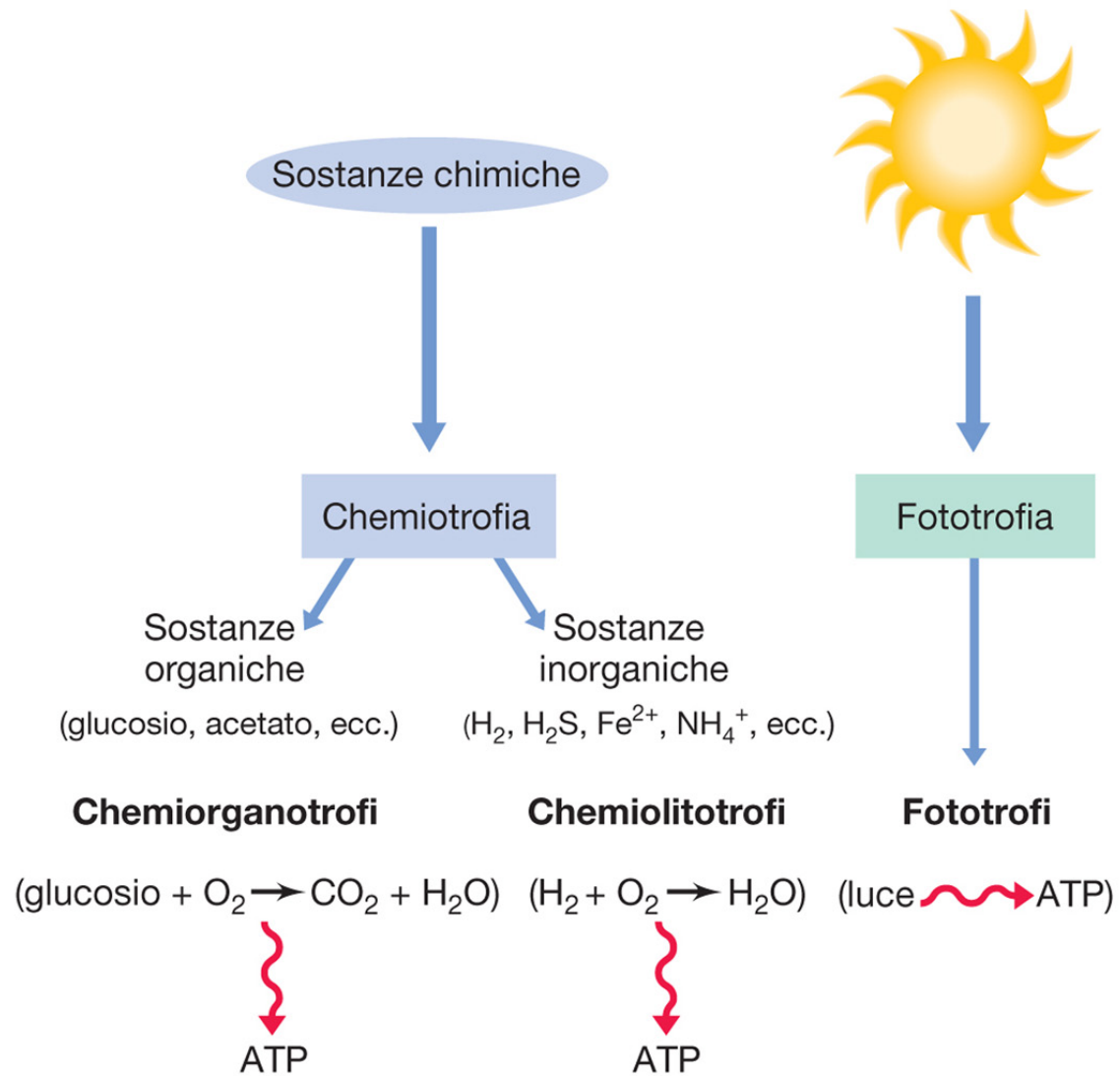


12. APPLIED BIOCHEMISTRY

Energy source

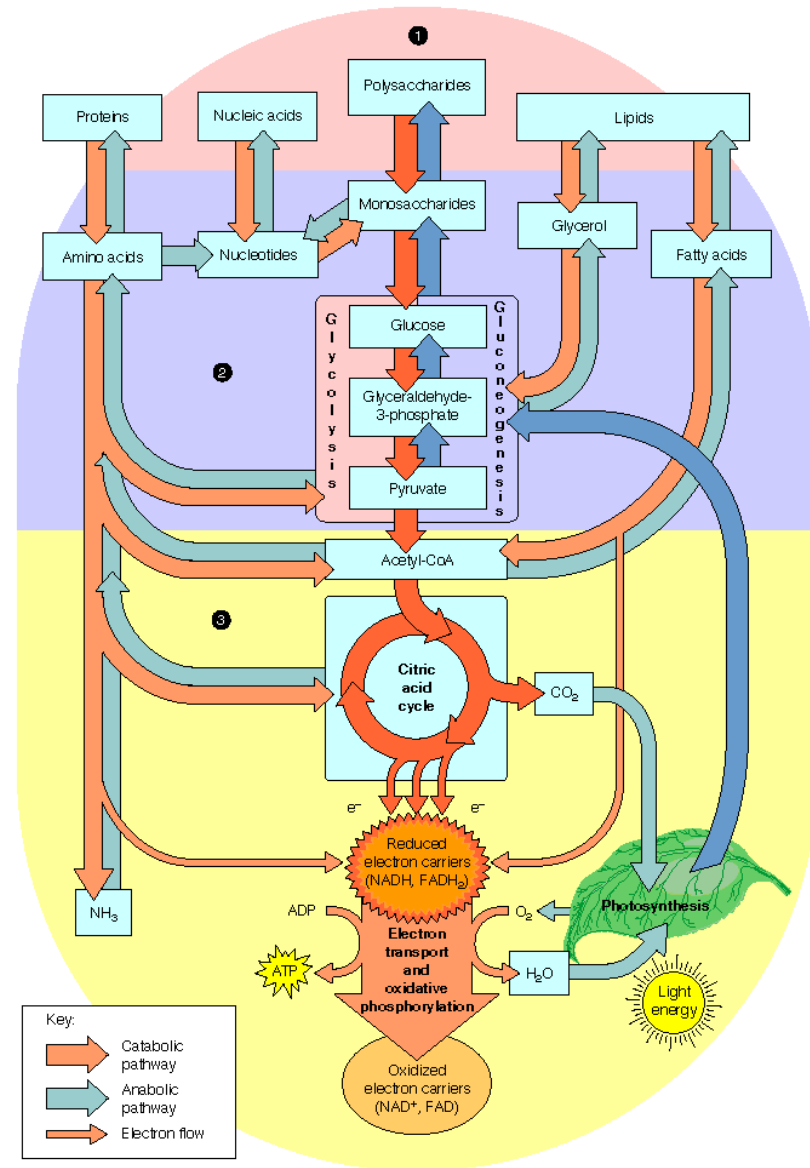
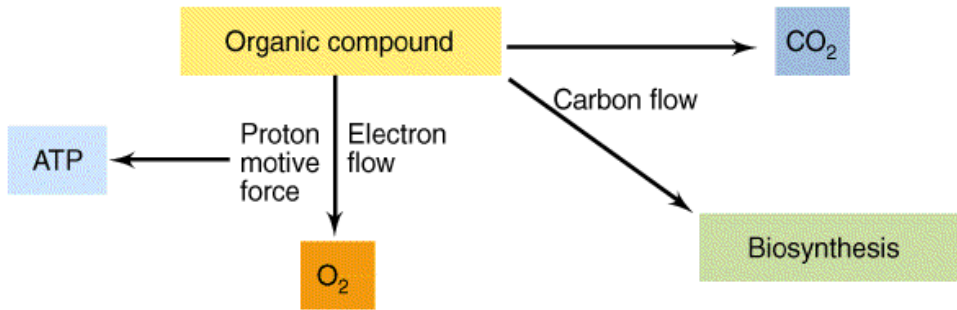


Energy and carbon source

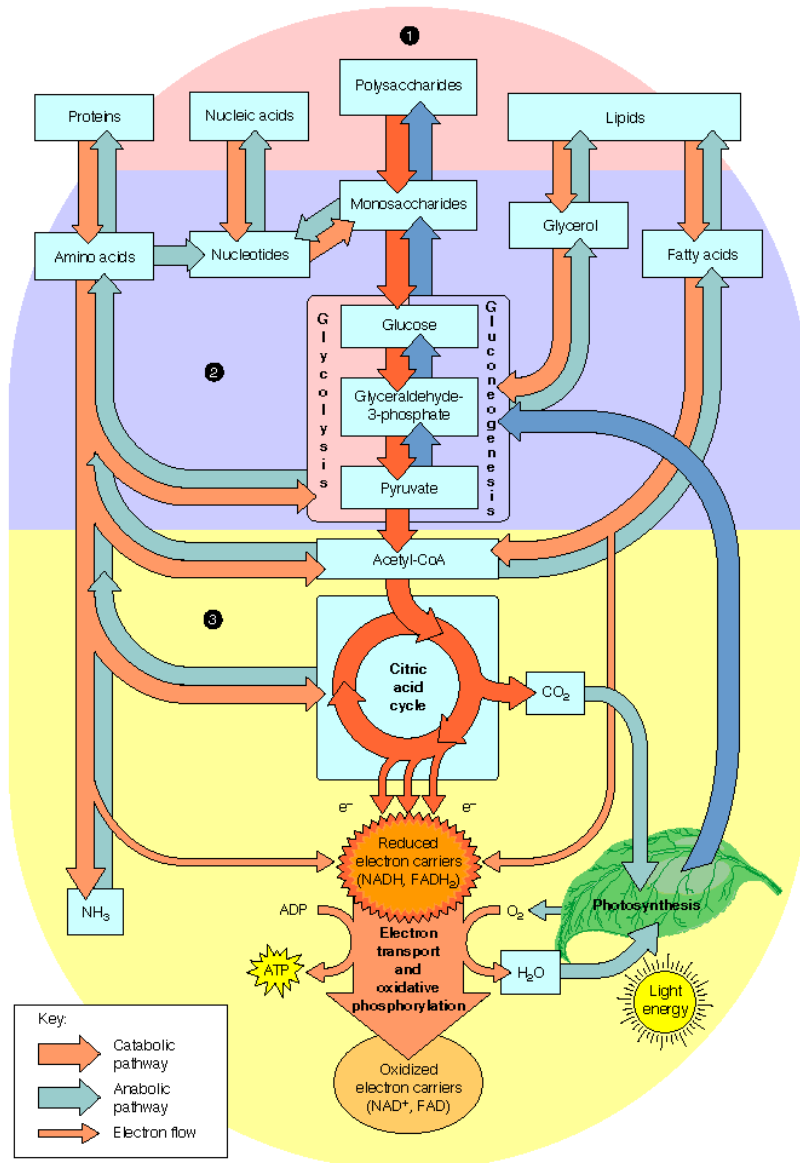
		Energy Source	
		Light (<i>photo-</i>)	Chemical compounds (<i>chemo-</i>)
Carbon Source	Carbon dioxide (<i>auto-</i>)	<p>Photoautotrophs</p> <ul style="list-style-type: none"> Plants, algae, and cyanobacteria use H₂O to reduce CO₂, producing O₂ as a byproduct Photosynthetic green sulfur and purple sulfur bacteria do not use H₂O nor produce O₂ 	<p>Chemoautotrophs</p> <ul style="list-style-type: none"> Hydrogen, sulfur, and nitrifying bacteria
	Organic compounds (<i>hetero-</i>)	<p>Photoheterotrophs</p> <ul style="list-style-type: none"> Green nonsulfur and purple nonsulfur bacteria 	<p>Chemoheterotrophs</p> <ul style="list-style-type: none"> Aerobic respiration: most animals, fungi, and protozoa, and many bacteria Anaerobic respiration: some animals and bacteria Fermentation: some bacteria and yeasts

Alternative energy generating patterns

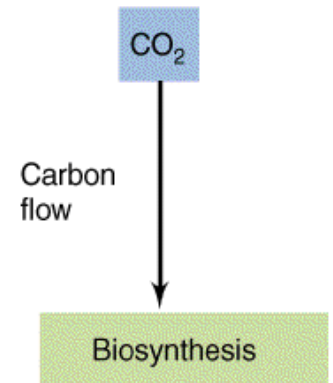
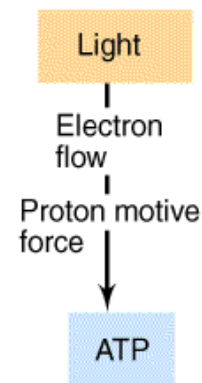
Aerobic respiration



