From entropy to information: Biased typewriters and the origin of life

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So much has been written about the possible origins of life on Earth (see, e.g., the popular books (Deamer, 1994; deDuve, 1995; Koonin, 2011; Morowitz, 2004)) that it sometimes seems that—barring an extraordinary breakthrough in experimental biochemistry (for example (Patel et al., 2015)), or else the discovery of the remnants of an ancient biochemistry (Davies et al., 2009)—nothing new can be said about the problem. But such a point of view does not take into account that perhaps not all the tools of scientific inquiry have been fully utilized in this endeavor to unravel our ultimate origin on this planet. Indeed, Origin-of-Life research has historically been confined to a fairly narrow range of disciplines, such as biochemistry and geochemistry. Today, a much broader set of tools is being unleashed on this problem, including mathematical (England, 2013; Smith, 2008; Vetsigian et al., 2006) and computational approaches (Mathis et al., 2015; Nowak and Ohtsuki, 2008; Segré et al., 2000; Vasas et al., 2012; Walker et al., 2012). Computational approaches to the study of possible origins of life are often derided because they lack a particular feature of biochemistry, or “because they do not take into account the specific properties of individual organic compounds and polymers” (Lazcano and Miller, 1996). Such a point of view ignores the possibility that life may not be a feature that is dependent on a particular biochemistry (Benner et al., 2004), but could instead be a feature of any chemistry that is capable of encoding information.

If the one invariant in life is information (information about how to replicate, that is), it then becomes imperative to understand the general principles by which information could arise by chance. It is generally understood that evolution, viewed as a computational process (Adami, 1998; Mayfield, 2013) leads to an increase in information on average. The amount of information that evolution has accumulated to date differs from organism to organism of course, and precise numbers are not known. A rough estimate of the amount of information stored in an organism’s genome can be obtained by calculating the amount of functional DNA in an organism’s genome (Adami, 2002, 2012). For humans (assuming a functional percentage of about 8% (Rands et al., 2014)), this means that our DNA codes for about half a billion bits.

Almost all of the information contained in our genome (and any other organism’s)

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1 It is not necessary to consider epigenetic variation in the estimate of information content, as all epigenetic changes are performed by enzymes whose information is already stored within DNA.

2 This number is (given the functional percentage of 8%) an upper limit on the information content, as protein coding regions display considerable variation and redundancy, which lowers information. However, as open reading frames only account for 1% of the human
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owes its existence to the evolutionary process. But the algorithm that is evolution cannot
be at work in the absence of replication, and therefore cannot explain the origin of life.
It is in principle possible that the first replicator did not originate on Earth but rather ar-
rived on Earth from extra-terrestrial sources (Arrhenius, 1908; Hoyle and Wickramas-
inghe, 1981; Wickramasinghe, 2011). Even if that was the case, such an origin story
does not obviate the need for emergence somewhere, so we may ask generally: “What
is the likelihood of spontaneous emergence of information?”. The question in itself is
not new, of course. Howard Pattee asked as early as 1961, shortly after the discovery
of the structure of DNA (but before the discovery of the genetic code) (Pattee, 1961):

(1) How did a disordered collection of elements which forms sequences with no restric-
tions produce, with reasonable probability, enough initial order to result in the general
property of self-replication? (2) Assuming the general property of self-replication for
all sequences, how did those particular sequences which now exist arise, with reason-
able probability, from the set of all possible sequences?

In order to estimate the likelihood of spontaneous emergence of a self-replicator, it
is necessary to estimate the minimal information necessary to replicate, because the
length of the sequence is not a good indicator of fitness. A quick gedankenexperiment
can clarify this. Imagine that a symbolic sequence (written using ASCII characters) can
replicate if and only if anywhere on the string the exact sequence origins appears. This
is a 7 letter sequence, and the total number of possible sequences of length 7 is $26^7$, or
about 8 billion. The likelihood to find this sequence by chance if a billion sequences are
tried is, obviously, about 1 in 8. But suppose we try sequences of length 1,000. If we
only ask that the word appears anywhere in the sequence, increasing sequence length
obviously increases both the number of possible sequences and the number of self-
replicators. Thus, the likelihood to find a self-replicator does not scale exponentially
with the length of the sequence (it does not become $26^{1000}$), but rather with the of the
sequence (as we will see momentarily). In the present example, the information content
is clearly 7 letters. But how do you measure the information content of biomolecules?

