given to each generation fully specify all but $m$ of the $n$ parameters, with the $m$ unknown parameters chosen uniformly at random without replacement. Then, $p(d|h)$ is zero for all strings $d$ with a 0 or 1 not matching the binary vector $h$ or that do not have exactly $m$ question marks (i.e. those consistent with $h$). Moreover, we assume that $p(d|h)$ is equal for all strings consistent with $h$. There are $\binom{n}{m}$ strings consistent with any hypothesis, so $p(d|h) = \frac{m!(n-m)!}{n!}$ for all $d$ consistent with $h$ (see Fig. 2).

Applying Bayes’ rule (Equation 1) using this hypothesis space and likelihood, the posterior
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The distribution is

\[
p(h|d) = \begin{cases} 
  \frac{p(h)}{\sum_{h':d|P(h')} h':d} & h' \vdash d \\
  0 & \text{otherwise}
\end{cases}
\]  

(2)

where \( h' \vdash d \) indicates that \( h \) is consistent with \( d \). This follows from the fact that \( p(d|h) \) is constant for all \( h \) such that \( h \vdash d \), meaning that the likelihood cancels from the numerator and denominator and the posterior is the prior renormalized over the set of consistent hypotheses. For a uniform prior, the posterior probability of a consistent hypothesis is simply the reciprocal of the number of consistent hypotheses. In the uniform case, \( 2^m \) of our hypotheses are valid, so \( p(h|d) = \frac{1}{2^m} \).

**Convergence Bounds**

We now seek to bound the time to convergence of the Markov chain formed by iterated learning. Bounds on the time to convergence are often expressed using *total variation distance*. This is a distance measure between two probability distributions \( \mu \) and \( \nu \) on some space \( \Omega \) that is defined as

\[
\| \mu - \nu \| \equiv \frac{1}{2} \sum_{x \in \Omega} |\mu(x) - \nu(x)| = \max_{A \subseteq \Omega} |\mu(A) - \nu(A)|
\]  

(Diaconis & Saloff-Coste, 1996). We seek to bound the rate of convergence of the distributions of the \( h^{(t)} \) to the stationary distribution, expressed via the number of iterations for the total variation distance to fall below a small number \( \xi \). This allows us to analytically determine how many iterations the Markov chain must be run to conclude that the current distribution is within \( \xi \) of the stationary distribution.

**Asymptotic Convergence Bounds**

To establish bounds on the convergence rate, we show that the Markov chains associated with iterated learning are reducible to Markov chains for which there are known bounds. As described above, we assume each learner receives sufficient data to set all but \( m \) of the \( n \) parameters in the hypothesis. We first consider the case where there is only one unknown parameter (\( m = 1 \)). In this case, each generation of iterated learning can change only one parameter at a time. This is equivalent to a random walk on a hypercube, where the hypercube has vertices with binary labels and each vertex is connected by an edge to only those vertices that differ in exactly one digit (see
We also assume that each vertex can transition to itself; this accounts for the case where a learner chooses the same parameter values as the previous generation. Previous analyses show that this Markov chain converges at the rate of $O(n \log n)$ (i.e. at a rate upper-bounded by some constant multiplied by $n \log n$) (Diaconis & Saloff-Coste, 1996). The multiplicative constant and lower order terms absorb the value of $\xi$, indicating the desired distance to convergence.

Now, we incorporate the fact that at each generation, there are $m$ parameters that the learner does not know, and thus up to $m$ parameters may change at each iteration. We assume the value of $m$ is fixed and constant across learners. Changing $m$ parameters at each step is equivalent to redefining an iteration as a collection of $m$ successive steps, each of which changes one parameter. Consider choosing which parameter to change at each step independently; this means that we might change a single parameter multiple times in one iteration. This process must converge in $O(\frac{n}{m} \log n)$ iterations, since each iteration in which we can change up to $m$ parameters is equivalent to $m$ steps in our original Markov chain. In our situation, however, we choose the $m$ parameters without replacement, so no parameter changes more than once per iteration. Since the net effect of changing the same parameter twice in one iteration is similar to changing it once (from the original value to the final value), changing $m$ different parameters brings us at least as close to convergence as changing fewer than $m$ different parameters. Thus, the Markov chain corresponding to our model converges in $O(\frac{n}{m} \log n)$ generations.

Simulation 1: Assessing the Bounds

To demonstrate the dependence of the convergence rate on $n$ and $m$, we simulated iterated learning for $m \in \{1, 2, 3\}$ and $n$ ranging from $m + 1$ to 10. For each simulation, the distribution $p(h^{(t)})$ over languages $h$ learned at generation $t$ can be calculated explicitly via the recursion

$$p(h^{(t)}) = \sum_{h^{(t-1)}} p(h^{(t)}|h^{(t-1)})p(h^{(t-1)})$$

We then define the convergence criterion as the number of iterations until $\|p(h^{(t)}) - p(h)\| < \xi$, with $\xi = 0.01$. We assume a worst case starting point; for the
case of the uniform prior, this is any state. Fig. 3 shows both the actual time to convergence as a
function of $n$ and $m$, and theoretical upper bounds on the time to convergence, equal to
$$\frac{n}{m} \log n + n \log \left( \frac{1}{\xi} \right)$$
(see Levin, Peres, & Wilmer, 2009, for incorporation of $\xi$ into the bound). As expected, we see that the time to convergence increases with $n$, and decreases proportional to $m$.
For relatively small $n$, the fact that time to convergence is increasing more than linearly is masked
by the size of the linear terms, but one can see that the results are consistent with the theoretical
upper bounds for the random walk on the hypercube.

