

Bioorthogonal non-canonical amino acid tagging

- BONCAT -

BONCAT in microbial ecology, as of September 2020, part I

Hatzenpichler R et al. *In situ visualization of newly synthesized proteins in environmental microbes using amino acid tagging and click chemistry*

Environ Microbiol, 16: 2568-2590 (2014)

- first application of BONCAT to uncultured microbes; development of BONCAT-FISH; correlation of BONCAT with nanoSIMS

Samo TJ et al. *Broad distribution and high proportion of protein synthesis active marine bacteria revealed by click chemistry at the single cell level*

Front Microbiol, 1: 48 (2014)

- application of BONCAT to seawater; correlation of BONCAT with microautoradiography

Hatzenpichler R and Orphan VJ *Detection of protein-synthesizing microorganisms in the environment via bioorthogonal non-canonical amino acid tagging (BONCAT)*

Book chapter for Hydrocarbon and Lipid Microbiology Protocols, Springer Protocols Handbooks, doi 10.1007/8623_2015_61 (2015)

- description of how to design and protocols for how to perform BONCAT-experiments using AHA and HPG

Hatzenpichler R et al. *Visualizing in situ translational activity for identifying and sorting slow-growing archaeal-bacterial consortia*

Proc Natl Acad Sci USA, 113: E4069-E4078 (2016)

- development of activity-based cell-sorting via bioorthogonal labeling (BONCAT-FACS); applied BONCAT-(CARD)FISH and BONCAT-FACS to deep-sea sediment consortia catalyzing the anaerobic oxidation of methane with sulfate

Leizeaga et al. *Using Click-Chemistry for Visualizing in Situ Changes of Translational Activity in Planktonic Marine Bacteria*

Front Microbiol, 8: 2360 (2017)

Pasulka AL et al. *Interrogating marine virus-host interactions and elemental transfer with BONCAT and nanoSIMS-based methods*

Environ Microbiol, 20: 671-692 (2018)

- first application of BONCAT to environmental phages and viruses; estimate of marine viral production rates by BONCAT and nanoSIMS

Couradeau et al. *Probing the active fraction of soil microbiomes using BONCAT-FACS*

Nat Comm, 10: 2770 (2019)

- first application of BONCAT to soil samples; reports that a surprisingly high proportion of soil microbes is translationally active

Sebastian et al. *High Growth Potential of Long-Term Starved Deep Ocean Opportunistic Heterotrophic Bacteria*

Front Microbiol, 10: 760 (2019)

Steward et al. *Metabolic Implications of Using BioOrthogonal Non-Canonical Amino Acid Tagging (BONCAT) for Tracking Protein Synthesis*

Front Microbiol, 11:197 (2020)

BONCAT in microbial ecology, as of September 2020, part II

Valentini et al. *Bioorthogonal non-canonical amino acid tagging reveals translationally active subpopulations of the cystic fibrosis lung microbiota*

Nature Comm, 11: 2287 (2020)

- Case study that applies BONCAT and BONCAT-FACS to cystic fibrosis patients' lung microbiomes

Lindivat et al. *Bioorthogonal Non-canonical Amino Acid Tagging Combined With Flow Cytometry for Determination of Activity in Aquatic Microorganisms*

Front Microbiol, 11: 1929 (2000)

Reichart et al. *Activity-based cell sorting reveals responses of uncultured archaea and bacteria to substrate amendment*

The ISME J, accepted (2020)

- Uses BONCAT-FACS to detect changes in single cell activity of a hot spring microbial community incubated in the presence of various growth substrates or under changing physicochemical conditions

BONCAT is a Next-generation physiology approach

Definition: ...any combination of techniques that analyze the phenotype of an individual cell in a microbiome in a non-destructive way, which enables the physical separation of this cell based solely on its phenotype for subsequent, downstream applications

Specifically, BONCAT is a type of substrate analog probing (SAP)

SAP uses molecules that carry either a fluorescence tag or a side group amenable to azide–alkyne click chemistry to obtain information on the overall biosynthetic activity or specific enzymatic function of the cell.

