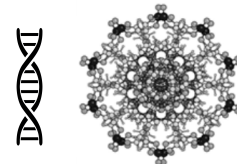


# DNA as Nanotechnology: Reassessing Life's Origin Through the Lens of Information and Genomic Intelligence

## ABSTRACT

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Version XII - Major Mathematical Additions - Major Tone Revisions - Updated References



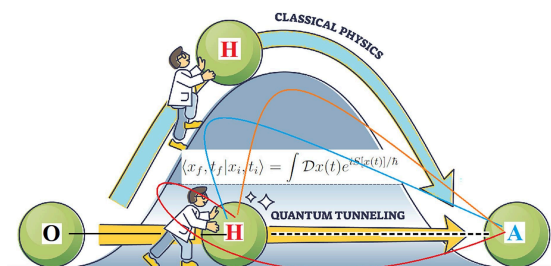
We undertake a comprehensive examination of the complex interplay between deoxyribonucleic acid (DNA), nanotechnology, and the origin of life, critically engaging with prevailing abiogenetic models. We advance the hypothesis that DNA functions at the quantum scale or exhibits quantum-mechanical characteristics, demonstrating a level of structural stability and informational complexity that challenges the assumptions underpinning theories of spontaneous molecular evolution. Central to the critique is the recognition of the indispensable role of enzymatic machinery in DNA replication—enzymes that, paradoxically, require DNA for their synthesis—thereby presenting a classic instantiation of the "chicken-and-egg" paradox. We further interrogate the significance of molecular chirality and evaluate the environmental prerequisites for biogenesis, contending that early Earth conditions were inherently unfavorable for the natural formation of either DNA or RNA. By synthesizing insights from molecular biology, quantum physics, and information theory, this analysis supports alternative frameworks. Ultimately, we call for a fundamental reassessment of evolutionary mechanisms and reposition DNA not merely as a passive genetic substrate, but as an advanced, self-organizing system for information storage and processing—one that challenges conventional biological paradigms. We propose a Mathematical proof utilizing Minimal Genome formation and the Universe's limit of Genetic generative capacity.

## Section A: The Scale and Quantum Implications of DNA: Proton Tunneling and Molecular Stability

DNA, with a diameter of approximately 2 nanometers (and a radius of 1 nanometer), and a base pair separation of merely 0.34 nanometers, operates at dimensions comparable to, or smaller than, the most advanced nanoscale transistors in modern CPUs. These physical characteristics place DNA within the threshold of the quantum domain, where subatomic phenomena such as quantum fluctuations and proton tunneling become relevant. At such scales, the structural integrity and informational fidelity of DNA would, under standard quantum expectations, be vulnerable to stochastic disturbances, particularly variations in hydrogen bonding driven by isotopic shifts.

Proton tunneling, in particular, is predicted to introduce a high frequency of random base pair mutations. Over time, these quantum-induced mutations should theoretically accumulate to a degree that compromises the functionality of the genetic code. However, empirical observations contradict this prediction. DNA exhibits a remarkable degree of structural and

informational stability, suggesting that it either mitigates or exploits quantum mechanical effects in a regulated manner.



(<https://www.sciencedirect.com/science/article/abs/pii/S0301010423002367> Luca Nanni)

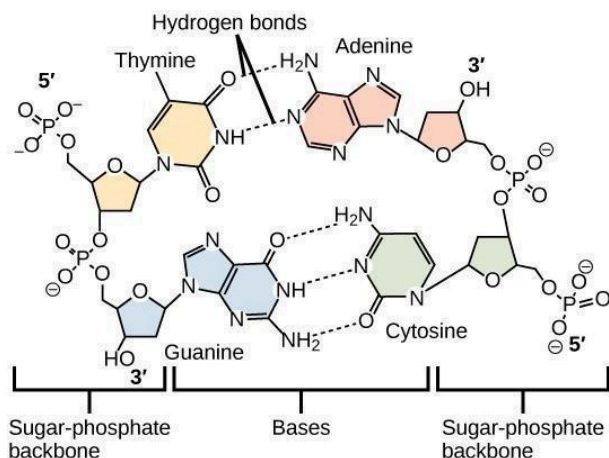
This perspective is supported by findings published in *Communications Physics*, which state:

“We determine that the quantum tunnelling contribution to the proton transfer rate is several orders of magnitude larger than the classical over-the-barrier hopping. Due to the significance of the quantum tunnelling even at biological temperatures, we find that the canonical and tautomeric forms of G-C inter-convert over timescales far shorter than biological ones and hence thermal equilibrium is rapidly reached. Furthermore, we find a large tautomeric occupation probability of  $1.73 \times 10^{-4}$ ,

suggesting that such proton transfer may well play a far more important role in DNA mutation than has hitherto been suggested. Our results could have far-reaching consequences for current models of genetic mutations” (Slocombe et al.).

The implications of such findings are profound. They suggest that DNA possesses inherent features capable of stabilizing its informational content against disruptive quantum phenomena. In the absence of such stabilizing mechanisms, the accumulation of deleterious mutations via quantum tunneling would render DNA evolutionarily unviable, as it would fail to preserve the integrity required for replication and functional biological development. Thus proton tunneling presents a distinctive challenge for any initial polymer in regards to Origin of Life.

This would render formation of tangible polymers with information preservation susceptible to high mutations that, if not mitigated immediately & from instantiation, will prohibit the formation of replicable molecules as they require a high threshold of informational stability, manifested through a replicator’s molecular and structural stability. The distinction of random crystallization and high fidelity information encoded in DNA/RNA is important to point out. The structural stability could not arise, as before that threshold has reached, the molecules integrity to preserve information would have already been compromised. This is distinctly different from crystallization as it doesn’t result in the same latent outcomes associated with DNA or RNA that create exponentially more **complex information systems** in comparison to crystallization. Crystallization does not encode high fidelity information like DNA or RNA.



(<https://raider.pressbooks.pub/biology1/chapter/4-dna/>)

## Section B: Error Catastrophe and the Threshold of Genomic Fidelity

One of the most critical requirements for the viability of DNA as a genetic medium is its extraordinary level of replication accuracy. Empirical models indicate that DNA/RNA or early replicators must achieve an initial replication fidelity exceeding 99.999%, derived from viral mutagenesis & Eigen’s paradox; by contrast, accuracy rates of even 90–95% are insufficient and would inevitably result in what is known as an **error catastrophe**. This phenomenon refers to the progressive and cumulative loss of genetic integrity within a population, driven by excessive mutation rates or the failure of error-correction mechanisms. Such degradation impairs cellular processes and division, ultimately leading to functional collapse. Furthermore, high mutation burdens are increasingly linked to aging and senescence in complex organisms.

Imagine a simple 100 nucleotide polymer with a 90% starting copying accuracy. The 1st generation would lose 10 nucleotides, for simplicity we assume with each following generation losing a further 10 nucleotides to errors. (static 10 bp/generation loss model) Within 5 generations, half or 50% of the information in the polymer would have degraded to noise/errors. This poses a hard problem for the first polymer, which in absence of error correction from its instantiation, would degrade unless error correction mechanisms mitigate such disturbances.

In reality, errors multiply and compound. Therefore the realistic problem of error accumulation in absence of error correction mechanisms is far more pronounced.

As summarized in a *PLOS Pathogens* figure article, “lethal mutagenesis” has been identified as a viable strategy for controlling RNA viruses precisely because of their susceptibility to error catastrophe—a vulnerability that underscores the fine balance necessary for genetic stability

“In RNA viruses, a small increase in their mutation rates can be sufficient to exceed their threshold of viability. Lethal mutagenesis is a therapeutic strategy based on the use of mutagens, driving viral populations to extinction. Extinction catastrophe can be experimentally induced by promutagenic nucleosides in cell culture models.” (Hassine)

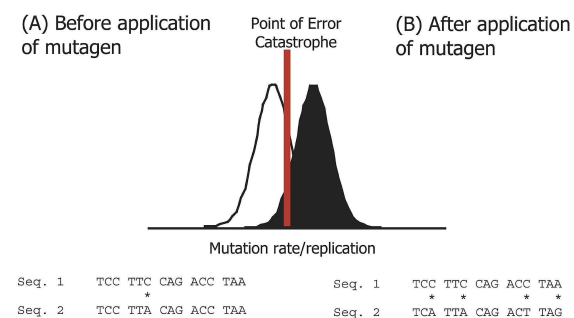
This principle poses a significant challenge to any abiogenesis model. In the earliest stages of life, how could primitive genetic systems have achieved such high fidelity in the absence of evolved proofreading mechanisms? Replication at the molecular level requires a suite of enzymatic functions—such as polymerase fidelity, exonuclease proofreading, and repair enzymes—all of which are themselves encoded by DNA or RNA. Thus, the question becomes deeply paradoxical: how could accurate replication arise in a prebiotic world where the very tools needed to ensure fidelity could not yet exist?

As pointed out in *Nucleic Acids Research* that mutation rates must be kept below an error threshold... otherwise the genetic information degenerates. Without a robust framework for error detection and correction, early genetic systems would have succumbed to exponential error accumulation, rendering stable inheritance—and thus evolution—impossible. This presents a formidable obstacle for any theory proposing a spontaneous, self-organizing origin of life.

The layered complexity of these enzymatic systems increases with ever increasing cellular processes. This increasing complexity of the information creates more susceptibility to error-catastrophe or corruption. Therefore you need a set of error correction mechanisms evolving ahead or simultaneously with a replicator's own ability to store information. This dual interdependence between error correction & information creates a paradoxical situation where the required information for both systems surpasses the information capacity of a replicator at any initial abiogenesis model. To resolve this paradox you may invoke the simultaneous formation of the Replicator, which is RNA/DNA and for it to spawn with its toolbox of error correcting enzymes, all of which themselves represent complex 3d structures. The accuracy rates are mandated to be as high as  $10^{-3}$  to  $10^{-4}$  in the initial stages of abiogenesis as RNA or DNA are at its most vulnerable state. A gradually evolving system with incremental improvements to error correction mechanisms simply can't evolve due to mathematical and logical constraints, the accumulative effect of errors would also render the error correction mechanisms themselves vulnerable to error catastrophe. These error correction mechanisms need to account not only for classical chemistry but also quantum effects such as proton tunneling that present a challenge to abiogenesis models that is not yet fully realized.

- A replicator (RNA/DNA) needs error-correction enzymes to persist.
- But those enzymes need to be encoded by the replicator.
- And the replication of both systems needs a fidelity higher than what's plausible without correction.

This forms a bootstrap problem: life can't start unless it's already highly ordered and protected from error. Simply put errors accumulate faster than they can be acted on by natural selection



(<https://journals.plos.org/plosbiology/article/figures?id=10.1371/journal.pbio.0020307>)

## Error Catastrophe simulation

Let's assume starting replication fidelity at 90% i.e the errors between generations are 10% - there is no error compounding and a static 1000bp/generation loss is assumed - if there are no error correction mechanisms then the whole 10000 bp long polymer will lose 1000 base pairs to mutations per generation - leading to complete informational degradation in 5 to 7 generations (50%-70%loss). This can be applied to the minimal genome of 543kbp and at 90% starting fidelity you lose - 54,300 base pairs to mutations in just 1 generation - this signifies how pivotal error correction is at any initial stage of abiogenesis & constitutes an unresolved physical constraint. Accounting for compounding errors i.e errors causing more errors - the realistic scenario is far more severe. Errors would accumulate exponentially, in a non linear amplification.

### \*Calculating 10% loss per generation for 100 bp polymer\*

Generation 1 - 100 bp to 90 bp

Generation 2 - 90 bp to 81 bp

Generation 3 - 81 bp to 72 bp

Generation 4 - 72 bp to 65 bp

Generation 5 - 65 bp to 59 bp

Generation 6 - 59 bp to 53 bp

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## Section C: The Information Paradox—DNA as the Ultimate Data Storage System

Among the most striking discoveries in molecular biology is the unparalleled information density and stability encoded within DNA. When compared to modern data storage technologies—such as magnetic tape, optical disks, or solid-state drives—DNA outperforms all in both volumetric efficiency and fidelity. Analogies often fail to convey the magnitude of this disparity.