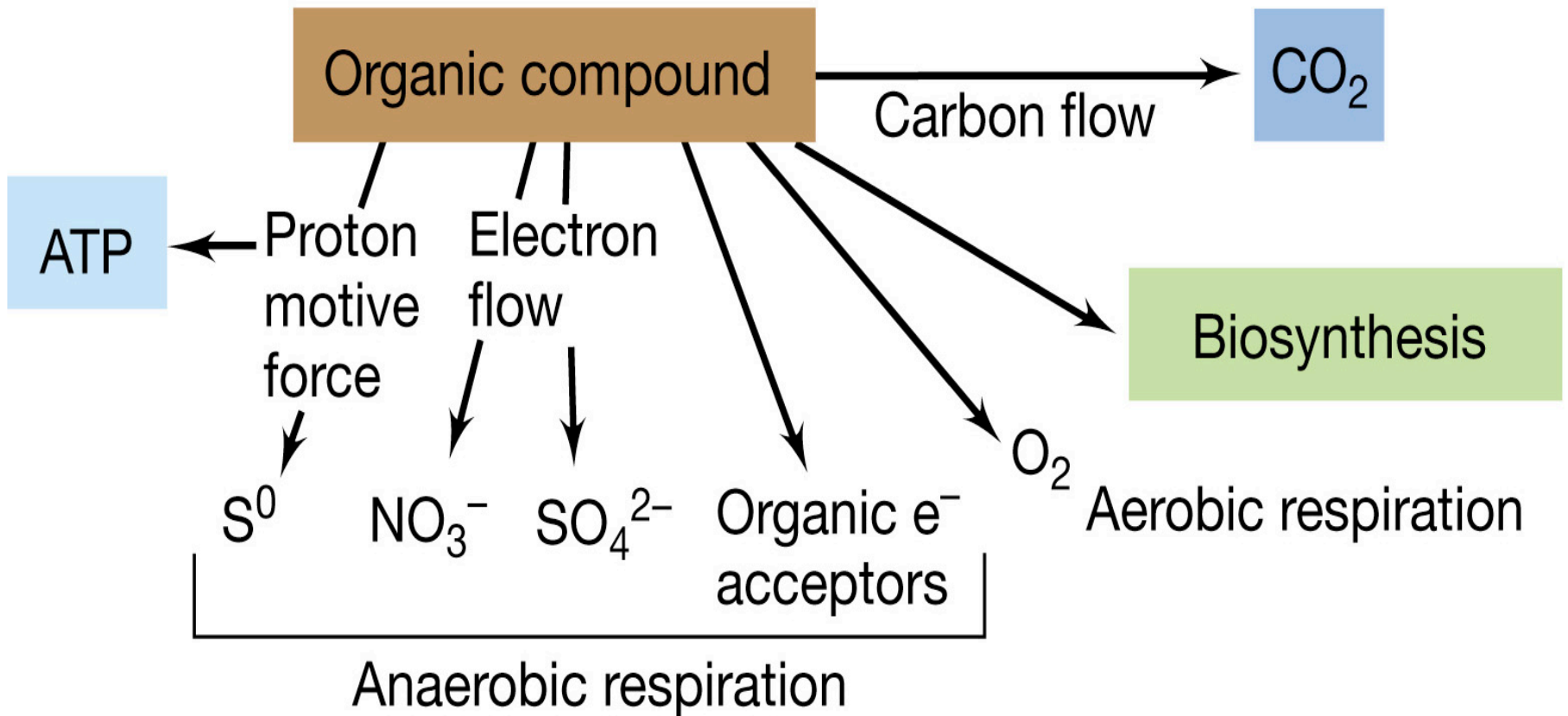
Alternative energy generating patterns



Phototrophic metabolism



Chemoorganotrophic metabolism

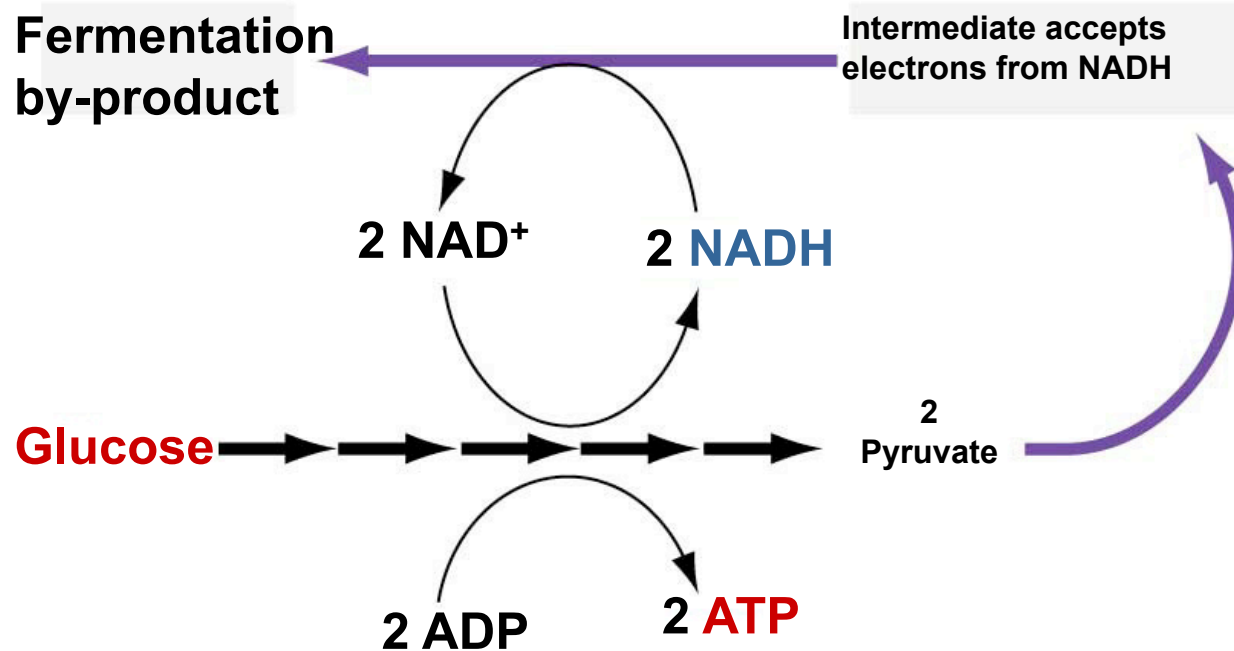


Chemoorganotrophic metabolism

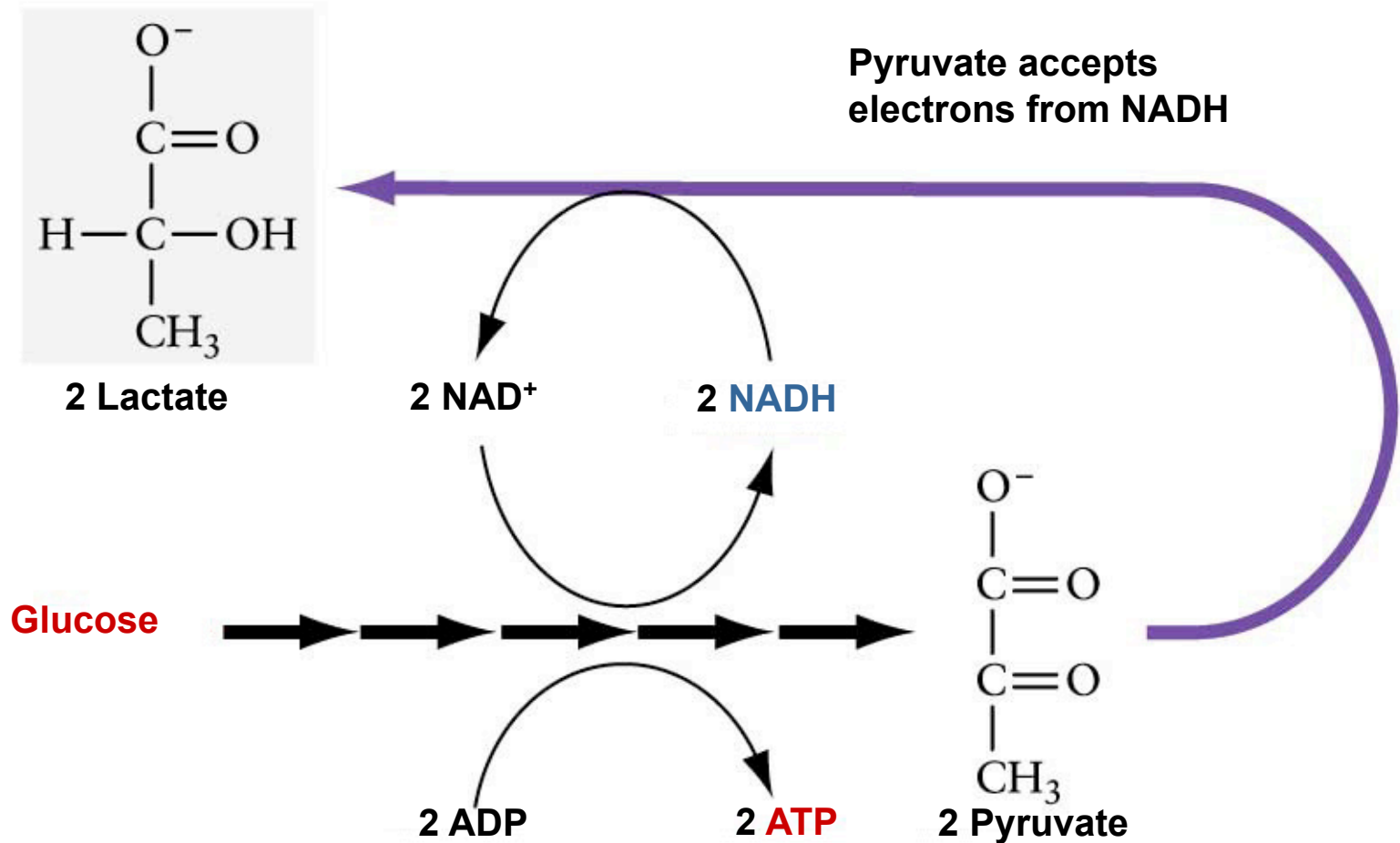
Electron acceptor	Final product	Name of the process	
O ₂	H ₂ O	Aerobic respiration	
NO ₃	NO ₂ , NH ₃ or N ₂	Anaerobic respiration: denitrification	<i>Bacillus</i> , <i>Pseudomonas</i>
SO ₄	S or H ₂ S	Anaerobic respiration: sulphates reduction	<i>Desulfovibrio</i>
fumarate	succinate	Anaerobic respiration: Organic e ⁻ acceptor	<i>E.coli</i>
CO ₂	CH ₄	Metanogenesis	(Archea)

Chemoorganotrophic metabolism

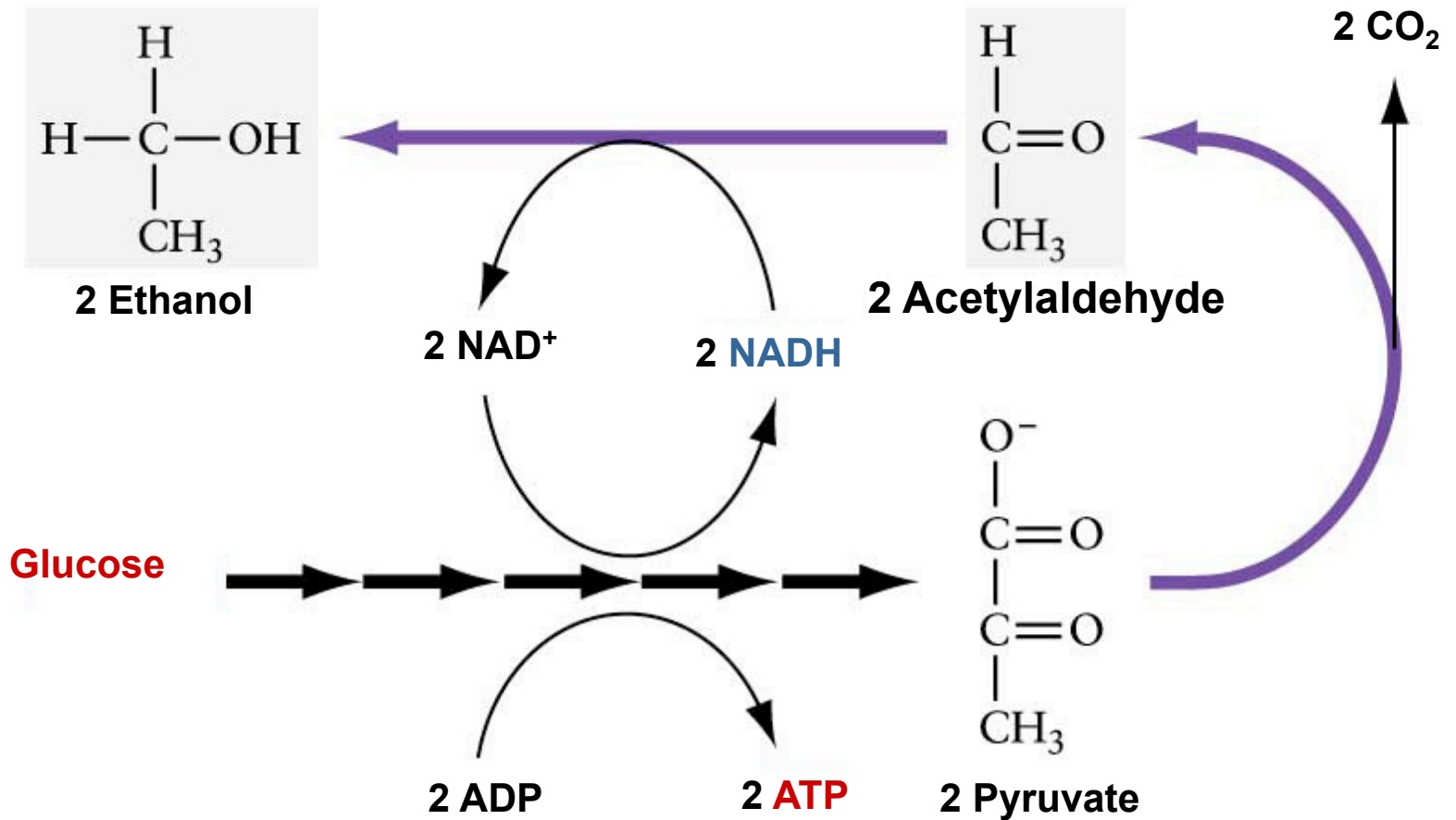
Fermentation pathways allows cells to regenerate NAD^+ for glycolysis