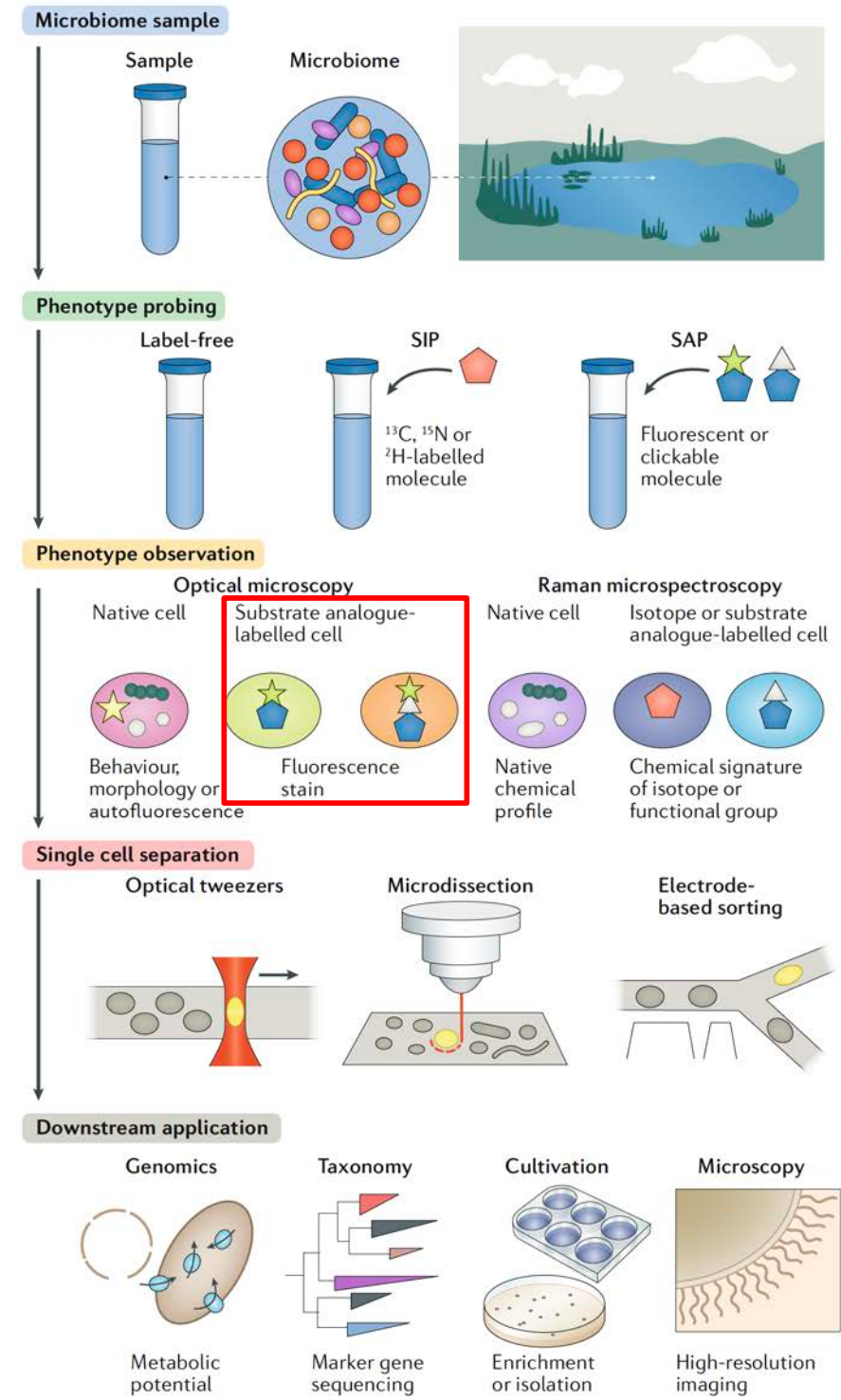


Fig. 2 | Next-generation physiology workflow to study microorganisms.

First, some definitions

bioorthogonal
non-interacting with cellular functionalities

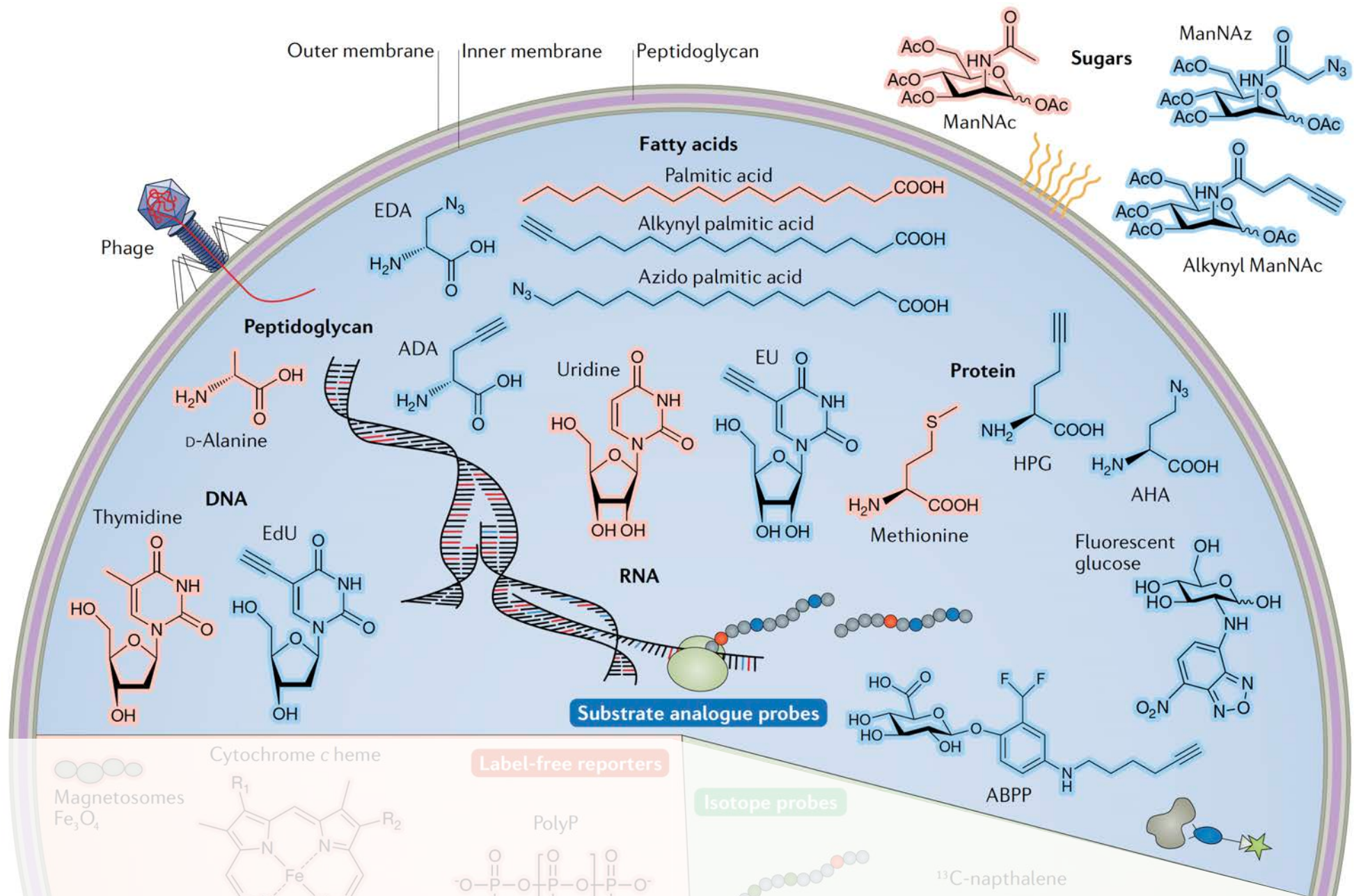
non-canonical
synthetic, not part of biological machinery