**Information content of biomolecules**

Generally speaking, the information content of a symbolic sequence is equal to the
amount of uncertainty (about a particular ensemble) it can reduce. This information can
be written mathematically in terms of the entropy of the ensemble (described by the
random variable $X$ that can take on states $x_1, ..., x_n$ with probabilities $p_1, ..., p_n$

$$H(X) = - \sum_{i=1}^{n} p_i \log p_i \quad (7.0)$$

and the conditional entropy $H(X|s)$, where $s$ is the sequence whose information content
we would like to measure, as

$$I(s) = H(X) - H(X|s) \quad (7.0)$$

The latter entropy is given by the conditional entropy distribution $p_{|s}$ instead. So, for
example, the sequence Colonel Mustard reduces the uncertainty about the identity of the
genome and regulatory sequences (the other 7%) are much less redundant, the true
information content of human DNA is likely not much lower.
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murderer in a popular board game from log₂ 6 ≈ 2.83 bits to zero (as there are a priori six suspects, and the sequence fingers the perpetrator), so the information content is 2.83 bits. The sequence length, on the contrary, is 15 (counting the space as a symbol), which translates to 15 log₂(27) ≈ 71.3 bits. Thus, sequence length and information content can be very different: information is about something, while sequence length is just entropy.

Unfortunately, we cannot measure the information content of biomolecules in the same manner, because we do not know the entropy of the ensemble that the biomolecular sequence is information about. Let us call this random variable $E$ (for "environment"), as it represents the environment within which the sequence is functional, in the same sense that $X$ above was the environment within which the sequence Colonel Mustard is functional. However, an information-theoretical "trick" allows us to make progress. Let $s$ be a functional biomolecule (a polymer of length $L$), and its information content (per the formula above)

$$ I(s) = H(E) - H(E|s) \tag{7.0} $$

that is, it is the entropy of the “world” minus the entropy of the world given that we know $s$. We can also define the average information content as

$$ \langle I \rangle = \sum_s p(s) I(s) = H(E) - H(E|S) = H(E : S) \tag{7.0} $$

where $H(E : S)$ is the shared entropy between environment and sequences, but again that formula is not useful because we do not know $H(E)$. However, the formula can also be written as

$$ \langle I \rangle = H(S) - H(S|E) \tag{7.0} $$

in terms of the entropy of sequences $H(S)$ and the conditional entropy of the sequences given an average environment. This is also not useful, as the world is not an average of environments, but one very particular one $E = e$. Could we write this in terms of a difference of entropies as in (7)? We then would guess that

$$ I(s) = H(S) - H(S|e) \tag{7.0} $$

but equation (7) is not mathematically identical to (7), as the identity only holds for the averages. However, Eq. (7) can be derived from an approach embedded in Kolmogorov complexity theory (Adami, 1998, 2002; Adami and Cerf, 2000), where that equation represents the “physical complexity” of the sequence. Furthermore, (7) is practical to the extent that its value can be estimated. For example, as $S$ is the ensemble of sequences, its entropy is simply given by $\log N$, where $N$ is the total number of sequences of that length (it is possible to extend this formalism to sequences of varying length). Sequences with an arbitrary function in environment $E = e$ have an entropy smaller than $\log N$. Let us imagine that the number of polymers with that function (in $e \in E$) is $N_e$ (with $N_e \ll N$). Then (here we specify the base of the logarithm by the number of possible monomers $D$)

$$ I(s) = -\log_D N_e \frac{N}{N} \tag{7.0} $$

which, it turns out, is identical to Szostak’s “functional complexity” measure (Szostak, 2003). It allows us to quantify the information content of a biomolecular sequence if the “density” of functional sequences $N_e/N$ is known, and makes it possible to calculate the likelihood of emergence (by chance), of a molecule with information content $I$. As
the likelihood must be given by the density of molecules of that type within the set of all molecules of that length, we find

$$P = \frac{N_e}{N} = D^{-I},$$ (7.0)

where the relationship to information content follows directly from (7). Thus we see (as advertised earlier), that this likelihood only depends on the information content of the sequence, but not on its length. Below, we will test this prediction using the digital life system Avida and find it violated. However, the origin of this apparent violation is easily tracked down, and we are confident that the equality holds exactly in principle.