Chemoorganotrophic metabolism

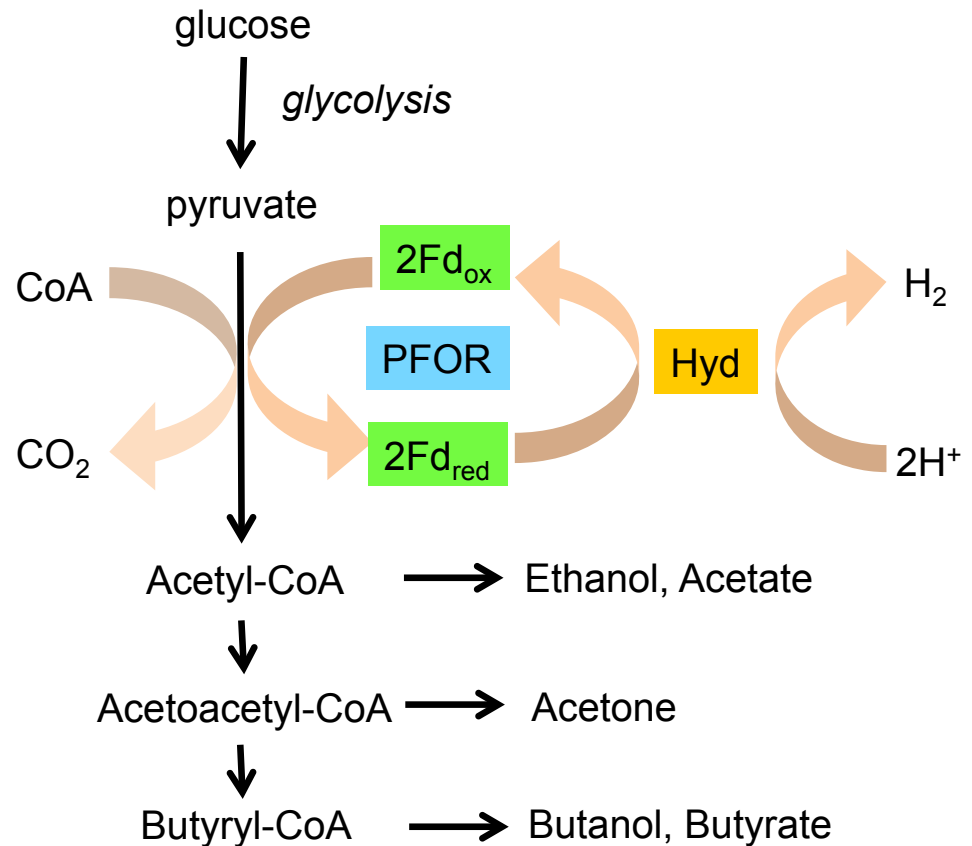


Chemoorganotrophic metabolism



Anaerobic Chemoorganotrophs: -Fermentors

- Genus *Clostridium*
 - Gram-positive rods found in soil
 - Endospores
- Ferment wide variety of compounds
- Representatives:
 - *C. tetani*,
 - *C. perfringens*,
 - *C. botulinum*



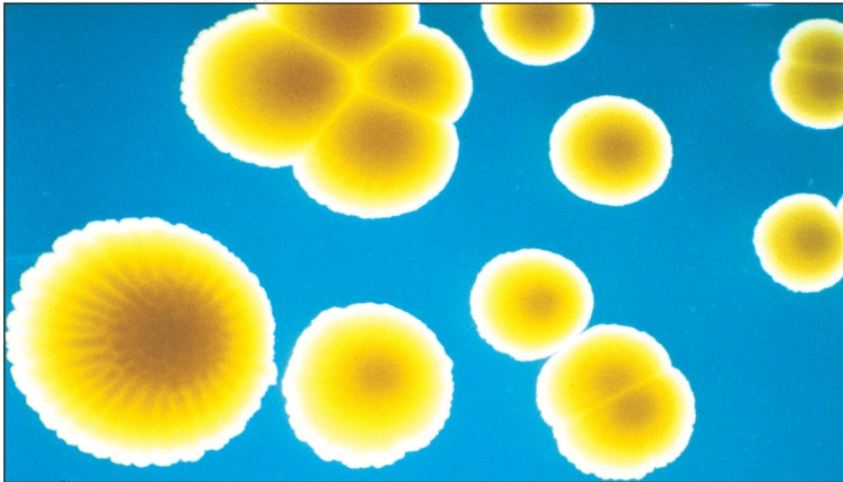
Anaerobic Chemotrophs

- *Propionibacterium* species are Gram-positive rods
- Organisms produce propionic acid as end product of fermentation
 - Found in anaerobic micro environments
 - Essential in the production of Swiss cheese
 - Also ferment lactic acid



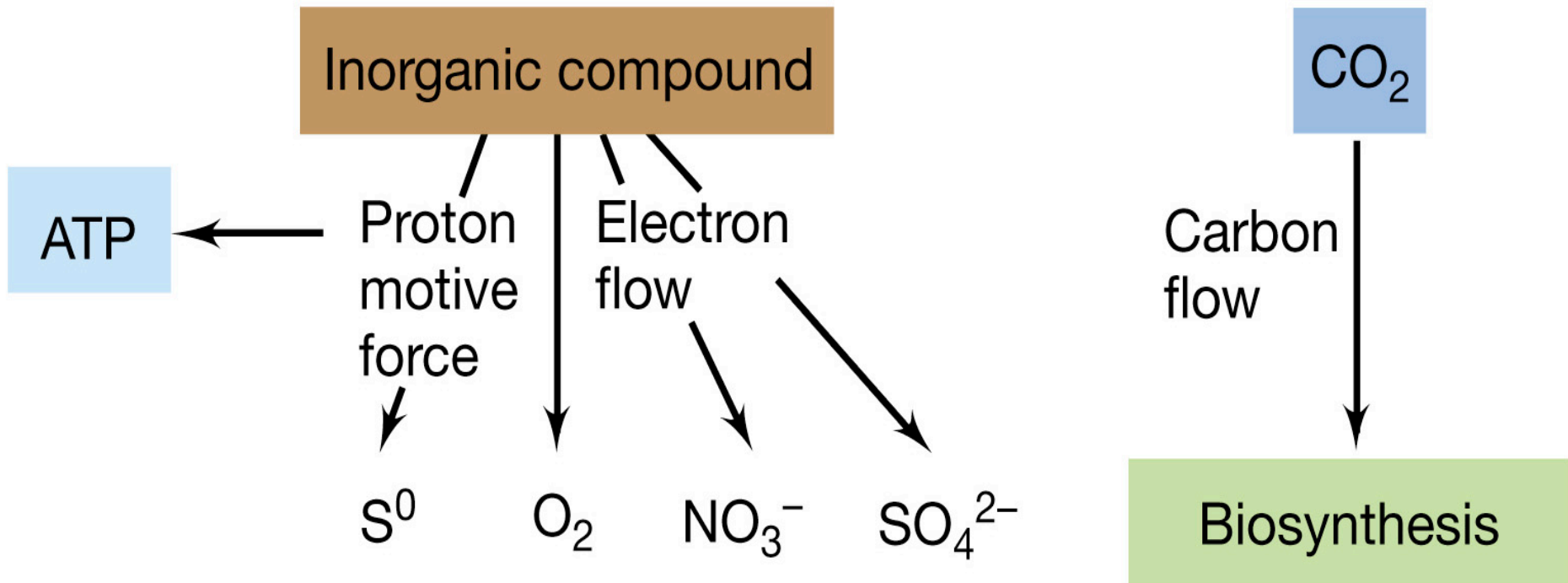
Aerobic Chemoorganotrophs: Obligate Aerobes

- Obligate aerobes obtain energy using aerobic respiration exclusively
- Characteristic genera include
 - > *Micrococcus*
 - Gram-positive cocci found in soil and dust
 - Produce yellow pigmented colonies



- *Mycobacterium*
 - > Gram-positive bacterium
 - > Live on dead and decaying matter
- *Pseudomonas*
 - > Gram-negative rods
 - > Motile and often pigmented
 - > Common opportunistic pathogen
- *Thermus* and *Deinococcus*
 - > Both have scientific and commercial uses
 - *Thermus* produces *Taq* polymerase
 - *Deinococcus* used to clean up radioactive contamination

Chemolithotrophic metabolism



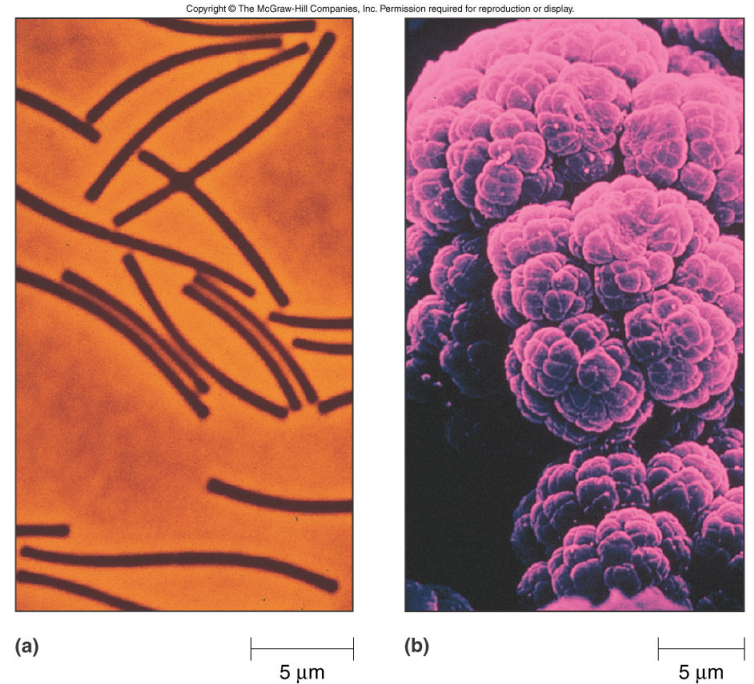
Chemolithotrophs

- Chemolithotrophs oxidize reduced inorganic chemicals (e.g. H₂) to produce energy
 - Rare organisms
 - Not O₂ tolerant
 - Terminal electron acceptor usually carbon dioxide or sulfur
 - Members of the domain *Archaea*

Group	Energy source	Final products	Microrganism
Hydrogen bacteria	H ₂	H ₂ O	<i>Alcaligenes, Pseudomonas</i>
Metanogenes	H ₂	H ₂ O	<i>Methanobacterium</i>
Carboxydobacteria	CO	CO ₂	<i>Rhodospirillum, Azotobacter</i>
Nitrifying bacteria	NH ₃	NO ₂	<i>Nitrosomonas</i>
Nitrifying bacteria	NO ₂	NO ₃	<i>Nitrobacter</i>
S-oxidizing bacteria	H ₂ S or S	SO ₄	<i>Thiobacillus, Sulfolobus</i>
Fe-bacteria	Fe ⁺⁺	Fe ⁺⁺⁺	<i>Gallionella, Thiobacillus</i>

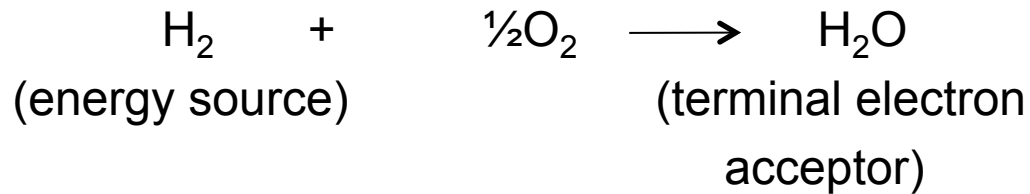
Chemolithotroph: Methanogens

- Members of Domain *Archaea*
- Found in sewage, swamps, marine sediments and digestive tract of mammals
- Highly sensitive to oxygen
- Produce energy (ATP) the reaction:
$$4\text{H}_2 + \text{CO}_2 \rightarrow \text{CH}_4 + 2\text{H}_2\text{O}$$



Aerobic Chemolithotrophs: Hydrogen-Oxidizing Bacteria

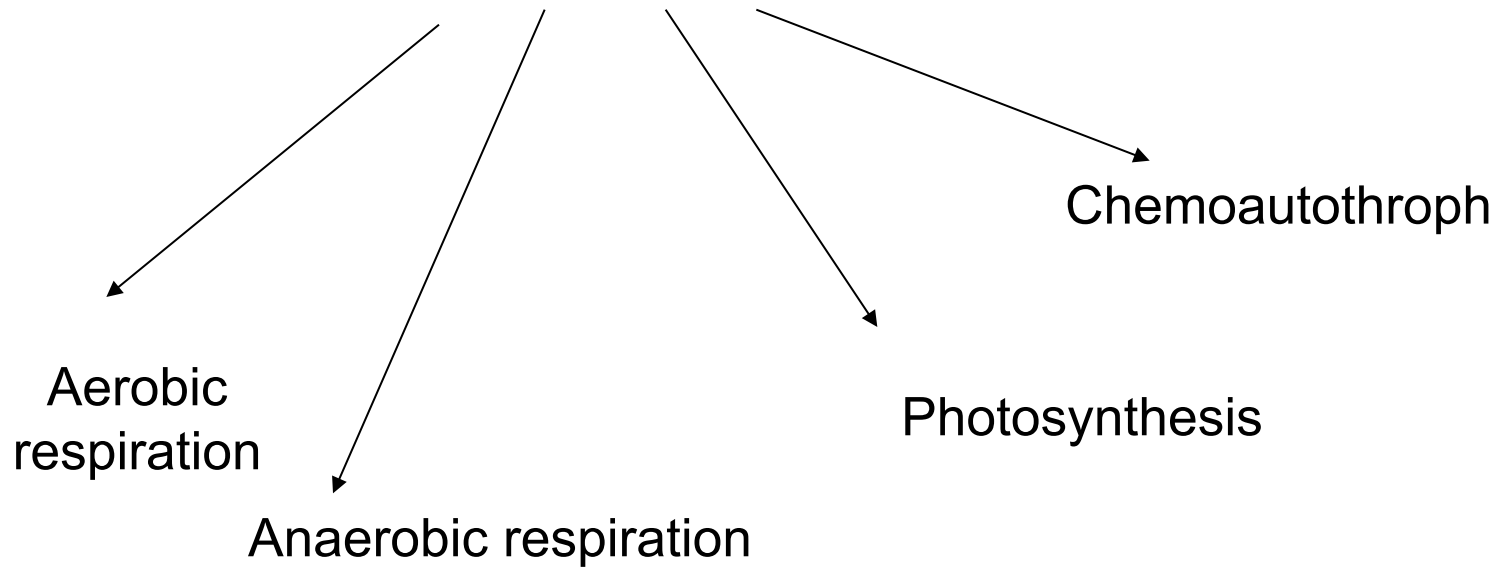
- Gram-negative bacteria
- Tend to be thermophilic
 - Found in hot springs, up to 95°C



Metabolic versatility

Rodospirillum rubrum

(Negative Gram, Proteobacteria)



BIOREMEDIATION

Bioremediation relies largely on the enzymatic activities of living organisms, usually microbes, to catalyze the destruction of pollutants or their transformation to less harmful forms.

