

**212 SM L03a**

# Core Concept

**01:** Evolution, Thermodynamics, Habitat diversity, Ecology, Physiology their integration define Microbiology

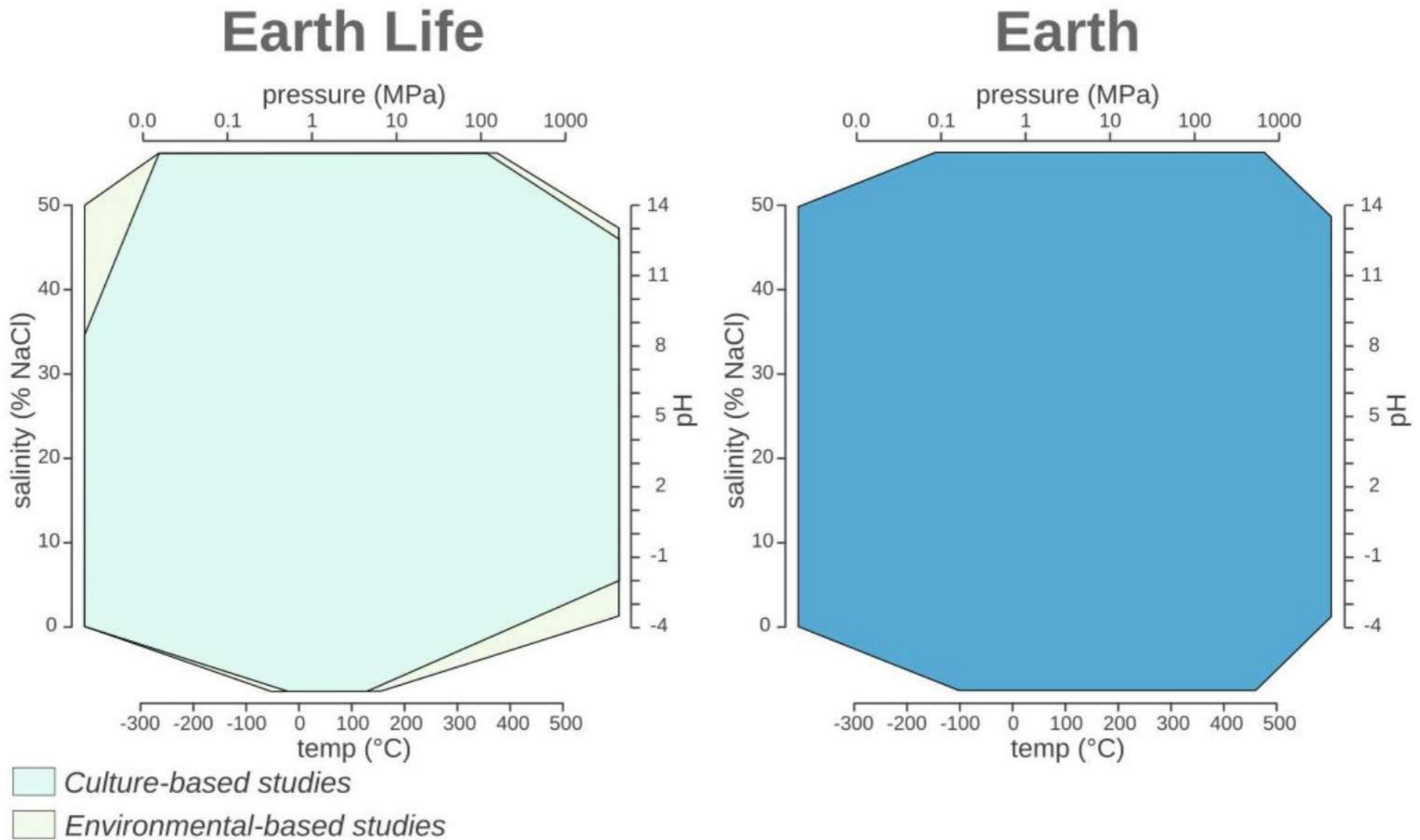
**02:** Unique goal of microbial life: survival, maintenance, generation of ATP, growth of new cells

**03:** Planet's habitat diversity results in genetic, molecular, metabolic and physiological microbial diversity

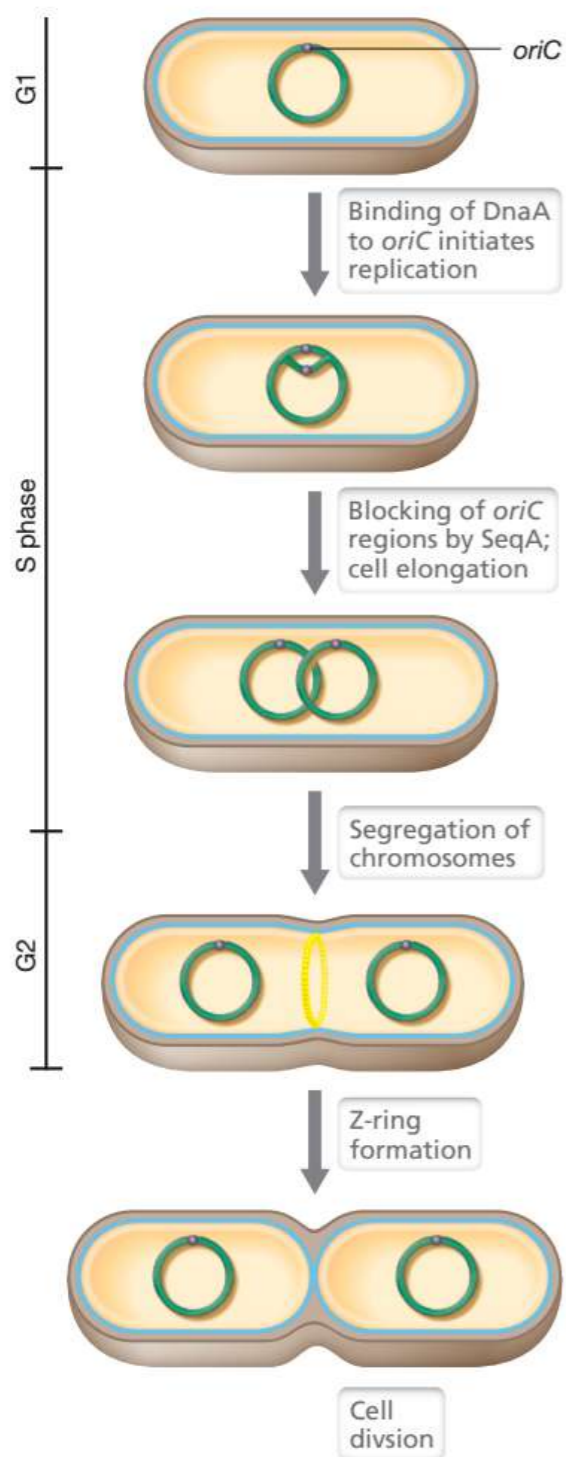
**How do microbes growth?  
How do microbes structure the  
environment?**

# Microbial Life on Earth

Merino et al., 2019

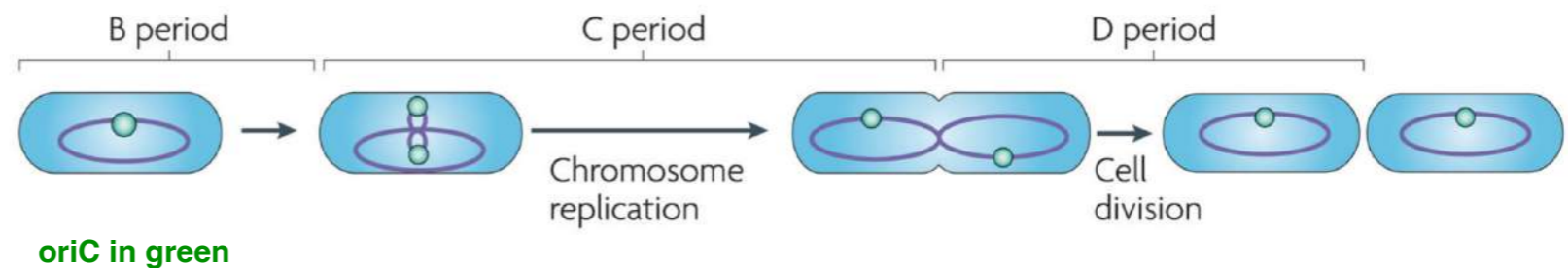


# Cellular growth



Growth is intimately connected with nutrient availability and energy status of the cell

Spatial and temporal coordination between the DNA replication and cell elongation, DNA segregation into new cells

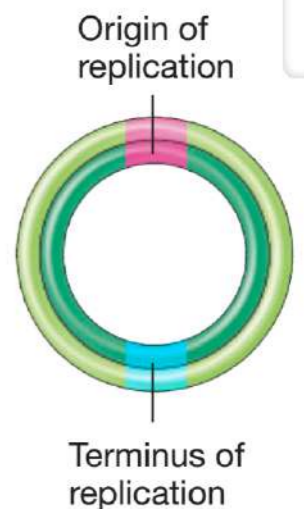


B period: time between division (birth) and the initiation of chromosome replication

C period: time window for chromosome replication

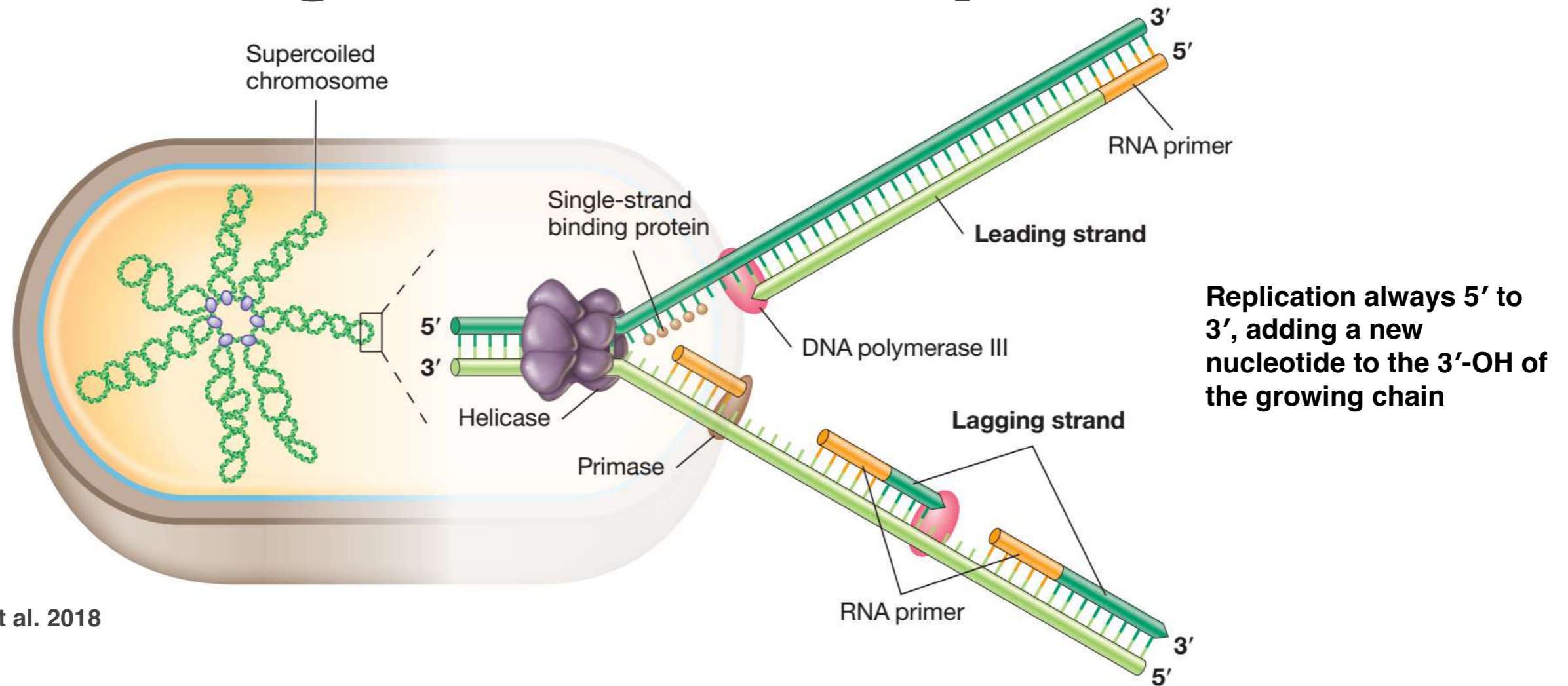
D period: time between the completion of chromosome replication and cell division

**Protein Tus binds to terminus site and stop replication**



Wang & Levin, 2009

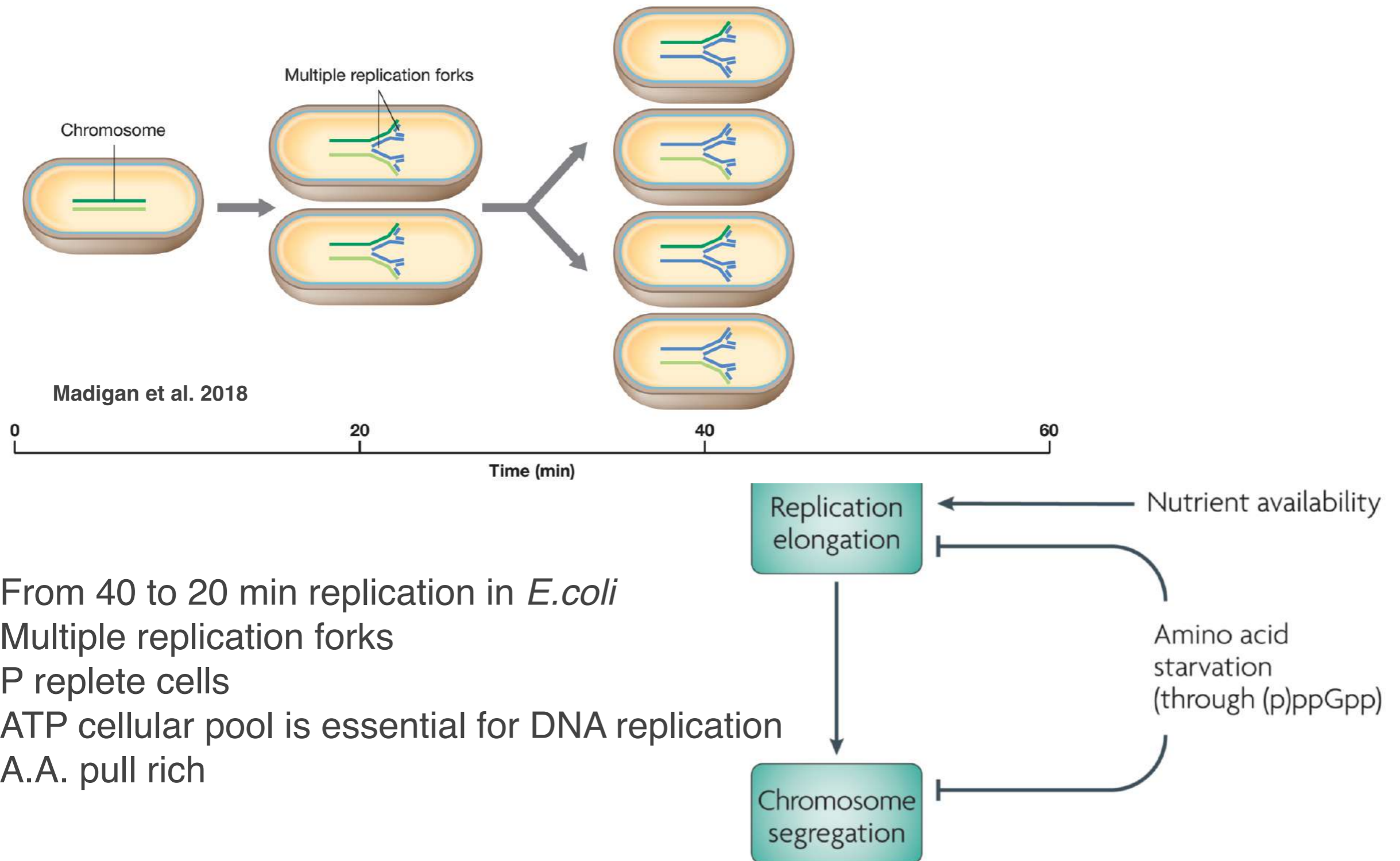
# Zooming into DNA replication



Madigan et al. 2018

1. DNA synthesis begins at a single site on chromosome, origin of replication (oriC), where DnaA binds and opens up double helix
2. Stabilization of strands by helicase (DnaB), and its helper loader protein (DnaC)
3. Two helicases are loaded, one onto each strand, facing in opposite directions
4. Two primase and two DNA polymerase III enzymes are loaded onto the DNA behind helicases and initiation of DNA replication begins
5. As replication proceeds, replication fork appears to move along the DNA