**Implications for Continuing Influence of a Common Ancestor**

In the previous sections, we used iterated learning as a model of language evolution, and showed that the number of generations required to reach convergence given a representation of languages inspired by Principles and Parameters is $O\left(\frac{n}{m} \log n\right)$, where $n$ represents the complexity of the hypothesis space and $m$ represents the degree to which incoming data (language) limits the language adopted by the learner. This result has two intuitively sensible implications. First, the fidelity with which languages are transmitted between generations – reflected in the value of $m$ – has a direct effect on the rate of convergence. With higher fidelity, languages diverge from one another more slowly. While we have only considered integer values of $m$, our results also apply to fractional values, which indicate the expected number of parameters that will be selected at each generation. From historical records, we can estimate how frequently parameters change in order to empirically estimate a fractional $m$. For example, if a parameter is changed on average every ten generations, then the convergence time is bounded by estimating $m$ to be $0.1$.\(^5\) Thus, historical records provide a way of gaining insight into how accurately languages are transmitted and estimating the value of $m$. Second, this bound implies that when we vary $n$, the time to convergence increases a little more than linearly but the size of the hypothesis space increases exponentially. Thus, relatively rapid convergence should occur even with very large hypothesis spaces.
Our results illustrate how convergence rates can be obtained for models of language evolution. These convergence rates can then be used to draw conclusions about the nature of linguistic universals. For example, we can use the results obtained above to identify constraints on the size of the hypothesis space and the fidelity of learning necessary for human languages to have completely lost the influence of a common ancestor. This indicates how assumptions about language learning (within this particular simple model) translate into claims about the origins of similarities across modern languages. In the case of the Principles and Parameters model that has been our focus so far, the exact (rather than asymptotic) bound is known, being $\frac{4}{n} \log n$, and the bound itself tightly tracks the point at which convergence occurs in the model (Diaconis & Saloff-Coste, 1996). Consequently, we can identify exactly which values of $m$ and $n$ would guarantee convergence, given some assumptions about the number of generations over which languages have been evolving.

Obtaining good estimates of the number of generations since the origin of language, the number of parameters appropriate in a Principles and Parameters model, and the rate of language change is challenging. However, we can illustrate how the mathematical results can be used by selecting a reasonable estimate for the number of generations and then examining what values of $m$ and $n$ are required for convergence to take place in this interval. Estimates for the time since the origin of human languages range from 50,000 to 160,000 years (e.g., Holden, 1998; Atkinson, 2011), so we focused on a period corresponding to 8,000 generations (approximately 160,000 years if each generation is 20 years); we ignore issues of multiple languages arising spontaneously, which would lead to multiple iterated learning chains, as well as questions of how the initial language emerged. Fig. 4 shows how times to convergence that fall within this interval vary given various plausible values of $m$ and $n$; note that this bound slightly underestimates the time to convergence due to omitting terms of order lower than $n$. With small $n$ and relatively large $m$, the graph suggests that it is possible that language has been transmitted for a sufficient number of generations to allow convergence; conversely, if $m$ is much smaller or $n$ is too high, the figure
suggests that language may not have existed for sufficient time to have converged.

**Insert Figure 4 about here**

Several authors have estimated the number of parameters that might be required for a
Principles and Parameters model; these estimates range from as low as 20 to as high as 50-100
parameters (Kayne, 2000; Lightfoot, 1999). With \( n = 20 \) parameters, the number of parameters
that changes each generation has to be \( m > 0.0019 \) to guarantee convergence; \( n = 100 \) requires
\( m > 0.014 \). We can also obtain an estimate for \( m \) by observing the frequency of parameter changes
in historical records, and ask what values of \( n \) are required for convergence. For example, Taylor
(1994) found that ancient Greek changed word orders over about 800 years, and Hroarsdottir
(2000) provides evidence that it took 600-700 years for Icelandic word order to change. Such
estimates provide a lower bound on the time for a single parameter to be changed. The Greek word
order occurred in about 40 generations, and the Icelandic word order change took 30 – 35
generations. These rates correspond to \( m = 0.025 \) and 0.029 – 0.033 respectively, implying that \( n \)
would have to be less than 158 and 177 – 201 for convergence to take place within 8,000
generations. Based on these values, it is plausible that human languages could have lost the
influence of a common ancestor. However, if the time language has existed is a bit smaller than
8,000 generations, the actual number of parameters needed to define human language a little larger
than the estimates given above, or the rate of change a little slower, then it could be the case that
the influence of a common ancestor still affects the similarities we see across languages.

While the graph in Fig. 4 gives some understanding of how our analysis applies to actual
human languages, the model we have used is extremely simple. This model illustrates how the
approach of bounding convergence rates can be useful to the debate about the origins of linguistic
universals, but drawing stronger conclusions will require considering more realistic models of
language evolution. In the remainder of the paper we provide some preliminary steps in this
direction, relaxing a number of the simplifying assumptions in our model and assessing
convergence time via simulation in order to determine the effect that these assumptions have on
Relaxing Simplifying Assumptions

We consider a variety of modifications to the assumptions of the original model to ascertain how these assumptions contribute to our predictions about the rate of language change. For each modification, further technical details can be found in the Appendix, and simulation results are shown in Fig. 5.