Click chemistry

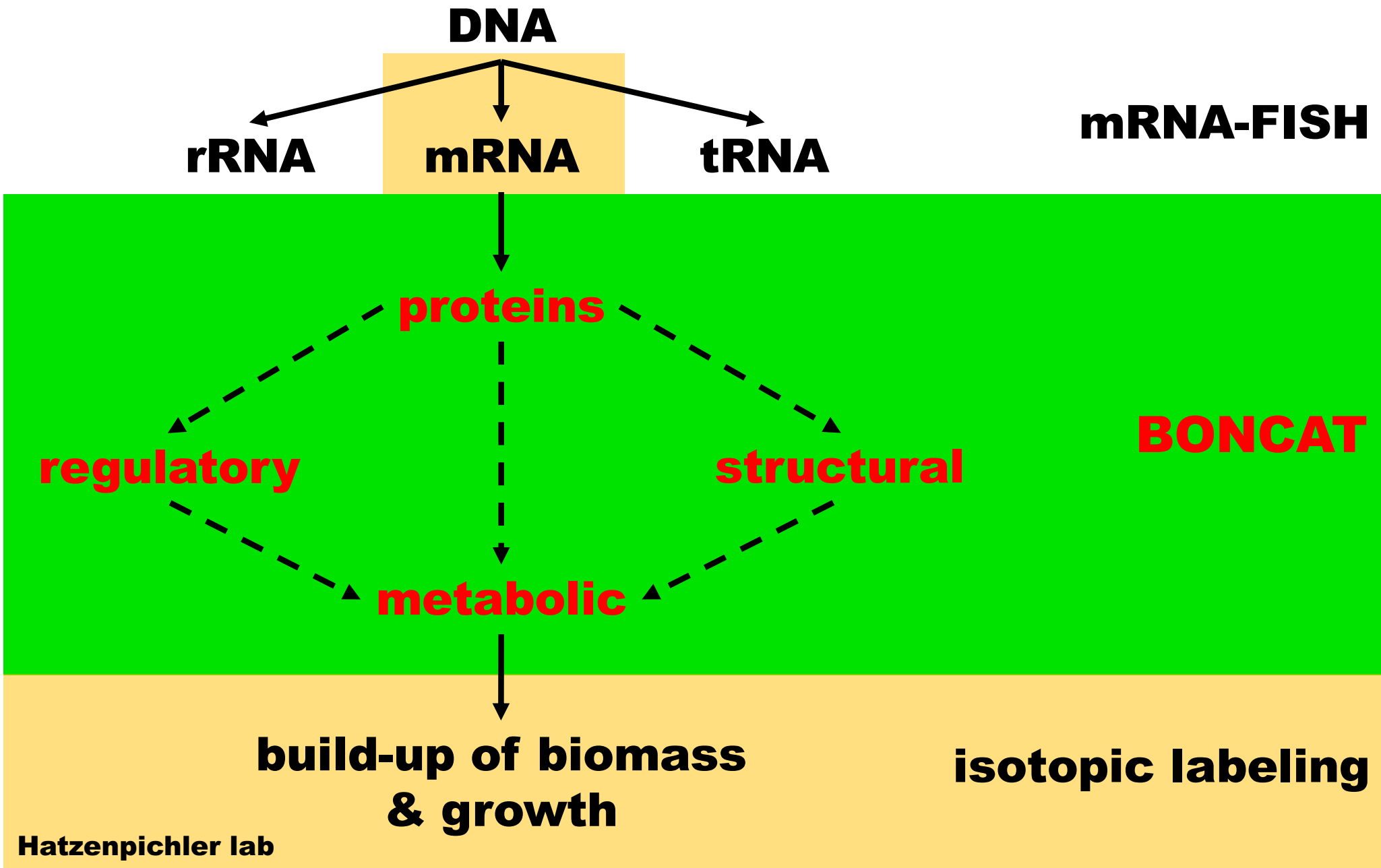
complete conversion of reagents to single product

- + mild conditions**
- + very fast**
- + in water**

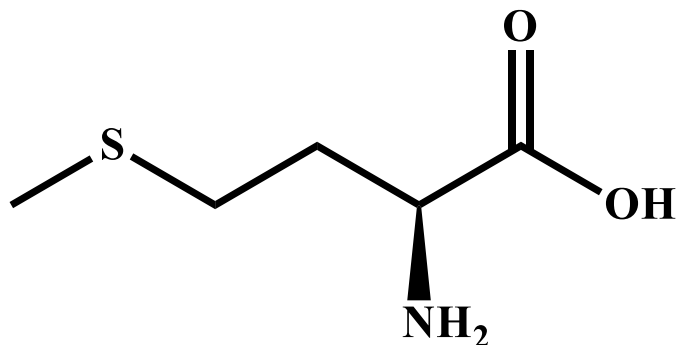
Examples for clickable substrate analogs