As reported in *IEEE Spectrum*, the informational capacity of DNA is extraordinary:

“Converted to digital media, a diploid genome can store 1.5 gigabytes of data... Since DNA has the ability to encode 2 bits per nucleotide, one gram of dried DNA can store 455 exabytes of data... Moreover, DNA exceeds by many times the storage density of magnetic tape or solid-state media. It has been calculated that all the information on the Internet—which one estimate puts at about 120 zettabytes—could be stored in a volume of DNA about the size of a sugar cube, or approximately a cubic centimeter”

“DNA is one of the most promising next-generation data carriers, **with a storage density of  $10^{19}$  bits of data per cubic centimeter**, and its three-dimensional structure makes it about eight orders of magnitude denser than other storage media. DNA amplification during PCR or replication during cell proliferation enables the quick and inexpensive copying of vast amounts of data. In addition, DNA can possibly endure millions of years if stored in optimal conditions and dehydrated, making it useful for data storage. Numerous space experiments on microorganisms have also proven their extraordinary durability in extreme conditions, which suggests that DNA could be a durable storage medium for data. Despite some remaining challenges, such as the need to refine methods for the fast and error-free synthesis of oligonucleotides, DNA is a promising candidate for future data storage.”  
<https://pmc.ncbi.nlm.nih.gov/articles/PMC10296570/>

This comparison is not hyperbolic; DNA is not only orders of magnitude denser in terms of storage but also vastly more stable. The natural error rate associated with DNA replication is extraordinarily low ( $\sim 10^{-9}$ ), making it the most reliable information storage medium known. It retains its informational integrity across generations and for millions or billions of years under favorable conditions. Such stability is unprecedented in any human-made system.

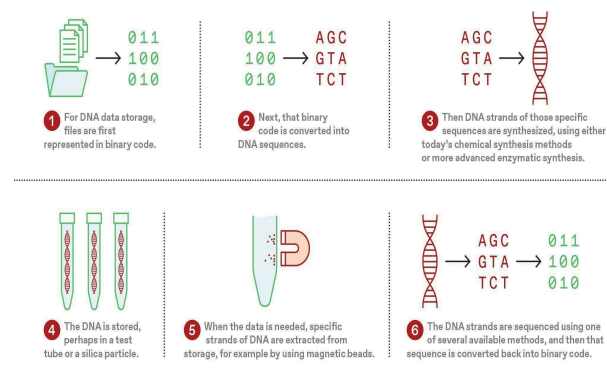
Yet DNA is far more than a static storage device. Its utility is entirely contingent on a suite of complex molecular machinery that enables transcription, translation, and replication. These processes require a host of highly specific enzymes: helicases to unwind the double helix, polymerases to read and replicate the sequence, ligases to bind fragments, and ribosomes to translate codons into functional proteins. Without this intricate biochemical infrastructure, a DNA molecule is biologically inert. As the biochemistry course materials at Western Oregon University indicate DNA replication and repair require a highly coordinated interaction of multiple enzymes and protein complexes.

The spontaneous, concurrent emergence of DNA alongside these enzymes in a prebiotic context is highly improbable and logically incoherent. As articulated in the *Structure and Function of Nucleic Acids*, the complexity and interdependence of nucleic acids and proteins make their simultaneous origin through unguided chemical processes appear extraordinarily implausible.

In terms of global presence, DNA is not a rare molecule but it permeates the biosphere. It is estimated that Earth harbors approximately  $5.3 \times 10^{31}$  ( $\pm 3.6 \times 10^{31}$ ) megabases (Mb) of DNA. Given conservative assumptions regarding transcription rates, this implies that the biosphere processes information at rates exceeding yottaNOPS ( $10^{24}$  nucleotide operations per second -Landenmark et al.), suggesting that life on Earth operates as an immense, decentralized computational substrate.

The broader implication of this information density and processing capability is that DNA embodies not merely a chemical scaffold for heredity but a computational and regulatory system of staggering sophistication. It orchestrates ecosystems, drives atmospheric transformations (such as oxygenation via cyanobacteria and plants), and underpins the interdependent feedback loops that sustain life on the planet. With every inhalation, human beings interact with the genetic

signatures of millions of microorganisms—testimony to the pervasive and dynamic role of DNA in shaping the biosphere.



(How DNA data storage works - Chris Philpot - <https://spectrum.ieee.org/dna-data-storage>)

## Section D: The Genetic Qubit—Quantum Behavior in DNA Base Pairing and Structural Forces

An intriguing and counterintuitive phenomenon emerges when examining the biophysical behavior of DNA at the quantum scale. Traditionally, it is understood that guanine-cytosine (GC) base pairs, which are connected via three hydrogen bonds, form stronger and more stable interactions than adenine-thymine (AT) pairs, which are stabilized by only two hydrogen bonds. From a conventional chemical standpoint, this increased bonding in GC pairs should confer enhanced stability and, by extension, a reduced susceptibility to mutation.

However, empirical data diverge significantly from this expectation. In reality, GC-rich regions within the genome exhibit **higher mutation rates** than AT-rich regions, suggesting that quantum mechanical phenomena—specifically proton tunneling—**may override classical chemical predictions**. Proton tunneling allows subatomic particles to traverse energy barriers, leading to tautomeric shifts and base mispairing events that can manifest as mutations during DNA replication.

“Several studies compare the probability of proton tunneling for the A-T and G-C base pairs. Tautomerization of the G-C pair has a large reverse reaction barrier than A-T, and therefore, the lifetime of G\*-C\* is longer than A\*-T\* and it can be maintained

throughout the replication process; consequently, G-C tautomerization can be a source of spontaneous point mutations (Slocombe et al., 2021)” <https://doi.org/10.1016/j.pbiomolbio.2022.05.009>

This paradox is succinctly articulated in findings published in *Genetics*

“Mutations that are introduced into the genome are generally AT-biased, i.e., the mutation rate from strong (GC) to weak (AT) nucleotides is higher than in the opposite direction. However, the proportion of fixed sites in the genome is GC-biased compared to the expectation based on mutation rate differences” (Bergman et al.).

These findings suggest that although GC bonds should be chemically more stable, quantum tunneling increases their mutagenic potential. Yet paradoxically, genomic regions with higher GC content are not only prevalent but appear to be selectively retained, implying a deeper layer of genomic organization and function that is not yet fully understood.

“Recently however, two independent analyses have shown that in virtually all Bacteria, independently of their genomic GC-content, there is an excess of G/C→A/T mutations [22,23]. This suggests that an unknown process, selective or neutral, is opposing this universal mutational bias by favouring the fixation of G/C alleles. Previously, an analysis of a large number of *E. coli* genomes had suggested a possible role of gBGC, based on the link between GC-content, recombination and the organization of the chromosome in this species [24]. However Hildebrand et al. [23] observed that the excess of G/C→A/T mutations was still present after removing datasets with evidence of recombination. Moreover they found no correlation between GC-content and recombination rate across bacterial species. They therefore concluded that this force could not be gBGC and hence that selection was driving an increase of genomic GC in Bacteria. The nature of this selective advantage remains however mysterious, though various hypotheses have been proposed [25,26].” <https://doi.org/10.1371/journal.pgen.1004941>

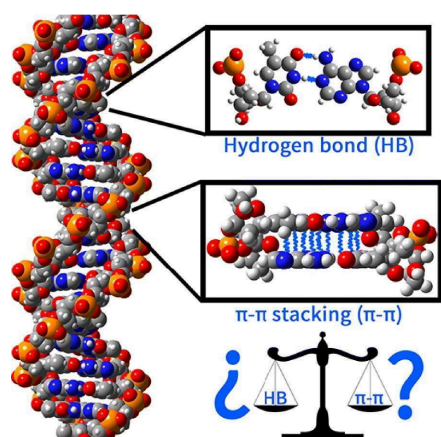
A second, equally critical force contributing to DNA's structural integrity is  **$\pi$ -stacking**. Because DNA base pairs are separated by only ~0.34 nanometers, their electron clouds overlap, generating a stacking force between the planar aromatic bases. This  $\pi$ -stacking results in a non-covalent interaction that enhances



molecular stability through hydrophobic and van der Waals forces. As described in *Oxford Reference*,

“The orientation of adjacent base pairs with their planes parallel and with their surfaces nearly in contact, as occurs in double-stranded DNA molecules. Base stacking is caused by hydrophobic interactions between purine and pyrimidine bases, and results in maximum hydrogen bonding between complementary base pairs.”

<https://www.oxfordreference.com/display/10.1093/oi/authority.20110803095449932>



(<https://www.sciencedirect.com/science/article/abs/pii/S0301462217303599>)

Together, these interactions suggest an exquisitely controlled nano scale system in which the sugar-phosphate backbone anchors nitrogenous bases that not only form hydrogen bonds subject to quantum tunneling, but also engage in  $\pi$ -stacking interactions that further reinforce the helical structure.

To summarize, the DNA double helix is stabilized through a complex interplay of covalent bonding, hydrogen bonding,  $\pi$ -stacking, and quantum mechanical dynamics. The expectation that such a system—with its structural elegance and informational precision—could have emerged by random molecular processes needs deeper evaluation.

The argument is now about the formation of a nano polymer that has the Informational stability and causative capacity to not only sustain its own existence whilst fighting environmental hazards over billions of years, but to maintain and evolve its informational complexity to an extent to terraform a whole planet and eventually create brains that are able to contemplate this monumental achievement. DNA watches itself through “your” eyes and “your” brains. With metaphor aside, it

represents something extremely profound about the nature of complex evolving information systems.

## Section E: Infodynamics and Informational Entropy—Toward a New Physical Law of Information

The traditional second law of thermodynamics asserts that the entropy of an isolated system tends to increase over time, leading to the dissipation of usable energy and the emergence of disorder. However, a new emerging theoretical framework—*the second law of infodynamics*—posits an alternative trajectory for systems characterized by information. Specifically, this law suggests that the informational entropy of systems containing encoded information, such as digital storage media or biological genetic material, must either remain constant or decrease over time, approaching a minimum entropy state at equilibrium.

As described in a recent AIP Advances publication:

“In contrast to the second law of thermodynamics, the second law of infodynamics states that the information entropy of systems containing information states must remain constant or decrease over time, reaching a certain minimum value at equilibrium... Remarkably, this indicates that the evolution of biological life tends in such a way that genetic mutations are not just random events as per the current Darwinian consensus, but instead undergo genetic mutations according to the second law of infodynamics, minimizing their information entropy” (Vopson).

The implications of this paradigm shift are profound. It challenges the long-held Darwinian assumption that mutations are inherently stochastic and directionless, instead suggesting that biological evolution may be governed by an underlying informational optimization principle. Such a framework could significantly reshape the fields of genetics, evolutionary biology, virology, pharmacology, and beyond—providing a new lens through which the structure and evolution of complex systems may be understood.

This ties neatly to homeostasis. As the external environment becomes more disorderly - the intra cellular environment has to maintain its *informational entropy* to maintain functionality and survival.

This theoretical backdrop becomes especially compelling when applied to the human genome. In terms of information generation—particularly in the memetic sense, as conceptualized by Richard Dawkins—human beings produce information at a rate dramatically disproportionate to their relatively small genomic divergence from non-human primates. Despite sharing approximately 98–99% of our genome with chimpanzees, humans generate an exponentially larger volume of cultural, technological, and linguistic data. This vast asymmetry has been referred to as the human “Datome,” a term encapsulating the full spectrum of human information output.

As noted in Caleb Scharf’s *The Ascent of Information*, this memetic explosion has created a negative ecological feedback loop. The anthropogenic acceleration of extinction rates, habitat destruction, and ecological destabilization - often referred to as the “Sixth Mass Extinction” - demonstrates a tension between information generation and biospheric sustainability. The New York Times reports that human activity is “altering the natural world at an unprecedented pace,” contributing to rapid biodiversity loss.

Among the most striking consequences is the alarming decline of global insect populations. A recent study by the University of California, Riverside, confirms a widespread and ongoing collapse in insect biodiversity—threatening ecosystems that rely on pollination, decomposition, and trophic stability (UCR Entomology). These changes raise a critical question: Is the human genome, with its unprecedented rate of information production, beneficial or detrimental to the global genomic system?

This inquiry intersects with broader discussions in information theory and evolutionary biology. From a purely genetic standpoint, the relatively small difference between the genomes of humans and apes belies the massive informational, technological, and societal disparity between the two species. Such a discrepancy defies conventional models of evolution by random mutation and selection alone. It invites scrutiny of phenomena such as the HSA2 anomaly - the fusion of two ancestral ape chromosomes to form human chromosome 2 which challenges gradualist accounts of speciation and evolutionary divergence, as discussed in detail later.

This section lays the foundation for a critical reassessment of evolutionary theory, not as a purely

random or competitive process, but as a potentially **directed flow of information** shaped by deeper physical laws. The notion that genetic mutations may occur along entropy-minimizing trajectories introduces a compelling alternative to standard Darwinian models.