Insert Figure 5 about here

Modifications to Learning

The first class of changes to the model retain the basic structure of iterated learning, but modify the assumptions about how learning takes place. In these models we can still calculate stationary distributions, and determine the total variation distance in simulations.

Optimality Theory. To determine whether linguistic representations other than Principles and Parameters produce asymptotic convergence rates similar to our original results, we consider another popular approach to modeling language learning. In Optimality Theory (OT), learning a language is learning the rankings of various constraints (Prince & Smolensky, 2004; McCarthy, 2004). These constraints are universal across languages and encode linguistic properties. For example, one constraint might encode that high vowels follow high vowels. Whether a construction is well-formed in a language is based on the ranking of the constraints that the construction violates. Specifically, well-formed constructions are those that violate the lowest-ranked constraints. Producing well-formed constructions thus requires determining how constraints are ranked in the target language. As we prove in the Appendix, using this linguistic representation gives the same asymptotic rate of convergence as for the Principles and Parameters representation; Fig. 5(a) shows this result in simulation.
Non-Uniform Priors. In our model, we have assumed that all languages are considered equally likely by learners. However, one might assume that there are certain languages that are easier to learn than others, making some languages more probable \textit{a priori}. We can examine the effect of this assumption by changing the entropy of the prior distribution: i.e., changing whether the prior makes all hypotheses equally likely or puts almost all weight on only a few hypotheses. Our previous analyses show how convergence time varies with the size of the hypothesis space, but non-uniform priors provide another way to model constraints on learning that might influence convergence. We examine the effects of using non-uniform priors by defining a distribution in which one language is designated a “prototype”, and the probability of all other languages decreases with distance from this prototype. As the entropy of the prior decreases, the upper bound on the time to convergence increases (see the Appendix for details).

Choosing a Hypothesis from the Posterior. In conducting our analyses, we assumed that learners sample from the posterior distribution over hypotheses, but there are other psychologically plausible methods of selecting hypotheses from the posterior. Alternative methods of selecting a hypothesis, such as selecting the hypothesis with the maximum posterior probability (MAP) and exponentiated sampling, have been considered in previous work (Griffiths & Kalish, 2007; Kirby et al., 2007). In the case of a uniform prior, both methods are equivalent to the sampling method we considered since all hypotheses with non-zero probability have the same probability in the uniform case; thus, our analyses of convergence time hold. In the non-uniform case, exponentiated sampling is equivalent to a variation on the simulation above concerning non-uniform priors (see the Appendix for details). For MAP in the non-uniform case, convergence to the prototype hypothesis (that with the highest probability in the non-uniform prior) will occur. The time in which this occurs is still $O(\frac{n}{m} \log n)$.\textsuperscript{6}

Non-Uniform Parameter Dropping. Another assumption in our original model was that each parameter of the grammar was equally likely to be dropped at each generation. However, one
might imagine that particular features of a language are less likely to be identified in the data than
others, which can be implemented by varying the probability of dropping each parameter. In this
model the distribution over languages converges to the prior distribution, but as shown in Fig. 5(c),
the time to convergence increases as the entropy of the distribution over dropping parameters
decreases (see the Appendix for details). Returning to the intuition that it is sufficient for
convergence to have had the opportunity to change each parameter once, it is reasonable that
making some parameters less likely to be dropped, and thus increasing the expected time to the
first iteration in which those parameters may be changed, will slow convergence.

 Modifications to Population Structure

One assumption of our original model that is clearly false is that there is a single learner at
each generation. In actuality, children learn from multiple people, and there is a population of
learners at each generation. One might believe that with this added complexity, the behavior of the
chains will differ significantly, making our analysis of the single learner chain less relevant. Thus,
we consider several ways that an iterated learning model might incorporate interaction between
multiple learners in order to see how this change affects convergence time. The additional
complexity of these models means that we cannot calculate the total variation distance exactly.
Instead, we use an empirical approximation of total variation distance described in the Appendix.
Additionally, these models do not necessarily have the prior as the stationary distribution when the
prior is non-uniform. The stationary distribution still contains information for understanding
inductive biases, however, as it is a combination of the prior and transmission artifacts of the
model. Convergence results remain important for these models because if we know that
convergence has not occurred in the model, we can conclude (as before) that based on the model,
common ancestor effects could still contribute to similarities across languages.

 Incorporating Communicative Success. The first way in which we consider interactions
between multiple learners is in a population fitness model similar to that used in other work on
language evolution (Komarova & Nowak, 2003). In this model, multiple learners exist at each generation, and each learner learns from a single parent. The learner samples a parent to learn from based on the parents’ fitness, which is a measure of the communicative efficacy of a speaker based on the languages spoken by those in the population. We considered two fitness models: the first lowers fitness additively for deviations in the language, and the second lowers fitness multiplicatively. Learners’ languages are more correlated in the latter model than the former, and learners in the latter model are more likely to choose the same parent as one another. As shown in Fig. 5(d), using either fitness function increases the time to convergence, but there is a much stronger slowing effect for the multiplicative function than for the additive function.