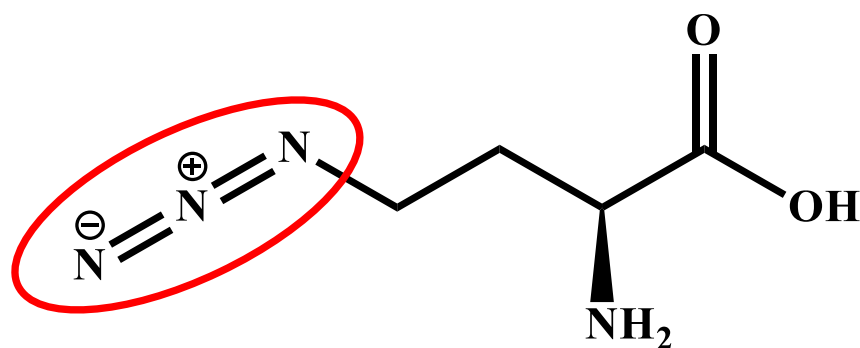
Activity assays on individual cell level



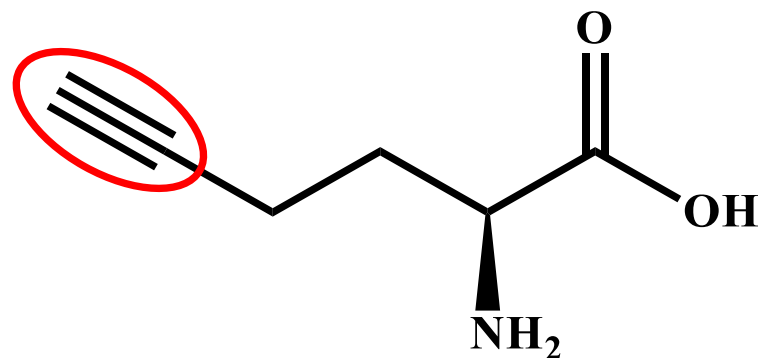
Bioorthogonal non-canonical amino acids



L-Methionine
Met



L-Azidohomoalanine
AHA

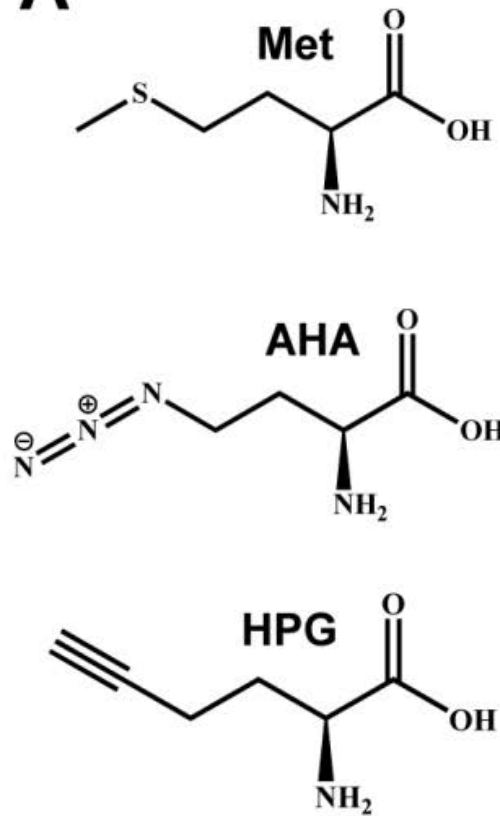


L-Homopropargylglycine
HPG

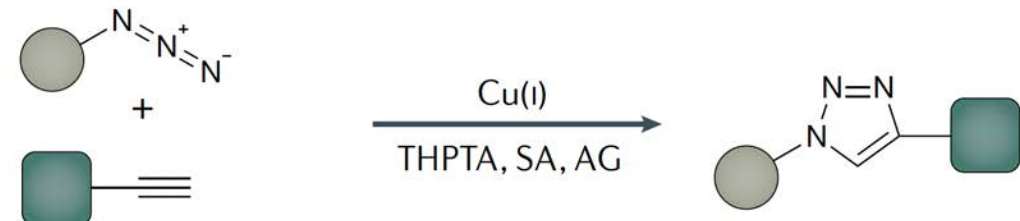
**synthetic amino acids
incorporate into new proteins instead of Met**