Why are microorganisms so important in this process?

They have extraordinary metabolic diversity!

A complex process depending on many factors including:

- ambient environmental conditions (pH, temperature, lack of nutrients & molecular oxygen)
- composition of the microbial community
- nature and amount of pollution present

BIOREMEDIATION

"use of living organisms (e.g., bacteria) to clean up oil spills or remove other pollutants from soil, water, and wastewater."

Source: United States Environmental Protection Agency, Office of Compliance and Assurance

"clean-up of pollution from soil, groundwater, surface water and air, using biological, usually microbiological processes"

Source: Philp et al., 2001

BIOREMEDIATION

Types of pollutants

Organic pollutants → catabolized

- Naturally occurring
- Xenobiotics - substances foreign to an entire biological system, i.e. artificial substances, which did not exist in nature before their synthesis by humans
- Metals from ore extraction and manufacturing

CONTAMINANTS FOR BIODEGRADATION

Readily
degradable

—
fuel oils, gasoline

ketones and
alcohols

monocyclic
aromatics

bicyclic aromatics
(naphthalene)

Somewhat
degradable

creosote, coal
tars

pentachloro-
phenol (PCP)

Difficult to
degrade

chlorinated
solvents (TCE)

some pesticides
and herbicides

Generally
recalcitrant

dioxins

polychlorinated
biphenyls (PCB)

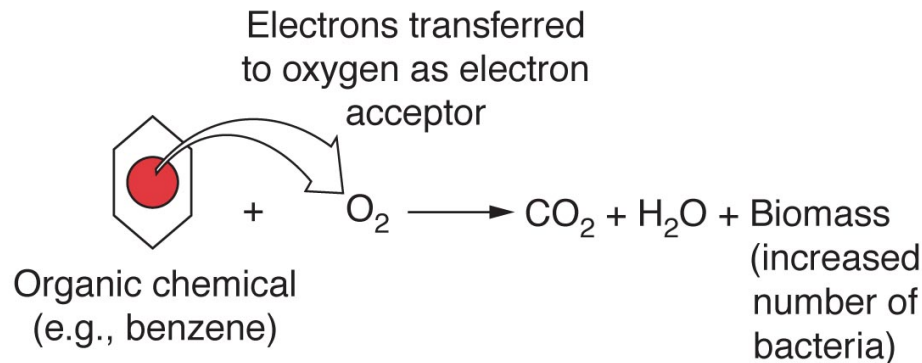
How Microbes Use the Contaminant

- Contaminants may serve as:
 - Primary substrate
 - enough available to be the sole energy source
 - Secondary substrate
 - provides energy, not available in high enough concentration
 - Cometary substrate
 - fortuitous transformation of a compound by a microbe relying on some other primary substrate

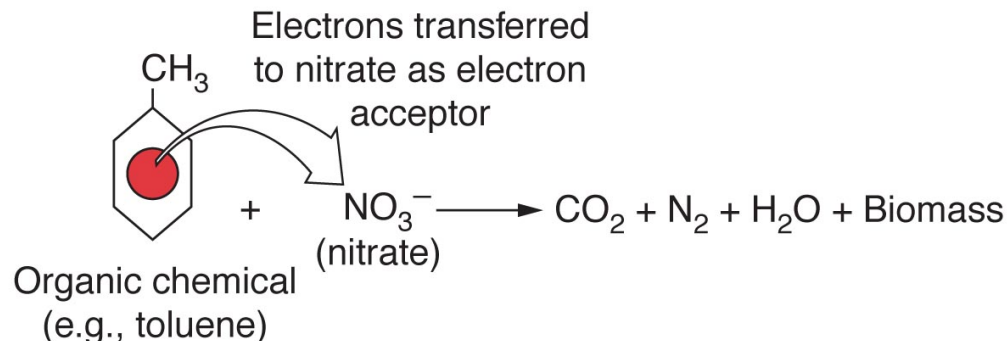
USE OF CONTAMINANTS AS PRIMARY SUBSTRATE

- Aerobic metabolism
 - Microbes use O_2 in their metabolism to degrade contaminants
- Anaerobic metabolism
 - Microbes substitute another chemical for O_2 to degrade contaminants
 - Nitrate, iron, sulfate, carbon dioxide, uranium, technicium, perchlorate

Aerobic biodegradation

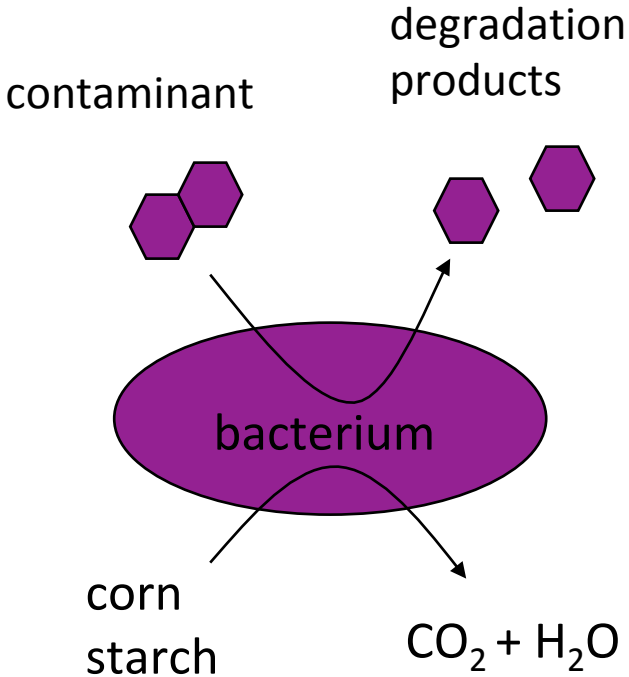


Anaerobic biodegradation

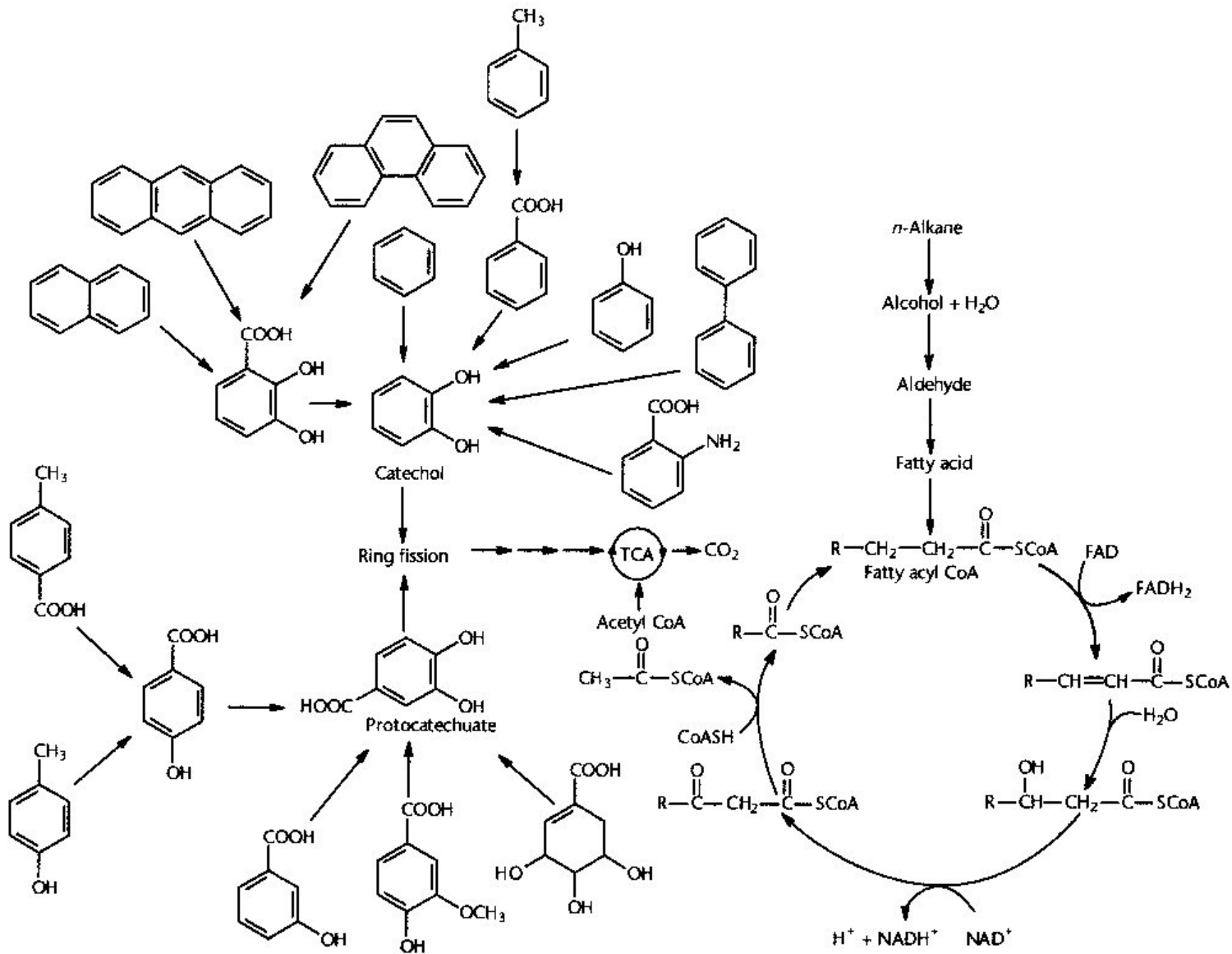


Cometabolism

Bacterium uses some other carbon and energy source to partially degrade contaminant (organic aromatic ring compound)