**Learning from Multiple Parents.** Another way of considering a population of learners is to assume that all learners learn from all parents in the previous generation, regardless of fitness. This also results in a longer convergence time. This occurs because learners tend towards the same hypotheses at each generation, meaning any given chain tends to have one dominant hypothesis that does not change for many iterations. The increase in convergence time becomes significantly greater with each additional parent; this is due to each parent reducing the chance that the dominant hypothesis will change at the next iteration. The slowing effect of multiple parents is also much greater than the slowing effect of the population fitness models.

**Learning from Multiple Generations.** Finally, our model assumes that learners learn only from the immediately preceding generation. However, real learners might be expected to be influenced by writing, which represents language produced by a previous generation, and by members of older generations still present in the population. Thus, we consider a model in which learners learn from multiple preceding generations, and we again find that the time to convergence slows. Fig. 5(f) shows that results are similar for this simulation and the previous simulation, although time to convergence is slightly smaller for learning from \( k \) generations rather than \( k \) parents.
General Discussion

Understanding the rate at which languages change can help explain whether similarities across languages may be partially attributable to a common ancestor, and for determining the implications of different assumptions about how languages are acquired and transmitted. To illustrate the utility of this approach, we analyzed the rate at which the influence of a common ancestor is lost in an iterated learning model in which each generation learns using data from the previous generation. This model forms a Markov chain, allowing us to use existing mathematical techniques to analyze how long the chain takes to converge to its stationary distribution, becoming independent of its initialization. For the simplest iterated learning model we considered, we showed that that the model converges in time $O\left(\frac{n}{m} \log n\right)$, where $n$ defines the size of the space of languages under consideration and $m$ defines the fidelity of transmission. By relaxing some of the simplifying assumptions of the model, we showed that in some cases, the rate of language change is unaffected; in other cases, language change is slowed. Using estimates of $n$ and $m$ drawn from the literature, this analysis suggests that some similarities across languages could be caused by the influence of a common ancestor. Such a result does not imply that many similarities are not due to constraints on learning; rather, the model suggests that if convergence has not occurred, then both a common ancestor and constraints on learning may influence at least some of the properties of modern languages. The conclusions we can draw from these models are limited by the simplicity of their assumptions, but our analyses illustrate how obtaining convergence rates for models of language evolution can be used to draw conclusions about the origins of similarities across languages, paving the way for the investigation of these questions using more realistic models.

In the remainder of the paper, we situate the models we have used within the broader literature, consider the limitations of our analysis, and highlight some directions for future work.
Relation to Other Models

We analyzed the rate of convergence of models of language evolution in order to understand how quickly the influence of a common ancestor is lost. Niyogi (2006) have considered this question for several other language learning algorithms. There also exist a variety of other approaches that can assist in answering this question. One such approach is the reconstruction of earlier languages based on their descendant languages. Such approaches often additionally examine how quickly the descendant languages diverged (Swadesh, 1952; Evans, Ringe, & Warnow, 2004). While these efforts remains controversial, the fact that they are even partially successful suggests that at least some evidence of the common ancestor can be traced back for a significant period. As mentioned at the beginning of this paper, reconstruction work has also found that current patterns across languages can be used to recapture some information about a common ancestral language (Dunn et al., 2011; Gell-Mann & Ruhlen, 2011). Again, this supports the claim that there are similarities across languages that are not solely caused by biases in human learning or innate linguistic universals. We see our results as a complement to this existing work that can help to clarify how modifications to a model of language evolution can affect time to convergence.

Limitations of the Analysis

We have already discussed some of the challenges posed by connecting our mathematical results to real processes of language change and considered some of the ways in which our models can be modified to be more realistic. In this section, we consider some of the more technical limitations of our analysis. For the mathematical analysis of convergence rate, we gave asymptotic bounds, which ignore proportionality constants. Such constants are independent of the relationship between the space of possible languages and the rate at which errors in learning occur (as defined by $n$ and $m$) but can have practical implications for applying the bound to understanding the rate of language change in the world. In the case of the bound we showed for the Principles and Parameters model, this proportionality constant is known; this fact was used to make Figure 4
showing convergence rates based on one’s beliefs about the values of $n$ and $m$ and to interpret the consequences of plausible estimates of $n$ and $m$ drawn from the literature. For the extensions of the model, the proportionality constants are not known, and in many cases, we have not provided explicit bounds. Instead, one must use simulation results to assess how quickly a particular model of language change will converge. Establishing bounds with known constants is a significant challenge, but one that we hope may ultimately be addressed through advances in the mathematical analysis of Markov chains.

A second limitation of our analysis is that we have only considered language representations that result in a discrete and finite set of possible languages. By focusing on hypothesis spaces based on Principles and Parameters and Optimality Theory, our analysis assumes that languages that cannot be represented by these systems are unlearnable. This is a strong assumption, and necessarily dictates that certain linguistic universals are due to constraints on human learning. One could relax this assumption by considering hypothesis spaces with an infinite number of languages. However, our analyses relied on results from the analysis of discrete Markov chains to assess the rate of convergence, and these results do not necessarily generalize when we consider language representations that allow infinite numbers of languages. In our formulation of the problem, converging to a stationary distribution is an instance of a random walk over a space of possible languages. It has been shown that discrete random walks will always converge to a stationary distribution given relatively simple technical conditions, while continuous random walks do not always converge to a stationary distribution (Woess, 2000). Thus, it is possible that such an analysis would predict that the influence of the common ancestor would never be completely lost. The results of Griffiths and Kalish (2007) do generalize to continuous Markov chains, indicating that we should expect models of language evolution based on iterated learning with Bayesian learners to have stationary distributions. However, obtaining convergence rates for these models remains a challenge, except in the simplest cases.
Future Work