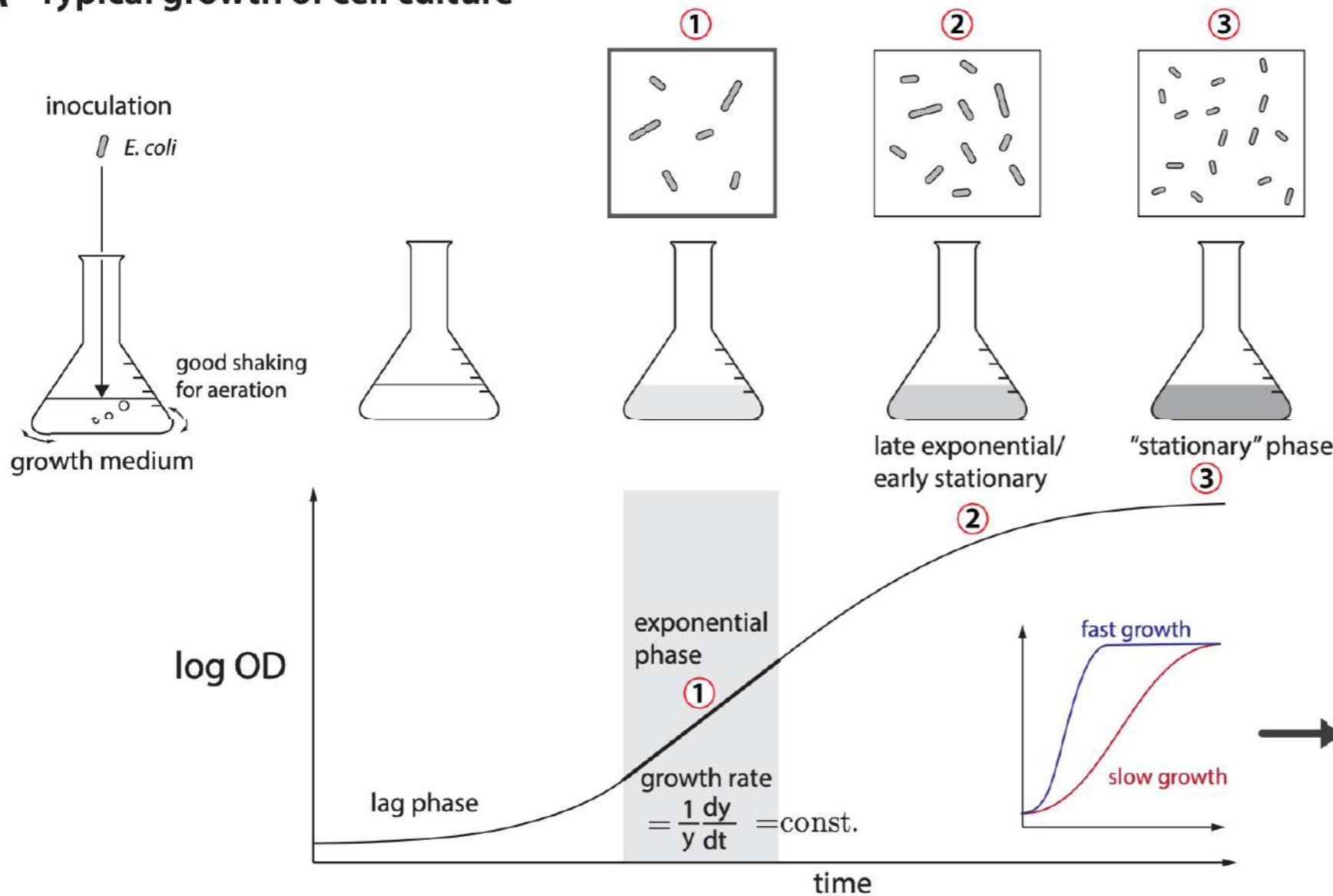
# Nutrient & energy status controls growth rate



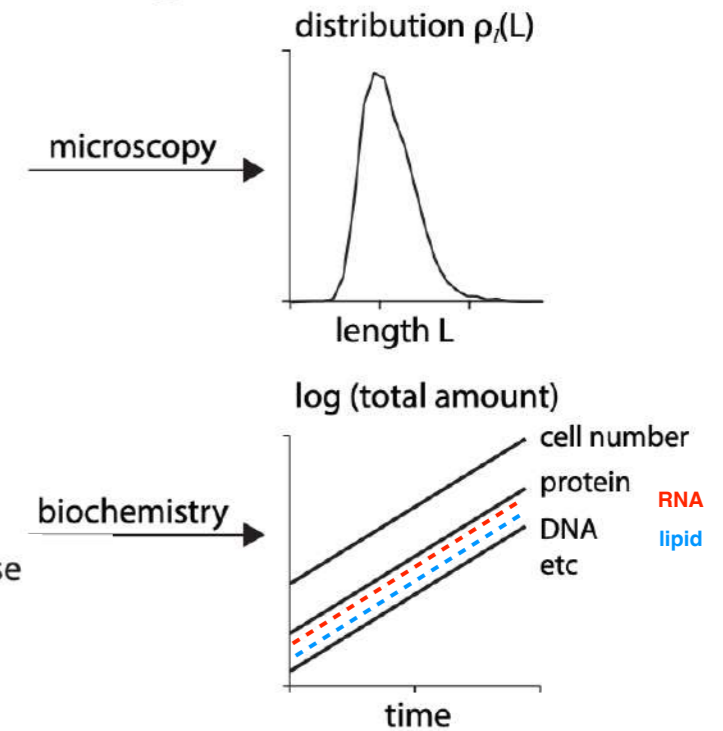


# Growth in an homogeneous-predictable-low diversity environment, I

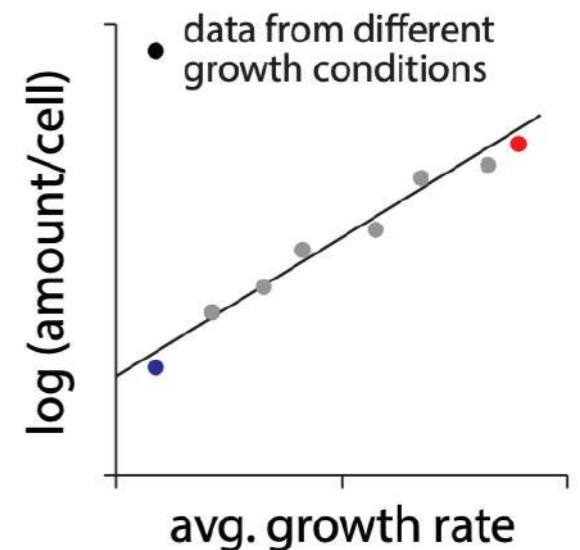
## A Typical growth of cell culture



## B What is measurable



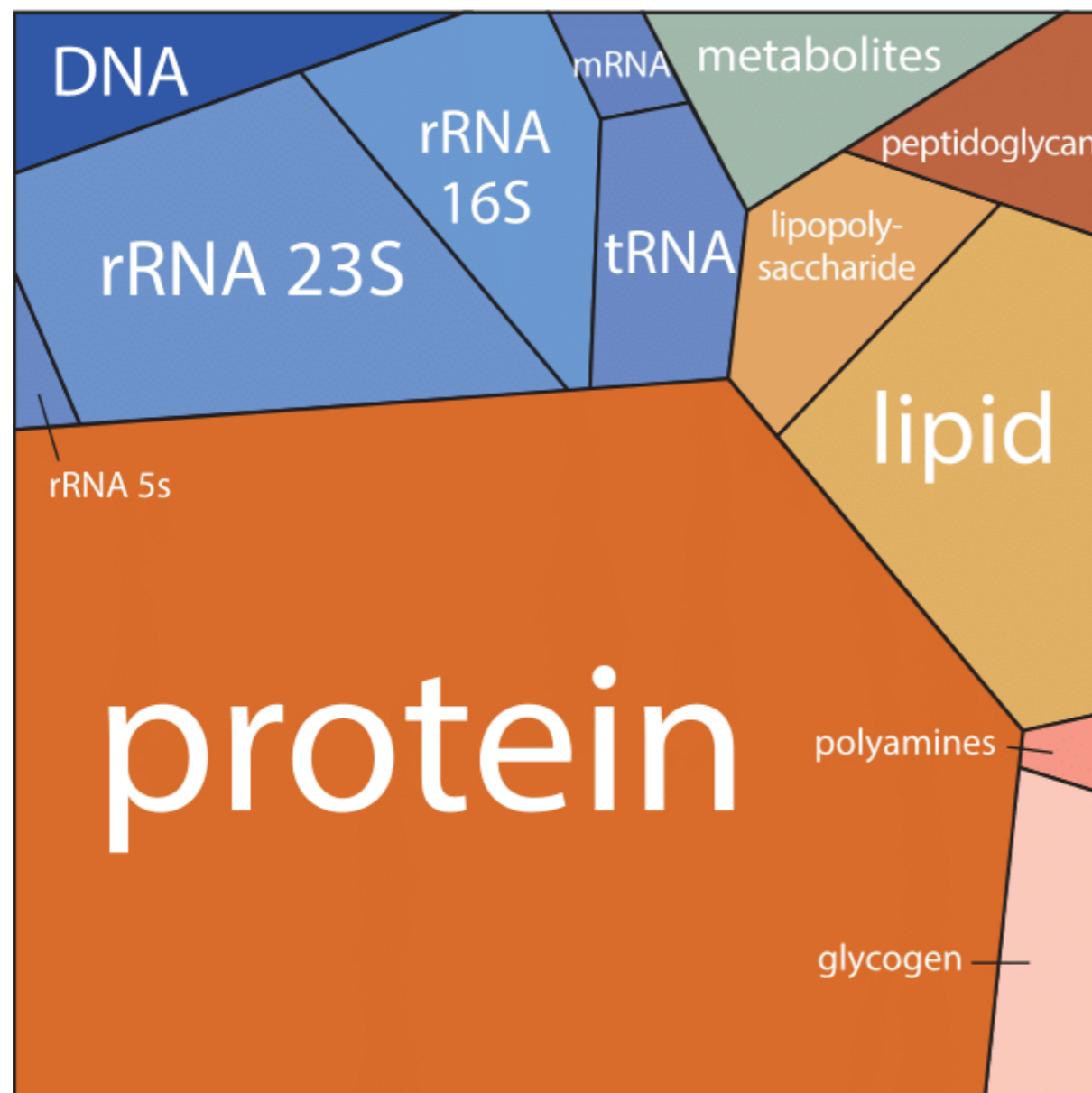
## C "(nutrient) growth law"





# Growth in an homogeneous-predictable-low diversity environment, II

A Voronoi tree diagram of *E.coli* composition



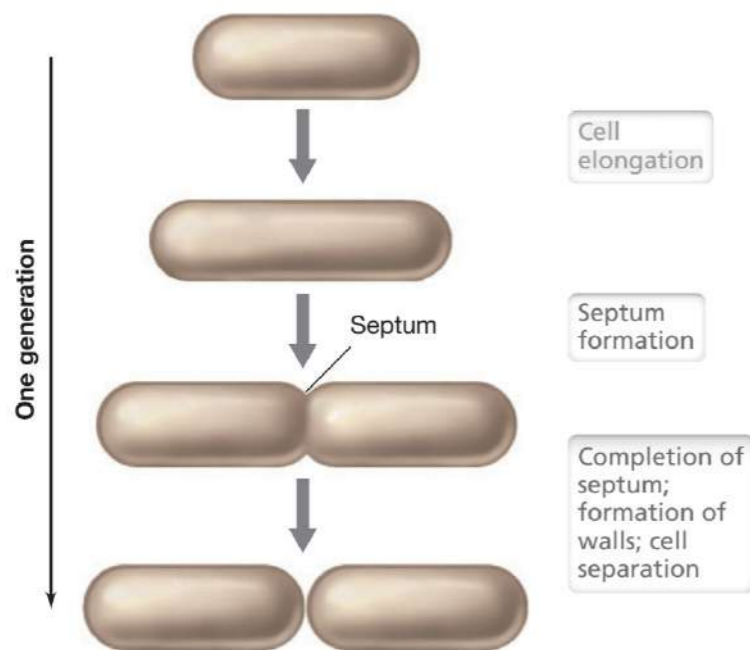
40 min

- Each polygon area is the relative fraction of the corresponding cellular constituent (dry mass)
- Similar colors = related functional role
- Steady-state mean cell size (large circles) scales exponentially with nutrient-determined growth rate

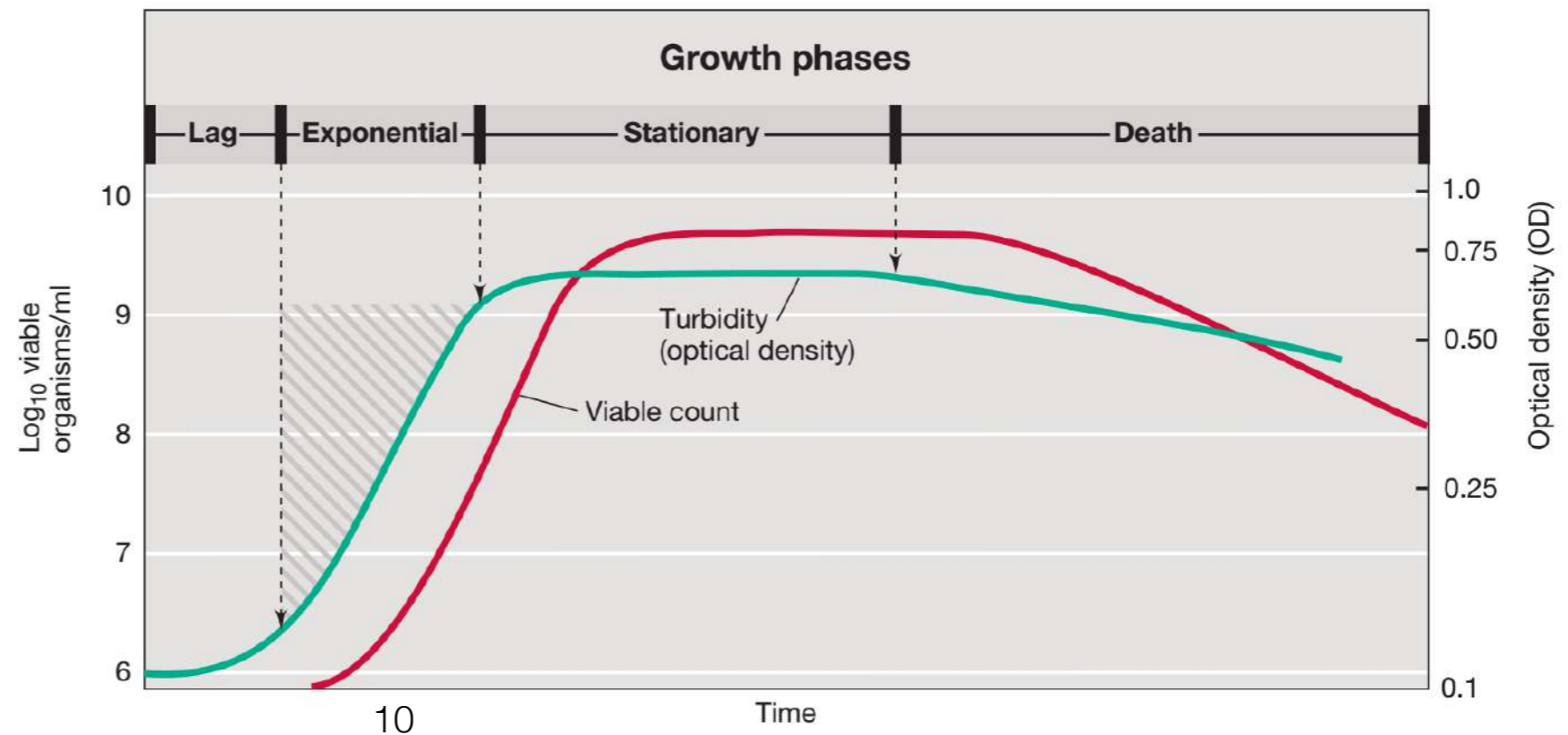
**Defined growth conditions**  
**What is missing?**