**Testing the likelihood of emergence by chance**

We first tested the likelihood to find the sequence origins by creating random ASCII polymers of length 7 using an alphabet of $D = 26$ (no spaces or other punctuation), and where each symbol was drawn from a uniform distribution over the letters a-z. When testing a billion sequences we did not find origins, which is in accord with the probability $P = 26^{-7}$ calculated above. Note that for ASCII strings (unlike the biomolecules) there is never any redundancy, so that $N_e = 1$ always. We then randomly searched for self-replicating sequences within the digital chemistry of the Avida Artificial Life system. (Adami, 1998; Adami and Brown, 1994; Ofria et al., 2009; Ofria and Wilke, 2004). In Avida, ASCII sequences can self-replicate, but only because these sequences are translated to instructions that are executed on virtual CPUs. In this sense, the sequences are really self-replicating computer programs, and because these sequences can mutate as they are copied, they evolve in a strictly Darwinian manner (see Table 1 for the arbitrary assignment of ASCII letters to avidian instructions). The Avida system has been used for over 20 years to test evolutionary dynamics (see, for example, the review (Adami, 2006) covering mostly the first ten years), and the likelihood of emergence of functional information (but not self-replication) has been studied in this system before (Hazen et al., 2007). (See also (Pargellis, 2003) for an investigation of spontaneous emergence of digital life in a related digital system).

The likelihood that any particular sequence coded within 26 instructions can replicate depends strongly on the meaning of each instruction. If a single letter (monomer) were to be interpreted as “replicate the entire sequence it is in”, then self-replicators would be very easy to find. Over the years of development of Avida, the meaning of each symbol has changed as the instruction set itself has changed over time, so the absolute values for the information content of self-replicators may also change in the future. We are here only interested in the rate at which self-replicators can be found in relationship to the information content, and how this rate depends on other factors in the environment that can be modified. Translated to a search for the origins of life, we are interested in how local (environmental) conditions can favorably increase the likelihood to find a self-replicator with information content $I$ purely by chance.

We first focused on avidian sequences constrained to length $L = 15$, as there already is a hand-written standard replicator of that length in Avida, given by the string wzacgzwvfcaxgah. If every instruction in this replicator were information, the likelihood of finding it by chance would be $26^{-15} \approx 6 \times 10^{-22}$. Even if we tested a million sequences per second per CPU (central processing unit), on 1,000 CPUs running in parallel, we only would expect to find a single self-replicator in about 50,000 years
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<table>
<thead>
<tr>
<th>Instruction</th>
<th>Description</th>
<th>Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>nop-A</td>
<td>no operation (type A)</td>
<td>a</td>
</tr>
<tr>
<td>nop-B</td>
<td>no operation (type B)</td>
<td>b</td>
</tr>
<tr>
<td>nop-C</td>
<td>no operation (type C)</td>
<td>c</td>
</tr>
<tr>
<td>if-n-eq</td>
<td>Execute next instruction only if <code>BX</code> does not equal complement</td>
<td>d</td>
</tr>
<tr>
<td>if-less</td>
<td>Execute next instruction only if <code>BX</code> is less than its complement</td>
<td>e</td>
</tr>
<tr>
<td>if-label</td>
<td>Execute next instruction only if template complement was just copied</td>
<td>f</td>
</tr>
<tr>
<td>move-head</td>
<td>Move instruction pointer to same position as flow-head</td>
<td>g</td>
</tr>
<tr>
<td>get-head</td>
<td>Write position of instruction pointer into register CX</td>
<td>h</td>
</tr>
<tr>
<td>set-flow</td>
<td>Move the flow-head to the memory position specified by <code>CX</code>'s</td>
<td>i</td>
</tr>
<tr>
<td>shift-r</td>
<td>Shift all the bits in <code>BX</code> one to the right</td>
<td>j</td>
</tr>
<tr>
<td>shift-l</td>
<td>Shift all the bits in <code>BX</code> one to the left</td>
<td>k</td>
</tr>
<tr>
<td>inc</td>
<td>Increment <code>BX</code></td>
<td>l</td>
</tr>
<tr>
<td>dec</td>
<td>Decrement <code>BX</code></td>
<td>m</td>
</tr>
<tr>
<td>push</td>
<td>Copy value of <code>BX</code> onto top of current stack</td>
<td>n</td>
</tr>
<tr>
<td>pop</td>
<td>Remove number from current stack and place in <code>BX</code></td>
<td>o</td>
</tr>
<tr>
<td>swap-stk</td>
<td>Toggle the active stack</td>
<td>p</td>
</tr>
<tr>
<td>swap</td>
<td>Swap the contents of <code>BX</code> with its complement</td>
<td>q</td>
</tr>
<tr>
<td>h-copy</td>
<td>Copy instruction from read-head to write-head and advance both</td>
<td>r</td>
</tr>
<tr>
<td>h-alloc</td>
<td>Allocate memory for <code>BX</code></td>
<td>s</td>
</tr>
<tr>
<td>h-divide</td>
<td>Divide off an offset located between read-head and write-head</td>
<td>t</td>
</tr>
<tr>
<td>h-search</td>
<td>Find complement template and place flow-head after it</td>
<td>u</td>
</tr>
</tbody>
</table>