Azide-alkyne click chemistry reactions

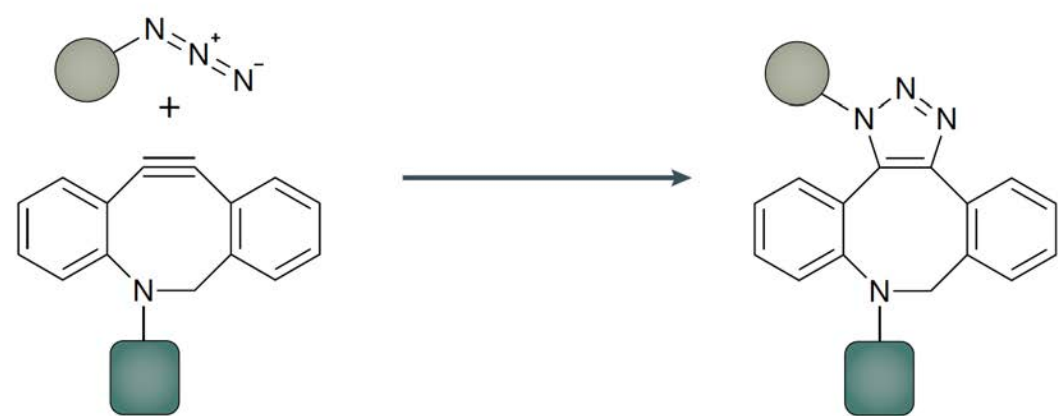
A



b Copper catalysed



c Strain promoted

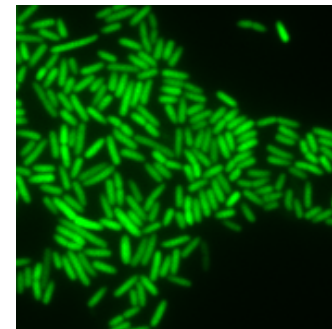
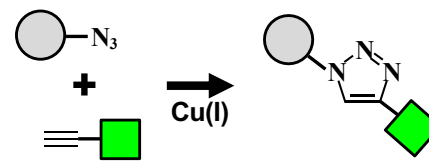
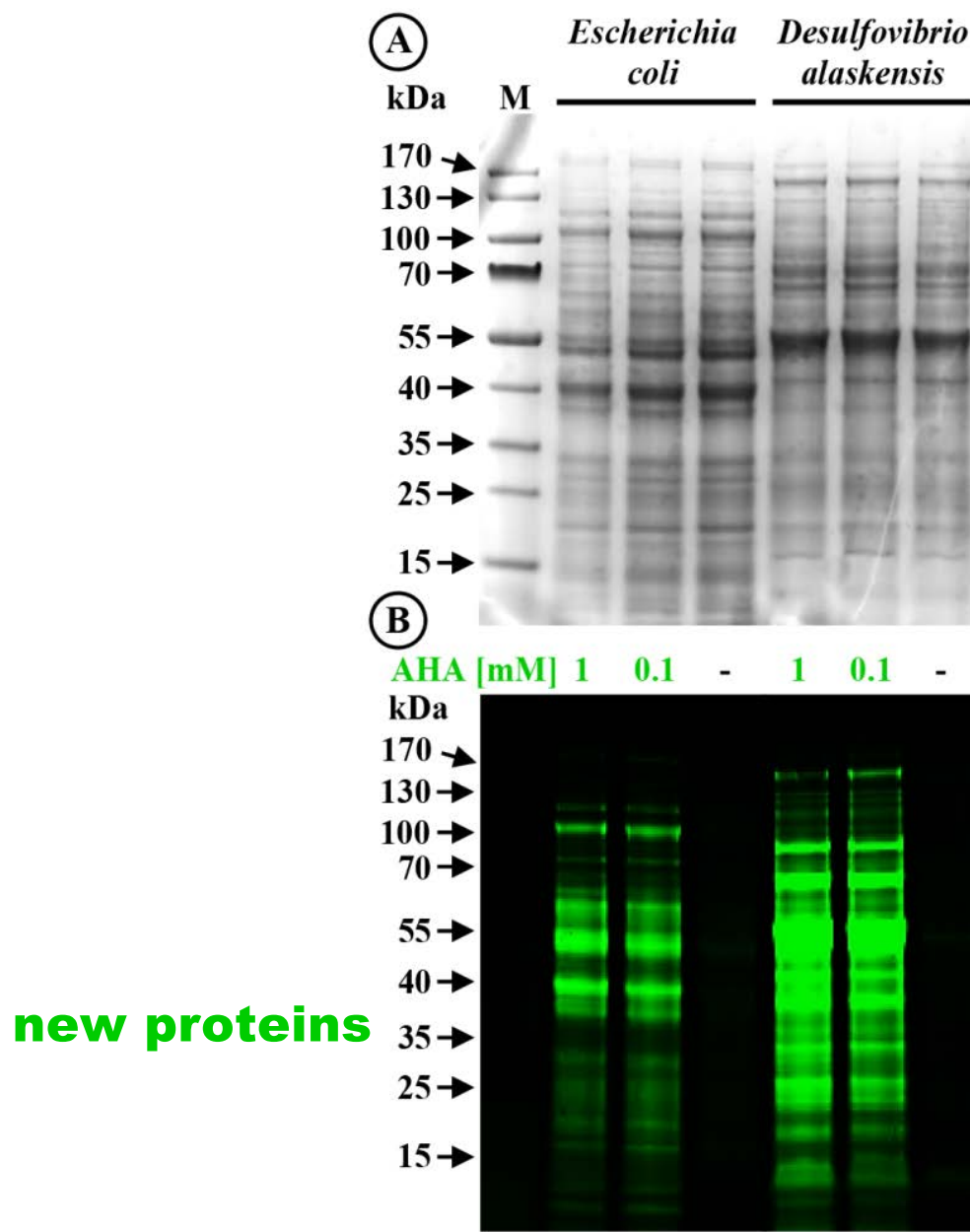


A. Structures of Met and its surrogates AHA and HPG, which compete with Met during translation.

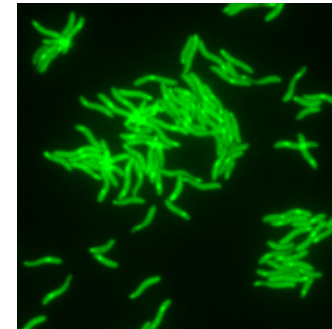
B. In Cu(I)-catalyzed click chemistry an azide group (N_3) is linked to a terminal alkyne residue, yielding a triazole conjugate.