# Growth

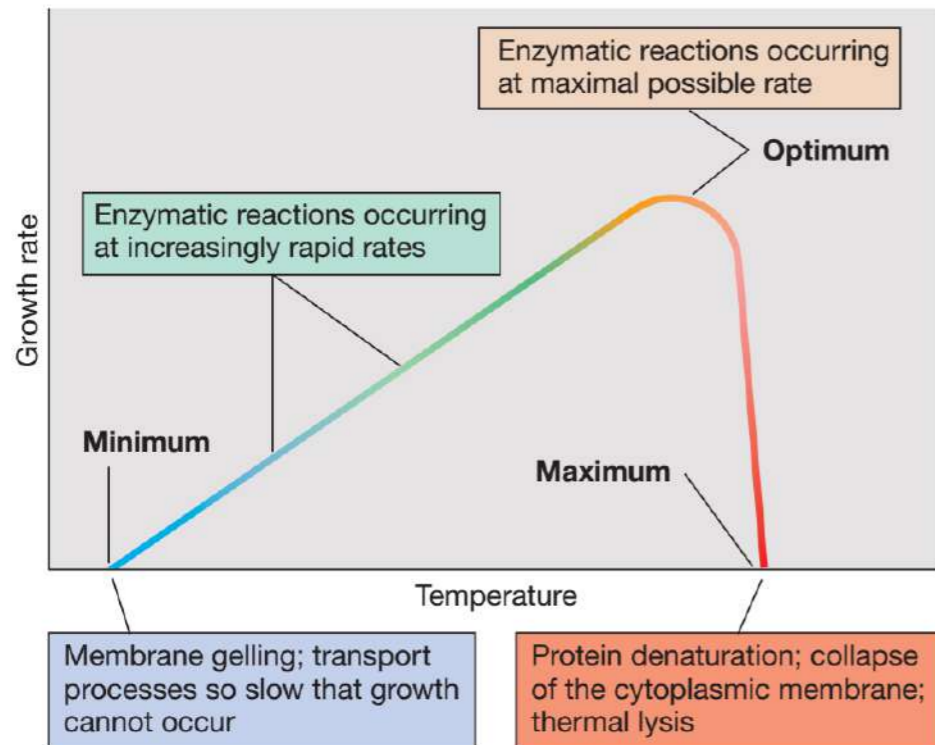
- A fixed relationship exists between initial cell number in a culture and cell number present after a period of exponential growth:  $N = N_0 2^n$  where  $N$  is final cell #,  $N_0$  is initial cell #, and  $n$  # of generations during period of exponential growth
- **Generation time (g)** of the exponentially growing population is  $t/n$ , where  $t$  is the duration of exponential growth in days, hours, or minutes (**g is the time from 1 cell to 2 cells**)
- Equation  $N = N_0 2^n$  can be expressed in terms of  $n$  by taking the logarithms of both sides:  $n = [3.3(\log N - \log N_0)]$
- **Instantaneous growth rate constant** expresses the rate at which the population is growing at any instant (by contrast,  $g$  is the mean time required for the cell population to double);  $k$  is expressed in units of reciprocal hours ( $h^{-1}$ ):  $k = 0.693/g$



Madigan et al. 2018

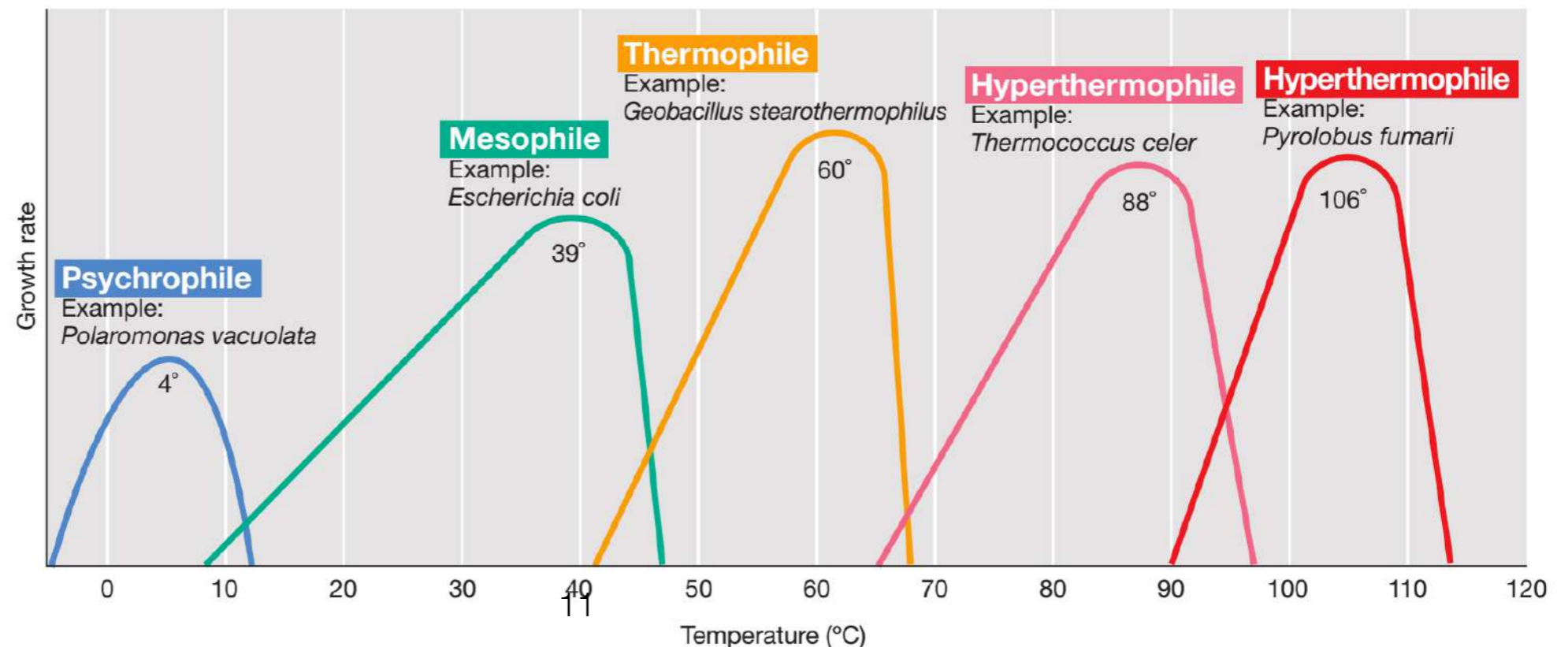


# Temperature affects growth



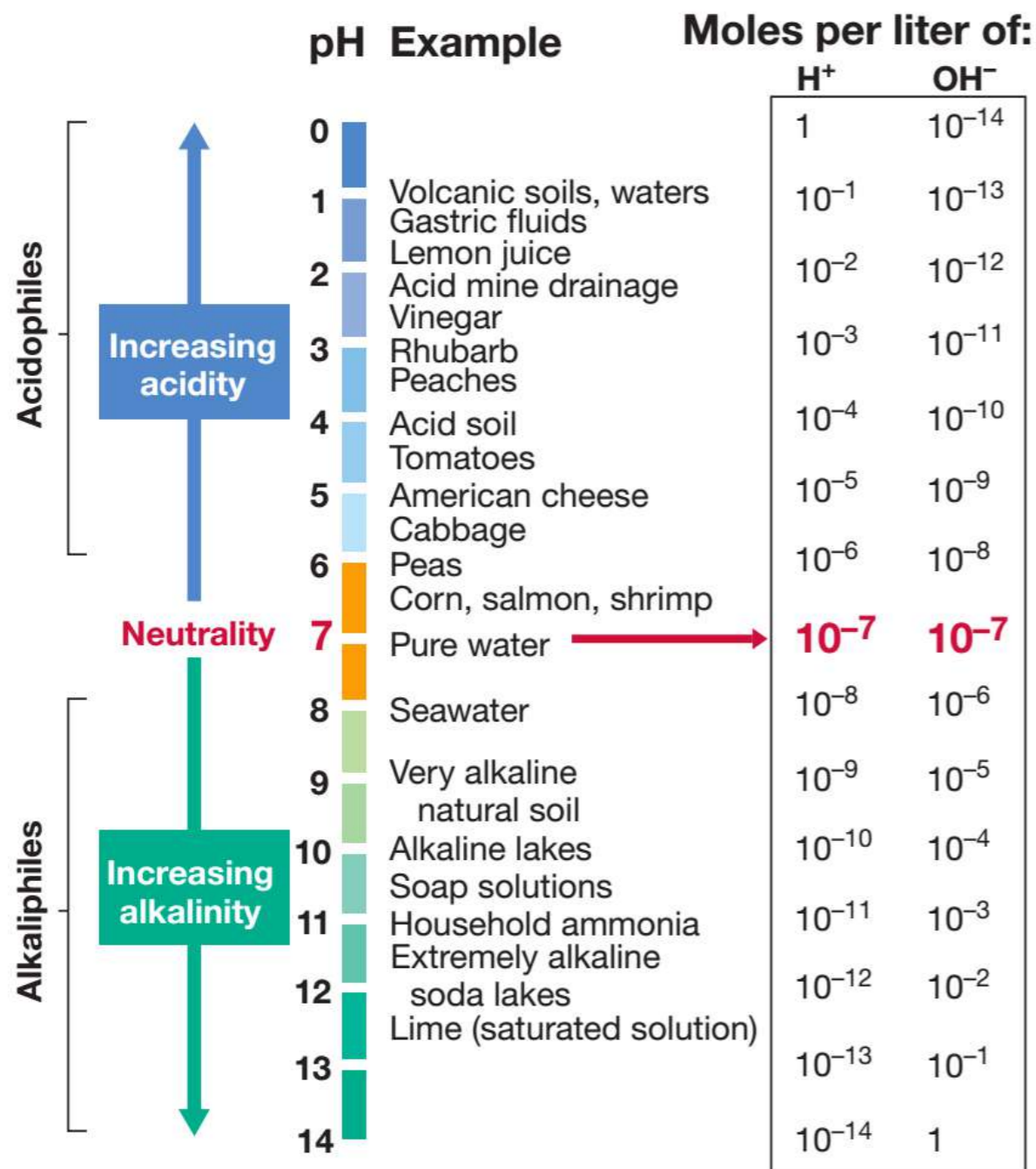
- **Reactions** occur **faster** at warmer temperatures b/c rate of collisions between molecules increases
- If temperature gets too hot the enzyme may denature and fail to function
- **Arrhenius activation energy**, which describes temperature effect on **catalytic rate constant**,  $k_{cat}$ , and thermal stability, which describes temperature effect on **thermal inactivation rate constant**,  $k_{inact}$

Peterson et al., 2007



Madigan et al. 2018

# pH affects growth



Optimal pH for growth refers to **extracellular environment only**

**Intracellular pH** must be maintained in 5 - 9 range

Extreme acidophiles and alkaliphiles maintain cytoplasmic pH values ~neutrality

**Extreme pH** affects macromolecule structures

H-bonds holding together strands of DNA break up at high pH

Lipids are hydrolyzed by an extremely basic pH

PMF responsible for production of ATP in cellular respiration depends on concentration gradient of H<sup>+</sup> across membrane



# pH adaptive strategies

Active (**proton exclusion, exchange, pumping, consumption and neutralization**) and passive (**cytoplasmic buffering**) mechanisms of pH homeostasis as well as damage mitigation strategies (**DNA repair, synthesis of acid stable proteins**)

Quatrini & Johnson, 2018

- *Lactobacillus plantarum* is an anaerobic bacterium that **produces lactic acid** as metabolic product and thus lowers pH but also prefers low pH values
- *Corynebacterium ammoniagenes* produces **urease that cleaves urea into ammonia** and thus increases pH at the same time, it prefers higher pH values
- *Pseudomonas veronii* also **increases medium pH** but prefers low pH values for growth
- *Serratia marcescens* **strongly lowers pH** but better tolerates comparably higher pH values, with a slight optimum at around pH 8

*In summary, we find that microbial growth often leads to dramatic changes in the pH of the environment, and this pH change can promote or inhibit bacterial growth*

Ratzke & Gore, 2018

# Osmotic pressure affects growth

- **Water availability** is expressed in terms of **water activity** ( $a_w$ ): vapor pressure of air in equilibrium with a substance or solution / the vapor pressure of pure water
- Values of  $a_w$  vary between 0 (no free water) and 1 (pure water)
- **H<sub>2</sub>O diffuses** from regions of high water concentration (**low solute concentration**) → regions of lower water concentration (**higher solute concentration**) in the process of **osmosis**
- Cytoplasm has a higher solute concentration than the environment, so H<sub>2</sub>O → into cell
- Under such conditions, cell is said to be in **positive water balance**, normal cell state

**TABLE 5.4 Water activity of several substances**

Water activity ( $a_w$ )	Material	Example organisms <sup>a</sup>
1.000	Pure water	<i>Caulobacter</i> , <i>Spirillum</i>
0.995	Human blood	<i>Streptococcus</i> , <i>Escherichia</i>
0.980	Seawater	<i>Pseudomonas</i> , <i>Vibrio</i>
0.950	Bread	Most gram-positive rods
0.900	Maple syrup, ham	Gram-positive cocci such as <i>Staphylococcus</i>
0.850	Salami	<i>Saccharomyces rouxii</i> (yeast)
0.800	Fruit cake, jams	<i>Zygosaccharomyces bailii</i> (yeast), <i>Penicillium</i> (fungus)
0.750	Salt lakes, salted fish	<i>Halobacterium</i> , <i>Halococcus</i>
0.700	Cereals, candy, dried fruit	<i>Xeromyces bisporus</i> and other xerophilic fungi

Madigan et al. 2018



# Osmotic pressure strategies

Madigan et al. 2018

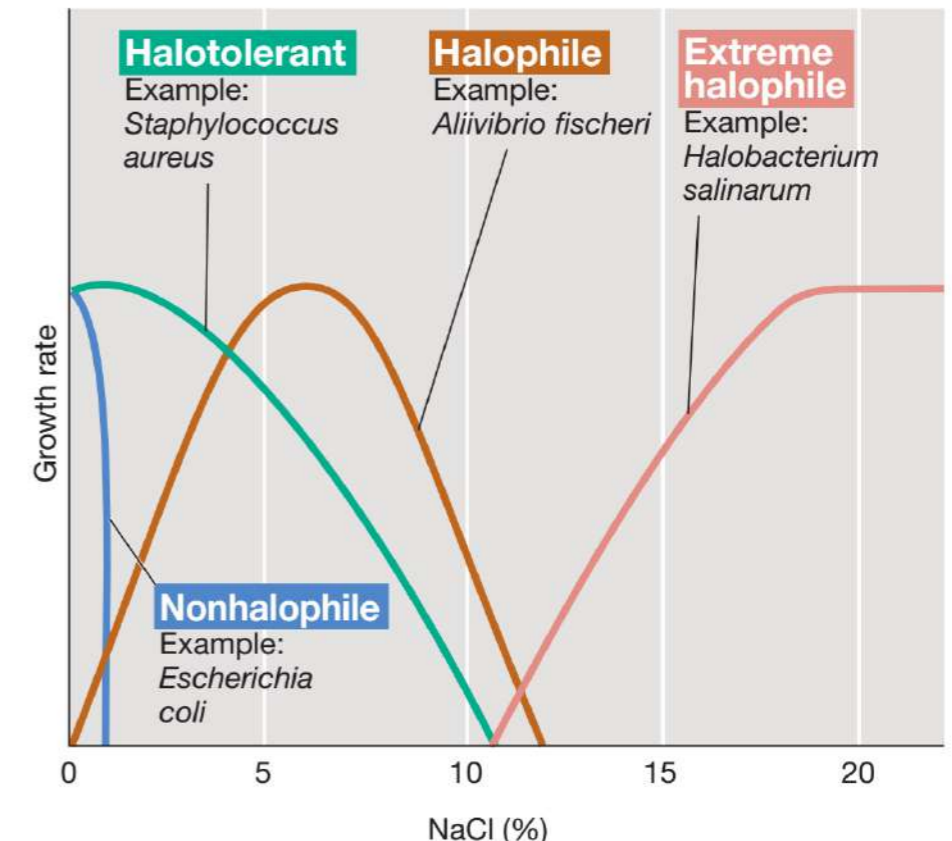
From high aw → low aw: cells maintains positive water balance **by increasing its internal solute (compatible not interference with metabolism) concentration:**