Cometabolism



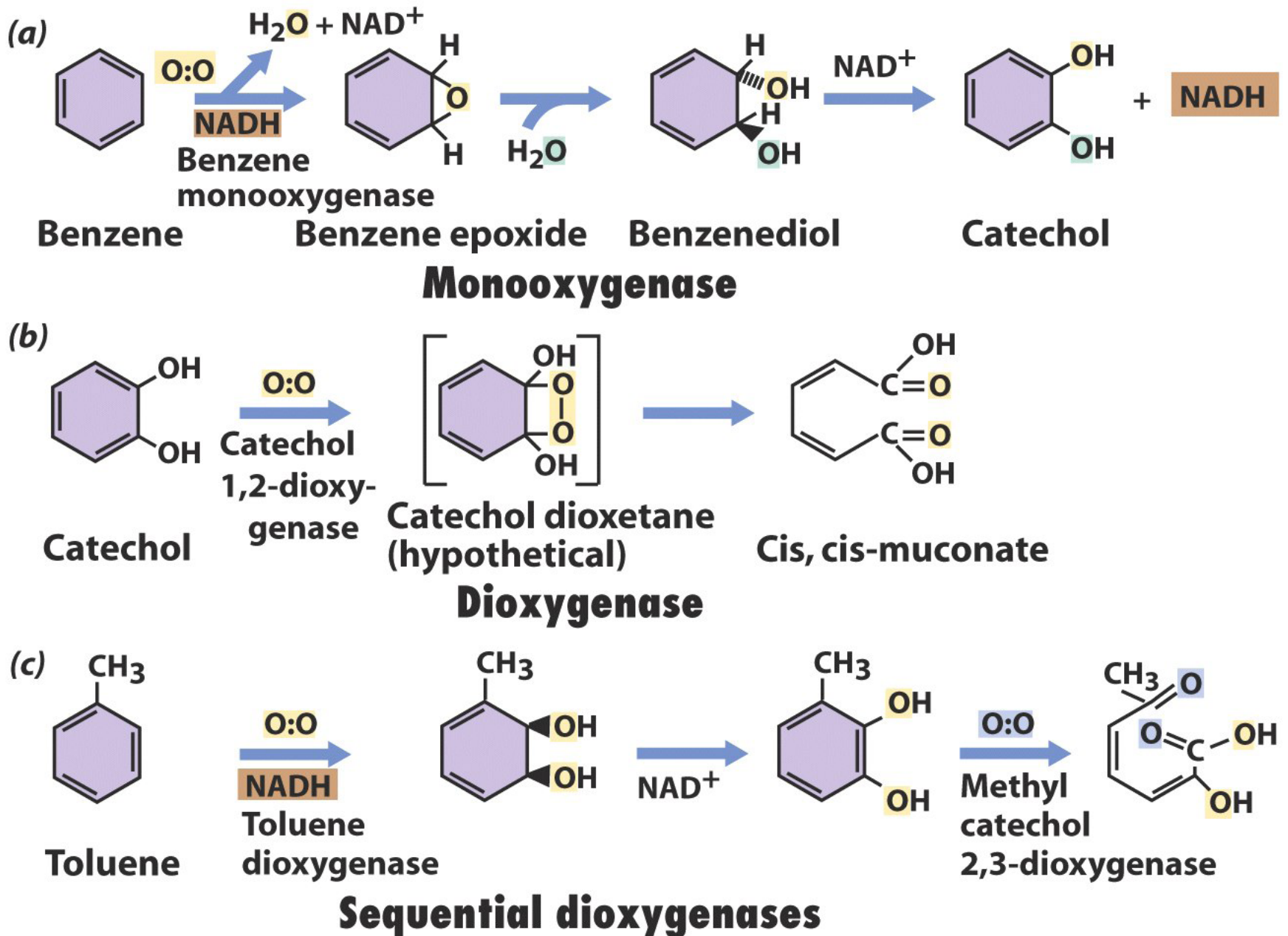


Figure 17-56 Brock Biology of Microorganisms 11/e
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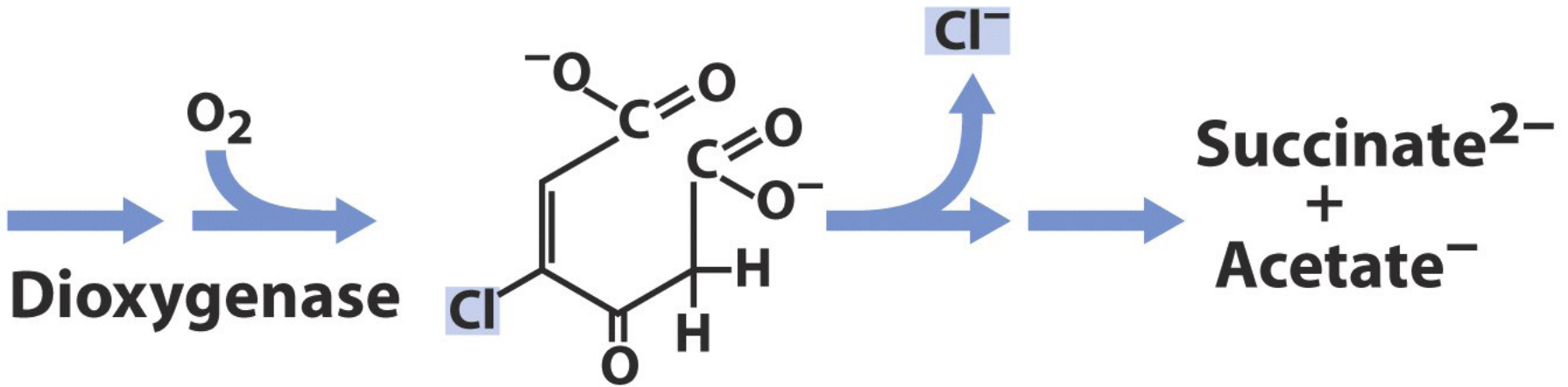
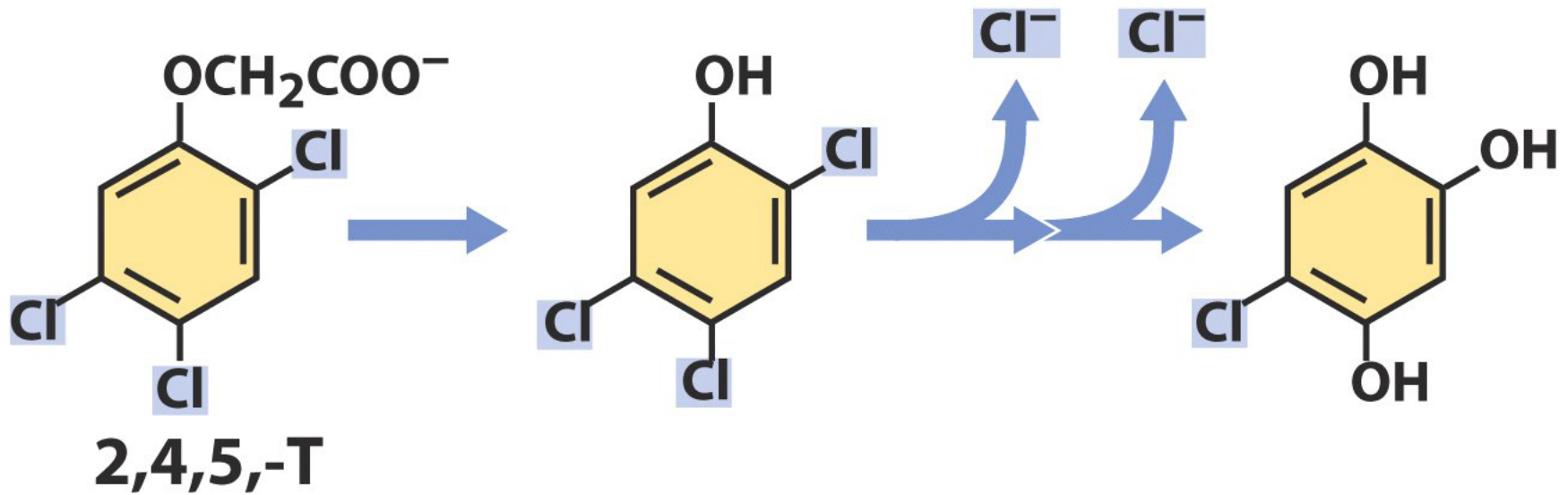


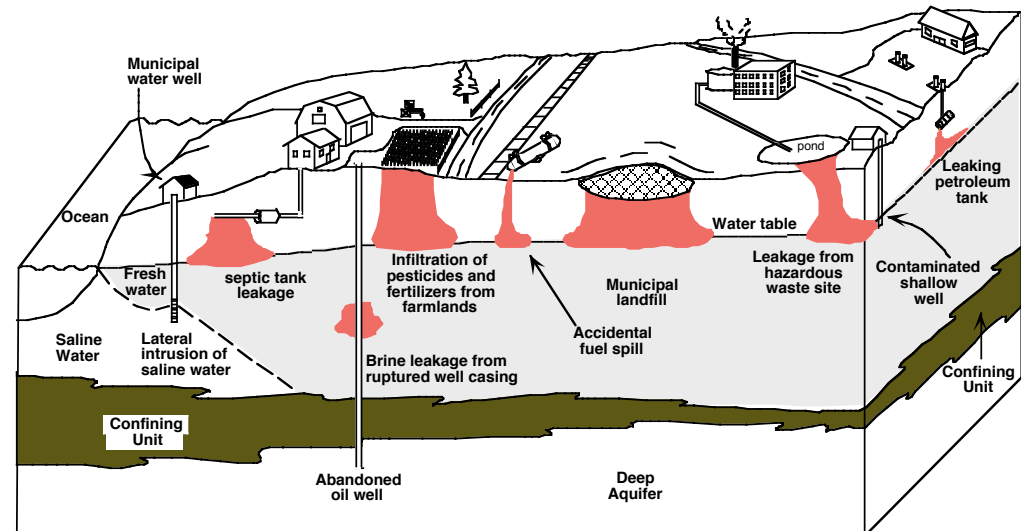
Figure 19-47b Brock Biology of Microorganisms 11/e
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Soil and Subsurface Contaminants

- Benzene and related fuel components (BTEX)
- Pyrene and other polynuclear aromatics
- Chlorinated aromatics and solvents
- Herbicides and pesticides
- Nitroaromatic explosives and plasticizers

Sources of Contamination

- Industrial spills and leaks
- Surface impoundments
- Storage tanks and pipes
- Landfills
- Burial areas and dumps
- Injection wells



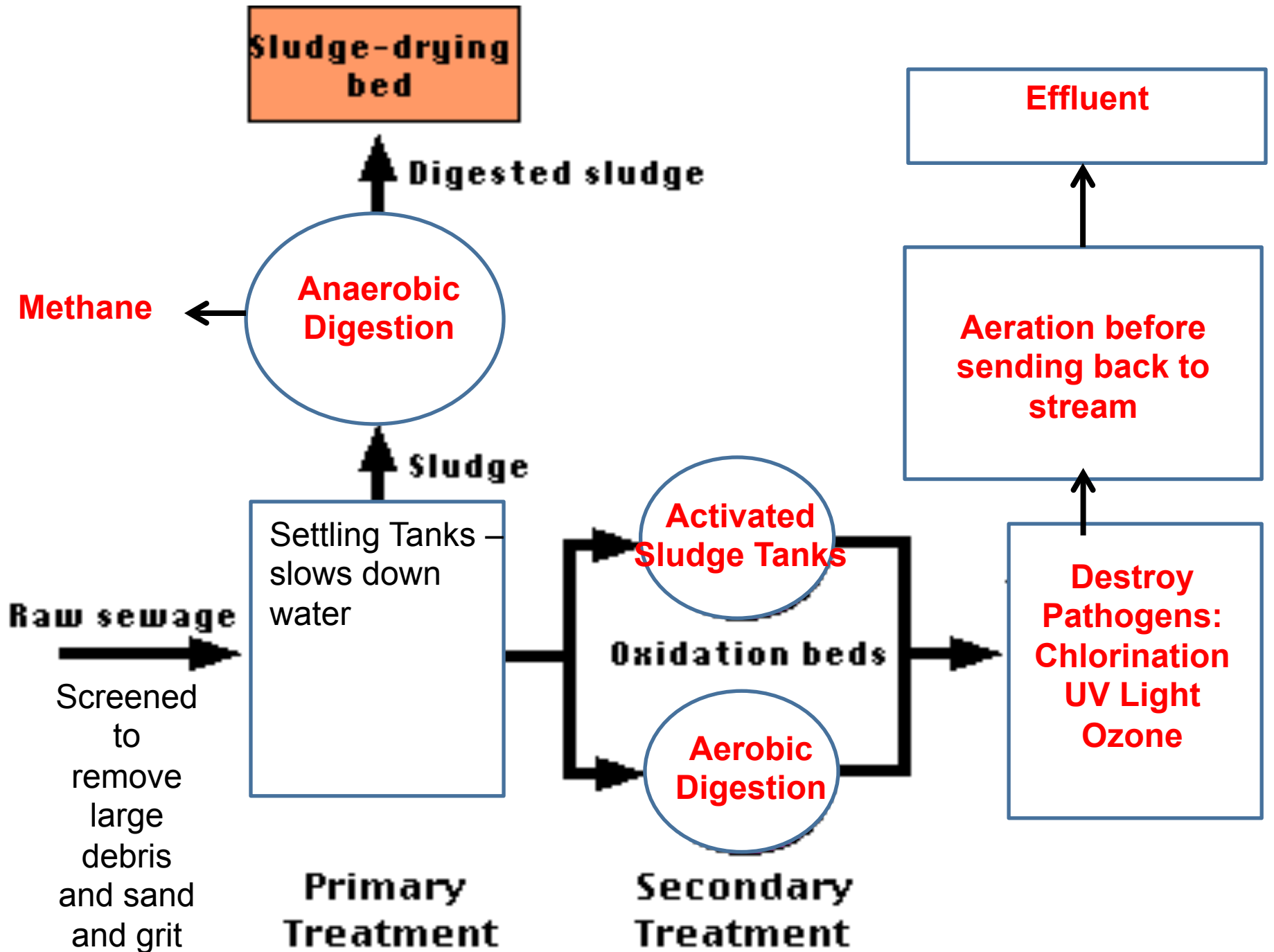
Wastewater Treatment

Treatment depends on three factors:

- 1) Slow water down - removes larger particles
- 2) Aerobic decomposition of organic material: natural bacteria decompose organic material IF enough dissolved oxygen is present in the water
- 3) Destroy pathogens (disease causing bacteria)

Biotic Transformations

- Result of metabolic activity of microbes
- Aerobic and anaerobic biodegradation
- Reduces aqueous concentrations of contaminant
- Reduction of contaminant mass
- Most significant process resulting in reduction of contaminant mass in a system



Wastewater Treatment

Secondary Treatment

- Secondary treatment is a biological process
- Utilizes bacteria and algae to metabolize organic matter in the wastewater



SHARE

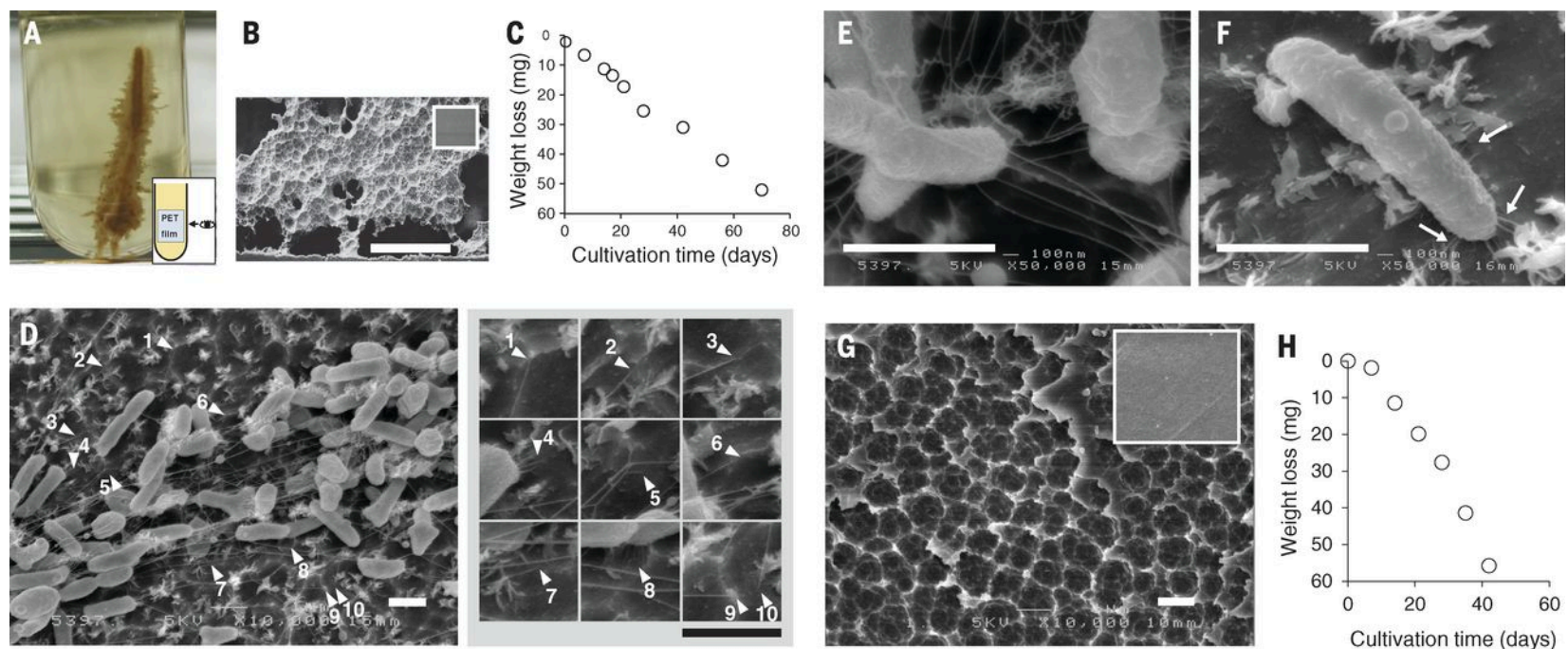
REPORT



A bacterium that degrades and assimilates poly(ethylene terephthalate)