Table 7.1 Instruction set of the avidian programming language used in this study. The notation `<BX>` implies that the command operates on a register specified by the subsequent nop instruction (for example, `nop-A` specifies the AX register, and so forth). If no nop instruction follows, use the register BX as a default. More details about this instruction set can be found in (Ofria et al., 2009).

of continuous search. We tested one billion sequences of $L = 15$ and found 58 self-replicators (all of them unique) by chance, indicating that the information content of self-replicators is vastly smaller than 15 mers. Indeed, we can estimate the information content as

$$I(15) = -\log_2(58 \times 10^9) \approx 5.11 \pm 0.04 \text{ mers},$$

with a one-σ error. Here, the ‘mer’ is a unit of information obtained by taking logarithms to the base of the alphabet size, so that a single monomer has up to one mer of entropy (Adami, 2002, 2012). This means that, within the replicating 15-mers, only about 5 of those 15 mers are information.

We next tested the information content of sequences constrained to several different lengths. Among a billion random sequences of $L = 30$, we found 106 replicators, which translates to

$$I(30) = -\log_2(106 \times 10^9) \approx 4.93 \pm 0.03 \text{ mers},$$

which is significantly different from $I(15)$. In fact, the calculated information content suggests that perhaps replicators of length five or six might exist, but an exhaustive search of all $11,881,376$ $L = 5$ sequences and all $308,915,776$ $L = 6$ sequences reveals this not to be the case. When searching a billion sequences of $L = 8$ we found 6 unique self-replicators, implying an information content

$$I(8) = -\log_2(6 \times 10^9) \approx 5.81 \pm 0.13 \text{ mers}.$$  

The six sequences we found are qxrchcwv, vxfgjgb, wxvxggb, vhfgwgb,
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Figure 7.1 Sketch of the avidian CPU, executing a segment of code. The CPU uses three registers (AX, BX, CX) and an instruction pointer (IP) that reads the program into the CPU. A read-head, a write-head, and a flow-head are used to specify positions in the CPU’s memory. For example, the ‘copy’ command reads from the read-head and writes to the write-head, while ‘jump’-type statements move the instruction pointer to the flow-head. The CPU uses two stacks to simulate an “infinite Turing tape”, and input/output buffers to communicate with its environment (reproduced from (Ofria et al., 2009), with permission).

We can understand this trend of decreasing information content with increasing length (violating Eq. (7)) as a consequence of the way we treat avidian sequences, namely as having a beginning and an end. Indeed, while the genome itself is circular, execution always begins at a marked instruction. We can see this effect at work using the example origins sequence that we used before. If we add a single letter to the 7-mer origins, the number of sequences that spell the word increases by 52 (adding the letter to the beginning or the end of the word), while the total number of possible sequences only increases by 26. Thus, the density of self-replicators increases with length, leading to a decrease of information.

We tested whether this decrease of information with increasing sequence length would continue, by testing 300 million sequences of length 100. We found 17 self-replicators among this set, which translates to \( I(100) = 5.10 \pm 0.09 \) mers and suggests that not only does the trend not continue (which of course would have been absurd), but may reverse itself. There is a subtle information-theoretic reason for an increase in information with increasing sequence length. Suppose that there is a single instruction that could abrogate self-replication if it is to be found anywhere within the sequence, when in its absence the sequence replicates (a ‘kill’ instruction, so to speak). Even though such an instruction is obviously not information about how to self-replicate,
its needed absence actually is information. When the sequence length increases, the presence of such a ‘kill’ instruction becomes more and more likely, and therefore the absence of the instruction over the increasing sequence length represents an increase in information. This is the trend suggested in Fig. 7.2.