E. coli Gene	Enzyme/Protein Function	Description
<i>dnaA</i>	Initiator Protein	Melts DNA at <i>oriC</i> , exposing two template ssDNA strands
<i>dnaB</i>	Helicase	Unwinds the DNA helix at the front end of each replication fork during replication
<i>dnaC</i>	Helicase Loader	Loads the DnaB Helicase onto the ssDNA template strands
<i>dnaG</i>	Primase	Synthesizes RNA primers used to initiate DNA synthesis
<i>dnaE</i>	$\alpha$ -Catalytic Subunit of DNA Polymerase III	Catalytic subunit of the main replicative polymerase during DNA replication
<i>dnaQ</i>	$\epsilon$ -Editing Subunit of DNA Polymerase III	Editing subunit of the main replicative polymerase during DNA replication
<i>dnaN</i>	$\beta$ -clamp subunit of DNA Polymerase III	Clamping subunit of the main replicative polymerase during DNA replication
<i>polA</i>	DNA Polymerase I	Processes Okazaki fragments and also fills in gaps during DNA repair processes
<i>polB</i>	DNA Polymerase II	Proofreading and editing, especially on lagging strand synthesis and some involvement in DNA repair
<i>ssb</i>	Single Stranded Binding Proteins (SSB)	Bind with single-stranded regions of DNA in the replication fork and prevent the strands from rejoining
A dimer encoded by <i>gyrA</i> and <i>gyrB</i>	DNA Gyrase	Type II Topoisomerase involved in relieving positive supercoiling tension caused by the action of Helicase
A dimer encoded by <i>parC</i> and <i>parE</i>	Topoisomerase IV	Type II Topoisomerase involved in decatenation of daughter chromosomes during DNA replication
<i>ligA</i>	DNA Ligase	Fixes nicks in the DNA backbone during DNA replication, DNA damage, and DNA repair processes

*Note: Only the genes involved in the formation of the catalytic domain of DNA polymerase III are listed*

(<https://wou.edu/chemistry/courses/online-chemistry-textbooks/ch450-and-ch451-biochemistry-defining-life-at-the-molecular-level/chapter-9-dna-replication-and-repair-2/>)

## Section F: Evolution as an Inherent Design Feature of DNA—Informational Adaptability and Systemic Feedback

Historically, the concept of evolution was approached from a top-down perspective, with a predominant focus on phenotypes—the observable traits of organisms—rather than the underlying genotypes that encode and determine those traits. Charles Darwin, formulating his theory of natural selection in the 19th century, did so without knowledge of molecular genetics. Consequently, evolutionary biology initially lacked insight into the molecular substrates—namely, DNA—that drive heritable variation and biological change.

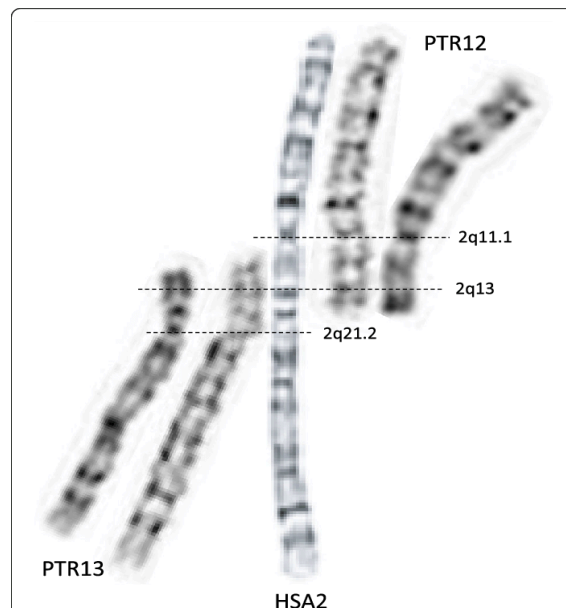
Modern understanding now reveals that evolution is not merely a descriptive framework of how organisms change over time, but a design feature intrinsic to DNA itself. At the molecular level, evolution describes how DNA maintains a finely tuned mutagenicity rate: low enough to preserve genomic integrity, yet flexible

enough to permit adaptation in response to environmental fluctuations. This dynamic allows DNA to not only respond to but also modulate its environment in a bidirectional feedback loop that enhances its own survivability.

Thus, evolution is more accurately conceptualized as a mechanism for optimizing the distribution of genetic information across the biosphere. The focus is not on the individual organism but on the persistence of DNA as an informational “entity”. Organisms/bodies are temporary vessels constructed by genes to propagate and optimize their own transmission. As Richard Dawkins famously articulated in *The Selfish Gene*, “We are survival machines—robot vehicles blindly programmed to preserve the selfish molecules known as genes.” Though not all of Dawkins’ conclusions are universally accepted, his foundational premise offers a compelling reorientation: evolution serves the gene, not the organism.

This perspective invites a critical reevaluation of evolutionary models. If an organism or species begins to threaten the integrity of the ‘global genomic system’, it may be phased out— via environmental pressures or population level genomic consequences. Humans, for example, have exerted unprecedented pressure on the biosphere, contributing to the ongoing sixth mass extinction and introducing existential threats such as nuclear weapons. These developments pose significant risks not only to individual species but to the long-term stability of DNA across the biosphere.

## F.1 HSA2 Anomaly - A Telomeric fusion event & it's implications for gradualist models



(Alignment of G-banded human (HSA2) and chimpanzee (PTR12 and PTR13) metaphase chromosomes - [https://www.researchgate.net/figure/Alignment-of-G-banded-human-HSA2-and-chimpanzee-PTR12-and-PTR13-metaphase-chromosomes\\_fig1\\_308663534](https://www.researchgate.net/figure/Alignment-of-G-banded-human-HSA2-and-chimpanzee-PTR12-and-PTR13-metaphase-chromosomes_fig1_308663534))

A particularly striking genomic anomaly is the fusion of human chromosome 2 (HSA2) - a merger of two ancestral ape chromosomes. This fusion results in a chromosomal count of 46 in *Homo sapiens*, compared to 48 in other great apes. This unique rearrangement poses a serious challenge to standard models of gradual speciation via random mutation, as individuals with different chromosome numbers typically cannot interbreed successfully. The HSA2 fusion stands as a genetic discontinuity that may signify a deeper, non-random influence guiding genomic evolution.

Mechanistically, this would involve the excision of the telomeric ends of chromosomes 2a and 2b—potentially requiring inversion for proper orientation—followed by their end to end fusion. This would initially result in a **dicentric chromosome**, which is typically unstable; thus, one centromere would need to be silenced to maintain mitotic viability. Dicentric chromosomes break during cell division. The mechanism of centromere inactivation is poorly understood.

“Erroneous DNA repair events linking two active centromeres generate dicentric chromosomes, whose



instability in mitosis is a major source of genome changes and a key contributor to oncogenesis. Dicentrics stem from double-strand break-induced and template switch-induced rearrangements, replication of hairpin-capped ends, and accidental telomere fusions (McClintock 1941; Bajer 1964; van Steensel et al. 1998; Lobachev et al. 2002; Mizuno et al. 2009, 2013; Stimpson et al. 2010, 2012; Marcand 2014). They can also be created by epigenetic neocentromere formation (Ishii et al. 2008; Stimpson and Sullivan 2010). Rare in normal contexts, double-strand break-induced dicentrics are frequent after exposure to ionizing radiation (Suto et al. 2013)."

<https://pmc.ncbi.nlm.nih.gov/articles/PMC4318148/>

The case for a single fusion event is compelling: for the chromosome to function and be heritable, all critical steps including telomeric truncation, fusion, and centromere inactivation would need to occur in close succession to establish a stably dividing chromosome. As one researcher notes: *"I propose that, unlike recurrent Robertsonian translocations in humans, the HSA2 fusion was a single nonrecurrent event that spread through a small polygamous clan population bottleneck. Its heterozygous to homozygous conversion, fixation".* (Stankiewicz)

For this fused chromosome to be both functional and heritable, the fusion site would require integration of a working gene complete with exonic and intronic architecture, minimal degradation in surrounding regions, and optimized satellite and telomeric sequences. Critically, such an event must occur in germline cells to be transmitted across generations.

What makes HSA2 particularly interesting is that it harbors critical genes associated with neural development & function. Based on accepted mutation rates ( $\sim 1$  in  $10^8$  per nucleotide per generation), the probability of  $\sim 20$ – $30$  specific, coordinated, functional mutations (encompasses structural and epigenetic changes) arising in a single generation is vanishingly small ( $10^{-160}$ – $10^{-240}$ ). Furthermore, successful reproduction would require a second individual with an identical chromosomal configuration, since 46 and 48 chromosome pairings typically result in reproductive barriers or inviability due to gametic mismatch. This necessitates either occurrence in two individuals with close temporal proximity or an alternate explanation.

*\*We are using a random mutation average to calculate the probability estimate - accounting for all structural and point mutations under one term for simplicity\**

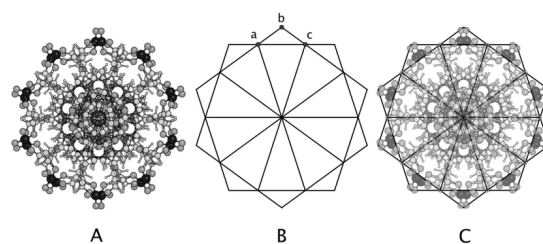
"We used human chromosome 2 as an example for the investigation of dicentric chromosome fate, since it is the most recent product of a chromosome fusion event fixed in the human lineage. It is largely known that the stability of the ancestrally dicentric chromosome 2 was gained by centromere inactivation....Previous studies showed that dicentric chromosomes overcame their instability via the inactivation of one of the two centromeres...In summary, our data support a model where human chromosome 2 likely overcame its instability by removing the chromosome IIq centromeric core located within the  $\alpha$ -satellite block C via a **one-step excision mechanism**."

<https://doi.org/10.1093/molbev/msx108>

Taken together, the precision, complexity, and improbability of this event challenge the sufficiency of unguided mutation and natural selection alone. It opens the door to reinterpreting this genomic transformation not as a random accident, but potentially as a highly coordinated, information-rich event - one that invites deeper inquiry into the informational architecture underlying biological systems.

## F.2. The Golden Genome: Emergent Order and Algorithmic Life

An additional layer of intrigue emerges when examining the structural and mathematical features of DNA and broader genomic systems. B-DNA, the canonical form of the double helix in most life on Earth, reveals a remarkable geometric property:



(Axial view of B-DNA (A) with a template of 10 "golden triangles" (B) shown overlain on the molecule (C). <https://www.mdpi.com/2073-8994/13/10/1949>)

"B-DNA, the informational molecule for life on earth, appears to contain ratios structured around the irrational number 1.618..., often known as the 'golden ratio'. This occurs in the ratio of the length/width of one turn of the helix; the ratio of the spacing of the two helices; and in the axial structure of the molecule which has ten-fold rotational symmetry. That this occurs in the information-carrying molecule for life is unexpected,

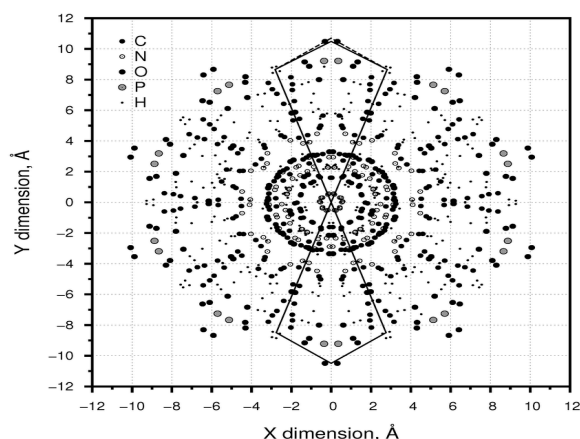
and suggests the action of some process. What this process might be is unclear, but it is central to any understanding of the formation of DNA, and so life." (Larsen)

This raises the possibility that DNA, rather than being an emergent product of purely random chemical evolution, may reflect deep mathematical constraints. The golden ratio is not only a mathematical curiosity—it is a number deeply embedded in biological structure, art, and natural design. Its consistent appearance in the spatial properties of DNA hints at a formative process.

This pattern extends from molecular architecture to genomic information systems. One study exploring codon-level genomic organization reports:

"We demonstrate that: (i) The whole Human Genome Structure uses the Universal Genetic Code Table as a tuning model. It predetermines global codons proportions and populations. The Universal Genetic Code Table governs both micro and macro behavior of the genome. (ii) We extend the Chargaff's second rule from the domain of single TCAG nucleotides to the larger domain of codon triplets. (iii) Codon frequencies in the human genome are clustered around 2 fractal-like attractors, strongly linked to the golden ratio 1.618." (Perez)

This implies the genome is not only chemically efficient but computationally elegant—its triplet code may be shaped by universal numerical laws not fully understood. Such tuning suggests that genome structure is not solely a product of adaptive selection but may reflect deeper algorithmic or mathematical principles that guide or constraint biological evolution.