We view our work as a first step in understanding how different assumptions in models of language evolution affect the rate at which the resulting languages lose the influence of a common ancestor. There are many ways in which this work could be extended. We have shown results for an arbitrary prior distribution. However, the structure of this distribution, which reflects cognitive biases, might be restricted in ways that shorten time to convergence. For example, if certain structures are unlearnable (as discussed in, e.g., Hunter, Lidz, Wellwood, & Conroy, 2010; Pinker & Jackendoff, 2009), then the number of possible languages is effectively decreased, lowering time to convergence.

While we considered a variety of relaxations of our original assumptions about language transmission, addressing what we saw as the most unrealistic assumptions, there remain a variety of possible relaxations that we did not consider. The framework we have presented allows one to consider how new variations might affect convergence time, and given the similarity of results across different relaxations, we believe that most other relaxations which bring the model closer to human language would also slow convergence time. We have examined relatively few relaxations that incorporate communicative pressures. There are a number of other ways of conceptualizing these pressures (e.g., Tily, 2010), that might lead to somewhat different results. Such pressures might decrease the time for an improbable feature of the common ancestor to disappear, but could conversely increase the time for improbable but possible variants to be attested.

There are also influences on language change that do not fit as easily into the iterated learning framework that we have presented. One such influence is geographic factors. Languages that are spoken by neighboring groups are likely to influence one another. To incorporate this factor into iterated learning, one might augment each learner with a location and have multiple descendants from each parent, each of which may migrate to a slightly different location. Learners may then learn both from their parent as well as from those who are geographically close to them. Such a model is likely to have different convergence properties than the iterated learning models
we considered and would be a combination of models of language learning and human migration.

Social factors, including deliberate language changes by social groups to differentiate from one another (Labov, 2001), are also outside the scope of the analysis we have presented. While these factors clearly result in language change, it is less obvious how to incorporate them into our mathematical model of language change and thus what effect they have on convergence time. Are the changes a straightforward application of cognitive biases, making them the equivalent of increasing $m$ and thus decreasing convergence time, or are the new languages derived from the old languages such that constraints on learning are less relevant than in cases where the fidelity of transmission is limited? Due to such complexities we did not incorporate social factors into our analyses, but this is clearly a direction that deserves further investigation.

Conclusion

Answering questions about the history of modern languages is challenging. Computational models provide a set of tools that can be used to explore the implications that different assumptions about language evolution have for this history. In this paper, we have shown how the question of whether the similarities between modern languages are partially due to the influence of a common ancestor can be expressed mathematically in terms of the convergence rates of stochastic processes that result from models of language evolution. We have illustrated how this approach can be used, analyzing the implications of a simple model of how languages are learned and transmitted. This analysis can be used to link theoretical and empirical results about language learning and language change to answers about the origins of similarities across languages. Drawing stronger conclusions will require analyzing more realistic models. We took a first step in this direction by exploring the effects of changing the assumptions behind our model, showing that these changes generally maintained the conclusions drawn from the original model. We hope that the methodology we have adopted allows others to take further steps along this path, ultimately yielding answers to some of the deep questions about the nature of linguistic universals.
References


Losing the influence of a common ancestor

Oxford: Blackwell.


Appendix

Relaxing Model Assumptions

We provide additional information below concerning the technical details of simulations. Additionally, Fig. 5 in the main text shows the results of simulations for each modification to the basic model.

Optimality Theory

To specify a Bayesian learner that uses a representation inspired by Optimality Theory (OT), we must identify the hypothesis space $H$, data space $D$, prior $p(h)$, and likelihood $p(d|h)$. In this representation, each hypothesis is an ordered list of $n$ constraints, with the order of constraints representing rank. The hypothesis space $H$ is thus the symmetric group of permutations of rank $n$, $S_n$, and is of size $n!$. We assume learners see sufficient data to specify the relative ordering of all but $m$ constraints. The data space is then strings over $\{1, 2, \ldots, n\}$ of length $n - m$, with no repeated elements, ordered from left-to-right in order of precedence (see Figure A1). The relative ordering of the $n - m$ specified constraints is maintained exactly from the generating hypothesis. We again see that the likelihood, $p(d|h)$, is 0 for all orderings not consistent with our hypothesis and equal for all consistent orderings. Analogously to the previous case, we select $m$ constraints to remove from the ranking randomly, giving $\binom{n}{m}$ possible data strings for each hypothesis. This gives $p(d|h) = \frac{m!(n-m)!}{n!}$. Thus, the posterior is the same as for the Principles and Parameters representation. Since we can freely permute $m$ of our parameters, we have $\frac{n!}{(n-m)!}$ consistent hypotheses for any data string $d$. If our prior is uniform, then $p(h|d) = \frac{(n-m)!}{n!}$ for all consistent $h$ and 0 otherwise.