### Biased typewriters

In a sense, the random search for self-replicators is very inefficient: it is known that functional molecular sequences cluster in genetic space, while vast regions of that space are devoid of function. Yet, the random generation of sequences searches all of genetic space evenly. Is there a way to focus the search more on sequences that are likely to be functional? It turns out there is, and this method only requires the generation of monomers using a biased probability density function that more resembles that generated by functional sequences (Adami, 2015). We first present a simple example (the biased typewriter), and then outline the theory behind the enhanced search.

Words in the English language have a very characteristic letter-frequency distribution that makes it possible to distinguish English text from random sequences of letters, and even text written in different languages. Fig. 7.3 (using data from (Lewand, 2000)) shows the frequency distribution of letters in English text, showing that ‘e’ appears more frequently than ‘t’, which itself is more frequent than ‘a’ and so on. As this is the expected frequency of letters in English, a focused search should generate words with these expected frequencies that is, the ‘monomers’ of English words should be

![Figure 7.2 Number of self-replicators per $10^8$ found for various genome sizes using an unbiased (uniform) probability distribution of monomers. The number of self-replicators per $10^8$ for $L = 100$ is estimated from sampling 300 million sequences only (all others used samples of $10^9$). Error bars are standard deviations.](image-url)
generated with the frequency distribution in Fig. (7.3), rather than uniformly. When we did this for 1 billion sequences of seven letters, we found origins twice. How large is the expected increase in likelihood?

We saw earlier that the information content of sequence \( s \) can be written as

\[
I(s) = - \log \frac{N_c}{N},
\]

which itself is an approximation of the form

\[
I(s) = H(S) - H(S|e),
\]

assuming that the distribution of functional sequences in genetic space is uniform\(^3\). The remaining entropy (given the current environment \( E = e \)) \( H(S|e) \) is not known a priori, but we can estimate it. This entropy of the polymer \( s \in S \) can be written in terms of the entropy of monomers, the shared entropy of all monomer pairs, triplets, and so on, using a formula that was first derived by Fano in a very different context (Fano, 1961, p. 58):

\[
H = \sum_i H(i) - \sum_{i<j} H(i : j) + \sum_{i:j:k} H(i : j : k) - \cdots
\]

where \( H(i) \) is the entropy of the \( i \)th monomer, \( H(i : j) \) is the shared entropy between

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\(^3\) The distinction between the entropy written as \( \log N_c \) or else as \( - \sum_s p(s|e) \log p(s|e) \) can viewed as the same distinction that is made in thermodynamics, where the former is known as the entropy in the "micro-canonical ensemble" whereas the latter entropy pertains to a "canonical ensemble" if \( p(s|e) \) is the canonical distribution, see, e.g. (Reif, 1965).
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Let us define the “average biotic entropy” $H_b$ as the average entropy-per-site for functional sequences (hence the name “biotic”)

$$H_b = \frac{1}{L} \sum_{i=1}^{L} H(i)$$

We distinguish this biotic entropy from the “abiotic” entropy $H_\star$, which is the entropy per-site within a sequence assembled at random. If each monomer appears with uniform probability, then the abiotic entropy is maximal: $H_\star = 1$. Using this definition, we can write (7) as

$$P_0 = D^{-L} = D^{-L \sum_{i=1}^{L} H(i)}.$$ (7.0)

If we were to generate ASCII sequences with a probability distribution obtained from English words (the equivalent of the biotic sample, see Fig.7.3), the abiotic entropy would be smaller than 1 (namely $H_\star \approx 0.89$, the entropy of the distribution in Fig. 7.3) while the biotic entropy must be zero, as there is only a single origins among 7-mers. Using the probability distribution of letters in English rather than the uniform distribution raises the probability to find the 7-mer origins to

$$P_\star = 26^{-70(0.89)}.$$ (7.0)

This seems like a small change, but the mean number of successes out of $10^9$ tries is increased from about 1 in 8 billion to 1.53 per billion. And indeed, we found the word twice when searching a billion sequences with the biased distribution shown in Fig. 7.3. Note, however, that the entropy of English is equal to the entropy $\frac{1}{L} \sum_{i} H(i)$ only if sequences cannot be aligned, and therefore that all $H(i) = H_\star$.