C. Strain-promoted click chemistry allows the copper-less conjugation of an azide group (N_3) with a cyclo-octyne-carrying molecule, yielding a triazole conjugate.

Incorporation into newly made proteins

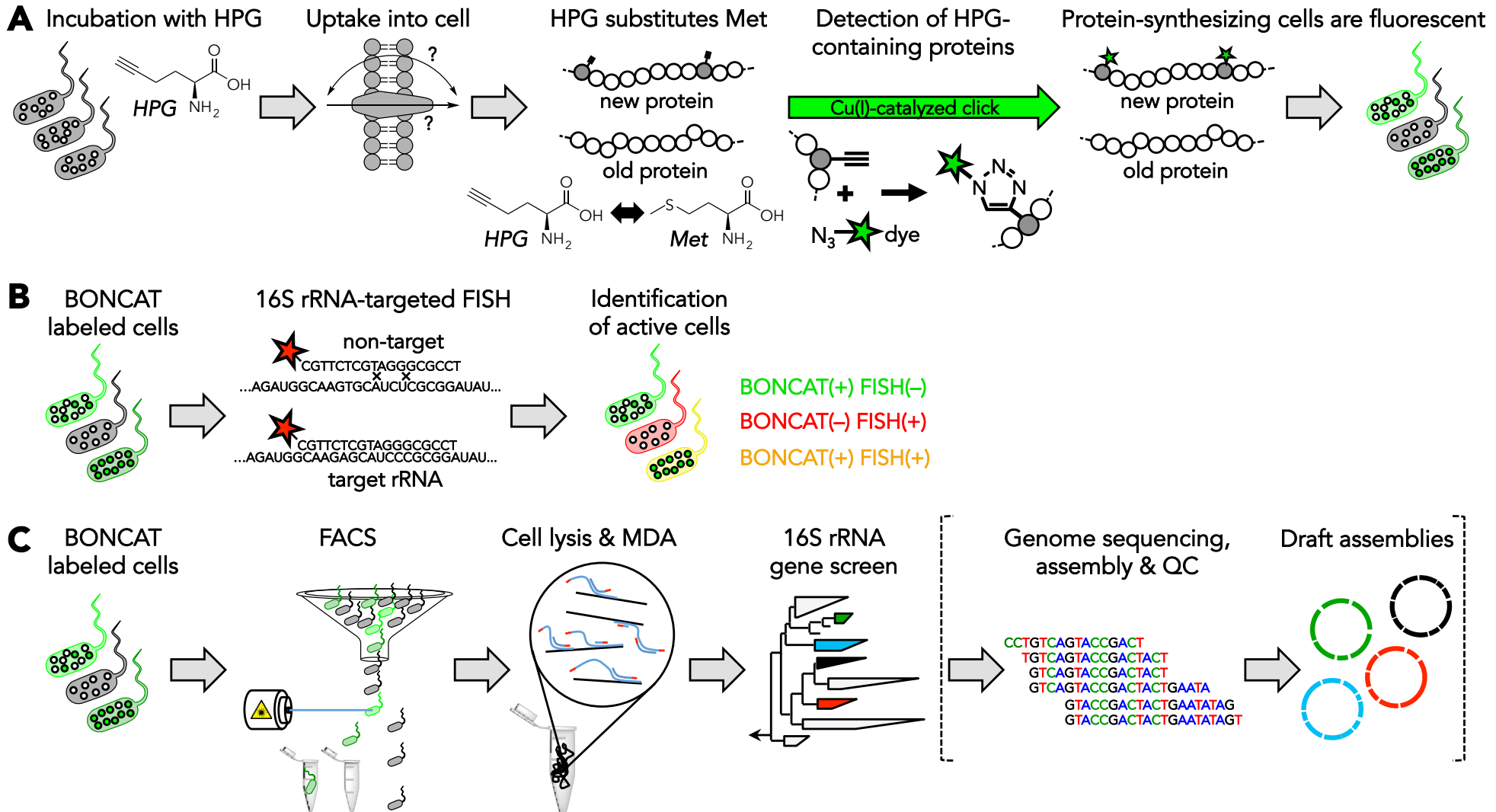