To bound the convergence of iterated learning with an optimality theory representation, we assume that, at each generation, the learner has sufficient data to rank all but $m$ of $n$ constraints. First consider the case where $m = 1$. The process of changing the ordering of one item in a
Losing the influence of a common ancestor

permutation while leaving the relative ordering of the other items unchanged has been studied previously in the context of a random-to-random shuffle (see Figure A1). The best bound for the random-to-random shuffle is $O(n \log n)$ by Diaconis and Saloff-Coste (1993), with the intuitive argument being similar to that given for the Principles and Parameters convergence bound. As before, we view each iteration as $m$ successive steps, making time to convergence $O\left(\frac{n}{m} \log n\right)$.

Insert Figure A1 about here

Non-Uniform Priors

We consider the case of learners who have a non-uniform prior over the languages in the hypothesis space. The uniform prior is the unique maximum entropy distribution for any hypothesis space. However, there is no unique solution for achieving a given entropy for a distribution with $k$ values. Thus, we altered entropy in the following, non-unique way. We define one hypothesis $h_p$ as the prototypical hypothesis. Then, we calculate the distance between $h_p$ and $h$ for each hypothesis $h$ using the Hamming distance $\Delta$. Then, for all $h$, $p(h) \propto \exp(-\beta \Delta(h, h_p))$.

Changing $\beta$ changes the entropy of the distribution. Changing entropy in this manner gives our priors a characteristic shape: $h_p$ has maximum probability, and the probability of other hypotheses decreases with distance from $h_p$. For an evenly spaced range of $\beta$ (from 0 to 6 in increments of 0.05), we calculated the time to convergence when the resulting prior was used, fixing $n = 6$ and $m = 2$. As the entropy of the prior decreases, the time to convergence increases, as mentioned in the main text. This result differs from previous simulations reported in earlier work because it is worst-case time bound rather than an expected time bound.

Choosing a Hypothesis from the Posterior

In the main article, we note that exponentiated sampling from a non-uniform distribution is equivalent to using a prior that has been transformed. In particular, raising the posterior to the power of $\gamma$ before sampling is equivalent to multiplying the $\beta$ parameter in the model we used to
construct our non-uniform priors by $\gamma$.

**Non-Uniform Parameter Dropping**

To model the situation in which certain parameters are more likely to be dropped at each iteration than others, we put a non-uniform distribution over the probability that any particular parameter will be dropped conditioned on the total number of parameters to drop. We consider the parameter in each place $i = 1, \ldots, n$, and define the probability of dropping the $i$th parameter given $m$ as proportional to $\exp(-\alpha(i - 1))$. Assuming that the probability of dropping each parameter is independent of the others given this conditioning (e.g., dropping the first parameter does not make dropping the second parameter more or less likely), then we can calculate the probability of a particular hypothesis given $m$ as $\prod_{i=1}^{n} p(d_{i})^{I(d_{i} = ?)}(1 - p(d_{i})^{1-I(d_{i} = ?)})$. With this definition, we can calculate transitions exactly, as in previous simulations. Time to convergence increases as the entropy of the distribution over dropping parameters decreases.

**Modifications to Population Structure**

We consider several ways that an iterated learning model might incorporate interaction between multiple learners in order to see how this change affects convergence time. The additional complexity of these models means that we cannot calculate the total variation distance exactly. Instead, we use an empirical approximation of total variation distance, described below.

Quantitatively, we analyze convergence for the following simulations by computing an empirical total variation distance. For each simulation, we run many parallel chains, all with the same starting point. We choose the starting point likely to cause the slowest convergence by starting all learners in the same language in those cases where there are multiple learners per generation. We then average the language distributions across chains at each iteration, giving an average language distribution for each iteration of the simulation. The total variation distance between this distribution and the prior can then be calculated. While the stationary distributions for these simulations may not be equal to the prior distribution in the general case, by symmetry the
prior is the stationary distribution in the uniform case. Thus, we are able to calculate the total variation distance between the language distribution at each iteration and the stationary distribution.

This empirical total variation distance raises another issue, however, due to the fact that we have a finite sample. Even if samples were drawn directly from the prior distribution, the total variation distance between a distribution over those samples and the prior distribution would be non-zero. To handle this issue, we developed the following empirical convergence criterion. We would like to find a cutoff point for the total variation distance such that when we reach this cutoff, we have converged. Given a simulation to be checked for convergence, samples are drawn from the stationary distribution at each iteration. The number of samples drawn per iteration is equal to the number of simulated chains. Then, after drawing all samples for all iterations, we can again average our distributions and calculate total variation distance. To ensure convergence, we choose a cutoff value such that for 95% of iterations based on draws from the prior, the total variation distance is less than the cutoff. By choosing such a cutoff value and starting all learners at the “worst” point for convergence, we find an upper bound for time to convergence.

There is one remaining issue: All of the simulations below involve multiple learners per generation, but the hypotheses of the learners are likely to be highly correlated. Thus, it would not be correct to sample the distribution as a collection of independent draws from the prior: Doing so would make many simulations fail to converge by this empirical criterion even after they have clearly converged by other measures. We thus treat the multiple correlated learners as equivalent to one learner. Consequently, we draw a sample for only one learner per iteration per simulated chain.