1. Pumping solutes into the cell from the environment
2. Synthesizing cytoplasmic solutes

**Compatible solutes are highly water-soluble organic molecules & electrolytes:** sugars, alcohols, and amino acid derivatives

Glycine betaine, an analog of the amino acid glycine, is widely distributed among halophilic bacteria

Other common compatible solutes include sugars such as sucrose and trehalose, dimethylsulfoniopropionate (produced by marine algae)



**H<sub>2</sub>O OUT** of cells as their medium becomes more concentrated (an osmotic upshift)

**H<sub>2</sub>O INTO** cells as their medium becomes more dilute (an osmotic downshift)

Bacterial membranes have high water permeabilities, so **cellular hydration is altered within seconds** of an osmotic shift

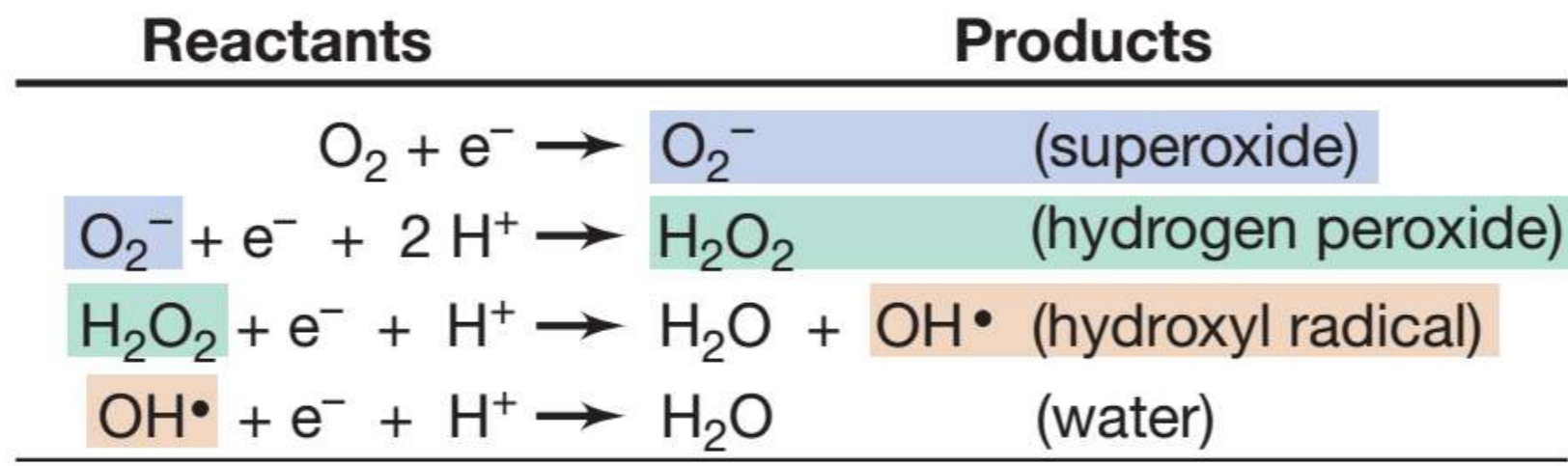
Approximately **0.5 g of water** is bound **per gram of cytoplasmic macromolecules**

At low osmolality, phosphate, the predominant inorganic anion, is present at a concentration of approximately 10 mM. Most metabolites are maintained at comparable or lower concentrations

At high osmolality, the amount of K<sup>+</sup> exceeds that of nucleic acid phosphate and glutamate accumulates as K<sup>+</sup> counterion (the concentration of glutamate rising from approximately 0.05 to 0.50 M). The trehalose concentration rises from approximately 0.04 to 0.4 M

# O<sub>2</sub> affects growth

- Molecular oxygen (O<sub>2</sub>) is not toxic
- O<sub>2</sub> can be converted to toxic oxygen by-products:
  - A. superoxide anion (O<sub>2</sub><sup>-</sup>)
  - B. hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) → damage cell components
  - C. hydroxyl radical (OH<sup>•</sup>) → oxidation macromolecules & other organic compounds
- All by-products of the reduction of O<sub>2</sub> to H<sub>2</sub>O in respiration
- Flavoproteins, quinones, and iron–sulfur proteins, electron carriers found all cells also catalyze some of these reductions



## Outcome:



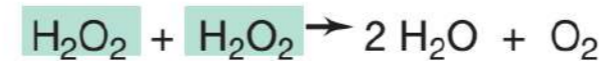
# O<sub>2</sub> adaptive strategies

**TABLE 5.6** Oxygen relationships of microorganisms

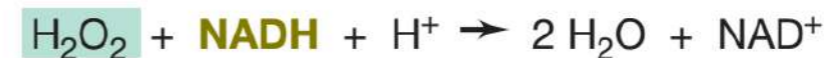
Group	Relationship to O <sub>2</sub>	Type of metabolism
<b>Aerobes</b>		
Obligate	Required	Aerobic respiration
Facultative	Not required, but growth better with O <sub>2</sub>	Aerobic respiration, anaerobic respiration, fermentation
Microaerophilic	Required but at levels lower than atmospheric	Aerobic respiration
<b>Anaerobes</b>		
Aerotolerant	Not required, and growth no better when O <sub>2</sub> present	Fermentation
Obligate	Harmful or lethal	Fermentation or anaerobic respiration

Specific niche

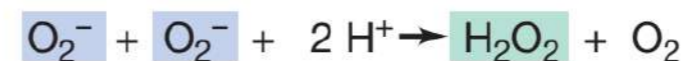
Metabolic machinery to detoxify



(a) **Catalase**



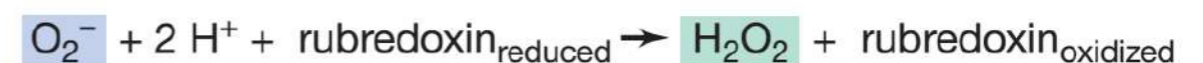
(b) **Peroxidase**



(c) **Superoxide dismutase**



(d) **Superoxide dismutase/catalase in combination**



(e) **Superoxide reductase**



# Elemental composition of Earth and microbes

Aston, 1924

- Universe, Earth, Life share important elements
- Majority of Universe is H and He and some others
- The essence of Life is the other elements

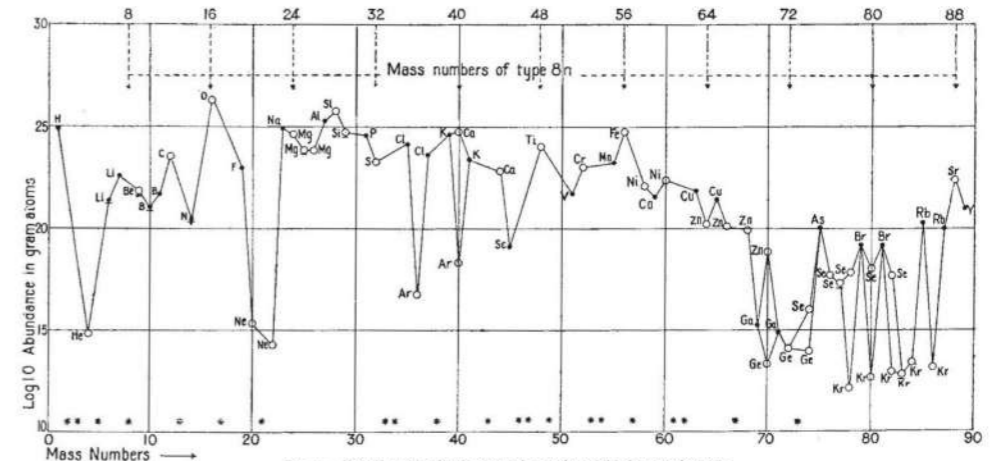


FIG. 1.—Relative abundance of atomic species of the first 39 elements.  
 ○ Even atomic number 41 } Total, 69 species.  
 ● Odd atomic number 28 }  
 \* Missing or doubtful mass numbers.

## The Elemental Composition of *E. coli*

Element	% dry	Substrate Source	Cellular Components
C	55	DOC, CO <sub>2</sub>	Main constituent of cellular material
O	20	O <sub>2</sub> , DOM, CO <sub>2</sub>	Constituent of cell material and cell water; O <sub>2</sub> primary electron acceptor in aerobic respiration
N	10	NH <sub>3</sub> , NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , DON, N <sub>2</sub>	Constituent of amino acids, nucleic acids, nucleotides, and coenzymes
H	8	DOM, H <sub>2</sub>	Main constituent of organic compounds and cell water
P	3	PO <sub>4</sub> <sup>3-</sup> , DOP	Constituent of nucleic acids, nucleotides, phospholipids, LPS, teichoic acids in gram positives
S	1	SO <sub>4</sub> , H <sub>2</sub> S, HS, DOM	Constituent of cysteine, methionine, glutathione, several coenzymes
K	1	Potassium salts	Main cellular inorganic cation and cofactor for certain enzymes
Mg	0.5	Magnesium salts	Inorganic cellular cation, cofactor for certain enzymatic reactions
Ca	0.5	Calcium salts	Inorganic cellular cation, cofactor for certain enzymes
Fe	0.002	Iron salts, DOM	Component of cytochromes and Fe-proteins; cofactor for many enzymes

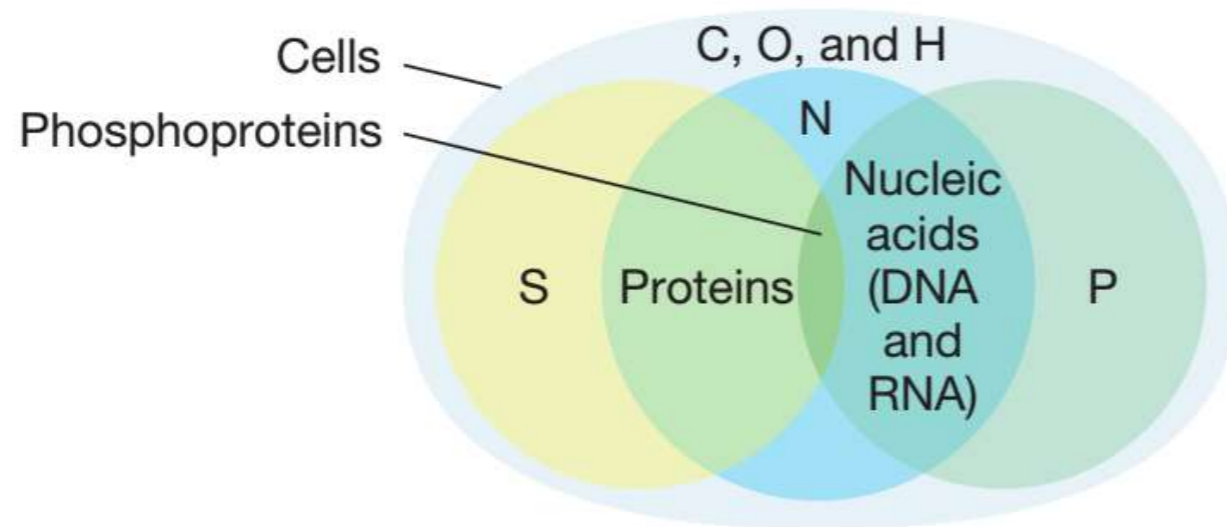
Group →

Period ↓	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
1	H																	He
2	Li	Be											B	C	N	O	F	Ne
3	Na	Mg											Al	Si	P	S	Cl	Ar
4	K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
5	Rb	Sr	Y	Zr	Nb	Mo	Tc	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	I	Xe
6	Cs	Ba	Lu	Hf	Ta	W	Re	Os	Ir	Pt	Au	Hg	Tl	Pb	Bi	Po	At	Rn
7			La	Ce	Pr	Nd												

- Essential for all microorganisms
- Essential cations/anions for most microorganisms
- Trace metals (Table 3.1), some essential
- Used for special functions
- Unessential, but metabolized
- Unessential, not metabolized

# From macromolecules to cell

**Elemental composition of informational macromolecules**



(b)

**Macromolecular composition of a cell**

Macromolecule	Percent of dry weight
Protein	55
Lipid	9.1
Polysaccharide	5.0
Lipopolysaccharide	3.4
DNA	3.1
RNA	20.5

(c)

- About 75% of microbial cell wet weight (a single cell of *Escherichia coli* weighs just  $10^{-12}$  g) is water
- The remainder ~ 25% is primarily macromolecules—proteins, nucleic acids, lipids, and polysaccharides
- The building blocks of these macromolecules are the amino acids, nucleotides, fatty acids, and sugars, respectively



# Basic Cellular Metabolism

1. Energy source to generate ATP
2. Carbon source of assembling cellular building blocks
3. For maintenance of existing cells/ for growth of new cells

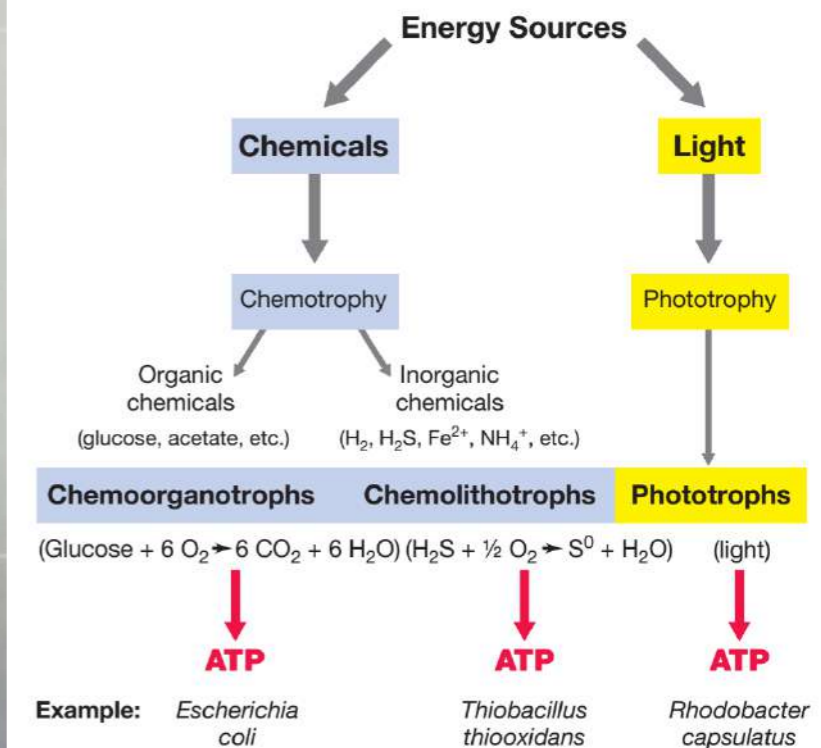
- > Successful exploitation well-defined energy and carbon source
- > Physical, biological and chemical properties vary in space and time

Carbon source	Energy source		
	Chemical, organic	Chemical, inorganic	Light
Fixed organic	<b>Chemosynthetic organoheterotroph</b> (Example: humans, fungi, <i>Pseudomonas</i> )	<b>Chemosynthetic lithoheterotroph</b> (Example: <i>Beggiatoa</i> sp.)	<b>Photosynthetic heterotroph</b> (Example: purple and green bacteria; <i>Rhodospirillum</i> )
Gaseous CO <sub>2</sub>		<b>Chemosynthetic lithoautotroph</b> (Example: ammonia-, hydrogen-, and sulfur-oxidizing bacteria; <i>Nitrosomonas</i> , <i>Aquifex</i> )	<b>Photosynthetic autotroph</b> (Example: plants, algae, <i>Prochlorococcus</i> )

**Terminology:**

- Autotroph: carbon from CO<sub>2</sub> fixation
- Heterotroph: carbon assimilated from (fixed) organic compounds
- Photosynthetic: energy from light
- Chemosynthetic: energy from oxidizing reduced chemicals
- Chemolitho: energy from oxidizing inorganic reduced chemicals
- Chemoorgano: energy from oxidizing organic reduced chemicals.