(B-DNA viewed axially with super-imposed polygons as described in the text-<https://www.mdpi.com/2073-8994/13/10/1949>)

The pervasiveness of Fibonacci and golden-ratio patterns further supports this hypothesis. A 2012 review surveying developmental biology and paleontology states:

"A survey of zoological literature affirmed the wide occurrence of Fibonacci numbers in the organization of acellular and prokaryotic life forms as well as in some eukaryotic protists and in the embryonic development and adult forms of many living and fossil remains of metazoan animals." This recurrence is not limited to complex organisms but appears across phylogenetic scales, from microscopic life to vertebrate morphogenesis. Fibonacci ratios emerge in patterns of growth, segmentation, and spatial organization, even in noncoding DNA. (Willie)

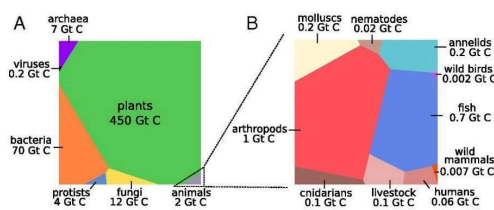
In botany, this same mathematical order governs the morphology of plants:

"Fibonacci phyllotaxis is commonly seen in all major groups of land plants. While a precise correlation is found between the internal pattern of the primary vascular system and the external pattern of appendages on the stem surface, it remains a big question how this regularity of Fibonacci phyllotaxis came into being in the course of evolution." (Okabe)

This widespread regularity across life forms cannot easily be explained as a side effect of natural selection alone, especially given the lack of immediate adaptive advantage in such geometric precision. Rather, these observations support a deeper view: that the emergence and structure of life are shaped by informational constraints and number-theoretic laws embedded into the substrate of biology.

Taken together, these insights converge & reinforce the argument that life's progression may not merely be a blind, stochastic walk across a fitness landscape, but instead a coordinated, information-rich trajectory, potentially orchestrated by principles not yet fully understood—principles that link mathematics, information theory, and biological form into a unified developmental framework.

Fig. 1.



Graphical representation of the global biomass distribution by taxa. (A) Absolute biomasses of different taxa are represented using a Voronoi diagram, with the area of each cell being proportional to that taxa global biomass (the specific shape of each polygon carries no meaning). This type of visualization is similar to pie charts but has a much higher dynamic range (a comparison is shown in [SI Appendix, Fig. S4](#)). Values are based on the estimates presented in [Table 1](#) and detailed in the [SI Appendix](#). A visual depiction without components with very slow metabolic activity, such as plant stems and tree trunks, is shown in [SI Appendix, Fig. S1](#). (B) Absolute biomass of different animal taxa. Related groups such as vertebrates are located next to each other. We estimate that the contribution of reptiles and amphibians to the total animal biomass is negligible, as we discuss in the [SI Appendix](#). Visualization performed using the online tool at [bionic-vis.biologie.uni-greifswald.de/](https://vis.biologie.uni-greifswald.de/).

(Graphical representation of the global biomass distribution by taxa" from Bar-On et al., 2018 - [https://www.researchgate.net/figure/Graphical-representation-of-the-global-biomass-distribution-by-taxa-from-Bar-On-et-al\\_fig3\\_334699717](https://www.researchgate.net/figure/Graphical-representation-of-the-global-biomass-distribution-by-taxa-from-Bar-On-et-al_fig3_334699717))

## Section G: The Neural Network of Genes—A Systems Biology View of Genomic Intelligence

At a systems level, genes collectively constitute a living information matrix—a vast, adaptive substrate through which information flows and evolves. Life, in this context, is best understood as information processing through the genome, with DNA acting as both the repository and executor of environmental feedback mechanisms. Organisms, in turn, function as biological nodes—autonomous sampling probes—through which DNA acquires data about the external world.

This networked model of genomic intelligence resembles an artificial neural network, in which distributed inputs from diverse phenotypes converge to influence a collective, system-level outcome.

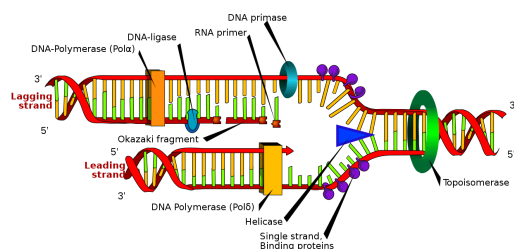
“Construction of gene regulatory networks (GRNs) is essential for elucidating the regulatory mechanisms underlying metabolic pathways, biological processes, and complex traits. In this study, we developed and evaluated machine learning, deep learning, and hybrid approaches for constructing GRNs by integrating prior knowledge and large-scale transcriptomic data from *Arabidopsis thaliana*, poplar, and maize.” (Gene regulatory network prediction using machine learning - <https://pmc.ncbi.nlm.nih.gov/articles/PMC12441907/>)

Genes may simultaneously influence multiple environmental variables via diverse phenotypic strategies across species. In this model, the genome's objectives may supersede the fitness of individual organisms. Traits that appear maladaptive at the individual level may serve broader genomic or ecological functions that benefit the gene pool as a whole.

As such, the conventional “organism-first” model of evolutionary biology becomes insufficient as Richard Dawkins envisioned in the *Selfish Gene*. The environment is not merely selecting for traits in a vacuum; DNA is actively probing and modulating the environment through its phenotypic outputs, orchestrating complex interactions across the biosphere. These interactions create layered feedback loops that mimic deep learning architectures, suggesting that the genome operates as a distributed computation system.

A study supports this paradigm shift. The authors describe how genomic interactions form complex structural and regulatory topologies that go beyond the organism-centric view of evolution and instead highlight large-scale, systems-level genomic behavior (Lieberman-Aiden et al.).

This model invites profound implications. Evolution, under this lens, is not solely governed by natural selection but by a multifactorial flow of information influenced by systemic genomic and environmental feedback. Natural selection remains a key player—but not the sole driver of evolutionary change.



(Components of DNA Replication. As DNA replication begins, Helicase separates the two strands of DNA to act as templates for replication. Single-strand binding proteins bind to each strand to stabilize and prevent them from reforming the double helix. Primase, an RNA polymerase, binds to the single-stranded DNA and synthesizes a short RNA primer in the 5' to 3' direction that is antiparallel to the parental strand. This RNA primer allows for DNA polymerase to begin replicating the DNA. Topoisomerase binds to the double helix upstream of the replication fork to prevent additional coiling by making small cuts in one of the DNA strands - <https://raider.pressbooks.pub/biology1/chapter/5-dna-replication/>)

## Section H: Abiogenesis Under the Microscope—A Probabilistic Impossibility?

The proposition that deoxyribonucleic acid (DNA)—the most sophisticated information storage and regulatory molecule known—could have arisen spontaneously through unguided, random processes not only strains the limits of biological plausibility but also the foundational principles of mathematics, logic, and probability theory. Whilst such statements are avoided in scientific nomenclature, the underlying methodological Naturalism assumed in biology needs to be examined deeply. When the complexity of DNA is analogized to human technology, the paradox is direct - DNA is exponentially more complex than any computer yet we entertain theories regarding DNAs spontaneous emergence seriously when anywhere else it would not be the case. This asymmetry between the mathematical constraints and the continued serious treatment of abiogenesis warrants examination as the data points in the opposite direction.

The core of this critique lies in the **overwhelming improbability** associated with the spontaneous generation of life's essential molecular machinery, given numerous hurdles facing incremental assembly models. If abiogenesis is to be upheld as a serious scientific hypothesis, then one must also concede the plausibility of virtually all events that are currently deemed impossible by probability theory, given the astronomical Improbabilities associated with abiogenesis. In this light, accepting abiogenesis on probabilistic grounds would require abandoning the methodological rigor that underpins science itself. To borrow from Occam's Razor—where the simplest explanation is usually preferred—abiogenesis, requiring

a cascade of highly specific and unlikely events, becomes the least parsimonious explanation.

## The Probability Problem: Borel's Law and the Limits of the Possible

The late mathematician Émile Borel articulated a guiding principle in probability theory: any event with a likelihood smaller than 1 in  $10^{50}$  can be considered practically impossible within the physical universe. While not an absolute physical law, Borel's threshold is widely used to contextualize the feasibility of events. According to this framework, **abiogenesis is not simply improbable but mathematically unjustifiable**.

For instance, the probability of forming a single functional protein through random amino acid assembly ranges from approximately 1 in  $10^{70}$  to 1 in  $10^{160}$ , depending on length and sequence specificity. The spontaneous emergence of a functional replicator carries an estimated probability as extreme as 1 in  $10^{10^{18}}$  (Koonin). These values render abiogenesis not merely unlikely, but **mathematically untenable** by standards of statistical reasoning alone.

Borel said that such events, while not technically impossible, are so improbable that it is irrational to expect them to occur in our universe

**Comparative Probability Benchmarks:** To further contextualize this improbability, let us consider several hypothetical scenarios that are objectively more probable than the spontaneous emergence of DNA, yet are universally dismissed as fantasy:

### 1. The Infinite Monkey Theorem:

If a monkey were to randomly type letters on a keyboard, the likelihood of it producing the full text of *Hamlet*, stripped of punctuation, is approximately 1 in  $3.4 \times 10^{183,946}$ . For comparison, the observable universe contains only about  $10^{80}$  atoms.

Now consider that the average human genome contains roughly 6.1 billion bases—each akin to a character in a coherent biological “text.” The probability of randomly assembling even the DNA sequence of a minimal viable cell (~500,000 base pairs or 1 million bases) far exceeds the improbability of typing *Hamlet* by chance. DNA is not merely a sequence—it is a functional script encoding enzymes, regulatory logic, and complex interactions that govern cellular life.



## 2. Cryptographic Analogy – 256-bit Encryption Key:

The number of possible 256-bit encryption keys is  $2^{256}$  ( $\sim 10^{77}$ ). Randomly guessing such a key is considered computationally infeasible and forms the basis of modern digital security. Yet this probability pales in comparison to the odds associated with spontaneously assembling a functional replicator.

## 3. 100 Heads in a Row:

Flipping a fair coin and landing on heads 100 times consecutively has a probability of  $\sim 7.9 \times 10^{-31}$ . This event is already considered effectively impossible in practice. Still, abiogenesis requires the successful convergence of far more improbable conditions under vastly more complex biochemical constraints.

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## The Case Against Random Origin

Given the probabilistic landscape, it is clear that abiogenesis stands **outside the bounds of probabilistic rationality**. The spontaneous emergence of life's molecular complexity requires such a series of improbable events that its inclusion in serious scientific discourse remains, at best, a profound inconsistency. It represents a philosophical contradiction in the foundations of biology. Despite knowing the informational density & complexity of DNA and its associated systems, abiogenesis appears to have garnered disproportionate traction that merits its own investigation. How did we overlook the numerous Mathematical, Biological & Chemical constraints to accommodate a model, which would not satisfy the scientific threshold of plausability.

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## Section I: Read, Write, Store, Execute, and Fabricate—DNA as a Complete Computational and Fabrication System

### I.1: DNA as a Self-Contained Information Processor and Architect of Life

DNA stands alone among natural molecules in its ability to **store, read, write, execute**, and even **fabricate**—serving as a complete, self-contained biological computing system. More than a static repository of data, DNA actively **generates** and **orchestrates** the formation of three-dimensional

molecular architectures: namely, proteins. These proteins, themselves extraordinarily complex polymers, are assembled through precise sequences encoded within DNA and interpreted through RNA transcription and translation mechanisms. The implications are staggering—DNA performs the computational equivalent of software writing its own hardware.

Here is a thought: You are not just learning about genomic ‘intelligence’; you’re experiencing its output.

The informational density and stability of the genome surpass even the most advanced digital storage technologies by orders of magnitude. It has been estimated that each human cell contains approximately **six feet of DNA**, and with a conservative estimate of **10 trillion cells per person**, the total length of DNA within a single human body is roughly **60 trillion feet**—or about **10 billion miles** (KQED). For reference, this is over **100 times the distance from Earth to the Sun**, which is approximately 93 million miles. When this is multiplied by the global human population—and further extended to include all animal, plant, fungal, and microbial life on Earth—the cumulative length of DNA forms an **information architecture of astronomical proportions**.

“Each human cell has around 6 feet of DNA... meaning each person has around 10 billion miles of DNA inside of them” (*KQED Science*).

“The current human reference genome includes 3.1 billion base pairs (3.1 Gb)” (*BioNumbers Database*, Harvard Medical School).

### I.2: One Functional Protein—An Impossibility by Random Assembly

The improbability of abiogenesis becomes even more pronounced when examined at the level of **individual proteins**. A functional protein is not simply a random string of amino acids—it is a folded, thermodynamically stable three-dimensional structure, often comprising hundreds of residues with precise hydrophobic patterns, active sites, and domain-level interactions. According to estimates published in *Journal of Molecular Biology* 2004 - the prevalence of random amino acid sequences yielding a functional protein fold is as low as **1 in  $10^{77}$** .