Population Fitness Model

In the main paper, we consider a population fitness model in which multiple learners exist at each generation. Each learner has a single parent from whom language is learned, and parents are sampled based on their fitness. Fitness is a measure of the communicative efficacy of a speaker
based on the languages spoken by those in the population. For each generation, we define the
fitness of learner $i$ as $f_i = \sum_{j=1}^{n} p(\text{parameter}_j)$ where $p(\text{parameter}_j)$ is the proportion of
the population with the same value for parameter $j$ as speaker $f_i$. For example, if we have three
learners speaking languages 1010, 1000, and 0010, the first will have $f_1 = 2/3 + 1 + 2/3 + 1 = \frac{10}{3}$,
while the other two will have $f_2 = f_3 = 2/3 + 1 + 1/3 + 1 = 3$. We call this the additive function for
population fitness. We also consider a multiplicative function for population fitness. In this case,
$f_i = \prod_{j=1}^{n} p(\text{parameter}_j)$. Intuitively, the multiplication function penalizes learners more strongly
for having a parameter value that differs from the norm. In the additive example, the first learner
has only slightly higher fitness than the other two. Using the multiplicative function, the first
learner would have $f_1 = \frac{4}{9}$ while for the other two, $f_2 = f_3 = \frac{2}{3}$. Since the probability of learning
from parent $i$ is proportion to $f_i$, this means that for the additive function subsequent learners
would be $\frac{10}{9}$ times as likely to choose parent 1 as to choose parent 2 or parent 3, while with the
multiplicative function the learners would be twice as likely to choose parent 1. Iterated learning
then proceeds as in the original case except that each learner in a generation may be learning from
a different parent.

For this simulation and the following two simulations, the qualitative behavior of the
simulations is affected by the expected rate of parameter change ($m$). Asymptotically, Komarova
and Nowak (2003) show that assuming the noise level in learning is not too high, learners will
converge to a single language with high probability. In our case, this is equivalent to saying that if
$m$ is not too high for our population fitness chain, as compared to $n$, then learners will show
linguistic coherence. This simply means that after sufficient iterations have passed, members of the
same generation in later iterations will share the same language as one another.

For each fitness function, multiplicative and additive, we simulated how the time to
convergence was affected by the number of learners at each generation (ranging from 2 to 20). All
simulations used $n = 5$ and $m = 1$. The empirical total variation distance was calculated by
determining the distribution over languages at each generation averaged across 1000 chains, as
described above. The worst case starting point for these simulations is to set all learners at the initial generation to have the same language, although any particular language can be chosen due to symmetry.

Both population fitness simulations have some slowing effect on time to convergence as the number of learners increases. However, this effect is extremely small when using the additive function: when the number of learners is increased from 2 to 20, on average only 8 more iterations are required for convergence. We see a much stronger effect for the multiplicative function, and the effect in this case appears to be exponential. The reason for the slowing is that the dominant language in one generation is closely coupled with the dominant language in the next generation. This coupling is more extreme when the multiplicative fitness function is used. Qualitatively, we found that the dominant language of the population stays the same over long periods. Thus, the Markov chain for this simulation is slower to converge due to the tendency to remain at a language favored by the starting point.

**Learning from Multiple Parents**

Another way of considering a population of learners is to assume that all learners learn from all parents in the previous generation, regardless of fitness. This requires defining a likelihood \( p(d|h) \) where \( d \) is a collection of \( \ell \) data vectors, one from each parent. To account for the fact that different parents may be producing data from different hypotheses, leading to the possibility of data with conflicting settings for a single parameter, we add a noise parameter \( \varepsilon \) to our model. We can then calculate \( p(d|h) \) as:

\[
p(h|d) = \prod_{i=1}^{\ell} p(h|d_i) \quad (3)
\]

\[
\propto \prod_{i=1}^{\ell} p(d_i|h) p(h) \quad (4)
\]

\[
\propto \prod_{i=1}^{\ell} \prod_{j=1}^{n} e^{I(h_j \neq d_i, d_j \neq ?)} (1 - \varepsilon)^{I(h_j = d_i)} \quad (5)
\]
where \( I(\cdot) \) takes the value 1 when its argument is true and 0 otherwise and \( d_{ij} \) is the \( j \)th parameter of data vector \( i \). With this definition of \( p(d|h) \), we can simulate iterated learning steps as in the original case.

We varied the number of learners per generation, as well as \( \varepsilon \), to determine the effect on time to convergence. For all simulations, we set \( n = 5 \) and \( m = 1 \), and calculated the empirical total variation distance using 1000 chains per simulation. As in the population fitness case, we used the worst-case starting point in which all members of the initial generation share the same language. Simulation results showed that learning from multiple parents increases time to convergence and as \( \varepsilon \) decreases, time to convergence increases further.

**Learning from Multiple Generations**

The technical details of the model for learning from multiple generations closely follow the details of the model for learning from multiple parents. We tested how varying the number of generations from which each learner learned as well as adjusting the noise parameter \( \varepsilon \) affected the time to convergence. Fixing \( n = 5 \) and \( m = 1 \), we ran 1000 chains per simulation to calculate the empirical total variation distance. The chains began with all previous generations sharing the same language, as this is the worst-case starting point. Learning from multiple generations also slows convergence.
Author Note

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Footnotes

1 Note that each generation may consist of a single learner or of a population of learners. Similar analytic results apply in both cases (Griffiths & Kalish, 2007); we primarily consider the case of a single learner at each generation.

2 Throughout this paper, we assume a finite space of possible languages in order to make computation tractable. In the Discussion, we consider possible implications of an unbounded space of languages.