Madsen, 2016





# Metabolism & Growth in a limited environment

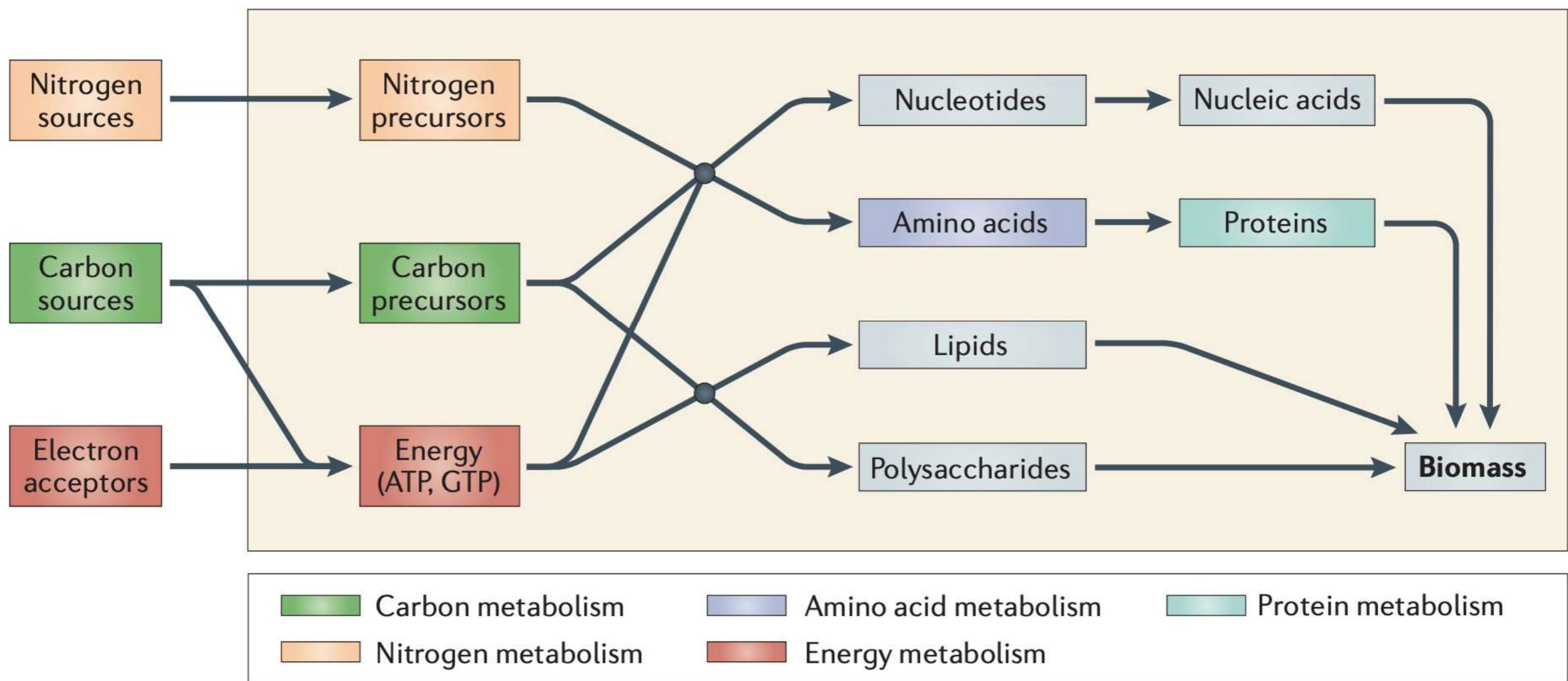
**Habitat characteristics and nutrient limitations faced by three physiological classes of microorganisms**

Habitat type	Photoautotroph	Chemolithotroph	Chemoorganoheterotroph
Ocean water	Daily light cycle, light penetration depth; scarce iron	Flux of reduced inorganic compounds, especially $\text{NH}_3$ , $\text{H}_2\text{S}$ , $\text{H}_2$ , or $\text{CH}_4$ from nutrient turnover and hydrothermal vents	Carbon flux from phototrophs, dead biomass, and influent waters
Lake water	Daily light cycle, light penetration depth; scarce phosphorus	Flux of reduced inorganic materials, especially $\text{NH}_3$ , $\text{H}_2$ , and $\text{CH}_4$ from nutrient turnover	Carbon flux from phototrophs, dead biomass and influent waters
Sediment (freshwater and oceanic)	Daily light cycle, light penetration depth	Flux of reduced inorganic materials, especially $\text{NH}_3$ and $\text{H}_2$ from nutrient turnover or $\text{H}_2$ , $\text{H}_2\text{S}$ , or $\text{CH}_4$ from hydrothermal vents	Flux of organic carbon from phototrophs and dead biomass; flux of final electron acceptors to carbon-rich anaerobic strata
Soil	Daily light cycle, light penetration depth	Flux of reduced gaseous substrates, especially methane from nutrient turnover by anaerobes	Slow turnover of soil humus, dead biomass, plant root exudates; leaf fall from vegetation
Subsurface sediment	No light	Flux of reduced inorganic materials, especially $\text{H}_2$ and $\text{CH}_4$ from geothermal origin	Carbon flux from nutrient turnover

# Coordinated Metabolism

Beyond fuelling cellular activities with building blocks and energy, metabolism also integrates environmental conditions into intracellular signals

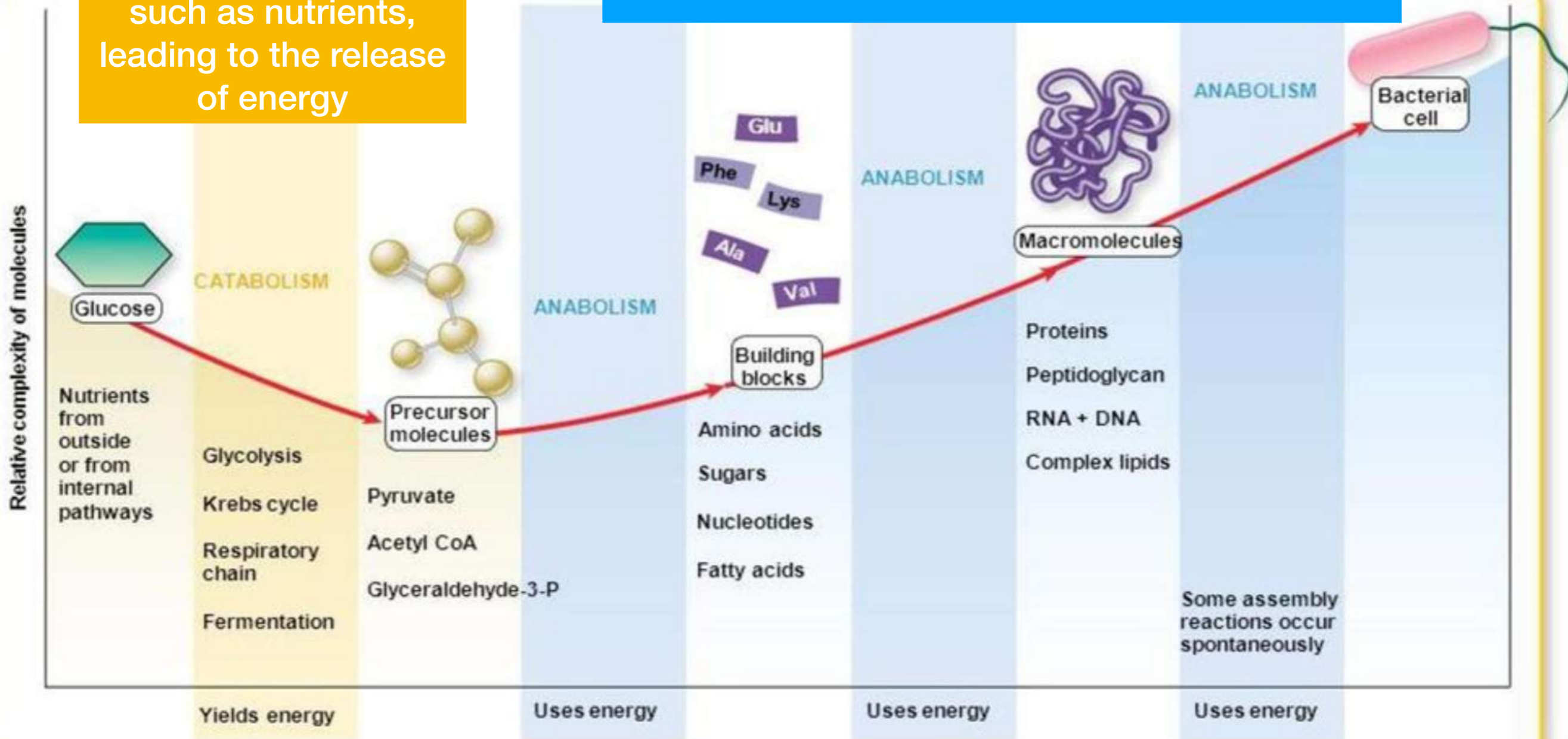
Metabolisms underlying regulatory network is complex and multifaceted



# Metabolism

**Catabolism:** The degradation of complex molecules, such as nutrients, leading to the release of energy

**Anabolism:** The energy-dependent formation of building blocks and macromolecules in a cell





# Real Growth in a limited environment

Rapid exponential growth is not real in the environment

In 48 h assuming that *E.coli* that doubles every 20 min, given its weight  $10^{-12}$  g, there will be  $2.2 \times 10^{31}$  g

In the environment:

- Sporadic rapid growth
- Slow growth
- Sporadic slow growth
- Dormancy

Estimates of microbial growth rate, dormancy, and survival time

Habitat	Organism	Doubling time (DT) or survival time (ST)	References
<b>Growth rate</b>			
Laboratory medium	<i>E. coli</i>	20 min DT	Koch, 1971
Human intestine	<i>E. coli</i>	12 h DT	Koch, 1971
Mouse	<i>Salmonella typhimurium</i>	10–24 h DT	Brock, 1971
Rumen	Heterotrophic bacteria	~12 h DT	Brock, 1971
Pond	Heterotrophic bacteria	2–10 h DT	Brock, 1971
Lake water	Heterotrophic bacteria	10–280 h DT	Jannasch, 1969
Ocean	Heterotrophic bacteria	20–200 h DT	Vaulot et al., 1995
Ocean	Autotroph, <i>Prochlorococcus</i>	~24 h DT	Gray and Williams, 1971
Soil	Heterotrophs: $\alpha$ Proteobacteria, rhizobia	100 days DT	Mailloux and Fuller, 2003
Shallow groundwater	Heterotrophs: <i>Acidovorax</i> , <i>Commamonas</i>	15 days DT	Hoehler and Jorgensen, 2013
Marine surface sediments	Sulfate reducers	1 year DT	Holmes et al., 2013
Shallow subsurface	<i>Geobacter</i>	46 h DT	Phelps et al., 1994;
Deep subsurface	Heterotrophs	100 years DT	Fredrickson and Onstott, 2001
Deep marine sediments	Sulfate reducers, heterotrophs	200–3000 year DT	Hoehler and Jorgensen, 2013

# Principle of Bioenergetics

Energy flows (radiation → chemical → heat)

Matter cycles (uptake → growth → death → recycle)

Energy is defined as the **ability to do work**, kilojoules (kJ), a unit of heat energy

All chemical reactions in a cell are accompanied by **changes in energy**, energy being either required or released as a reaction proceeds

$\Delta G^{0'} < 0$ , reaction will proceed with the release of free energy-  
**exergonic**

$\Delta G^{0'} > 0$ , reaction requires energy in order to proceed- **endergonic**

A  $\Delta G^{0'} > 0$  reaction under standard conditions can become exergonic under the actual conditions present in the microbial habitat

**Thermodynamics: the branch of chemistry that predicts what can happen based on the energy state of reactants and products**

In 1877, the American mathematician, Josiah Gibbs, sought to quantify the amount of "useful work" that can be harnessed during completion of chemical (or other) reactions.

$A + B \rightleftharpoons C + D$

**Types of reactions**

- Changes in state
- Dissolution/precipitation
- Complexation
- Acid/base
- Sorption/desorption
- Oxidation/reduction (key for biological systems to generate ATP)

Products – reactants =  $-\Delta G$  (free energy)

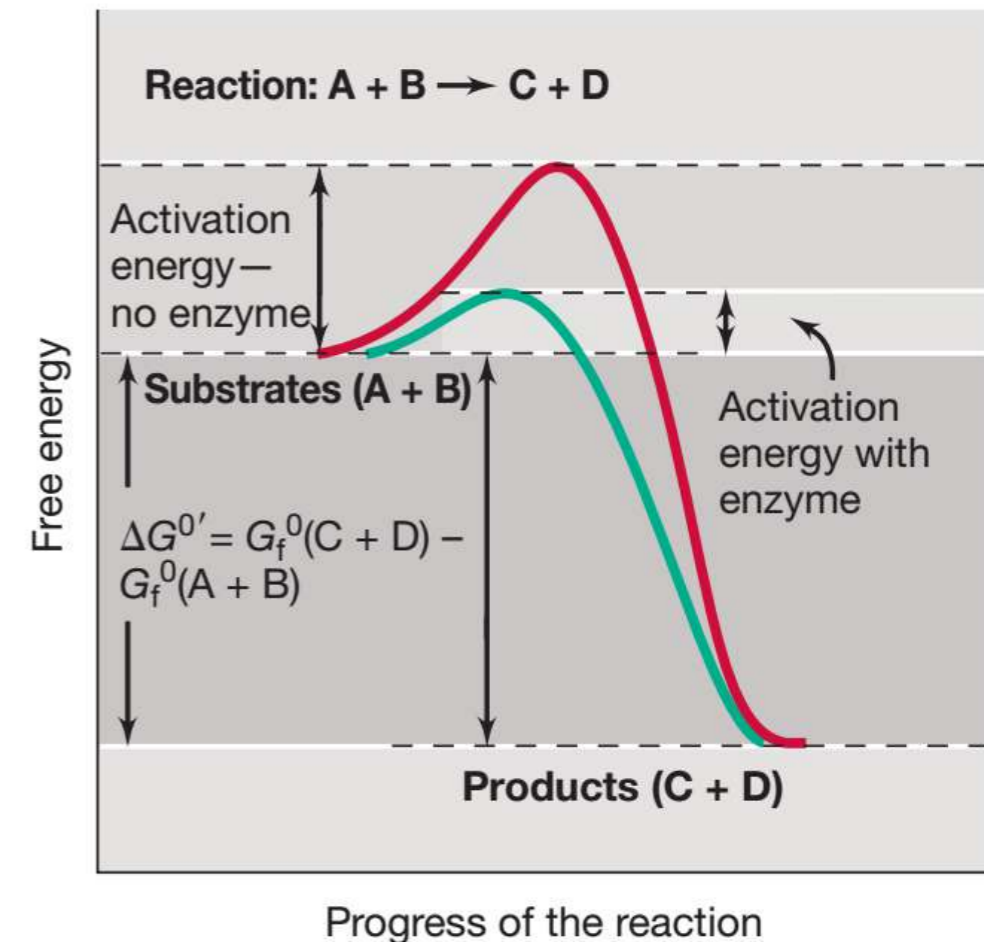
$$\Delta G = \Delta G^{\circ} + RT \frac{[C][D]}{[A][B]}$$

$\Delta G$  = free-energy change under conditions specified  
 $\Delta G^{\circ}$  = free-energy change under standard conditions of 1 atm pressure and 1 m concentration  
 $T$  = temperature (degrees K)  
 $R$  = universal gas constant (8.29 J/mol/K)