Can searching with a biased probability distribution increase the chance of finding a self-replicator in Avida? We first took the 58 self-replicators we found when searching $L = 15$ sequences, and created a monomer-probability distribution $p_\star$ out of them. This distribution in Fig. 7.4 shows that within these randomly created replicating sequences, the 26 instructions appear far from uniformly in the sequence (as of course is expected),
in the same way as English (because it conveys information) has a non-uniform letter
distribution. The entropy of the distribution shown in Fig. 7.4 is $H_\ast = 0.91$ mers.

Figure 7.4 The biased probability distribution $p_\ast$ of Avida instructions ob-
tained from the genomes of 58 randomly generated $L = 15$ replicators (the
meaning of each letter is described in Table ??). The solid black line repre-
sents the probability for a uniform distribution.

According to the approximation we made above, biasing the monomer creation process
using this particular probability distribution should lead to an enhancement $E$ of the
likelihood of finding a self-replicator

$$E = \frac{P_\ast}{P_0} = \frac{D^{-\Delta H_0 + H_\ast}}{D^{-\Delta H_0}} = D^\Delta(1 - H_\ast).$$

Eq. (7) suggests that the enhancement factor $E$ only depends on the bias in the distri-
bution and the length of sequence. However, we should not be fooled into believing
that any reduced entropy $H_\ast$ will lead to an enhancement in the probability to find self-
replicators by chance: the distribution $p_\ast$ needs to be close to the distribution of actual
replicators. For example, omitting the instruction ‘x’ (the $h$-divide instruction that
splits off a completed copy, see Table 7.1) certainly leads to an entropy less than one,
but using such a biased distribution cannot net a self-replicator as $h$-divide is required
for replication.

We proceeded to test Eq. (7), by searching for self-replicators using the biased dis-
tribution $p_\ast$ (see Methods). Among a billion sequences of $L = 15$ generated in this
manner, we found 14,495 self-replicators, an enhancement of $E = 14,495/58 \approx 250$,
while Eq. (7) predicted an enhancement of $E = 81.3$. We also tested whether changing
the probability distribution from uniform gradually towards $p_\ast$ leads to a gradual in-
crease in the $E$. The empirical enhancement factor shown in Fig. 7.5 indeed increases
with the bias, and is larger than the one predicted from the simple approximation (7).
This difference is likely due to a number of effects. On the one hand, we are neglect-
ing any higher order correlations in Eq. (7). On the other hand, we are assuming that $H_* \approx H(i)$ for all $i$, that is, that the entropy at each site is the same. This is not at all true for functional sequences that can be aligned (see, e.g., (Adami and Cerf, 2000; Adami et al., 2000; Gupta and Adami, 2014)). Sequences that are obtained from a random procedure (rather than from an evolutionary process) are likely difficult to align, and therefore $H_* \approx H(i)$ may hold.

Figure 7.5 The enhancement factor $E$ to find entropys for genomes of 15 instructions as a function of the bias, using an interpolated probability distribution $p(i,b)$. Here, $b = 0$ means unbiased, and $b = 1$ uses a fully biased distribution $p_*$. Black circles represent estimates (calculated as the number of self-replicators per $10^8$ for a biased distribution divided by the number of self-replicators per $10^8$ for a uniform distribution), while error bars are standard deviations. The solid line is the naive prediction given by Eq. (7).

The enhancement works for sequences of any length, but depends on how well the biased distribution represents actual functional replicators. For example, as we found only 6 self-replicators of length 8, the distribution $p_*(8)$ is fairly coarse (see Fig. 7.6A), while the distribution we obtained from the 106 $L = 30$ replicators has a significant uniform contribution (Fig. 7.6B), because among the 30 instructions only a handful need to carry information in order for the sequence to be able to replicate. We show in Fig. 7.7 the enhancement achieved by biasing the search for each of the three length classes $L = 8, 15,$ and 30.

Could we use the probability distribution for sequences obtained in one length group to bias the search in another length group? Such a procedure might be useful if the statistics of monomer usage is poor (as for the case $L = 8$), or if the distribution was obtained from a sequence with too much entropy (as for the case $L = 30$). It turns out that this is not the case; biasing the $L = 30$ search using $p_*(15)$ does not work well (144.3 replicators found per $10^8$ compared to biasing with the “native” $p_*(30)$ (297 per $10^8$). In the same manner, biasing the $L = 8$ search works best with the “native” bias $p_*(8)$, yielding 230 per $10^8$, as opposed to only 15.8 per $10^8$ biasing with $p_*(15)$. 
Figure 7.6 Probability distribution of instructions. A: $p_*(8)$ obtained from the replicators of length $L = 8$, giving rise to an entropy $H_*(8) = 0.71$ mers. B: $p_*(30)$ obtained from the replicators of length $L = 30$, giving rise to an entropy $H_*(30) = 0.98$ mers. The solid horizontal line denotes the uniform probability distribution $1/26$ in both panels.