Shosuke Yoshida^{1,2,*}, Kazumi Hiraga¹, Toshihiko Takehana³, Ikuo Taniguchi⁴, Hironao Yamaji¹, Yasuhito Maeda⁵, Kiyotsuna Toyohara⁵, Kenji Miyamoto^{2,†}, Yoshiharu Kimura⁴, Kohei Oda^{1,†}

Fig. 1 Microbial growth on PET. Microbial growth on PET. The degradation of PET film (60 mg, 20 × 15 × 0.2 mm) by microbial consortium no. 46 at 30°C is shown in (A) to (C). The MLE (modified lettuce and egg) medium (10 mL) was changed biweekly. **(A)** Growth of no. 46 on PET film after 20 days. **(B)** SEM image of degraded PET film after 70 days. The inset shows intact PET film. Scale bar, 0.5 mm. **(C)** Time course of PET film degradation by no. 46. PET film degradation by *Ideonella sakaiensis* 201-F6 at 30°C is shown in (D) to (H). The YSV (yeast extract–sodium carbonate–vitamins) medium was changed weekly. **(D to F)** SEM images of *I. sakaiensis* cells grown on PET film for 60 hours. Scale bars, 1 μm. Arrow heads in the left panel of (D) indicate contact points of cell appendages and the PET film surface. Magnifications are shown in the right panel. Arrows in (F) indicate appendages between the cell and the PET film surface. **(G)** SEM image of a degraded PET film surface after washing out adherent cells. The inset shows intact PET film. Scale bar, 1 μm. **(H)** Time course of PET film degradation by *I. sakaiensis*



Shosuke Yoshida et al. Science 2016;351:1196-1199



Fig. 2 ISF6_4831 protein is a PETase. ISF6_4831 protein is a PETase. Effects of PETase on PET film are shown in (A) and (B). (A) SEM image of the treated PET film surface. The inset shows intact PET film. Scale bar, 5 μm . (B) High-performance liquid chromatography spectrum of the products released from the PET film. (C) Unrooted phylogenetic tree of known PET hydrolytic enzymes. (D) Substrate specificity of four phylogenetically distinct PET hydrolytic enzymes (b/a indicates the ratio of the values in the middle-left panel to those in the leftmost panel). (E) Activity of the PET hydrolytic enzymes for highly crystallized PET (hcPET). (F) Effect of temperature on enzymatic PET film hydrolysis.

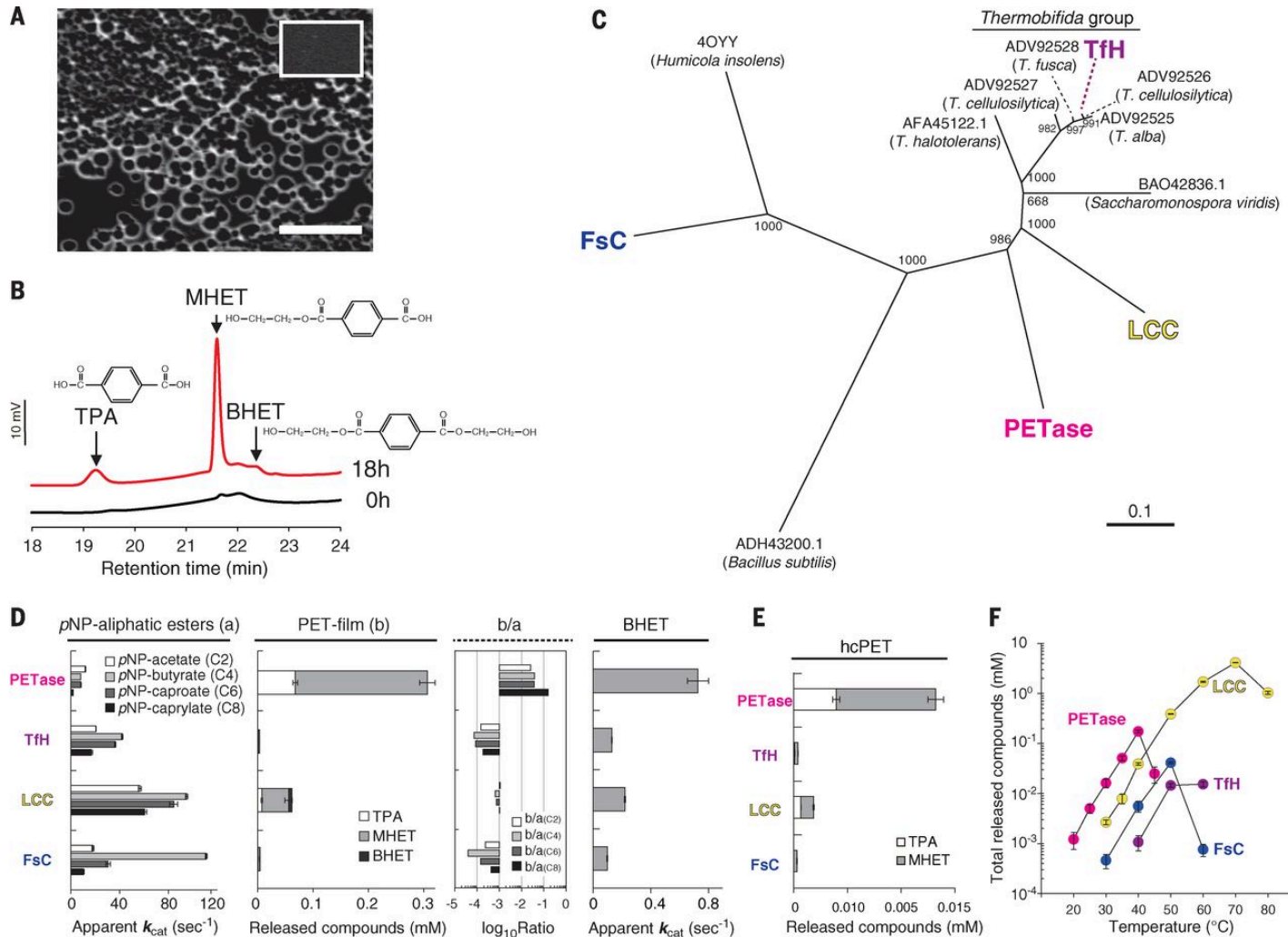
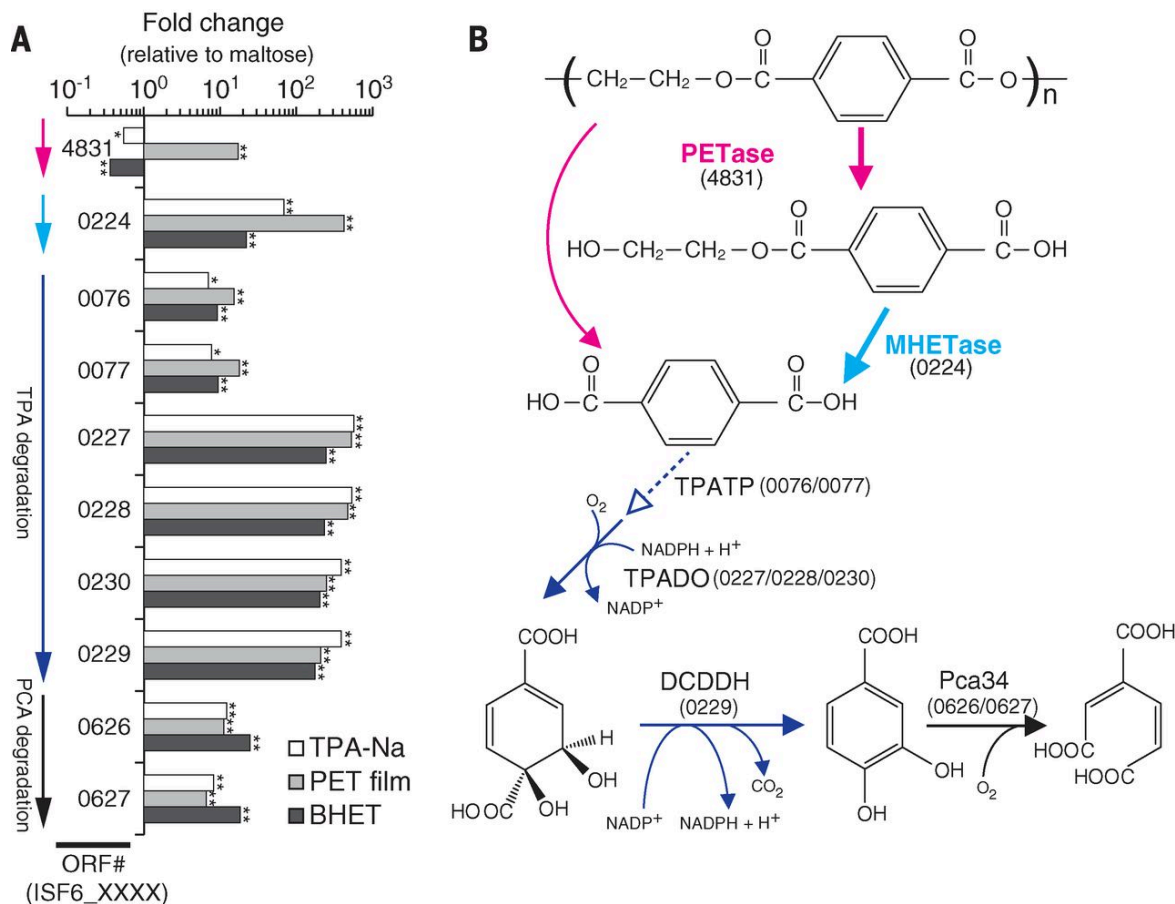
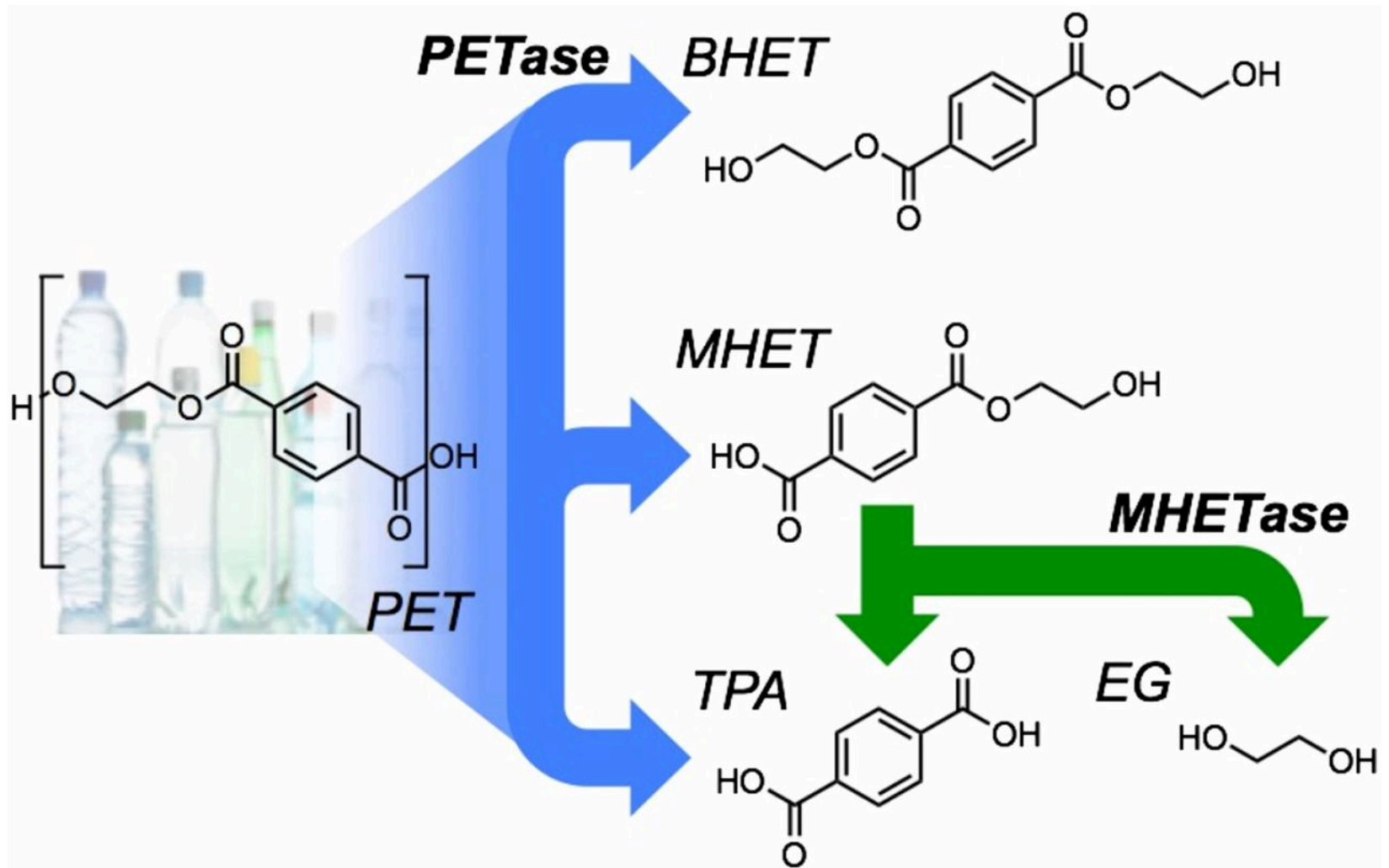


Fig. 3 PET metabolism by *I. sakaiensis*. (A) Transcript levels of selected genes when grown on TPA-Na, PET film, or BHET, relative to those when grown on maltose (B) Predicted *I. sakaiensis* PET degradation pathway. The cellular localization of PETase and MHETase was predicted first (supplementary text, section S1). Extracellular PETase hydrolyzes PET to produce MHET (the major product) and TPA. MHETase, a predicted lipoprotein, hydrolyzes MHET to TPA and EG. TPA is incorporated through the TPA transporter (TPATP) (17) and catabolized by TPA 1,2-dioxygenase (TPADO), followed by 1,2-dihydroxy-3,5-cyclohexadiene-1,4-dicarboxylate dehydrogenase (DCDDH). The resultant PCA is ring-cleaved by PCA 3,4-dioxygenase (Pca34). The predicted TPA degradation pathway is further described in the supplementary text (section S2).