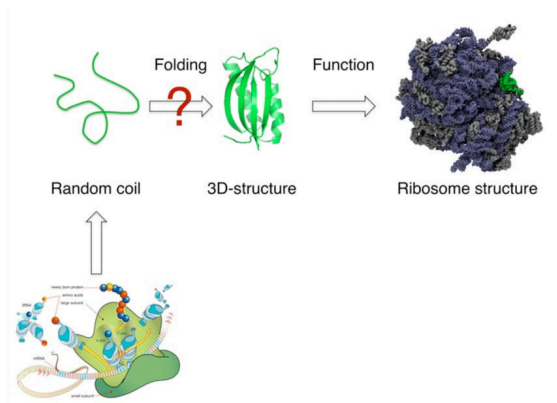
“This implies the overall prevalence of sequences performing a specific function by any domain-sized fold may be as low as 1 in  $10^{77}$ , adding to the body of evidence that functional

folds require highly extraordinary sequences” (Axe, 2004).

Moreover, studies such as the one by **Gauger and Axe (2011)** on the *Kbl2* and *BioF2* enzymes demonstrate that the evolutionary accessibility of new enzymatic functions—even between closely related homologs—may lie **beyond the temporal scope of Earth’s biological history**. Their findings suggest that even under optimal evolutionary assumptions, achieving a new functional enzyme would require **over  $10^{30}$  generations**, far exceeding the time available since life began.

“Successful functional conversion would... require seven or more nucleotide substitutions. Evolutionary innovations requiring that many changes would be extraordinarily rare... probable only on timescales much longer than the age of life on earth... This places the innovation well beyond what can be expected within the time that life has existed on earth” (Gauger and Axe, 2011).

Other estimates derived from biochemical modeling support similar conclusions. Folding simulations for ribosomal protein S6, for instance, reveal that sequence connectivity, overlapping foldons, and parallel folding pathways are essential to achieving functional conformation—further reducing the plausibility of random emergence.



(From RNA to a functional protein - [https://www.researchgate.net/figure/From-RNA-to-a-functional-protein-The-way-in-which-proteins-are-expressed-and-translated\\_fig1\\_265561300](https://www.researchgate.net/figure/From-RNA-to-a-functional-protein-The-way-in-which-proteins-are-expressed-and-translated_fig1_265561300))

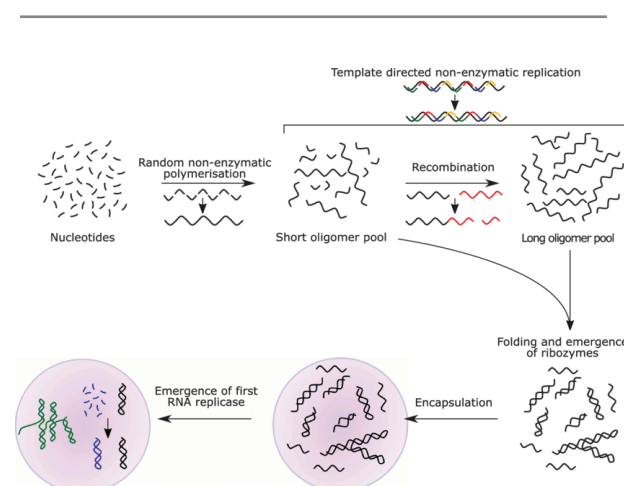
The folding of ribosomal protein S6 relies on coordinated sequence connectivity, overlapping foldons, and parallel folding pathways (Haglund).

When we combine these probabilistic assessments with the necessity for precise enzymatic coordination in DNA replication, transcription, translation, and protein folding, the case for **unguided molecular assembly is not supported by current probabilistic frameworks**.

## DNA as a Nano-Computer and Self-Assembling Fabricator

In sum, DNA operates as a **multifunctional nanotechnological system**, far surpassing even our most advanced human-engineered technologies. It **stores data** with unparalleled density, **executes instructions** via RNA intermediates, and **fabricates complex molecular machines** in three dimensions—at room temperature, under aqueous conditions, and with near-perfect error correction. The fact that it can do all of this within the confines of a living cell underscores a fundamental paradox: **such a system could not reasonably have arisen through chance-driven processes alone**.

The sheer volume, stability, and interdependence of genomic information systems demand **alternative frameworks for the origin of life**, ones that do not rely solely on chance but consider guided, informational, or perhaps even NHI mechanisms such as **directed panspermia**, an eventuality meriting discussion (Crick). As a computational and fabrication unit, DNA embodies the convergence of **software and hardware, design and execution, storage and synthesis**—all in one molecular architecture.



(A schematic representation of the classical RNA world hypothesis. Initially, synthesis and random polymerisation of nucleotides result in pools of nucleic acid oligomers, in which template-directed non-enzymatic replication may occur. Recombination reactions result in the generation of longer oligomers) <https://doi.org/10.1042/ETLS20190024>)

## Section J : The “RNA First” Hypothesis—A Critique of Its Scientific Viability

The hypothesis that RNA could have emerged prior to DNA and proteins as the first self-replicating molecule—the so-called “**RNA World**” hypothesis—has been widely circulated in the origin of life research. However, when subjected to rigorous mathematical scrutiny and biochemical feasibility, this model is not supported by current probabilistic frameworks. While the concept seeks to resolve the “chicken-and-egg” dilemma by positing that RNA preceded the appearance of both proteins and DNA, it only pushes the problem further upstream—without providing a viable mechanism for how such a system could emerge by chance.

The formation of even a single **self-replicating RNA molecule**, approximately 100 nucleotides in length, has been estimated to fall within a probability range of  **$10^{-120}$  to  $10^{-600}$**  depending on the assumptions involved. These estimates, derived from combinatorial models assuming random polymerization under idealized prebiotic conditions, place the **spontaneous emergence of RNA firmly outside the realm of statistical plausibility**.

“The probability of this latter process succeeding is so vanishingly small that its happening even once anywhere in the visible universe would count as a piece of exceptional good luck” (Shapiro, 2007).

“Many scientists believe life began with the spontaneous formation of a replicator. This idea has been supported by “prebiotic” syntheses carried out by chemists using modern apparatus and purified reagents. The probability that such reactions would take place spontaneously on the early Earth is minute” (Shapiro - <https://pubmed.ncbi.nlm.nih.gov/16776061/>)

As renowned NYU chemist **Dr. Robert Shapiro** emphasized in numerous papers and interviews, the idea that multiple nucleotide sequences could randomly assemble into both a non-catalytic RNA strand and a catalytic **ribozyme**—simultaneously and in the correct conformation—is not only improbable but **logically inconsistent**. A ribozyme is a specific RNA molecule folded into a tertiary structure, enabling it to catalyze biochemical reactions much like a protein enzyme. But such molecular folding itself depends on fine-tuned

thermodynamics, hydrogen bonding, and sequence-specific hydrophobicity—features that do not occur spontaneously.

To resolve the paradox of a self-replicating RNA molecule, some models simply add more RNA strands—each encoding catalytic activity—thus compounding the improbability problem with each addition. Indeed, as a 2022 review admits:

“However, no self-replicase has been identified to date. Alternatively, autocatalytic systems involving multiple RNA species capable of ligation and recombination may enable self-reproduction. However, it remains unclear how evolution could emerge in autocatalytic systems.” ([PubMed 36203246](https://pubmed.ncbi.nlm.nih.gov/36203246/)).

This concession highlights the **circular reasoning** and speculative layering that has plagued RNA-first models. The more one attempts to solve the catalytic and replication challenges by invoking auxiliary ribozymes or cooperative networks, the more astronomically improbable the entire framework becomes.

### Expert Testimonies: Shapiro, Hoyle, Crick, and Koonin

Dr. Robert Shapiro, a pioneer in the study of nucleic acids, consistently challenged the RNA World paradigm as deeply flawed. In his 2007 publication:

“Yet even if nature could have provided a primordial soup of suitable building blocks, whether nucleotides or a simpler substitute, their spontaneous assembly into a replicator involves implausibilities that dwarf those required for the preparation of the soup” (Shapiro, 2007).

No gradual or incremental processes have enough statistical robustness to withstand the challenge of informational entropy. Echoing this skepticism is **Sir Fred Hoyle**, a Cambridge astrophysicist and committed atheist, who famously calculated the improbability of life’s origin through random chance:

“The chance of obtaining all the necessary enzymes for life in a random trial is 1 in  $10^{40,000}$ ... an outrageously small probability that could not be faced even if the entire universe were an organic soup. The enormous information content of even the simplest living systems cannot... be generated by what are often

called ‘natural’ means” (Hoyle, *Evolution from Space*).

Even **Francis Crick**, co-discoverer of the DNA double helix and Nobel laureate, conceded the untenability of terrestrial abiogenesis, advocating instead for **directed panspermia**—the idea that life was seeded on Earth via extraterrestrial intervention.

Finally, **Dr. Eugene Koonin**, one of the foremost experts in evolutionary genomics, concluded that **abiogenesis is only remotely possible under a multiverse cosmological model**. In his analysis, the emergence of a primitive coupled translation-replication system—the minimum requirement for evolution—requires an improbability on the order of  $10^{-1018}$ .

“At a minimum, spontaneous formation of:

- two rRNAs with a total size of at least 1000 nucleotides
- ~10 primitive adaptors of ~30 nucleotides each, in total, ~300 nucleotides
- at least one RNA encoding a replicase, ~500 nucleotides (low bound) is required. In the above notation,  $n = 1800$ , resulting in  $E < 10^{-1018}$ .

In other words, even in this toy model that assumes a deliberately inflated rate of RNA production, the probability that a coupled translation-replication emerges by chance in a single O-region is  $P < 10^{-1018}$ . Obviously, this version of the breakthrough stage can be considered only in the context of a universe with an infinite (or, in the very least, extremely vast) number of O-regions.” (Koonin, *Biology Direct*, 2007).

Such a conclusion effectively **removes abiogenesis from the scope of empirical science**, placing it into the speculative domain of multiverse theory. In simpler terms, it becomes untestable and unrepeatable—violating core tenets of scientific methodology.

### The Intractable Paradox of RNA First

The “RNA First” hypothesis, though once hailed as a promising avenue for explaining life’s origin, has been increasingly revealed as **a theoretically appealing but**

**experimentally unsupported framework**. Its internal contradictions, dependence on unknown prebiotic conditions, and reliance on exponentially improbable molecular events place it squarely **outside the domain of scientific credibility**. It fails not only mathematically and chemically, but philosophically—offering no falsifiable mechanism and no viable model for the spontaneous emergence of complexity.

If one requires a ribozyme to form spontaneously to replicate RNA, and the ribozyme is itself RNA, then we arrive at yet another “chicken-and-egg” problem. The framework requires the product of the process it is attempting to explain.

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## Section K: Directed Panspermia and the Logical Paradoxes of Abiogenesis

### Directed Panspermia: A Serious Scientific Alternative

As our understanding of molecular biology, genetics, and biochemistry has advanced, traditional 19th-century models of abiogenesis—that life originated spontaneously from a prebiotic “soup”—have come under increasing scrutiny. In light of this, Nobel Laureate **Francis Crick** and chemist **Leslie Orgel** proposed an alternative hypothesis: **Directed Panspermia**. This theory suggests that life was deliberately seeded on Earth by an advanced Non Human Intelligence bypassing the immense improbabilities associated with spontaneous biogenesis on Earth.

In their seminal 1973 paper published in *Icarus*, Crick and Orgel wrote:

“We conclude that it is possible that life reached the Earth in this way, but that the scientific evidence is inadequate at the present time to say anything about the probability. We draw attention to the kinds of evidence that might throw additional light on the topic... Thus the idea of Directed Panspermia cannot at the moment be rejected by any simple argument. It is radically different from the idea that life started here ab initio without infection from elsewhere” (Crick & Orgel, 1973).

This proposal, while controversial, is not pseudoscientific. It arises from a sober assessment of the **informational, structural, and probabilistic**



**complexity** inherent in DNA, which—as later sections will demonstrate—renders spontaneous abiogenesis virtually untenable. While the probability of panspermia remains unquantified, its **logical consistency** and ability to account for the otherwise intractable hurdles of abiogenesis demand serious scientific attention. Many may view it as moving the problem from Earth to Planet x - the inevitability of infinite regress demands us to ask - who created the NHI?

## Logical Paradoxes of Abiogenesis

The spontaneous origin of life is undermined not only by improbability but by a suite of **fundamental logical contradictions** embedded in its theoretical framework. Below are two of the most critical:

### a) The Enzyme-DNA Paradox – The Ultimate Chicken-and-Egg Problem

For DNA or RNA to replicate, a complex ensemble of **enzymatic machinery** is required. This includes DNA polymerases, helicases, ligases, and proofreading enzymes. However, these very enzymes are themselves encoded by DNA (or, in some cases, RNA). This creates a closed loop of dependency:

- **DNA requires enzymes to replicate.**
- **Enzymes require DNA to exist.**

Such mutual dependency cannot arise incrementally, as **partial or non-functional intermediates would be useless**. Consequently, the entire system must be in place simultaneously for replication to occur—rendering the hypothesis of gradualistic, stepwise evolution from a simple molecule inherently flawed.