***E. coli*
respiring glucose**

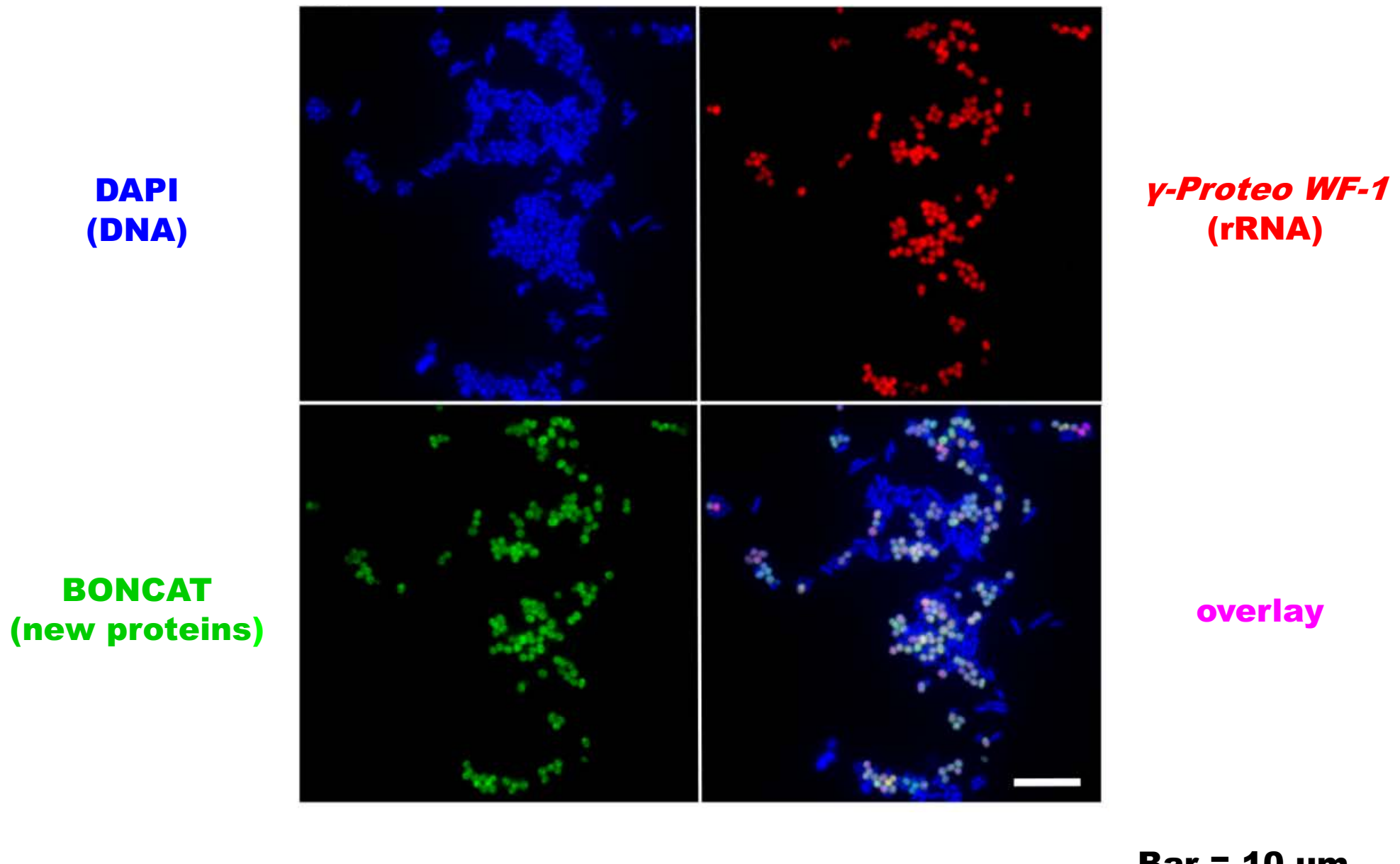


***D. alaskensis*
sulfate reducer**

Visualizing, identifying, and sorting translationally active microbes

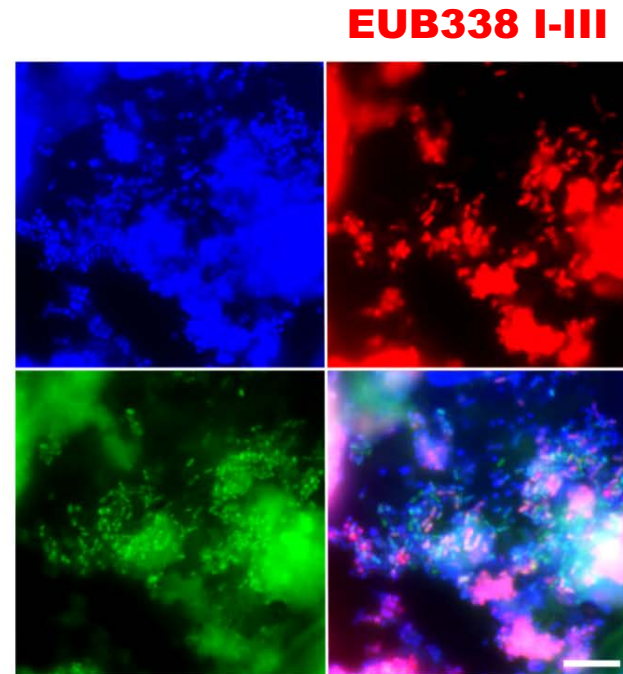
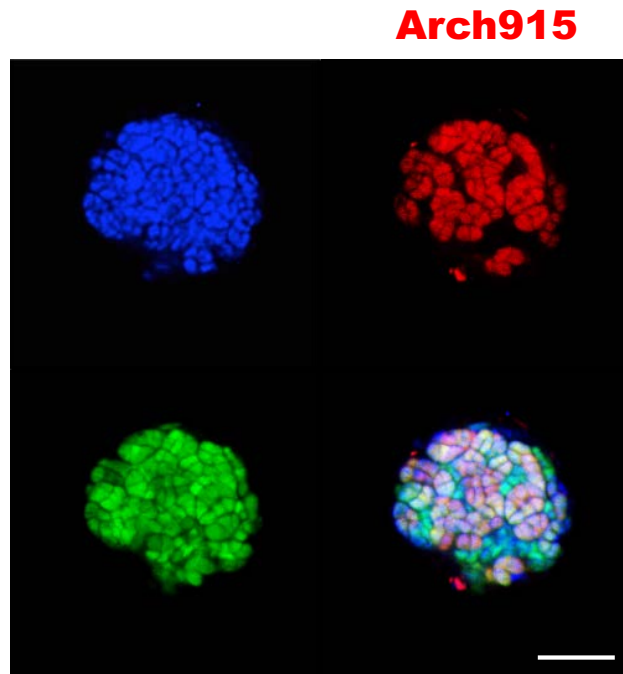