- PETase (PET-digesting enzyme) converts PET to mono(2-hydroxyethyl) terephthalic acid (MHET), with trace amounts of terephthalic acid (TPA) and bis(2-hydroxyethyl)-TPA as secondary products.
- A second enzyme, MHETase (MHET-digesting enzyme), further converts MHET into the two monomers, TPA and ethylene glycol (EG).

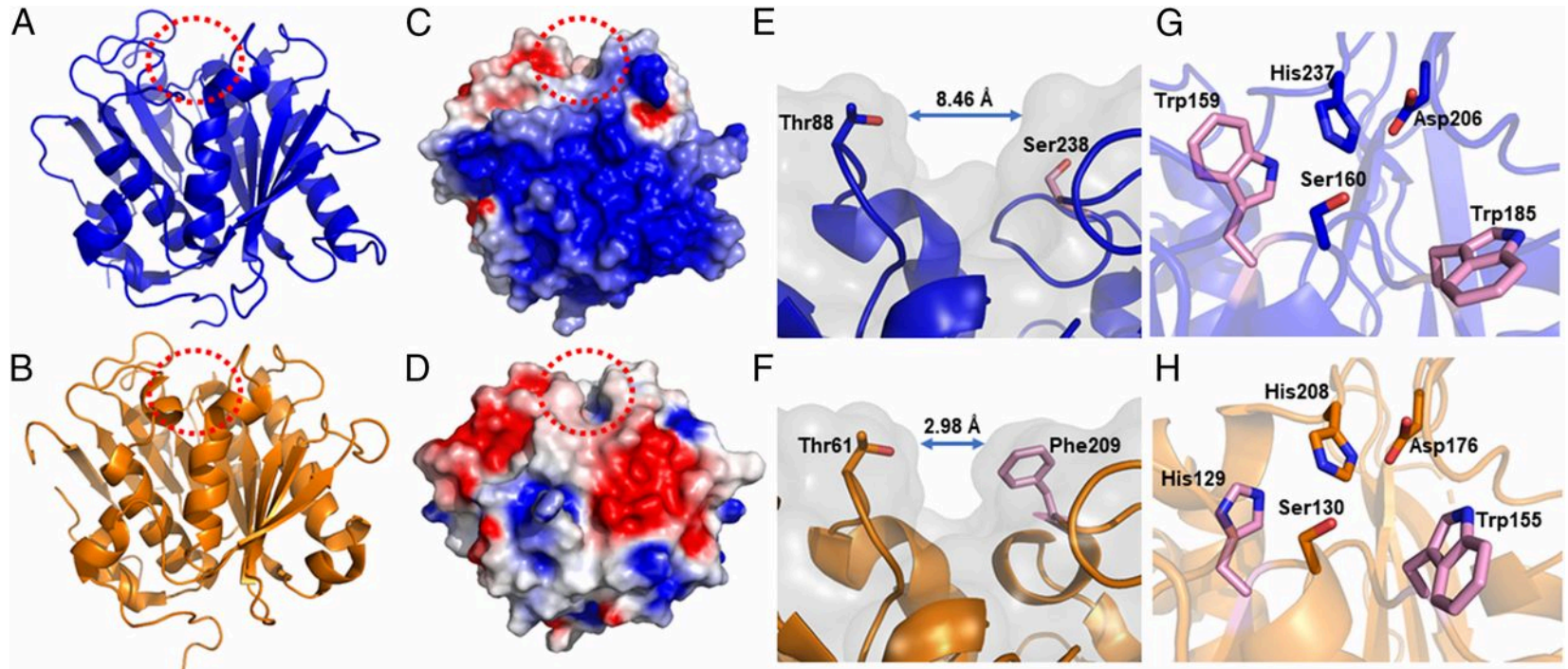
PETase catalyzes the depolymerization of PET to bis(2-hydroxyethyl)-TPA (BHET), MHET, and TPA. MHETase converts MHET to TPA and EG.



Harry P. Austin et al. PNAS 2018;115:19:E4350-E4357

PNAS

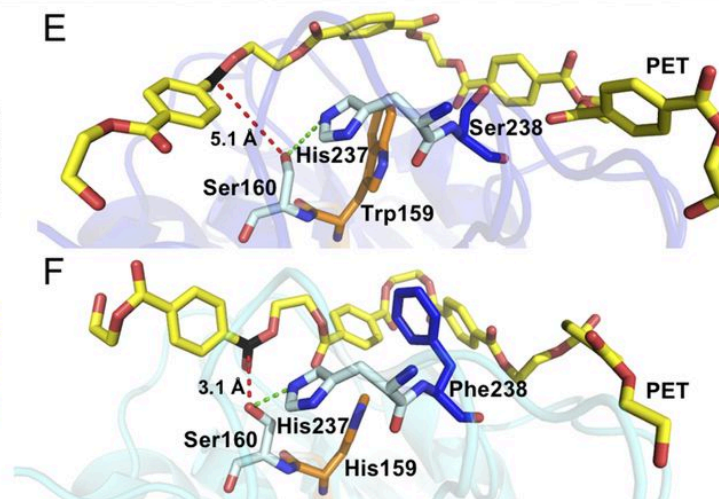
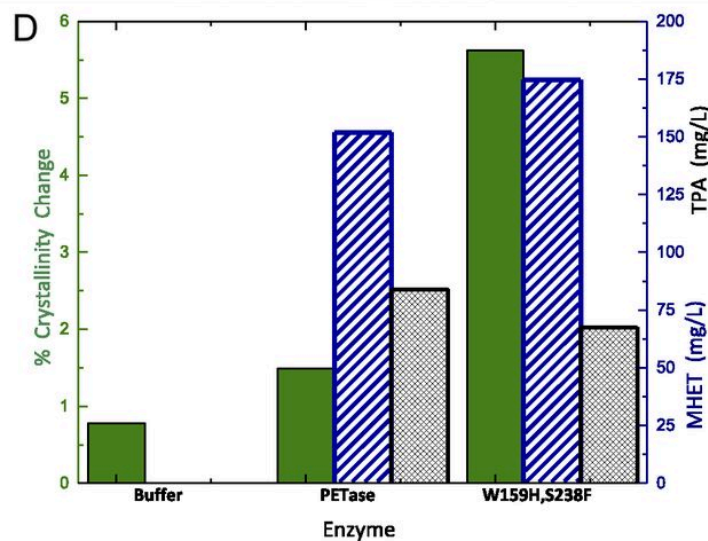
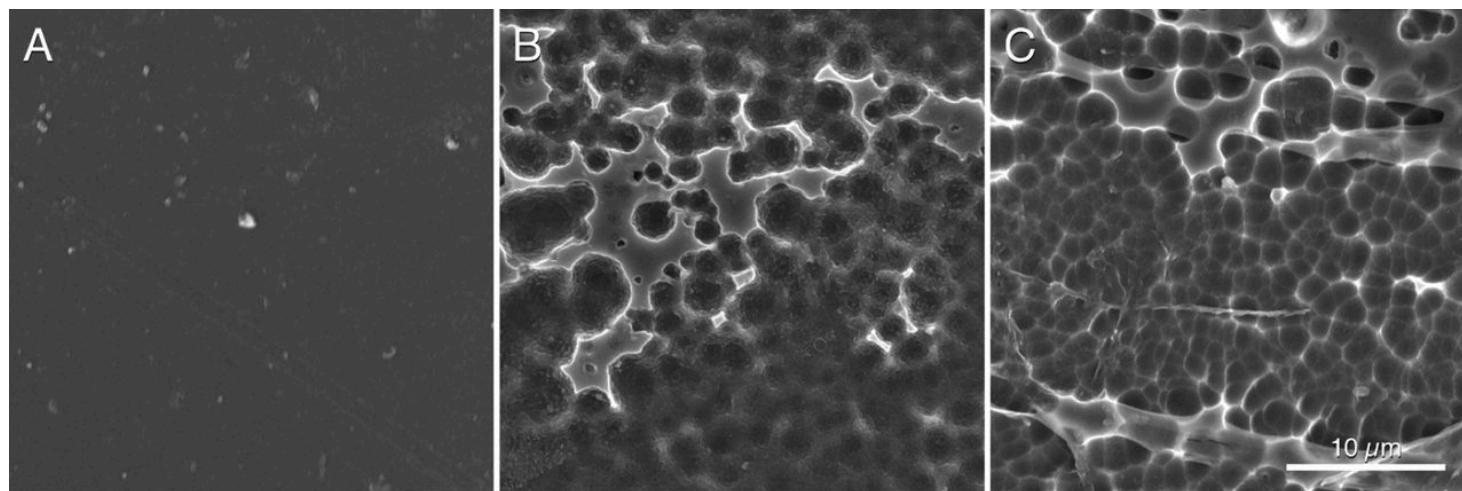
Structure of PETase.



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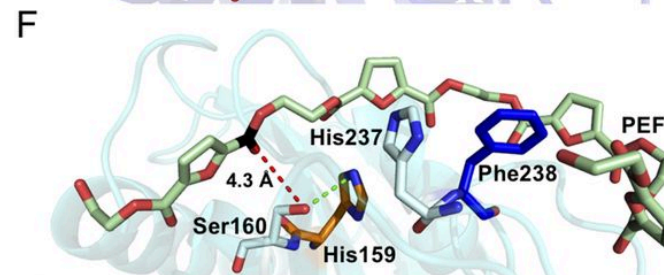
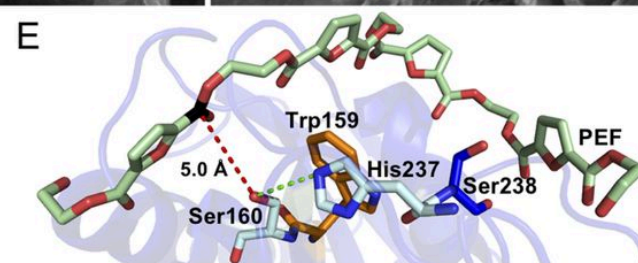
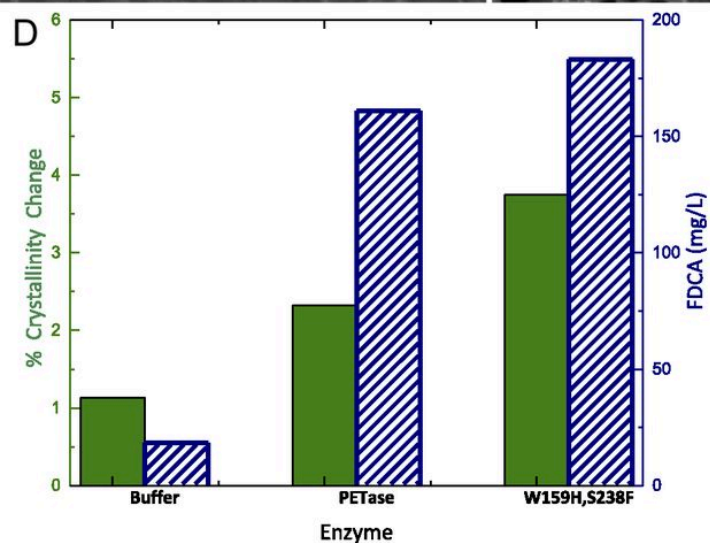
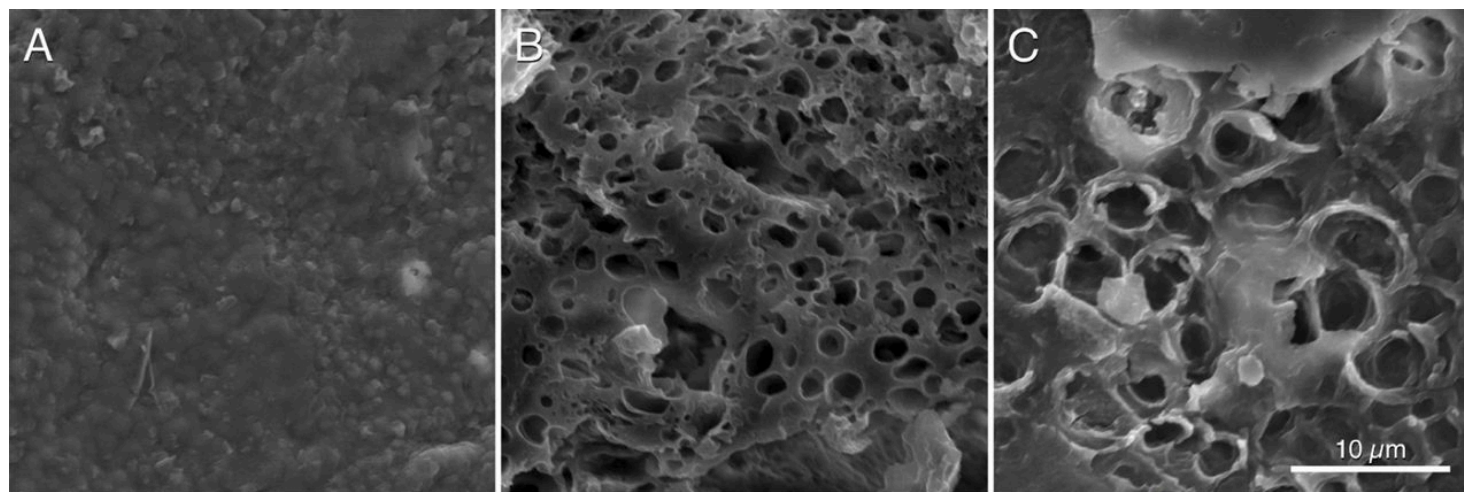
Comparison of PETase and the engineered enzyme S238F/W159H with PET. (A) Buffer-only control of PET coupon.