### b) Aqueous Instability of DNA – The Water Paradox

Contrary to the common narrative that life began in water, DNA is **chemically unstable in aqueous environments**. It is susceptible to **hydrolysis, oxidation**, and degradation by microbial activity. Experimental data demonstrates that:

“As is well known to forensic science, tissue and genetic material can persist for extended periods of time in conditions where oxygen and

microbial action are reduced or absent, such as DNA extracted from museum specimens or sediment and ice cores<sup>13</sup>. However, DNA can degrade rapidly (i.e., minutes) in aquatic environments due to hydrolysis, oxidation, and microbial activity.”(Seymour 2018).

“The covalent structure of DNA is unstable in aqueous solution... it tends to hydrolyze to its monomeric components... A single base transformation may be sufficient to cause a mutation or inactivate the DNA” (Shapiro 1981).

Even under mildly oxidative conditions, DNA’s half-life ranges between hours to days. The narrative that delicate nucleotides could spontaneously assemble in a pre biotic environment and remain stable long enough to evolve into complex life forms is **in direct contradiction** with known chemical kinetics.

This should not be confused with DNA’s half life in pure water at stable temperatures as those conditions are not reflective of realistic pre biotic chemistry

“Researchers examined the hydrolysis of dimethyl phosphate, as a model for the cleavage of phosphodiester linkages in DNA, and observed a half-time, pH-independent in the range near neutrality, of 140,000 years (primary source). Even at that level of stability, it can be inferred that a single DNA molecule from the human genome (with  $\sim 3 \times 10^9$  linkages of this kind) undergoes one backbone cleavage in 20 min. Accordingly, the genetic material would be highly unstable without enzymes to bring about DNA repair.”(Halftime of spontaneous DNA hydrolysis at 25°C-<https://bionumbers.hms.harvard.edu/bionumber.aspx?id=105355&ver=3>)

Thus, any abiogenetic scenario must explain not only the spontaneous formation of a vast and specific information-carrying polymer, but also how it was **shielded from rapid degradation** in the environment that allegedly enabled its creation. This presents a **double paradox**: the environment must be reactive enough to catalyze complex chemistry, yet gentle enough to preserve the results—a balance that is, as current science shows, **chemically contradictory**.

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## Section L: Homeostasis and the Cellular Paradox of Molecular Machinery

A central, yet often understated, requirement for the emergence and persistence of life is **homeostasis**—the capacity of a system to maintain internal stability amid external fluctuations. For DNA or RNA to function meaningfully within a biological context, it must be **encapsulated within a protective membrane**. This membrane does not merely serve as a passive boundary but operates as a dynamic, regulated interface that sustains cellular equilibrium. Without such regulation, nucleic acids would rapidly degrade due to oxidation, hydrolysis, and environmental volatility.

## The Membrane Challenge

Prebiotic models often propose that early life emerged inside **liposomes**—simple spherical vesicles composed of lipid bilayers. However, while liposomes can form spontaneously, they are **non-functional** in the biochemical sense. Real cell membranes are **highly specialized structures**, embedded with a vast array of **ion channels, proton pumps, transport proteins, and regulatory systems**. These components maintain essential electrochemical gradients and control osmotic pressure, ion flux, and nutrient uptake—each of which is vital for maintaining intracellular conditions conducive to life.

Without these regulatory systems, even a strand of intact DNA would be useless. Uncontrolled diffusion would rapidly collapse any emerging biochemical processes, and toxic buildup of ions would destroy nascent molecular structures.

## ATP Synthase: The Irreducibly Complex Proton Motor

Among the most astonishing components of cellular homeostasis is **ATP synthase**, a rotary molecular motor that synthesizes **adenosine triphosphate (ATP)**—the energy currency of all known life. This enzyme complex is embedded in the membrane and functions by exploiting the **proton gradient** across the membrane, a product of proton pumps and metabolic activity.

According to the *NCBI Molecular Cell Biology* reference:

“Detailed structural studies have established the mechanism of ATP synthase action, which involves mechanical coupling between the  $F_0$  and  $F_1$  subunits. In particular, the flow of

protons through  $F_0$  drives the rotation of  $F_1$ , which acts as a rotary motor to drive ATP synthesis” (NCBI Bookshelf).

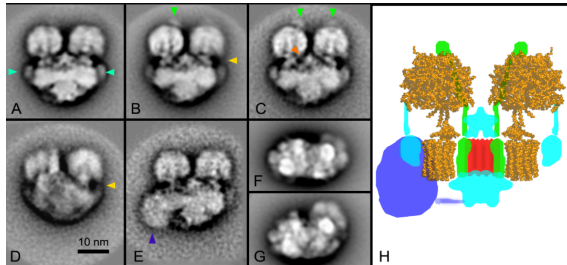
This motor, on average, produces approximately **40 kilograms of ATP per day per human**—a staggering feat of molecular coordination. The rotor spins at subatomic scales, converting proton motive force into mechanical rotation and then into chemical bond formation.

This essential cellular component is pre LUCA. Implying that the existence of ATP synthase dates to the earliest life forms. This temporal constraint is critical: if ATP synthase existed in Earth's first cells, then the circular dependency described in Section B—DNA requiring ATP, ATP requiring ATP synthase, ATP synthase requiring DNA—must have been resolved instantaneously at life's origin. No gradualistic evolutionary pathway could operate before functional ATP generation, yet ATP generation requires the complete genetic machinery. The presence of maximal complexity in minimal time further undermines naturalistic assembly scenarios. This is not merely a complex enzyme that evolved over billions of years—this complexity existed in Earth's first cells, before natural selection could refine simpler precursors.

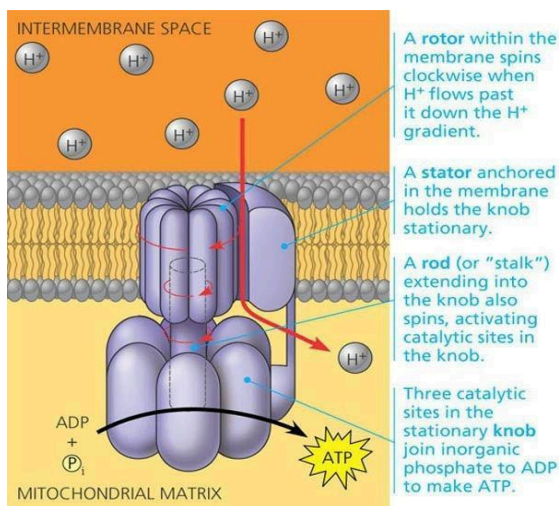
“The ATP synthase complex is thought to have originated prior to the Last Universal Common Ancestor (LUCA) and analyses of ATP synthase genes, together with ribosomes, have played a key role in inferring and rooting the tree of life. We reconstruct the evolutionary history of ATP synthases using an expanded taxon sampling set and develop a phylogenetic cross-bracing approach, constraining equivalent speciation nodes to be contemporaneous, based on the phylogenetic imprint of endosymbioses and ancient gene duplications. This approach results in a highly resolved, dated species tree and establishes an absolute timeline for ATP synthase evolution. Our analyses show that the divergence of ATP synthase into F- and A/V-type lineages was a very early event in cellular evolution dating back to more than 4 Ga, potentially predating the diversification of Archaea and Bacteria” <https://www.nature.com/articles/s41467-023-42924-w>

The architecture and mechanism of ATP synthase have led many to argue that it represents a clear case of **irreducible complexity**. Its function requires all parts—rotor, stator, proton channel, and catalytic head—to operate simultaneously and in perfect coordination. Any partial version of the enzyme would

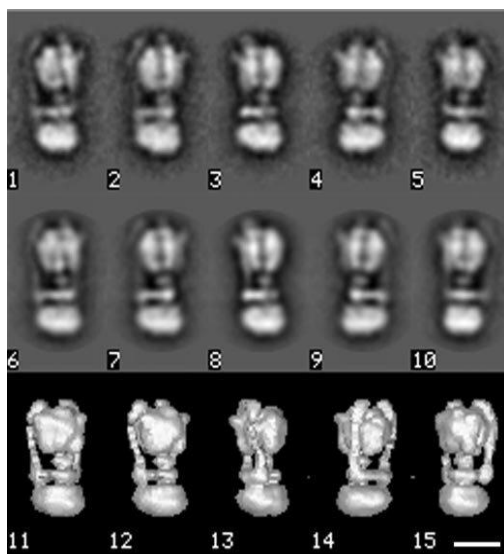
be non-functional and would not confer selective advantage. As such, the gradualistic pathway proposed by Darwinian evolution appears insufficient under current models to explain the emergence of such a system.



Highly Divergent Mitochondrial ATP Synthase Complexes in *Tetrahymena thermophila* - <https://doi.org/10.1371/journal.pbio.1000418>

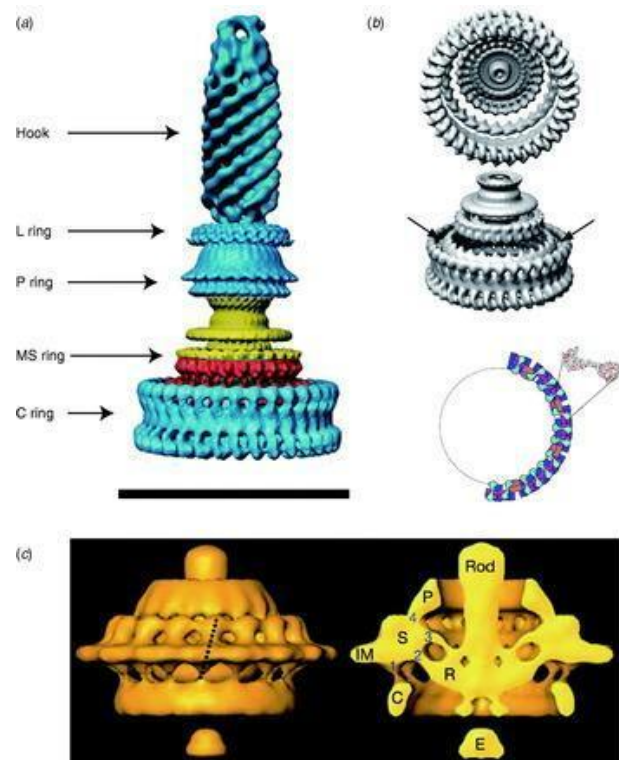


Comprehensive Diagram for the Structure of ATP Synthase - [https://www.researchgate.net/figure/Comprehensive-Diagram-for-the-Structure-of-ATP-Synthase-Taken-from-3\\_fig1\\_357759332](https://www.researchgate.net/figure/Comprehensive-Diagram-for-the-Structure-of-ATP-Synthase-Taken-from-3_fig1_357759332)



Three-dimensional Structure of A1A0 ATP Synthase from the Hyperthermophilic Archaeon *Pyrococcus furiosus* by Electron Microscopy - [https://www.jbc.org/article/S0021-9258\(20\)32273-0/fulltext](https://www.jbc.org/article/S0021-9258(20)32273-0/fulltext)

## Beyond ATP Synthase: Molecular Motors and Bioengineering Precision



(The structure of the flagellum extracted from *S. typhimurium* cells - <https://www.cell.com/current-biology/fulltext/S0960-9822%2806%2902286-X#fig1>)

ATP synthase is not the only rotary motor found in biology. The **bacterial flagellum**, another molecular rotary engine, operates like an outboard motor, with a drive shaft, rotor, and stator—all composed of protein subunits. It enables bacterial locomotion at remarkable speeds relative to cell size and is similarly dependent on ion gradients and highly coordinated assembly processes.

These biological machines function by **harnessing subatomic particles and exploiting quantum-level gradients**, a level of coordination and efficiency unmatched in any known human technology.

To suggest that these molecular motors **emerged spontaneously** via undirected processes not only lacks supporting evidence but contradicts our current understanding of information processing, engineering principles, and biochemical kinetics. The complexity, specificity, and interdependence of components defy the plausibility of sequential, random mutations producing these systems by natural selection alone.

## Conclusion: Cellular Homeostasis as a Hallmark of Non-Random Origin

The requirement for homeostasis reveals yet another fatal flaw in abiogenetic models. Life does not emerge in a vacuum—it requires containment, regulation, and energy conversion. A strand of DNA or RNA without a membrane is inert. A membrane without functional protein pumps and motors is a leaky container. And energy metabolism without ATP synthase is biologically non-viable.