Figure 7.7 Empirical enhancement factor (black dots, with 1σ counting error), along with the predicted enhancement factor using the entropy of the distribution based on Eq.(7) (grey dots) for $L = 8, 15, 30$.

Finally we asked whether taking the self-replicators obtained from a biased search (and that consequently nets many more replicators) gives rise to a more accurate probability distribution $p_*$, which then could be used for a more ‘targeted’ biased search. By “rebiasing” successively (see Methods), we did indeed obtain more and more replicators, albeit with diminishing returns (see Fig. 7.8).
Figure 7.8 A: Average per-site entropy $H_s$ for replicators in different length classes, at various stages of biasing. “Unbiased” reports the average per-site entropy obtained from the self-replicators that were found in an unbiased search, and whose biased distribution was used to find the self-replicators whose average per-site entropy is shown in “1st Bias”. Those in turn were used for a biased search that gave rise to replicators with bias shown in “2nd Bias”, and so on. B: Number of self-replicators (per billion) found at each biasing stage. Biasing the distribution with more “focused” probability distributions $p_x$ leads to an increasing yield of self-replicators, albeit with a diminishing return. In re-biasing with $L = 8$, some duplicate sequences were obtained, and those are not included in the count.

**Discussion**

One of the defining characteristics of life (perhaps the defining characteristic) is that life encodes information, and information leaves a trace in the monomer abundance distribution (a non-random frequency distribution) (Dorn and Adami, 2011; Dorn et al., 2011) of the informational polymers. As life evolves, the information contained in it increases on average (Adami et al., 2000), but evolution cannot explain where the first bits came from. Information can in principle arise by chance, just as an English word can appear by chance within an ASCII string that is created randomly, as per the “dactylographic monkeys” metaphor. The “infinite monkey theorem” posits that a million monkeys typing on a million keyboards, if given enough time (and typing randomly) could ultimately type out all of Shakespeare’s works. However, the theorem is misleading, as even correctly typing out the first 30 characters of Hamlet’s soliloquy (“To be or not to be...”) cannot occur during the time our universe has been around (about $4.36 \times 10^{17}$ seconds), as Hamlet’s 30-mer is one in about $3 \times 10^{42}$. Using biased typewriters will not allow the monkeys to finish either, as it is only accelerating the search by a factor $E \approx 46,700$.

We can ask whether more sophisticated methods of biasing exist. One look at Eq.(7) suffices to answer this question in the positive. We could begin by generating sequences biased in such a way that the more common 2-mers are generated with increased likelihood. In English text, for example, the “dimers” “th”, “he”, and “in” appear with frequencies 3.56%, 3.07%, and 2.43% respectively, which are significantly larger than the random dimer expectation = 0.15%. Indeed, as the frequency of ‘or’ is 1.28%, while ‘ig’ appears at 0.255%, our 7-mer origins would be found fairly fast. Likewise, in our 6 replicators of length $L = 8$ the dimer gb appears significantly more often than expected by the product of the likelihood of g and b.
Such biased search procedures can also accelerate the search for functional biomolecules where the target is a function other than self-replication. For example, when designing random peptide libraries (either for screening purposes or to perform directed evolution), researchers often bias the codons in such a way that the stop codon is rare (so-called NNB or NNS/NNK libraries (Barbas et al., 1992)). Hackel et al. (Hackel et al., 2010) went beyond such simple biases and constructed a protein library to screen for binding to a set of 7 targets. To bias the random sequences, they mimicked the amino acid distribution in human and mouse CDR-H3 loops (complementarity determining regions, which are found in antibodies), and found that such a library outcompetes even NNB libraries significantly: of the 20 binders that they found, 18 were traced back to the CDR-biased library.