Madsen, 2016

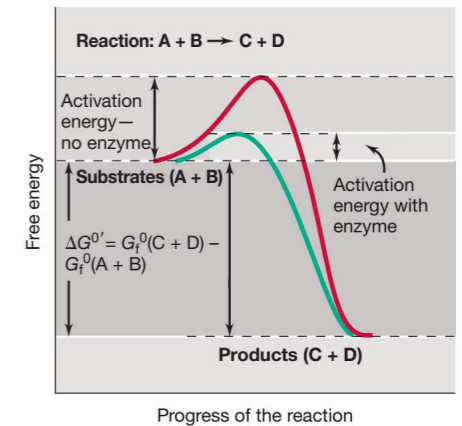
# Enzyme, I

- Free-energy calculations reveal only whether energy is released or required in a given reaction
- $\Delta G^{0'}$  says nothing about the rate of the reaction
- **Activation energy** can be viewed as the **minimum energy required** for a chemical reaction to begin
- **Catalysts** function **by lowering the activation energy** of a reaction thereby increasing the reaction rate





# Enzyme, II



- **Catalysts** have no effect on the energetics or the equilibrium of a reaction but **affect the rate at which a reaction proceeds**
- **Most cellular reactions will not proceed at significant rates without catalysis**
- The **major** catalysts in cells are enzymes, **proteins** (or in a few cases, **RNAs**) that are **highly specific**
- This **specificity** is a function of the precise **3D structure** of the enzyme.
- In an enzyme-catalyzed reaction, the enzyme combines with the reactant, called a substrate, forming an enzyme–substrate complex. Then, as the reaction proceeds, the product is released and the enzyme is returned to its original state, ready to catalyze a new round of the reaction
- **Prosthetic** groups bind **tightly** to their enzymes, usually covalently and permanently (e.g. heme group present in cytochromes such as cytochrome c)
- **Coenzymes**, with a few exceptions, **are loosely and often transiently** bound to enzymes
- Single coenzyme molecule may associate with a number of different enzymes (e.g. vitamins)

# Speed and specificity of a reaction

If the enzyme has more than **one possible substrate**, the  $k_{cat}/K_m$  values determine the **specificity** of the enzyme for each

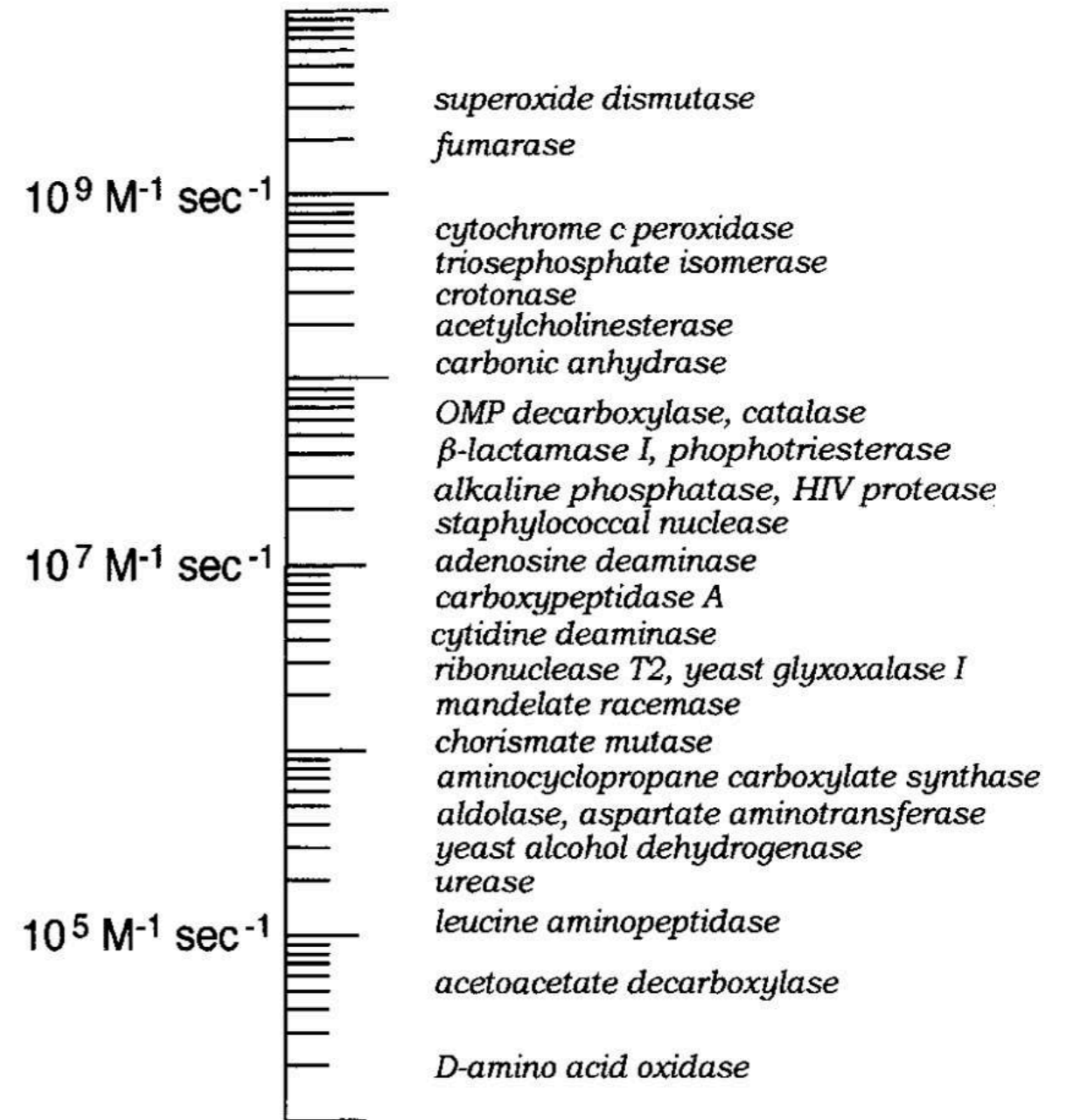
The **higher** this value the **more specific** the enzyme is for that substrate

This is because a high value of  $k_{cat}$  and a low value of  $K_m$  are expected for the best substrates

If  $k_{cat}/K_m$  – which is the apparent second-order rate constant for the **enzyme-catalyzed reaction** – approaches the **diffusion limit** ( $\sim 10^8$ – $10^9 \text{ M}^{-1} \text{ s}^{-1}$ ), the enzyme cannot catalyze the reaction any better and is said to have reached '**catalytic perfection**'

—> second-order rate constants that approach their rates of encounter ( $\sim 10^9 \text{ s}^{-1} \text{ M}^{-1}$ ) with the substrate in solution

*Triosphosphate isomerase*, *superoxide dismutase* and *carbonic anhydrase* are examples of perfect enzymes



**FIGURE 1.** Representative values of  $k_{cat}/K_m$  at 25 °C compiled from the literature.

# Evolution in action !

The **half-life** of a reaction: **amount of time needed for a reactant concentration to decrease by half compared to its initial concentration**

Enzymes allow organisms to **channel the flow of matter to their own advantage**, allowing some reactions to proceed rapidly compared with other reactions that offer no selective advantage

After a substrate is bound at an enzyme's active site, its half-life is usually a small fraction of 1 s

**Rapid turnover is necessary if any enzyme is to produce a significant rate of reaction at the limited concentration ( $<10^{-5}$  M) at which enzymes are present within the cell**

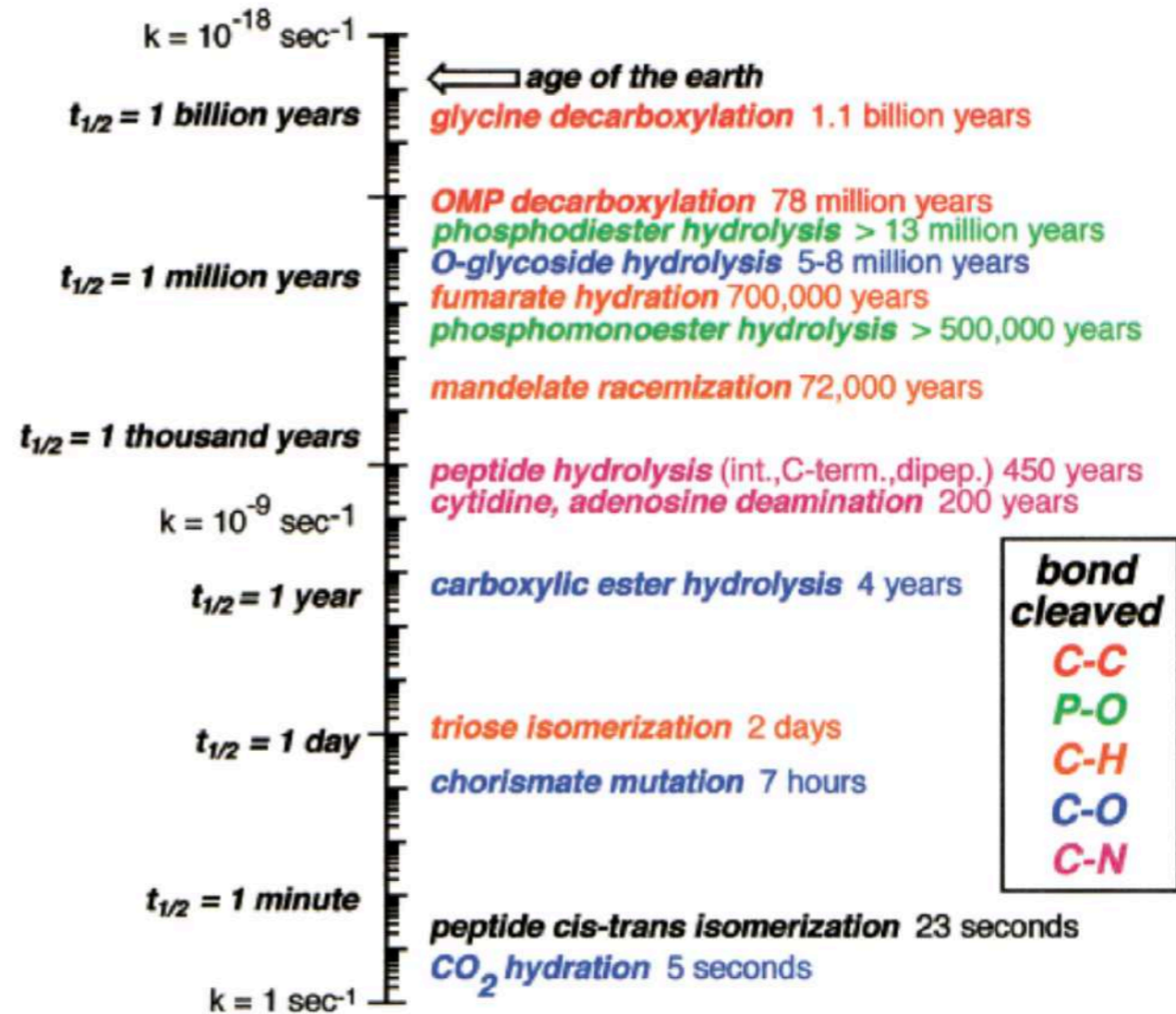


FIGURE 4. Natural half-times of some biological reactions in neutral solution at 25 °C.

WOLFENDEN and SNIDER, 2001

**Table 1. Cleavage of Polymers at 25 and 100 °C**

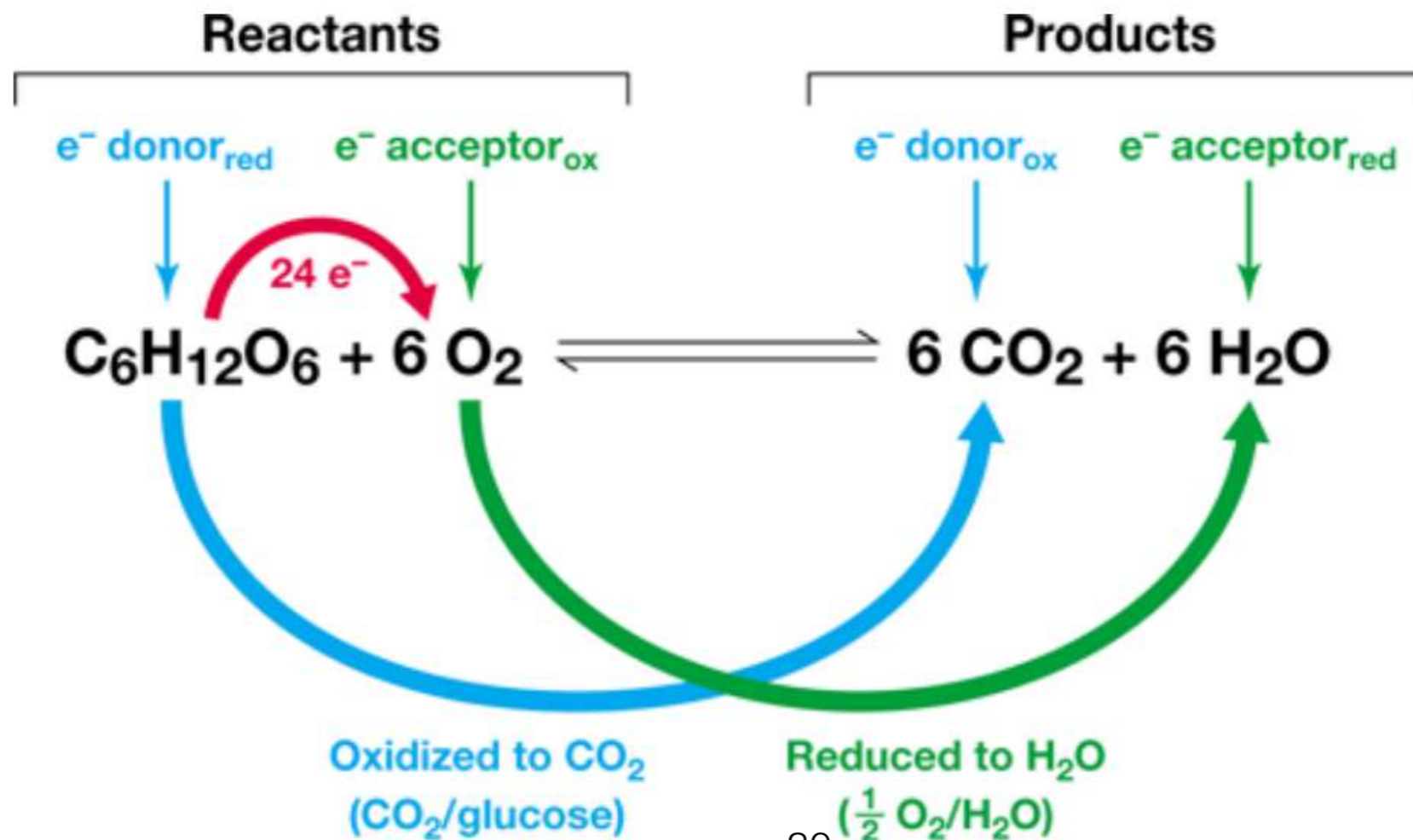
reaction	bond $t_{1/2}$		no. of bonds per polymer	$t_{1/2}$ per cleavage event	
	25 °C	100 °C		25 °C	100 °C
protein hydrolysis	400 years	5.5 weeks	123 (RNase A)	4 years	7 hours
polysaccharide hydrolysis	$4.7 \times 10^6$ years	160 years	$10^5$ residues (glycogen)	50 years	12 hours
RNA hydrolysis	4 years	9 days	70 residues (tRNA)	20 days	3 hours
DNA hydrolysis	140 000 years	22 years	$10^9$ residues (human DNA)	1 month	2 hours



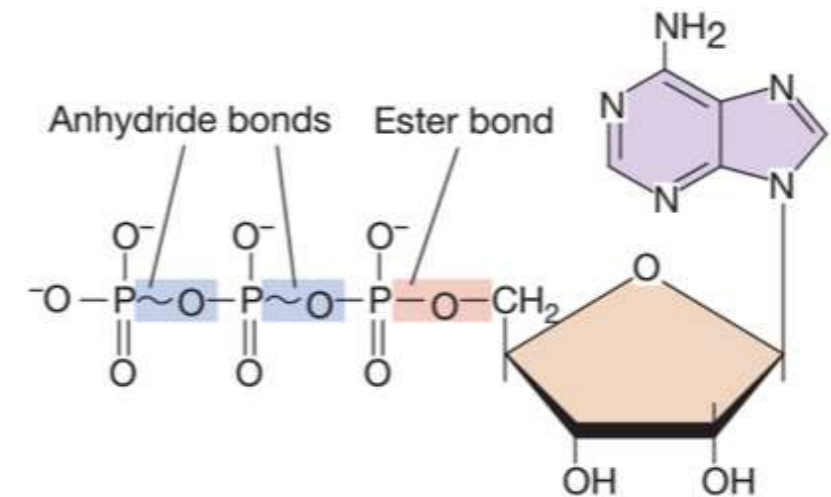
# Fundamentals in Metabolisms

- Transfer  $e^-$  and conserve energy
- Reactions are not performed in single-step  $\rightarrow$  consecutive reactions in different part of the cells
- Need of soluble  $e^-$  carriers:  $\text{NAD}^+/\text{NADH}$ ,  $\text{FAD}^+/\text{FADH}_2$