The emergence of such integrated systems—where **every part is essential from the outset**—suggests a level of foresight, coordination, and functional interdependence that naturalistic explanations have yet to account for. Whether one favors models of **abiogenesis**, **directed panspermia**, or alternative frameworks, what becomes increasingly clear is that **undirected evolution via natural selection alone is insufficient** to explain the precision machinery observed at the molecular level of life.

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## Section M: The Oxidation Dilemma – A No-Win Scenario for Prebiotic Chemistry

One of the most underappreciated yet critical obstacles to abiogenesis is the **oxidation paradox**. DNA and RNA are **highly sensitive to oxidative environments**, and yet, any atmosphere rich in oxygen—such as Earth's current atmosphere—would rapidly degrade nucleic acids through oxidation. Conversely, if we assume an early Earth atmosphere devoid of oxygen, we face another catastrophic challenge: **the absence of an ozone layer**, which would allow **unfiltered ultraviolet radiation** to bombard the planet's surface. This radiation is potent enough to break molecular bonds and obliterate nucleic acid structures.

In essence:

- **With oxygen:** nucleic acids oxidize and degrade.
- **Without oxygen:** nucleic acids are destroyed by UV radiation.

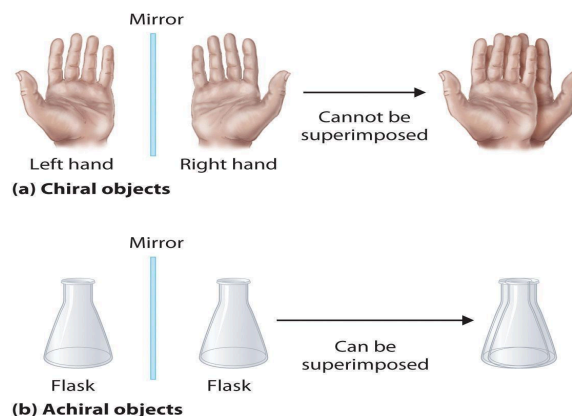
To reconcile this, abiogenesis models often speculate that nucleic acids formed in **non-aqueous, non-oxidizing environments**, possibly shielded within **volcanic vents** or **subsurface mineral matrices**. However, this hypothesis veers into speculative territory

with **minimal empirical support** and contradicts multiple observed chemical constraints.

Even if we imagine a perfect environment—one that simultaneously avoids oxygen, UV radiation, hydrolysis, and a plethora of biochemical chemical constraints — the spontaneous formation and stabilization of complex molecules such as DNA or RNA still **does not follow** from known prebiotic chemistry. The problem isn't just the environment—it's the **complexity and fragility** of the information inside the molecules, in addition to the required structural stability.

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## Section N: Chirality – A Critical Problem in Abiogenesis Theories



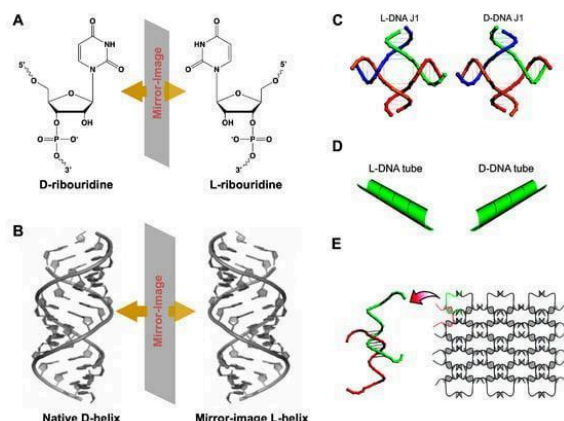
“The homochirality of biological molecules (the use of only left handed amino acids and only right handed sugars) has long been known to be an important characteristic of life. Current ideas on the origin of life do not explain the origin of homochirality, yet the widely accepted ‘RNA world’ model cannot work without it.”(Jeremy Bailey-<https://www.sciencedirect.com/science/article/abs/pii/S0094576500000242>)

A second, often overlooked issue that undermines the plausibility of abiogenesis is the problem of **molecular chirality**. Organic molecules, including amino acids and sugars, exist in **two mirror-image forms** known as **enantiomers**—designated as **left-handed (L)** and **right-handed (D)**. These forms are chemically identical but **non-superimposable**, akin to one's left and right hands.



Crucially, **all known life** on Earth exhibits **monochirality**:

- **Proteins** are composed exclusively of **L-amino acids**.
- **DNA and RNA sugars** are made entirely of **D-ribose or D-deoxyribose**.



[(A) Chemical structures of D-ribose, L-ribose, and (B) duplex of native D-DNA and L-DNA. (C) Structural models of L- and D-DNA J1 molecules. (D) The opposite chirality of the L- (left-column) and D- (right column) DNA nanotubes revealed. (E) Schematic drawings of two-dimensional nanoarrays self-assembled from L-DNA. (C-E) - <https://www.mdpi.com/2073-4425/13/1/46>]

This is not a trivial characteristic—it is absolutely essential. Even a **single misaligned enantiomer** in a protein or nucleic acid can **disrupt folding**, render the molecule nonfunctional, or even introduce toxicity. For abiogenesis to be viable, one must explain not just how complex molecules formed, but **why only one chiral form was selected across all life**, and **how that selection was enforced in prebiotic conditions**.

## Limitations of Prebiotic Experiments

The famous **Miller-Urey experiment**, often celebrated for producing amino acids under simulated early Earth conditions, failed to account for chirality. It produced a **racemic mixture** - equal parts of left and right handed amino acids. Such mixtures are **non-functional in biological systems**. Moreover, the experiment required **highly purified industrial reagents that do not reflect natural geological environments**, and it generated **toxic byproducts** that would be detrimental to DNA, RNA, and cellular stability.

The methodological contradictions inherent in the work of Sutherland, Szostak, and the investigators of the

QT-45 ribozyme reveal a fundamental divergence between experimental success and prebiotic relevance. In these frameworks, the transition from chemical chaos to biological information is not an emergent property of undirected matter; rather, it is the direct result of **researcher intelligent interventions** that bypass the stochastic hurdles and degradative pressures inherent in standard abiogenesis environments.

Every milestone in these studies is achieved by manually removing the "hard problems" facing abiogenesis. Rather than demonstrating the *de novo* emergence of complexity, these experiments rely on highly purified, specialized reagents, components essentially "bought pre-made" that possess a level of structural and chemical fidelity impossible to maintain in a primitive, unshielded environment.

“Experimentalists in the field of prebiotic chemistry strive to re-enact what may have happened when life arose from inanimate material. How often human intervention was needed to obtain a specific result in their studies is worth reporting....Organic chemists, if not all experimentalists in the field of prebiotic chemistry, are faced with a similar dilemma. We do our best to perform experiments that we believe re-enact possible steps of prebiotic evolution, but we know that we need to intervene manually to obtain meaningful results. Further, the ideal experiment does not involve any human intervention. A reaction or a reaction network is allowed to unfold, and the sample is only broken up when the experiment has been finished. Perhaps, samples are drawn, as in the famous Miller experiment<sup>L2</sup>, but there is no addition of new chemicals or an artificial change in conditions.” (Clemens Richert-<https://pmc.ncbi.nlm.nih.gov/articles/PMC6290120/>)

Sutherland, for instance, must stage sequential reactions, manually cleaning up toxic byproducts and controlling pH and UV exposure at necessary steps to prevent the system from degrading into non-functional chemical noise. (<https://pmc.ncbi.nlm.nih.gov/articles/PMC4568310/>)

Similarly, in the QT-45 study this intervention is manifest in the use of a pre-synthesized, trillion-member RNA library. By manually orchestrating the iterative selection process through cycles of amplification and reverse transcription, the researchers provide the necessary replication fidelity and error correction that are naturally absent in prebiotic environments. An illustration of directed or

guided selection by intelligent researchers, as an example:

“We initiated selections from three small, random sequence pools ( $\sim 6 \times 10^{12}$  unique RNA sequences) each containing a short (20, 30 or 40 nucleotides) randomized region constituted as a tandem repeat (fig. S1). We challenged the pools first to catalyze a single templated ligation of a primer to an adjacent triphosphorylated substrate that is covalently linked to the library via a flexible RNA linker (Fig. 1A, top). Once catalytic activity was observed, we challenged the pools to catalyze triplet polymerization” (<https://pmc.ncbi.nlm.nih.gov/articles/PMC7618777/>)

The reliance on stabilized precursors and researcher-directed environments masks the fundamental "chicken & egg" paradox. By providing the high purity inputs and environmental shielding that nature lacks, these models do not prove life can emerge without design. Instead they provide a precise quantification of the specific interventions required to overcome the natural tendency of undirected chemistry toward disorder. Viewed philosophically, the methods sections of these papers serve as the most powerful evidence against their own abstracts, documenting the necessity of specific, directed parameters to achieve functional molecular results.

“A critical question to ask when seeking to apply new results in origins is: how can this chemistry occur without direct intervention from a chemist?” (Do-Nothing Prebiotic Chemistry-<https://pmc.ncbi.nlm.nih.gov/articles/PMC11713876/>)

This general pattern permeates deeply in modern origin of life research, thus individually listing and critiquing each one specifically becomes redundant whilst the underlying concept of directed chemistry becomes apparent.

## Chirality in Biological Systems – A Case Study in Improbability

Consider **T4 DNA ligase**, an enzyme essential for DNA repair and recombination. It comprises **487 amino acids**, each of which must be of **precise chirality** and **correct sequence** to fold into a functional 3D structure. With average amino acids measuring just 0.4 to 1 nanometer, the margin for error is infinitesimal. A

single inversion in chirality would lead to catastrophic misfolding.

“The T4 DNA ligase is a single polypeptide with 487 deduced amino acids” (*ScienceDirect*). <https://www.sciencedirect.com/topics/pharmacology-toxicology-and-pharmaceutical-science/dna-ligase>

The exclusivity of chirality drastically reduces the already astronomically low probability of abiogenesis, **cutting the likelihood in half at every molecular junction**.

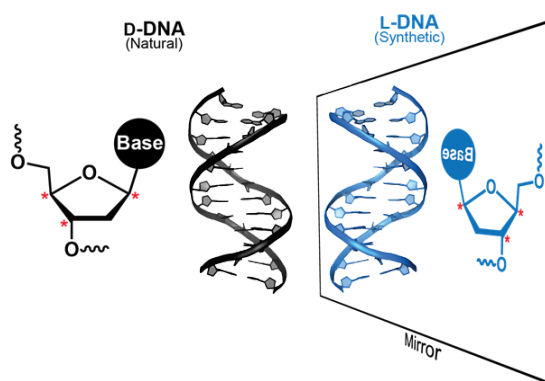
## D-DNA vs. L-DNA – A Biological Enigma

Ironically, DNA itself can exist in two chiral forms:

- **D-DNA**, which is the naturally occurring form in all known life.
- **L-DNA**, its **mirror image**, which is structurally stable and biochemically viable, yet **absent in nature**.

“L-DNA strand-displacement systems show superior intracellular stability and minimal leak... compared to D-DNA” (*ACS Synthetic Biology*, 2020).

“L-DNA is more stable than D-DNA to enzymatic degradation...” yet life propagates through the less stable form (*GeneLink*).



(The two enantiomers of DNA. Chirality centers are indicated with an asterisk-<https://www.glenresearch.com/reports/gr33-21>)

Despite its **greater biochemical resilience**, L-DNA has **never been observed as the basis for life**, raising profound questions about the origin and selection of chirality. Even more perplexing, L-DNA and D-DNA

**cannot hybridize**—they are completely incompatible, underscoring how deeply embedded chirality is in life's molecular foundation.

## The Impossibility of Random Assembly Under Natural Conditions

To produce a single viable cell, one would need:

- Hundreds of **chiral amino acids** arranged in a specific sequence.
- Hundreds of **chiral sugars and phosphates** to build the DNA backbone.
- Dozens of highly specific **enzymes** to replicate, splice, and repair DNA.
- Encapsulation within a **membrane** equipped with **ion pumps** and **proton channels**.
- An operational **ATP synthase** motor to generate energy.
- A stable environment devoid of oxidative damage and radiation.

This degree of coordination, orientation, and chemical selectivity is so extreme that even **one misstep** in chirality, folding, or assembly could render the entire system inoperative.

## Conclusion: Chirality and Oxidation as Foundational Challenges to Abiogenesis

Both the **oxidation paradox** and the **chirality problem** strike at the core of abiogenesis. These are not esoteric or edge-case issues—they are foundational hurdles that must be addressed to validate any naturalistic origin-of-life model.

- **Oxidation** renders DNA and RNA chemically nonviable in both oxygen-rich and oxygen-poor environments.
- **Chirality** demands a precision that defies all known random prebiotic chemistry.
- The use of less stable D-DNA over L-DNA, despite the latter's superior biochemical properties,

raises further doubts about the randomness of molecular selection.