Harry P. Austin et al. PNAS 2018;115:19:E4350-E4357

PNAS

Comparison of PETase and the engineered enzyme S238F/W159H with PEF. (A) Buffer-only control of PEF coupon.



Harry P. Austin et al. PNAS 2018;115:19:E4350-E4357

PNAS

[3 Biotech](#)

December 2016, 6:200 | [Cite as](#)

Biodegradation of endocrine disruptor dibutyl phthalate (DBP) by a newly isolated *Methylobacillus* sp. V29b and the DBP degradation pathway

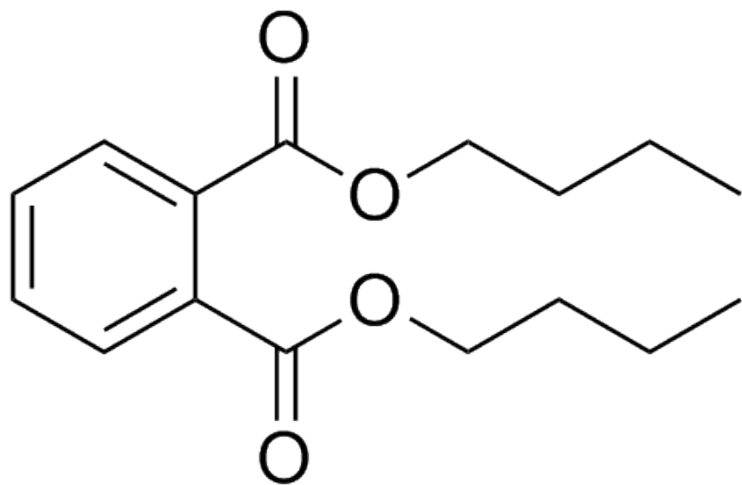
Authors

[Authors and affiliations](#)

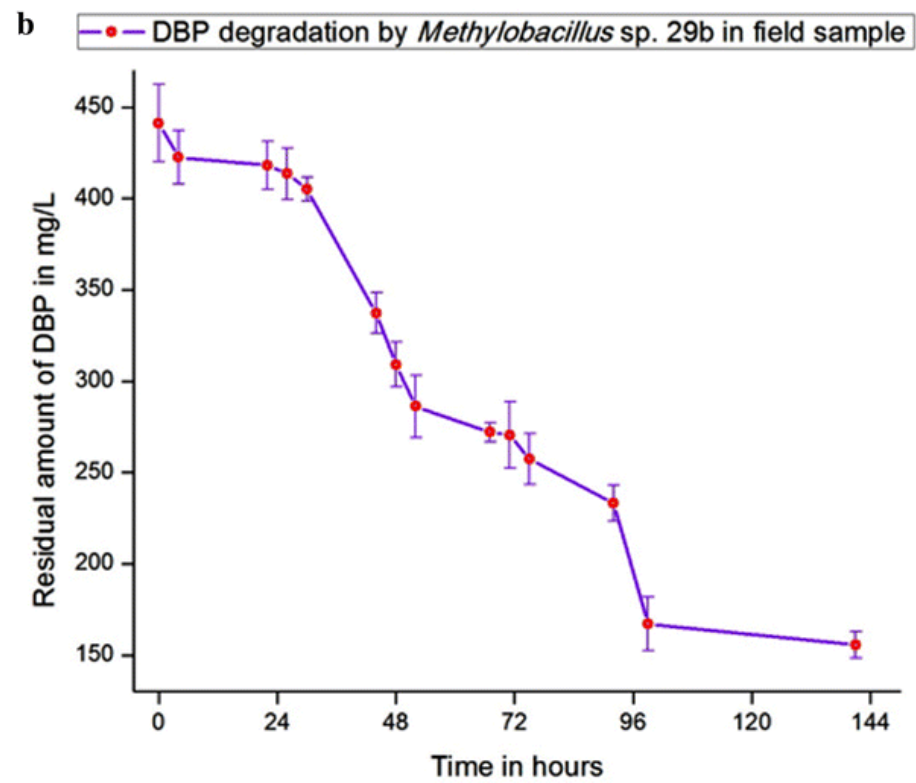
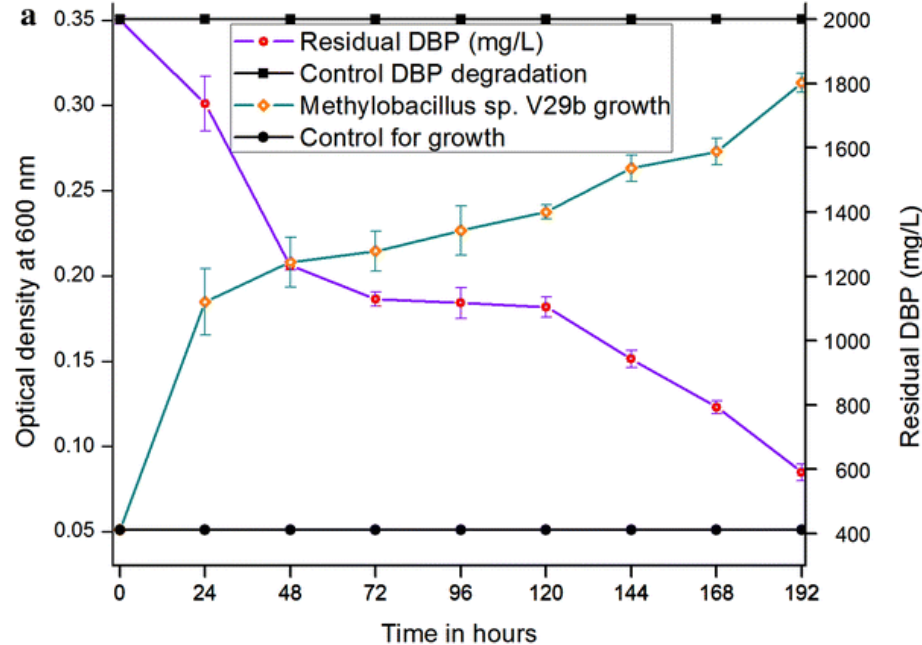
Vinay Kumar , S. S. Maitra

a) Degradation of DBP by *Methylobacillus* sp. V29b in MSM.

b) Degradation of DBP by *Methylobacillus* sp. V29b in the sample contaminated with DBP



DBP



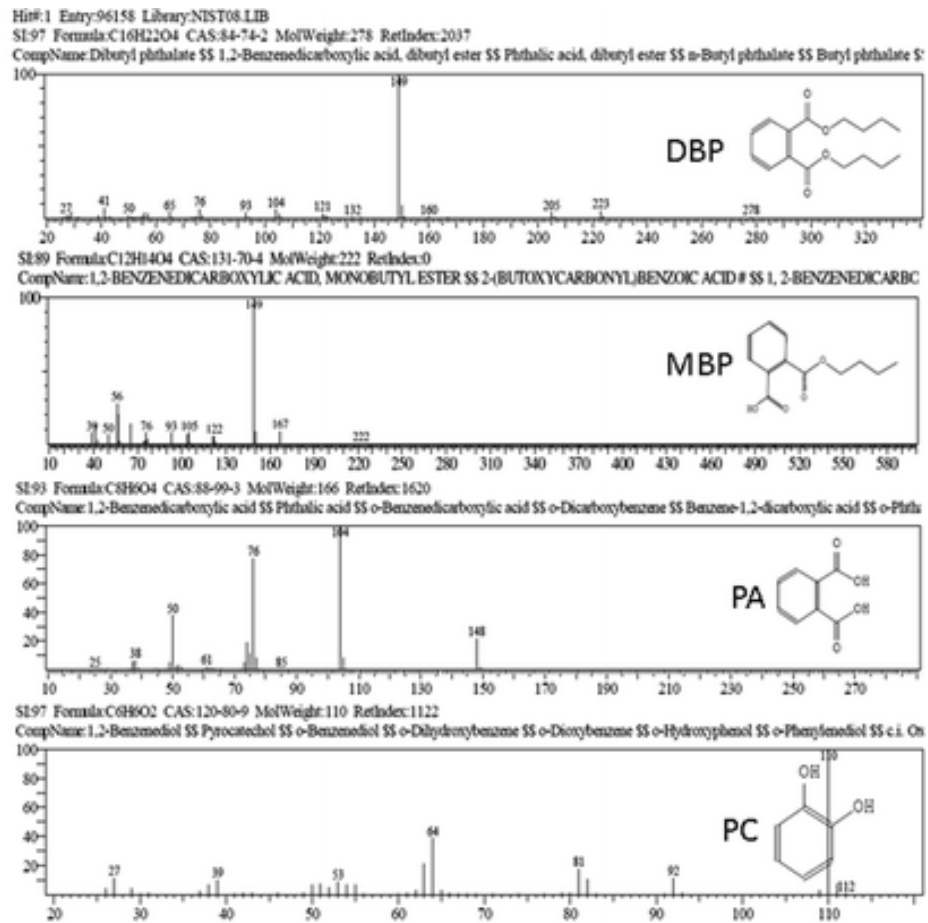
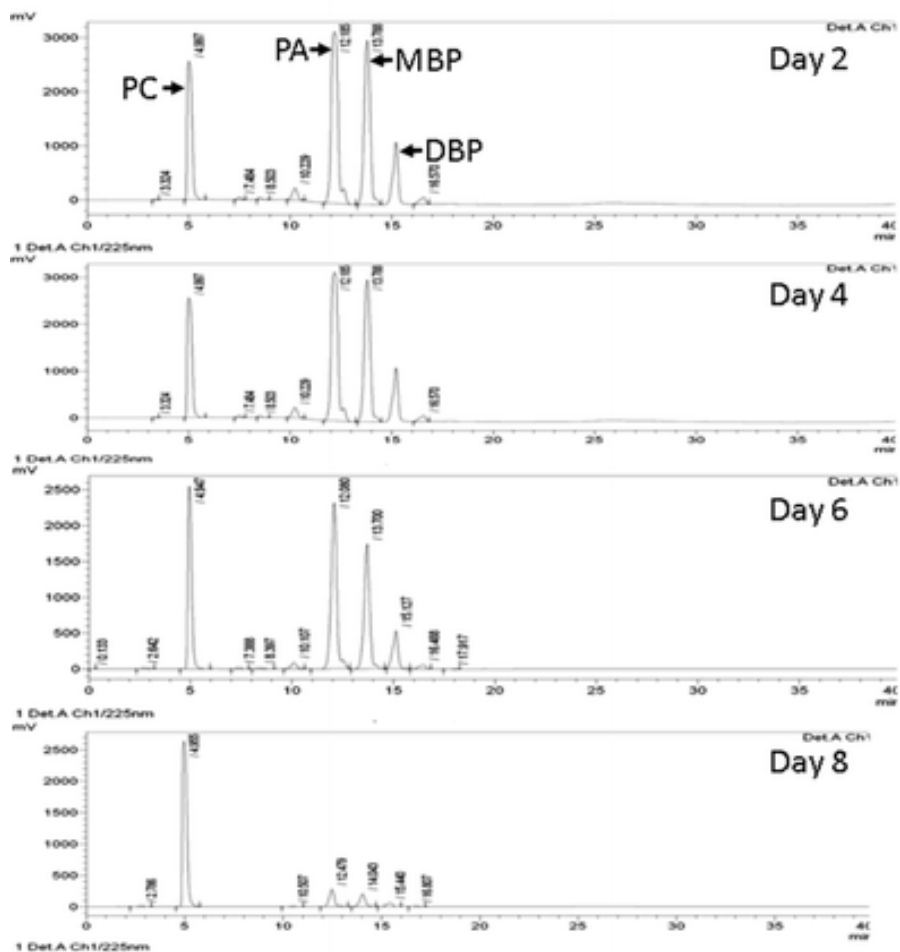


Fig. 5
DBP degradation metabolic intermediates identified by GC-MS. **a** HPLC chromatogram of the metabolic intermediates. **b** Structure and m/z of the identified metabolic intermediates. *DBP* dibutyl phthalate, *MBP* monobutyl phthalate, *PA* phthalic acid, *PC* pyrocatechol

A proposed biochemical pathway for DBP degradation by *Methylobacillus* sp. V29b. *DBP* dibutyl phthalate, *MBP* monobutyl phthalate, *PA* phthalic acid, *PC* pyrocatechol

