## The Transition to Life

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## Based on Simplest Life Now:

Need:

- 1. Nucleic Acids
- 2. Proteins
- 3. Lipids
- Carbohydrates (Pigments)

Replicatable Information Enzymes (Catalysts) Membranes (Enclosure) Energy Storage (Energy Conversion)

Too much to ask of chemical evolution  $\Rightarrow$  Protolife?

## <u>Protolife</u>

 "Virus" Free living but equivalent in complexity
 Protein + Nucleic Acid + Supply by Environment

Genetic Code

- 2. Protein Protolife
  Protein —— Self Replication?
- Nucleic Acid Protolife
   RNA → Self Catalysis?

4. Something Else
Minerals
Clay Layers
Mineral - Molecule
Pyrite
Thioesters

**Genetic Takeover** 

 $? \longrightarrow \mathsf{RNA} \longrightarrow \mathsf{DNA}$ 

## Protein-Based Protolife

1. Proteinoid microspheres - Sidney Fox Amino Acids + Dry Heat  $\longrightarrow$  Proteinoids (Hot Tidepool?)  $\downarrow$  H<sub>2</sub>O (Tide)

Protocells Protolife? Can Add Proteinoid Split Bud Form Chains

(Look like life) Grow Divide "Reproduce" Bud J Like Bacteria

**Microspheres** 

But "Reproduction" not exact Later incorporate Nucleic Acids Proteinoid  $\longrightarrow$  Cells  $\longrightarrow$  Genes

<u>Problem:</u> How to incorporate Nucleic acids?

### Nucleic Acid Based Protolife

RNA ----> Genes ---> Protein -----> Cells Self-replicating RNA molecules Experiment by Sol Spiegelman RNA from Q<sub>6</sub> Virus - parasite on bacteria Injects RNA - Bacterium makes replicase Enzyme to Replicate RNA RNA multiplies, using activated nucleotides in Bacterium ——— new viruses

In Test Tube: Template RNA, Replicase, Activated Nucleotides (ATP, CTP, GTP, UTP)

 $\Rightarrow$  RNA copied <u>without</u> machinery of cell

Variation: <u>No</u> template RNA

Replicase made RNA from nucleotides

Manfred Eigen - further experiments with RNA in test tube:

Mutant RNA strands compete Degrade to smallest (~ 200 nucleotides) RNA that replicase could recognize (Monster - Selfish RNA)

RNA can do self-catalysis in some cases Could this have led to self replication?

### Eigen scenario

- A replicating RNA molecule forms by chance (random replicator - not a gene) ribozyme (catalyst, made of RNA)
- 2. Family of **similar** RNA's develops (quasispecies)
- 3. Connection to proteins(quasispecies specialize to make parts of protein)

- 4. Complex interactions (hypercycles)
- 5. Use lipids to make protocells
- 6. Competition leads to biological evolution



# Problems with Nucleic Acid First Scenario

- 1. Hard to get monomers
- 2. Unlikely to link correctly
- 3. Need existing proteins and lipids
- 4. Hypercycles subject to instabilities

N = size of molecular population



⇒ Only narrow range of sizes works

## The Origin of the Genetic Code

- We need more than either protein or RNA protolife
- Need interaction via genetic code
- Need translation
- Consider first a scenario by R. Shapiro

## Shapiro's Fable

The case for the "chicken"

Protein first  $\Rightarrow$  replication problem

"interpreters" aminoacyl tRNA synthetases

Match tRNA &

Amino acids

Could an earlier version have copied proteins directly?



tRNA



synthetases. The tRNA is recognized in the second step (through the features designated paracodon in this article) by the enzyme carrying a bound aminoacyl-AMP intermediate. Participation of the aminoacyl group in the recognition process is thus an attractive possibility.

- Early Evolution: Start with 4-6 amino acid types, gradually add more enzymes increase in size and catalytic power
- 2. First use of phosphate as energy? (ATP) or sugar-phosphate chains for construction (Teichoic acids in membranes of some bacteria) (partial  $Q_{\beta}$  replicase)
- Bases added for structure
   Support for protein synthesis → ribosome

- 4. Begin to copy RNA (Full  $Q_{\beta}$  replicase) Natural selection  $\longrightarrow$  better ribosome
- 5. Specialized, Short RNA aided attachment of amino acids to proteins; became tRNA
- 6. Then mRNA to align tRNA's now a separate genetic system that evolves
- 7. DNA developed from RNA

Shapiro dates last step to prokaryote -eukaryote split (different ways of storing DNA info)

Tests:

- 1. Synthesize in lab? Not possible yet.
- 2. Molecular archaeology vestigial ability of interpreters to recognize amino acids in proteins
- 3. Survivors of protein era? prions?