## Section O: Minimal Genome – A Mathematical Refutation of Abiogenesis

Perhaps the most compelling, empirical challenge to the theory of abiogenesis arises not from philosophical speculation, but from **experimental science itself**. In a groundbreaking study, scientists at the J. Craig Venter Institute (JCVI) synthesized the **simplest known cell capable of sustaining life**, termed **JCVI-syn3.A**. By systematically removing all non-essential genes, researchers were able to isolate the **minimal genetic blueprint required for autonomous cellular function**.

The result? A genome consisting of **493 genes**.

According to standard molecular biology estimates, the **average coding sequence per gene is approximately 1,000 base pairs (bp)**. This places the minimal genome of JCVI-syn3.A at roughly **493,000 base pairs**—and this is under **highly optimized laboratory conditions**, with significant biochemical support and no environmental challenges.

“The average size of a protein molecule allows one to predict that there are approximately 1,000 nucleotide pairs of coding sequence per gene”

*(National Research Council (US) Committee on Mapping and Sequencing the Human Genome.)*

<https://www.ncbi.nlm.nih.gov/books/NBK218247>

Let us now consider the statistical implications of such a genome forming **spontaneously**.

## Probability Calculations for Minimal Genomes

DNA is composed of **four nucleotides**: A, T, C, and G. Therefore, the probability of correctly selecting one nucleotide by chance is **1 in 4**, or 0.25. The probability of randomly assembling a precise sequence of  $n$  base pairs is therefore  $(1/4)^n$ .

$(1/4)^n$  is generous to abiogenesis as additional constraints like chirality etc place the realistic probability  $(1/16)^n$  to  $(1/64)^n$  or worse\*

### Case 1: JCVI-syn3.A – 493,000 base pairs

Using the lower-bound estimate of 1,000 base pairs per gene:

- $P = (1/4)^{493,000} \approx 1 \text{ in } 10^{296,000}$

If we use the upper-bound estimate of 2,000 base pairs per gene (to reflect regulatory and intronic regions):

- $P = (1/4)^{986,000} \approx 1 \text{ in } 10^{592,000}$

These are not just small numbers—they are **orders of magnitude beyond anything conceivably occurring by chance** within the age or physical confines of the universe.

### Case 2: Hypothetical Reduction to 100 Genes

Assuming a radically reduced minimal genome of just **100 genes** (highly unrealistic), each consisting of 1,000 nucleotides:

- $P = (1/4)^{100,000} \approx 1 \text{ in } 10^{60,000}$

Even in this most charitable scenario, the probability remains **astronomically low**, still exceeding the estimated number of particles in the observable universe ( $\sim 10^{80}$ ).

*\*JCVI-syn3A is a genetically minimal bacterial cell, consisting of only of 493 genes on a single 543-kbp circular chromosome”  
<https://doi.org/10.1016/j.cell.2021.12.025>*

*1 kbp = 1000 base pairs therefore 543 kbp=543,000 base pairs*

*P=10<sup>-326,918</sup> - the calculation in the paper underestimated the improbability in its lower bound estimate but used a 1000 gene avg assumption for simplifying the calculation\**

## MATHEMATICAL INVERSION - WHAT CAN THE UNIVERSE GENERATE?

Now we present an inversion of the math to calculate the Universe's maximal upper bound capacity to generate a genome

Given that there are  $10^{80}$  particles

Time elapsed Age of universe =  $4.35 \times 10^{17}$  seconds

Max reactions per atom per second  $10^{13}$

$$\text{Total trials} = 10^{80} \times 10^{13} \times 4.35 \times 10^{17} = 4.35 \times 10^{(80+13+17)} = 4.35 \times 10^{110} \text{ total reactions}$$

**For a specific sequence of length n base pairs:**

$$\text{Probability} = (1/4)^n$$

$$\text{For at least one success: } 4.35 \times 10^{110} \times (1/4)^n \geq 1$$

**Solving for n:**

$$(1/4)^n \geq 1 / (4.35 \times 10^{110})$$

$$4^n \leq 4.35 \times 10^{110}$$

$$n \times \log(4) \leq \log(4.35 \times 10^{110})$$

$$n \leq 110.64 \div 0.602 = 184 \text{ base pairs}$$

**The universe's absolute maximum random sequence length: ~184 bp**

**For reference a single Gene is ~1000 base pairs on average**

**This is 18% of a single Gene and about 0.03% of the 543,000 base pairs of JCVI-syn3.A**

The observable universe lacks sufficient atoms, time, and reactions to randomly assemble even 200 base pairs of specific DNA sequence—yet the simplest known life requires over 500,000.

Let's further this idea by applying it to just the Earth - if assumed that the Earth's atoms are also reacting since the universe began, a planetary primordial soup, an extremely generous concession for abiogenesis, we would only maximally yield 134 Base pairs.

Atoms on Earth:  $1.33 \times 10^{50}$



Reaction rate:  $10^{13}$  reactions per atom per second  
 Time available:  $4.35 \times 10^{17}$  seconds (age of universe)  
 Total reactions:  $1.33 \times 10^{50} \times 10^{13} \times 4.35 \times 10^{17} = 5.79 \times 10^{80}$

For a specific sequence of  $n$  base pairs to form at least once:

Required:  $5.79 \times 10^{80} \times (1/4)^n \geq 1$   
 Solving:  $n \leq 80.76 \div \log_{10}(4) = 80.76 \div 0.602 = 134.1$  base pairs

Borel's Threshold Analysis:

Borel's Universal Probability Bound: Events with probability  $< 10^{-50}$  are considered mathematically impossible in our universe.

For DNA sequences:  $(1/4)^n = 10^{-50}$

Solving:  $n = 50 \div \log_{10}(4) = 50 \div 0.602 = 83$  base pairs

This represents a Mathematical firewall on random assembly of complex systems in the universe. The gap is so large no amount of concession or gradualist modelling resolves the discontinuity.

Despite the model appearing simple, if one were to calculate realistic earth bound scenarios, the probability space dramatically shrinks and your maximum base pair limit falls even lower.

Even if we assume only 1% of the minimal genome needs to be specified - 5,430 base pairs is still many magnitudes beyond 184 base pairs. The probability is  $10^{-3269}$ , lying far beyond Borel's Threshold.

## Can Selection Bias help?

Given that Universe trials =  $4.35 \times 10^{110}$  total reactions

$\log(4.35 \times 10^{110}) = 110.638$

P (corrected Probability) per position

Formula

$n = \log(\text{Total Trials}) / -\log(P)$

P (Corrected Probability)	$-\log(P)$	Base Pairs $110.638 / -\log(P)$	bias %
0.25 or 1/4 (random)	0.602	184	25
0.50	0.301	368	50
0.75	0.125	885	75
0.90	0.0458	2416	90
0.95	0.0223	4961	95
0.995	0.00218	50,751	99.5
0.999	0.000435	254,340	99.9
0.9995	0.000217	509,853	99.95
0.9999	0.0000434	2,547,406	99.99

The table indicates that the required selection bias to reach the minimal genome is ~99.95%. No known stochastic or selective process operates at this required fidelity in a pre biotic environment. Approaching merely 1% of the minimal genome (5430bp) requires selection specifying over 95%. This is outside the domain of known stochastic processes usually associated with abiogenesis. This calculation already assumes the totality of the universe as a Genome generator whilst any realistic calculation would yield remarkably reduced base pair numbers. This serves as the maximal upper bound concession in favor of abiogenesis. A bias so strong indicates what type of process may be required to explain the instantiation of life.

## Implications for Abiogenesis

The probabilistic landscape presented by the simplest life-supporting genome is **irreconcilable with any random or undirected process**. These calculations do not merely cast doubt on abiogenesis; they constitute a **mathematical untenability** of it under the assumptions of standard chemistry, thermodynamics, and information theory.

To uphold abiogenesis in light of these numbers requires one to **suspend known laws of biochemistry**,

ignore **entropy constraints**, and posit **unverified mechanisms** for spontaneous complexity. In essence, it amounts to **arguing for a miracle**—the very kind of explanation science typically seeks to avoid.

## The DNA Information Paradox

DNA is more than a molecule; it is an **information-processing system** of unparalleled sophistication. It performs:

- **Storage** with densities far beyond modern silicon-based technology.
- **Self-replication** with astonishing fidelity, often exceeding 1 error in  $10^9$  nucleotides.
- **Self-correction** through error-checking enzymes.
- **Execution** via transcription and translation into functional proteins.

Such performance places DNA in the domain of **algorithmic precision** and **computational design**. No known system—natural or artificial—approaches this level of integrated functionality and resilience.

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## Conclusion: The Inescapable Inference

The cumulative findings presented in this work offer a comprehensive reassessment of DNA as not merely a biochemical polymer, but as a nanotechnological system exhibiting properties that are both informationally dense and structurally optimized. By synthesizing insights from quantum physics, molecular biology, information theory, and thermodynamics, this paper challenges the prevailing paradigms that have traditionally underpinned abiogenesis and evolutionary theory.

DNA's stability at the quantum scale—despite being susceptible to proton tunneling and quantum decoherence—indicates an inherent capacity for preserving informational coherence that defies predictions made by classical chemistry. Its chirality, replication fidelity,  $\pi$ -stacking interactions, and capacity for error correction demonstrate an interdependent system of components that cannot be linearly reduced to simpler precursors without forfeiting function. The

informational paradox is perhaps most salient: DNA stores, transmits, and executes digital instructions with a density and reliability unmatched by any known human-engineered system. This informational functionality is coupled with the ability to produce complex 3D molecular structures such as proteins—thus acting as a read-write-execute substrate.

Furthermore, probabilistic analyses rooted in the mathematics of combinatorics and information entropy reveal that the spontaneous formation of a minimally viable genome, such as that of JCVI-syn3.A with 493 essential genes, lies far beyond the threshold of plausibility defined by Borel's Threshold. Even under optimistically reduced parameters, the probability of assembling functional nucleic acid sequences by chance remains effectively zero. These estimates do not merely cast doubt on abiogenesis—they render it mathematically and biochemically untenable.

The issue of biological homochirality further undermines the naturalistic synthesis of life. No known physical law mandates the exclusive emergence of left-handed amino acids and right-handed sugars, and yet life universally adheres to this asymmetry. Abiotic chemistry invariably yields racemic mixtures, which are biologically incompatible and often toxic. The absence of a known selection mechanism for chirality in prebiotic environments points again to an organizing principle beyond random thermodynamic drift.

Finally, the recursive dependency between DNA and the enzymatic machinery necessary for its own replication and expression introduces an ontological dilemma—one that cannot be resolved within the constraints of gradualist evolutionary theory. The spontaneous co-emergence of all essential components in a functional configuration requires either a fundamental rethinking of physical laws or the introduction of directed causality.

Taken together, these constraints—biochemical, informational, environmental, and probabilistic—demand the consideration of alternative frameworks for the origin of life. Among these, the theory of directed panspermia, as proposed by Crick and Orgel, offers a logically consistent alternative that circumvents the paradoxes intrinsic to terrestrial abiogenesis. More broadly, the evidence suggests that life is not the consequence of undirected chemical accidents but of processes governed by directed

informational causality, whether instantiated by natural laws not yet understood or by external agency.

In conclusion, DNA must be understood not simply as the product of evolution but as its architect—a molecular information system that encodes, preserves, and propagates life with a degree of precision that invites deeper investigation into the origins of information itself. The path forward in origin of life research must therefore shift from purely chemical narratives toward those that interrogate the informational and algorithmic foundations of biology. Only through such a paradigm shift can science hope to resolve the most foundational question it faces: not merely how life persists, but how it began.

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*Every conclusion here points backward to a question the reader hasn't asked.*

## Abiogenesis Models - Maximum Base Pair Limits $n \leq \log_{10}(\text{Total Trials}) \div \log_{10}(4)$

Model	Max BP
Entire Universe	184
Entire Earth	134
All Oceans	116
Primordial Soup	111
Deep Hydrothermal Vents	112
Shallow Hydrothermal Systems	108
Submarine Alkaline Vents	108
Tidal Pools (Wet-Dry Cycles)	106
Iron-Sulfur World	102
Lipid World	100
RNA World (Prebiotic)	100
Clay Mineral Surfaces	98
Deep Crustal Environments	98

Ice Eutectic Phase	96
Mineral Pore Spaces	96
Atmospheric Aerosols	96
Volcanic Glass Surfaces	95
Formamide Pools	95
Phosphate-Rich Environments	93
Cyanosulfidic Chemistry	93
UV-Irradiated Surfaces	93
Metabolic Cycles (Autocatalytic)	93

**\*All models assume constants generous to Abiogenesis. Minor variations in total trials affect the final results by  $\pm 10$  base pairs\***

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