3 Note that these various probabilities form our model of the learners. Learners need not actually hold them explicitly, nor perform the exact computations, provided that they act as if they do.

4 An intuition for this result comes from the following argument. A sufficient condition for convergence to the uniform prior we have assumed is that all parameters have been left unspecified in the data at least once. This is true because each time a parameter is left undefined, its new value is insensitive to its current value. The result after all parameters have been left unspecified at least once is then equivalent to drawing a vector of values uniformly at random. The time to convergence is thus upper-bounded by the time required to sample all parameters at least once. This is a version of the coupon-collector problem, being equivalent to asking how many boxes of cereal one must purchase to collect \( n \) distinct coupons, assuming each box contains one of the \( n \) coupons and coupons are distributed uniformly over boxes (Feller, 1968). The first box provides one coupon, but then the chance of getting a distinct coupon in the next box is \( (n - 1)/n \). In general, the chance of getting a distinct coupon in the next box after obtaining a total of \( i \) different coupons is \( (n - i)/n \). The expected time to find the next coupon is thus \( n/(n - i) \), and the expected time to find all coupons is \( n \sum_{i=1}^{n} \frac{1}{i} \), or \( n \) times the \( n \)th harmonic number. The bound of \( n \log n \) results from an asymptotic analysis of the harmonic numbers, showing that the largest term in the asymptotic approximation grows as \( \log n \) as \( n \) becomes large.

5 Since a parameter may be dropped but take a new value that is the same as the previous one,
the expected number of parameters to actually change per generation is upper bounded by rather than equal to the value of $m$.

6Intuitively, this rate can again be interpreted using the coupon-collector problem. At every step, the learner changes unknown parameters to match the prototype. The problem is thus still analogous to the coupon-collector problem, with the worst case being when all $n$ parameters differ from the prototype.
Figure Captions

Figure 1. Language evolution by iterated learning. (a) Each learner sees data, forms a hypothesis, and generates the data provided to the next learner. (b) The underlying stochastic process, with \(d_t\) and \(h_t\) being the data generated by the \(t\)th learner and the hypothesis selected by that learner respectively. (c) We consider the Markov chain over hypotheses formed by summing over the data variables. All learners share the same prior \(p(h)\), and each learner assumes the input data were created using the same \(p(d|h)\).

Figure 2. A simple model of language evolution. (a) Representation of a hypothesis and data item for Bayesian learning of linguistic parameters. On the left is a possible hypothesis for \(n = 3\); the center shows a possible data output derived from this hypothesis (with \(m = 1\)), and the right shows all hypotheses consistent with this data output. (b) Movement between hypotheses is analogous to a walk on the hypercube when \(m = 1\). Above, the corners (hypotheses) that could be reached after one step (iteration) beginning at 000 are shown.

Figure 3. Rate of convergence using a uniform prior and the Principles and Parameters representation. Time to convergence is proportional to \(\frac{n}{m} \log n\), and is measured in the number of generations to convergence. Gray lines show theoretical upper bounds for these simulations (based on convergence of a random walk on the hypercube), and black lines show simulation results.

Figure 4. Number of generations necessary for convergence, as a function of \(n\) and \(m\). The surface is shown for values that converge in 8,000 generations or fewer; the black line shows the two-dimensional projection of values of \(m\) and \(n\) for which convergence is estimated at 8,000 generations. This corresponds to 160,000 years, with 20 years per generation, which is an upper limit on the maximum amount of time language may have existed.

Figure 5. Effects of relaxing simplifying assumptions on time to convergence. (a) Optimality Theory representation: Results using this representation are similar to those for the Principles and
Parameters representation. (b) Entropy of prior distribution: As the entropy of the prior increases, indicating more uniformity, the time to convergence decreases. (c) Entropy of the distribution of the probability of dropping each parameter of the language vector: As the entropy of this distribution decreases, time to convergence increases dramatically. (d) Population fitness model: Convergence time is relatively constant with increasing population size if an additive fitness function is used, but increases with a multiplicative fitness function. (e) Learning from multiple parents: As the number of parents increases, the time to convergence also increases. (f) Learning from multiple generations: Showing a pattern very similar to that of learning from multiple parents, the time to convergence increases with the number of generations from which the learner learns.

*Figure A1.* Assumptions for model inspired by Optimality Theory (OT). (a) Representation of a hypothesis and data item for OT. The relative ordering of A and C is preserved in the data, but not the ordering of B. (b) Movement among hypotheses for the OT case is analogous to a shuffle in which a random card (in this case, the grey card) is removed and reinserted into a random spot.
Losing the influence of a common ancestor, Figure 1
Losing the influence of a common ancestor, Figure 2
Losing the influence of a common ancestor, Figure 3

![Graph showing the relationship between the number of parameters and convergence time for different values of m (1, 2, 3).]
Losing the influence of a common ancestor, Figure 4
Losing the influence of a common ancestor, Figure 5

ConsolidatedFigure.pdf
Losing the influence of a common ancestor, Figure A1

(a) \( h_{t-1} \quad d_{t-1} \quad h_t \)

\[
\begin{align*}
A & \quad B & \quad C \\
B & \quad A & \quad C \\
C & \quad B & \quad A
\end{align*}
\]

(b) Remove card

\[
\begin{align*}
A & \\
C & \quad B
\end{align*}
\]