Madigan et al. 2020

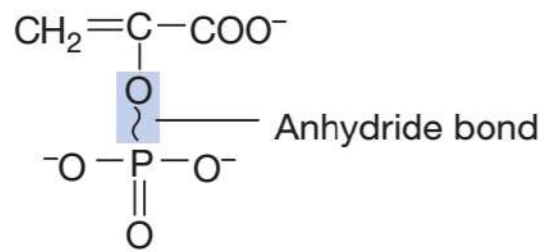


# Energy conservation

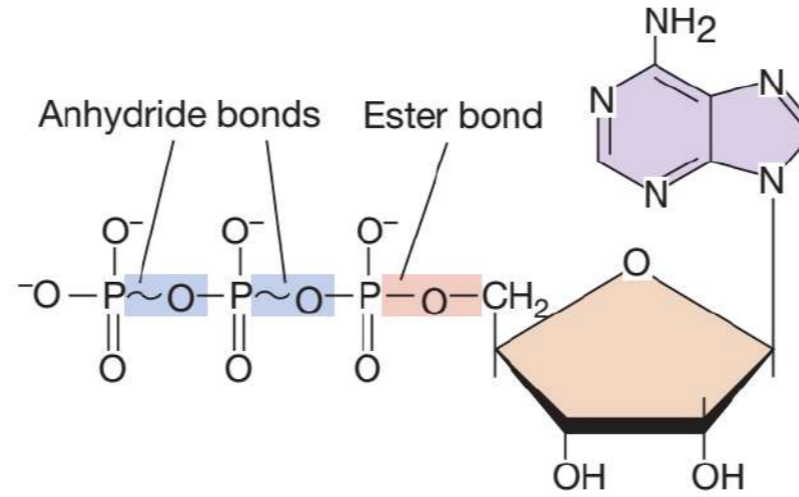


- ATP generation, adenosine triphosphate
- **ATP the energy currency of the cell** → cell motility, biosynthetic reactions, replication, cell growth and heredity, its generation relays:
  - 1. **Substrate-level phosphorylation**
  - 2. **Membrane-bound e<sup>-</sup> transport chain** → create **H<sup>+</sup> motive force** that drives ATP synthetase embedded in cytoplasmic membrane
- Earth including biota are a very heterogeneous and complex environment → thermodynamics (prediction of reactions that are energetically favorable)

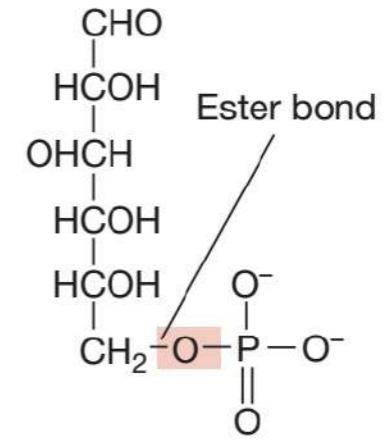
# Energy-rich compounds



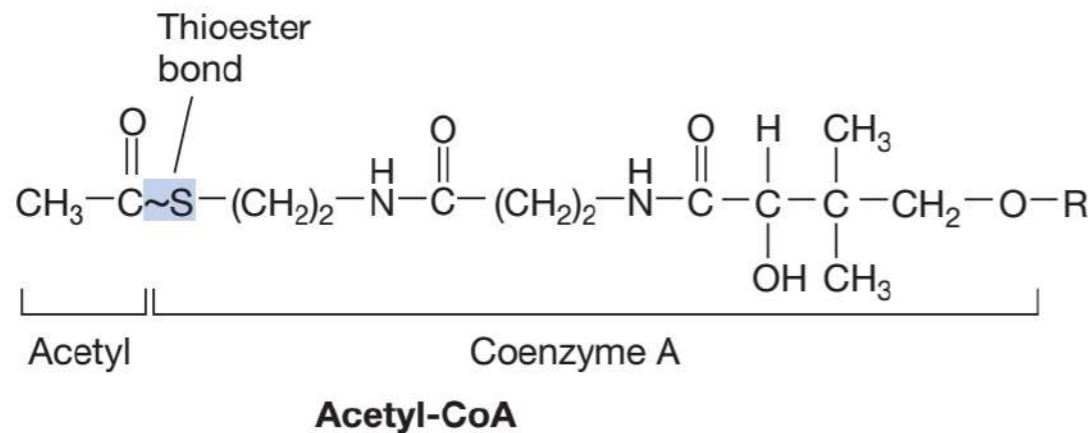
Phosphoenolpyruvate



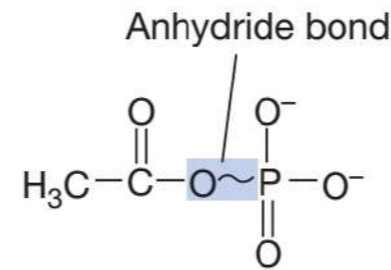
Adenosine triphosphate (ATP)



Glucose 6-phosphate



Acetyl-CoA



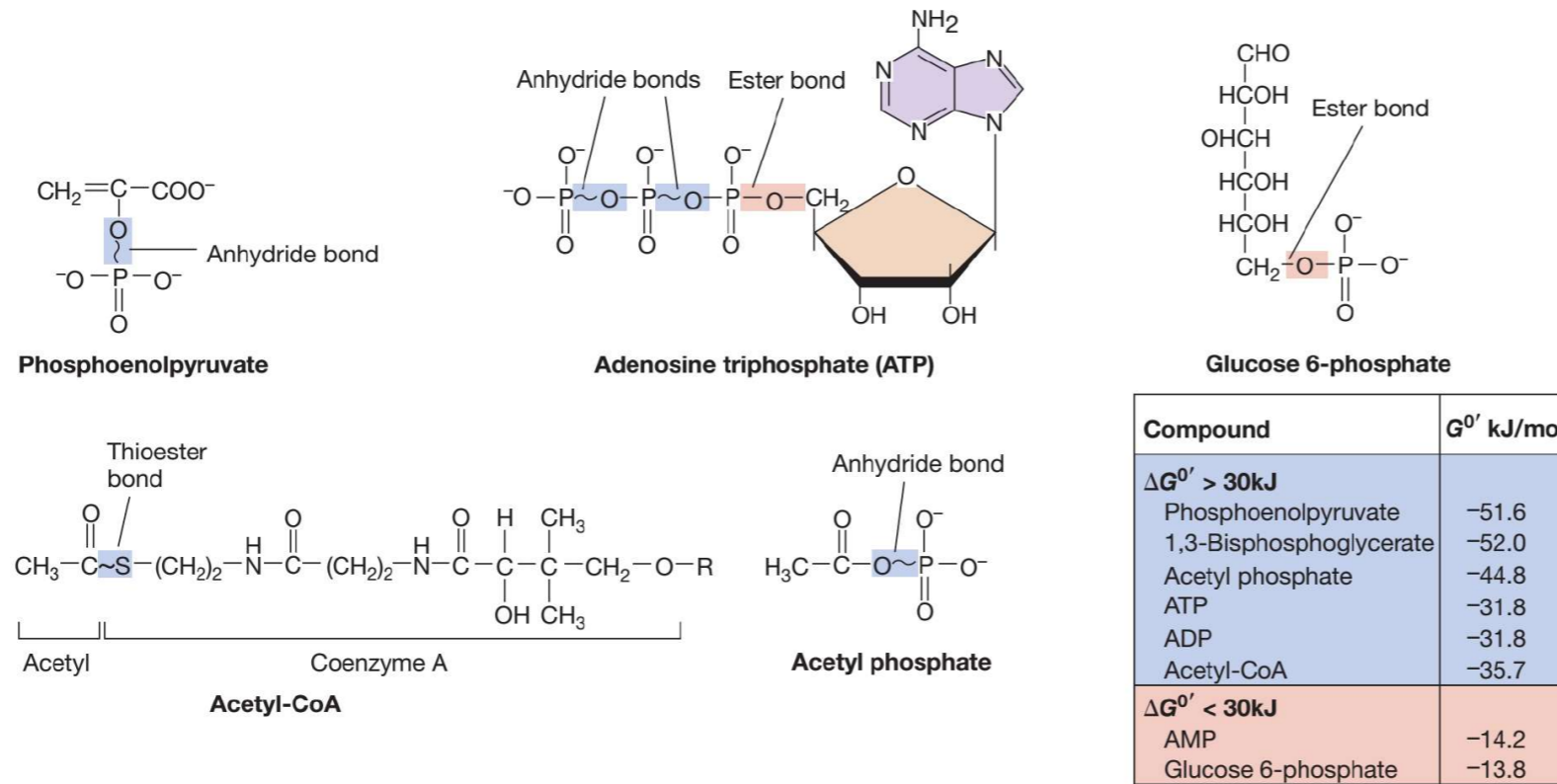
Acetyl phosphate

Compound	$G^{0'}$ kJ/mol
<b><math>\Delta G^{0'} &gt; 30\text{kJ}</math></b>	
Phosphoenolpyruvate	-51.6
1,3-Bisphosphoglycerate	-52.0
Acetyl phosphate	-44.8
ATP	-31.8
ADP	-31.8
Acetyl-CoA	-35.7
<b><math>\Delta G^{0'} &lt; 30\text{kJ}</math></b>	
AMP	-14.2
Glucose 6-phosphate	-13.8



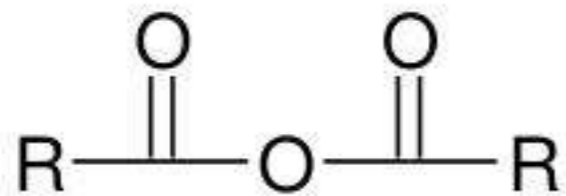
# Energy-rich compounds

Madigan et al. 2018

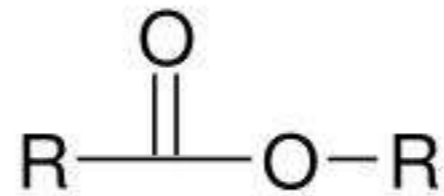


- The energy released from **redox** reactions **fuels energy-requiring cell functions**
- Free energy released in the coupled **exergonic redox** reaction **must first be trapped** by the cell and conserved
- Energy conservation in cells is accomplished through the formation of a set of compounds containing **energy-rich phosphate or sulfur bonds**
- The biosynthesis of these compounds functions as the **free-energy trap**, and their hydrolysis releases this energy **to drive endergonic reactions**

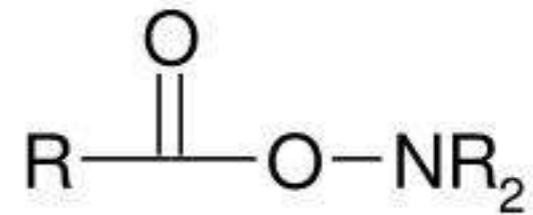
# C and P: Anhydrides and Esters



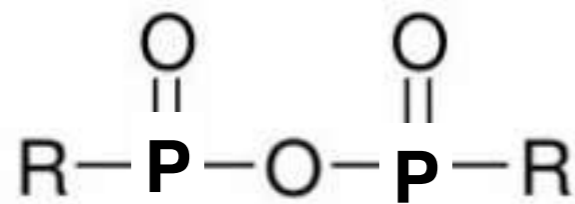
Anhydride



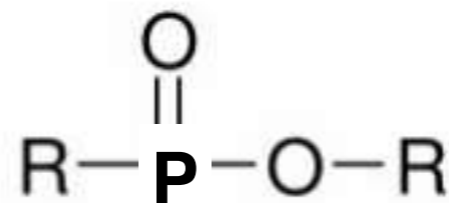
Ester



Amide



Anhydride

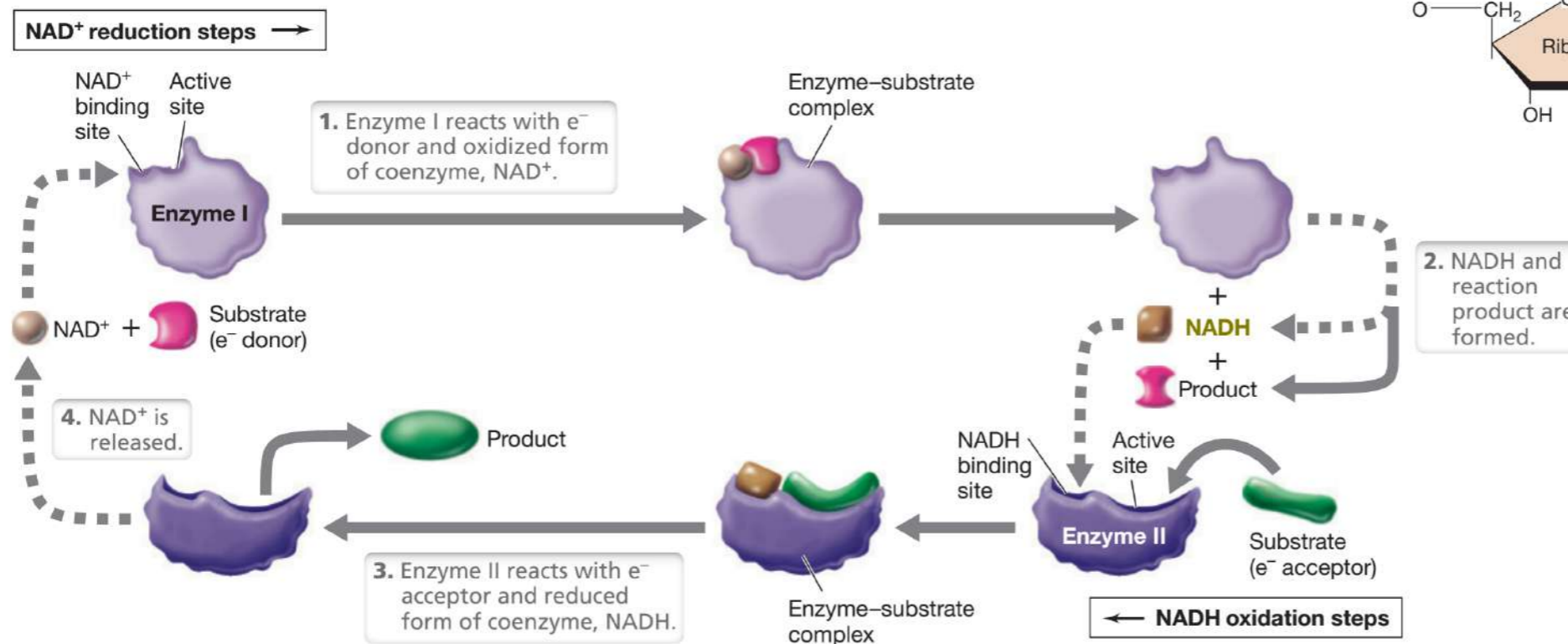
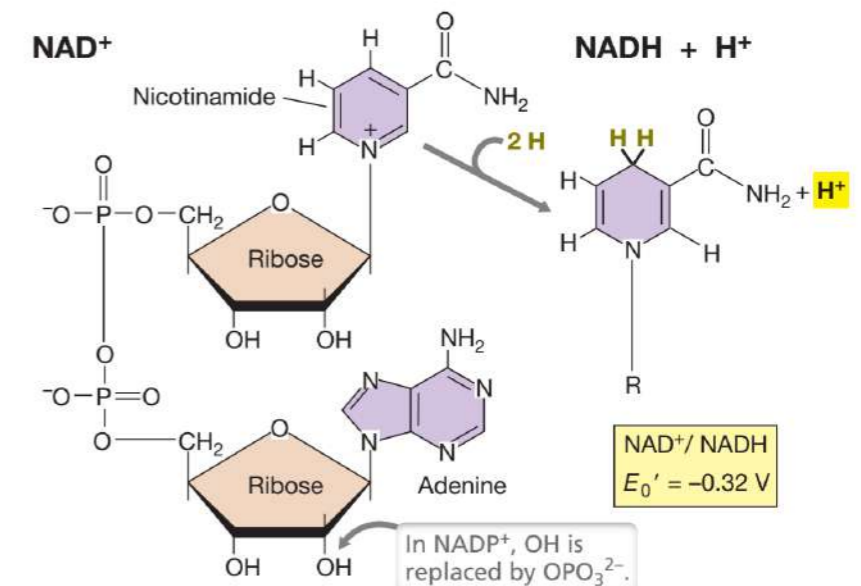