Support for the "chicken"

- 1988 discovery that interpreter does not use tRNA codon to recognize correct tRNA (in some cases) ~ 1/2
  - instead a single base pair at the other end of tRNA
  - ⇒ simpler, older code second genetic code
  - ⇒ connection of interpreter and tRNA more primitive than current code

2. Dyson modeling of molecular "populations"

Transition from disorder to order (non-life) (life)

Finds number of monomer types likely to be 9 - 11 (ok if used ~ 1/2 of modern proteins) But nucleotides (only 4) - not enough

Favors protein first



## The Egg Strikes Back

Other work shows some RNA can catalyze Non-RNA reactions

- 1. RNA in ribosome appears to be what catalyzes peptide bond formation Noller, et al. 1992, *Science*, **256**, 1416
- 2. RNA "ribozyme" catalyzes reactions between amino acids and tRNAs

→ First "interpreter" may have been RNA Piccirilli, et al. 1992, *Science*, **256**, 1420

## Origin of the Genetic Code

Crucial step in any theory

Early versions probably coded fewer amino acids - less specific

![](_page_24_Figure_0.jpeg)

#### Others added later

## **Evolution of Genetic Code**

Gaining specificity

If early tRNAs carries more than 1 kind of amino acid

![](_page_25_Figure_3.jpeg)

Evidence that code has evolved Freeland, et al. Tested 10<sup>6</sup> other codes

Only one better at minimizing bad effects of mutations

 $\Rightarrow$  Natural Selection

Still Evolving Some organisms have slightly different codes in mitochondria or in nucleus

### Other Ideas

- Neither the chicken nor the egg came first
- Transitional forms that were later discarded

## Or was it the "egkin"?

Some experiments with peptide nucleic acid (PNA).

PNA: <u>Peptide</u> backbone with bases

Can act as template for polymerization of RNA From activated nucleotides (Böhler, et al., *Nature*, **376**, 578 & comments by Piccirilli, pg. 548 17 Aug. 1995

PNA could be simpler to form under prebiotic conditions Main point is that a simpler thing (not necessarily PNA) could have preceded RNA

### Membranes

- Membranes provide enclosure
  - Also fundamental for metabolism
- Membranes never arise from scratch
  - Always passed down and added to
  - All derived from ancestral cell
- T. Cavalier-Smith proposes membranes
  - Plus nucleic acid formed "ob-cell"
  - Merger of 2 ob-cells formed first cell

## **Thioester World**

- 1. Need precursor to RNA world
- Need energy conversion Protometabolism Background:

![](_page_30_Figure_3.jpeg)

Hydroxyl + Carboxyl

C. deDuve In <u>Vital Dust</u>

![](_page_31_Figure_0.jpeg)

Thiol + Carboxyl

Thiols involved in metabolism, particularly in ancient pathways

Also can catalyze ester formation by group transfer Reactions e.g. peptide bonds

### **Catalytic Multimers**

C. deDuve

# "Multimer" short peptides and esters (NH<sub>2</sub>) (OH) of amino acids and hydroxy acids

### **Energy Sources**

Basic need is hydrogen atoms (or electrons in excited states)

In pure water	$\frac{H^+ + OH^2}{H_2O}$	more if acidic
e⁻ + H+ –	→ <b>H</b> *	excited H

Now chlorophyll + sunlight

![](_page_34_Figure_0.jpeg)

## Transition to Phosphate

Energy currency in life now is ATP Adenosine Triphosphate used to make bonds, remove H<sub>2</sub>O

Earlier, inorganic phosphate p–p diphosphate or polyphosphate still involved in ATP reactions

![](_page_35_Figure_3.jpeg)

age—will be followed by GUC. This AGUC sequence is complementary to the terminal sequence written in antiparallel fashion, and will cause the chain to double up as follows:

![](_page_36_Figure_2.jpeg)

Assume now that this folded chain is subject to elongation, by the addition of new nucleotides, from right to left, to the U end. The presence of G next to the A paired with the terminal U is likely to favor the addition of a complementary C over that of the other three possible nucleotides. Repeat the process and you get U added opposite A, G opposite C, G again opposite the next C, and so on. What you get is the formation of a stretch complementary over all its length to the other end of the molecule:

![](_page_36_Figure_4.jpeg)

### Summary of Proto-Life Development

<u>Stage</u>	<b>Proteins</b>	<u>Halfway # 1</u> Peptide Nucleic Acids	<u>Halfway # 2</u> RNA Ribozyme	<u>Nucleic Acids</u>	
Monomers	Amino Acids	Bases Amino Acids	Ribose Sugars Bases Phosphates Amino Acids	Ribose Sugars Bases Phosphate	
Polymerization	Proteinoids	Short strands of PNA's	Short strands of RNA + amino acids	Short strands of RNA	
Replication	?	Affinity for complementary bases + ease of peptide bonding	Affinity for complementary bases	Affinity for complementary bases	
Pre-life	Proteinoids + RNA?	Separation of proteins and nucleic acids	Separation of nucleic acids and protein parts	RNA adapts proteinoids as needed	
Life	Proteins	Disappears	Disappears	DNA and RNA	