The implications of the present theoretical and computational analysis of the emergence of informational “molecules” by chance for the problem of understanding the origin of life are straightforward. It is well known that monomers do not form spontaneously at the same rate. The abiotic distribution of amino acids is heavily skewed both in spark synthesis experiments as well as in meteorites (Dorn et al., 2011), and the same is true for other monomers such as carboxylic acids, and many other candidate alphabets in biochemistry. In many cases, the abiotic skew (often due to thermodynamic considerations) will work against the probability of spontaneous emergence of information, but in some cases it may work in its favor. In particular, we might imagine that in complex geochemical environments the biotic distributions can be significantly different in one environment compared to another, raising the chance of abiogenesis in one environment and lowering it in another.

We also immediately note that in chemistries where molecules do not self-replicate but catalyze the formation of other molecules, the abundance distribution of monomers would change in each catalysis step. If these monomers are recycled via reversible polymerization (Walker et al., 2012), then the activity of the molecules can change the entropy of monomers, which in turn changes the likelihood of spontaneous discovery. Should this process “run in the right direction”, it is possible that self-replicators are the inevitable outcome. This hypothesis seems testable in digital life systems such as Avida.

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Methods
In order to explore the spontaneous emergence of self-replicators in Avida, we generated random genomes of length $L$. These genome sequences were generated with different probability distributions for the avidian instructions (we used Avida version 2.14, which can be downloaded from https://github.com/devosoft/Avida). First, we generated $10^9$ random genomes for lengths $L = \{8, 15, 30\}$ and $3 \times 10^8$ sequences for $L = 100$ with an unbiased (that is, uniform) instruction distribution $1/26$ (because there are 26 possible instructions). In order to decide whether a genome could successfully self-replicate, we performed two tests. First, we checked whether the organism would successfully divide within its lifespan. Here, we used the traditional Avida parameters.
for an organism’s lifespan: it must divide before it executes $20 \times L$ instructions. While this indicates that an avidian could successfully reproduce, it does not imply that the avidian’s descendants could also reproduce. In our search we found many viable avidians that would successfully divide into two non-viable organisms. Therefore, we only counted avidians that could self-replicate and produce offspring that could also self-replicate as true self-replicators (in other words, they are “colony-forming”). This does not mean that every self-replicator would produce a perfect copy of itself in the absence of mutation; in fact, most of these replicators undergo implicit mutations solely due to their genome sequence, and their offspring differ in length from the parent (LaBar et al., 2015). In analyzing a genome’s ability to self-replicate, we used the default Avida settings, described for example in (Ofria et al., 2009).

Next, we generated random genome sequences with a biased instruction distribution. These biased distributions were calculated by altering the probability that each instruction was generated by our random search. The probability of an instruction $i$ being generated for a biased search was set at

$$p(i, b) = (1 - b)(1/26) + bp_*(i),$$

where $b$ is the bias, $0 \leq b \leq 1$, and $p_*(i)$ is the probability that instruction $i$ appears in the set of all genomes that were classified as self-replicators in the unbiased search. When $b = 0$, the distribution is the uniform distribution and when $b = 1$, the distribution is the frequency distribution for the instructions in the set of self-replicators $p_*$ found with the unbiased search for a given length. The parameter $b$ allows us to set the bias, and thus the entropy, of the distribution to detect the role of the instruction entropy in determining the likelihood of spontaneous self-replicator emergence. For genomes of $L = 15$, we generated $10^9$ random sequences with $b = 1$ and $10^8$ random sequences with $b = \{0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9\}$.

Finally, we performed searches where we iteratively biased the distribution of instructions. First, we generated self-replicators with an unbiased instruction distribution. We then created another set of self-replicators with a biased distribution of instructions using the above equation with $b = 1$ (referred to as “1st bias”). However, as opposed to stopping the self-replicator generation process, we then searched for self-replicators two more times (referred to as ‘2nd bias’ and ‘3rd bias’). Each time, we used the set of self-replicators from the previous bias: the distribution of instructions for the 2nd bias was derived from the set of self-replicators obtained from the 1st bias, and the distribution of instructions for the 3rd bias was derived from the set of self-replicators from the 2nd bias (in both of these we set $b = 1$). We generated $10^9$ random genomes using the 1st bias for $L = \{8, 30\}$ and $10^8$ random genomes using the 2nd and 3rd bias for $L = \{8, 15, 30\}$ with a biased instruction distribution. For $L = 15$, we used the $10^9$ random genomes described above to obtain the 1st bias.

References