Ester

**ATP**

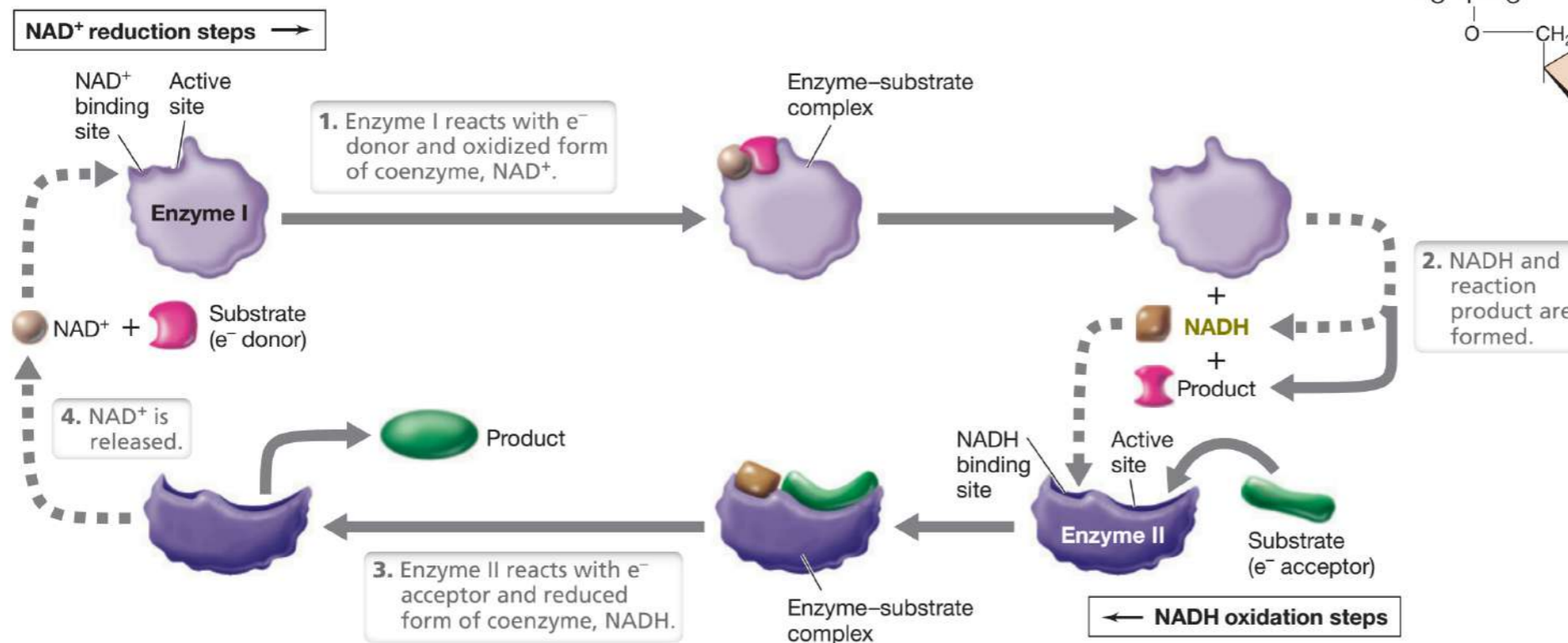
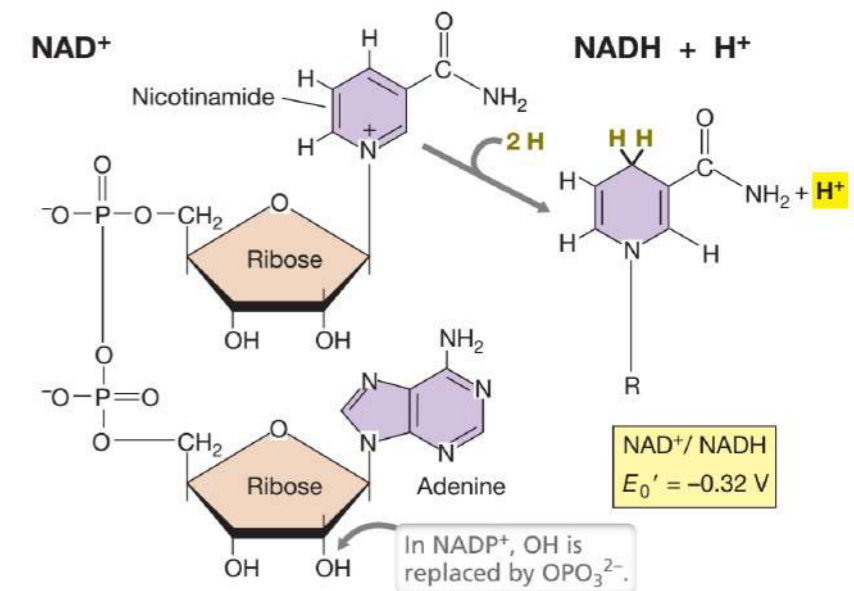
# Nicotinamide adenine dinucleotide, I

- Redox reactions are typically facilitated by coenzymes that associate with the redox enzymes that catalyze the reaction
- Reduction potential  $\text{NAD}^+/\text{NADH} = -0.32\text{V}$
- Electron plus proton carrier, transporting  $2\text{e}^-$  and  $2\text{H}^+$  simultaneously



# Nicotinamide adenine dinucleotide, II

- NADH is a good electron donor
- Nicotinamide adenine dinucleotide phosphate (NADP<sup>+</sup>) is made from NAD<sup>+</sup> by adding a phosphate molecule
- NADP<sup>+</sup> /NADPH participates in anabolic redox reactions (biosynthesis of cellular)
- NAD<sup>+</sup>/NADH participates in catabolic redox reactions





# Electron Donors and Electron Acceptors

- Cells conserve energy released from **exergonic reactions** by **coupling** the reaction to the **biosynthesis** of **energy-rich compounds**, such as ATP
- Reactions that **release** sufficient **energy** to form **ATP** require **oxidation–reduction** biochemistry
- An **oxidation** is the **removal of an electron** (or electrons) from a substance, and a **reduction** is the **addition of an electron** (or electrons) to a substance.
- In redox reactions, we refer to the **substance oxidized** as the **electron donor**, and the **substance reduced** as the **electron acceptor**
- By convention, in writing a redox couple, the **oxidized** form of the couple is always placed on the **left** (before the forward /slash) followed by the **reduced** form **after** the forward slash

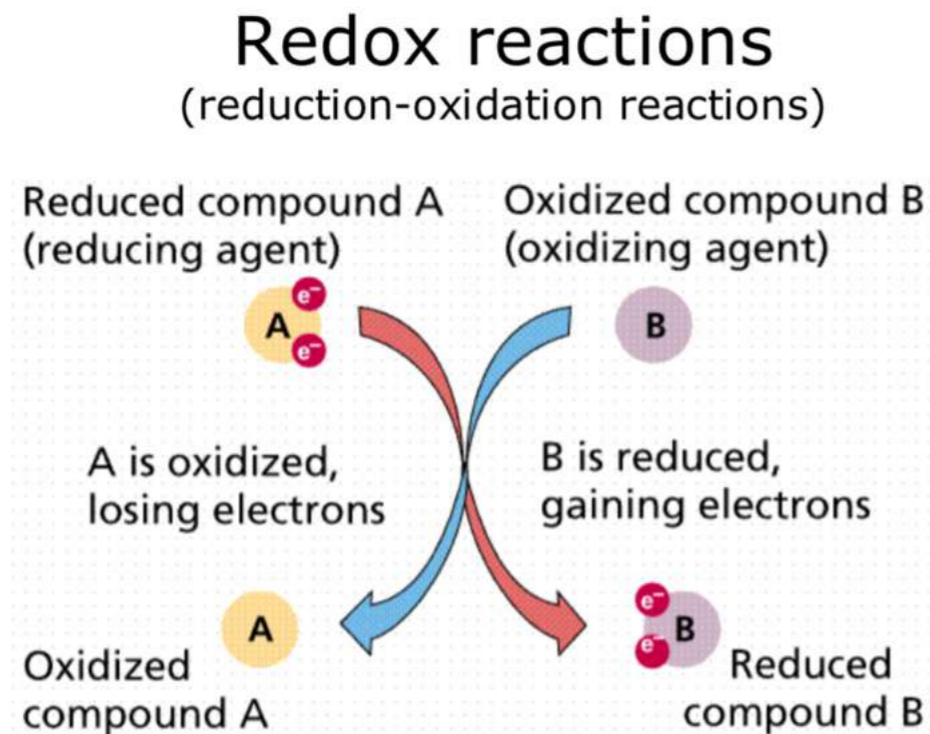
# Electron Donors and Electron Acceptors

- **Substances differ** in their tendency to donate or accept electrons
- This tendency is expressed as their reduction potential ( $E^{0'}$ , standard conditions), a value measured in volts (V) compared with that of a reference substance,  $H_2$
- When two redox couples react, the **reduced substance** of the couple ( $E^{0'}$  is  $< 0$ , negative) **donates electrons** to the **oxidized substance** ( $E^{0'}$   $> 0$ , positive)
- The half reaction with the more negative  $E^{0'}$  proceeds as an oxidation and is therefore written in the opposite direction

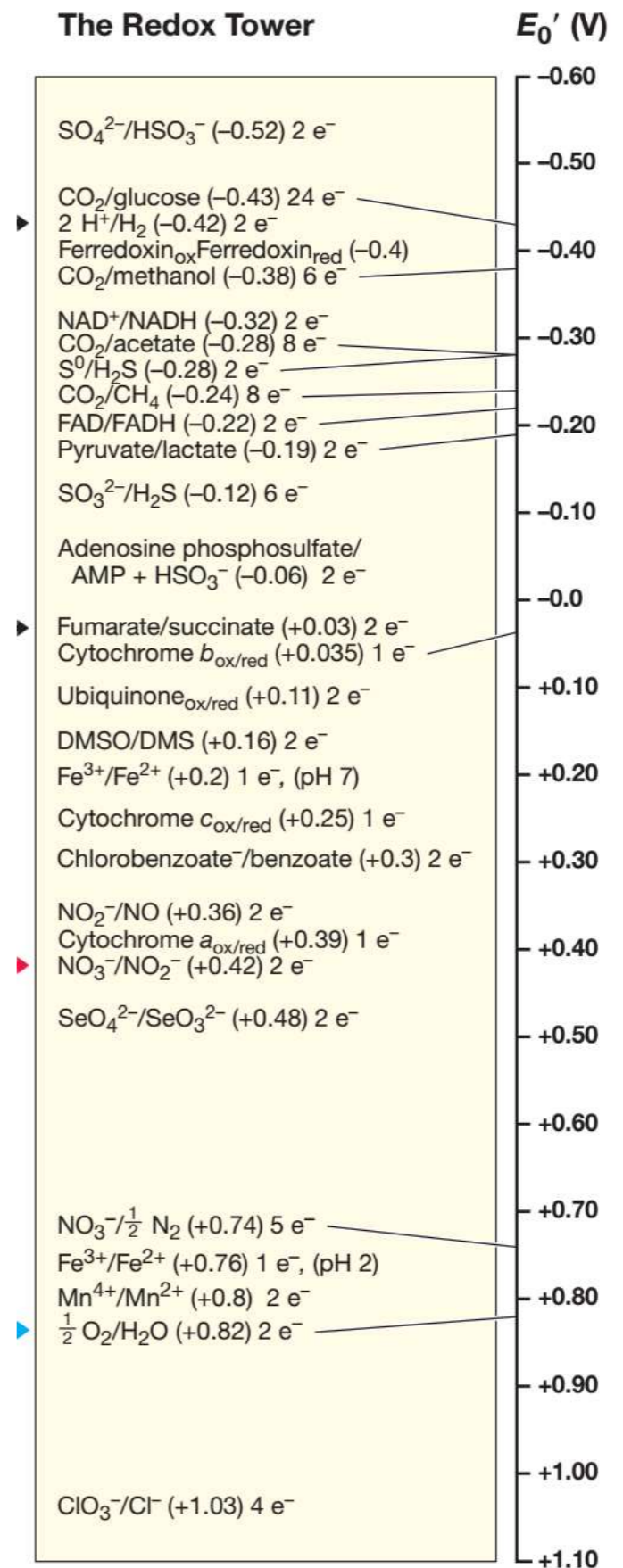
# Redox Tower

Reduced

- Redox couples are arranged from the strongest donors at the top ( $E^{\circ'} < 0$ ) to the strongest acceptors at the bottom ( $E^{\circ'} > 0$ )
- The larger the difference in reduction potential between electron donor and electron acceptor, the more free energy is released ( $\Delta G^{\circ'}$  can be computed via Nerst equation from reduction potential)



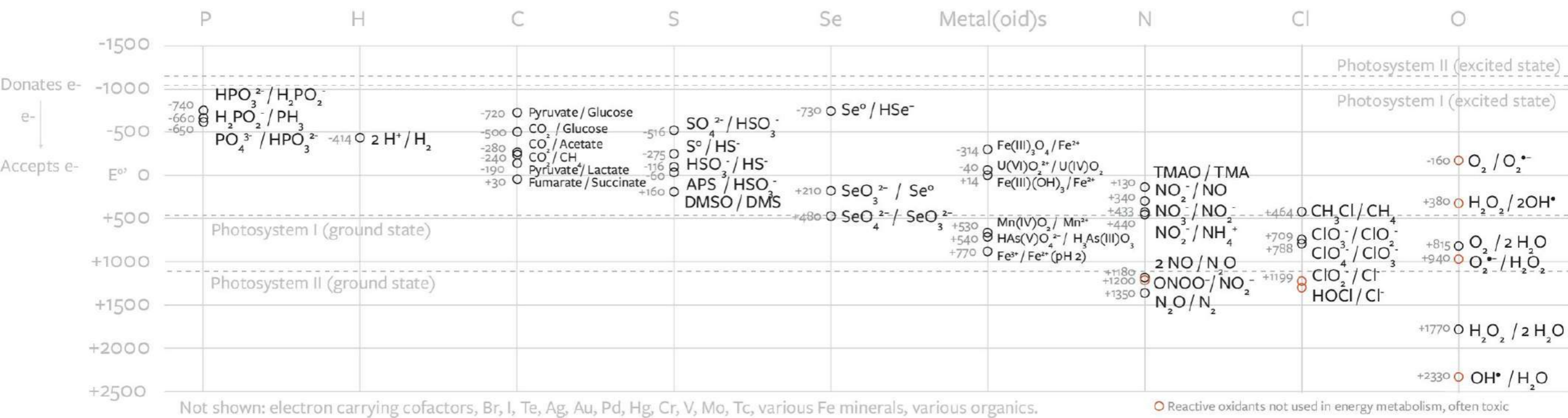
Oxidized



# Redox couples in the environment

## Redox couples and potentials (mV) for elements common in biology at pH 7 and temperature 25 C \*

Redox potential indicates the propensity for a compound to transfer electrons to another compound. A more-negative redox potential means a compound is more likely to donate electrons (e-).  
**All of life gets its energy by capturing the change in potential energy** from the transfer of electrons from the reducing compound to the oxidizing compound.



Across periodic table



# High diversity of key compound in the Electron Transport Chain



Standard redox potential ( $E_0'$  [mV, 25°C, pH = 7]) are indicated by dashed (- - -) lines

If physiological or environmental conditions are known to shift the potential from the  $E_0'$ , redox windows are indicated (solid lines)

- **Blue:** aerobes
- **Green:** facultative anaerobes
- **Red–yellow:** obligate anaerobes

c-type cytochromes can cover a broad range of redox potentials as indicated.

Not all bacteria mentioned will cover the whole range.