Identification of translationally active cells



Bar = 10 μ m

BONCAT-FISH of uncultured microbes



**Methane seep
ANME-SRB consortium**

Tongue biofilm and saliva

**Freshwater from Lily
pond on Caltech campus**

**DAPI
(DNA)**

**BONCAT
(new proteins)**

**FISH
(rRNA)**

Overlay

Hatzenpichler *et al.*, 2014; Hatzenpichler *et al.*, 2015; Hatzenpichler *et al.*, 2016

Hatzenpichler lab

Visualizing new proteins *in situ*

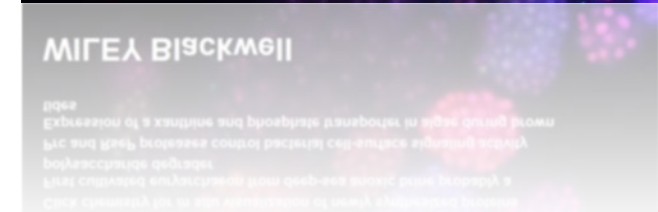
**generally applicable
(works for all taxonomies and
physiologies tested so far)**

**detectable after 2%
of generation time**

**FISH-BONCAT links function
and identity of a cell**

**BONCAT correlates with
 $^{15}\text{NH}_3$ incorporation (nanoSIMS)**

**no change in protein expression
(Bagert *et al.*, 2014)**



Hatzenpichler *et al.*, 2014

Limitations and advantages of BONCAT-FISH

uptake and incorporation

Methionine-rich samples are tough

hard to quantitate amount of new proteins in uncultured cells

potential for cell inactivation or community shifts

links cellular identity and function

fluorescence-based *in situ* activity studies

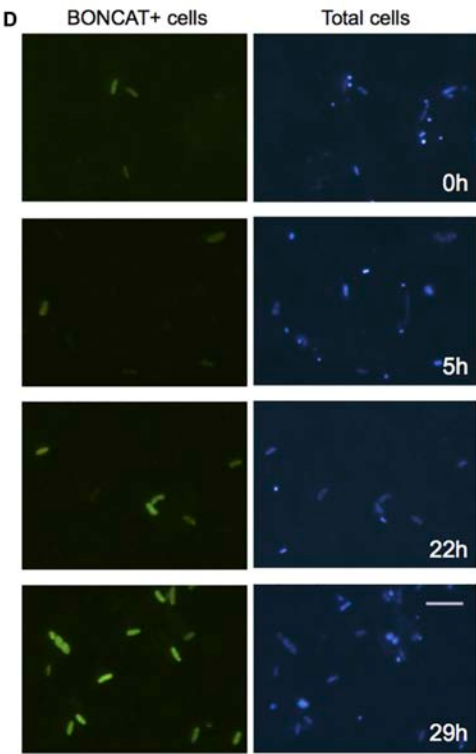
metabolic screening

activity-based cell-sorting

fast + highly selective + cheap + easily available

1 h azide-alkyne ~\$500 epi-scope

Research examples

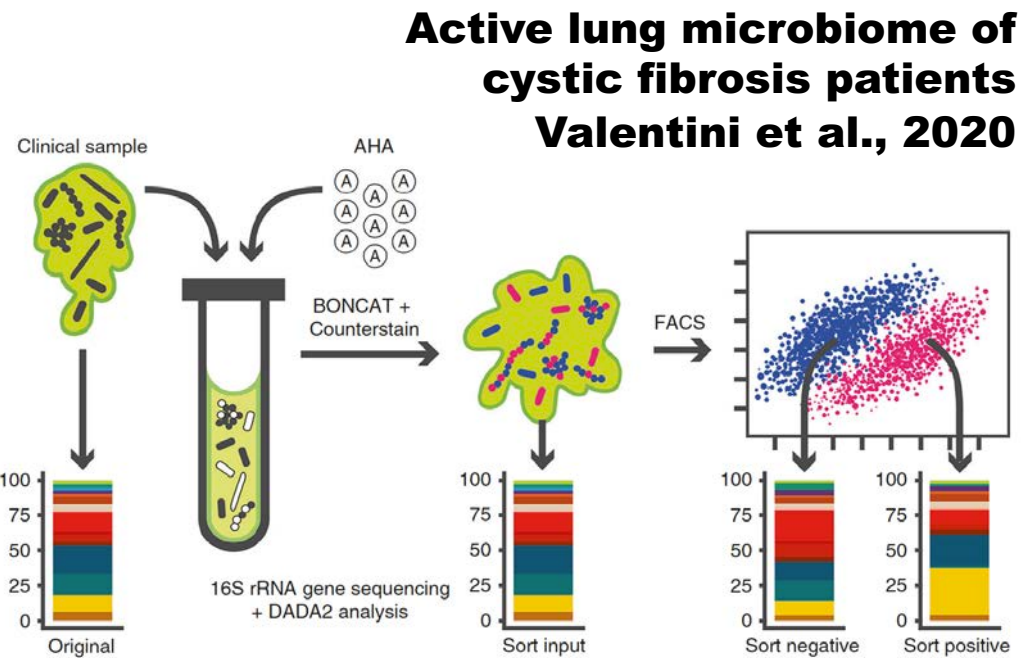
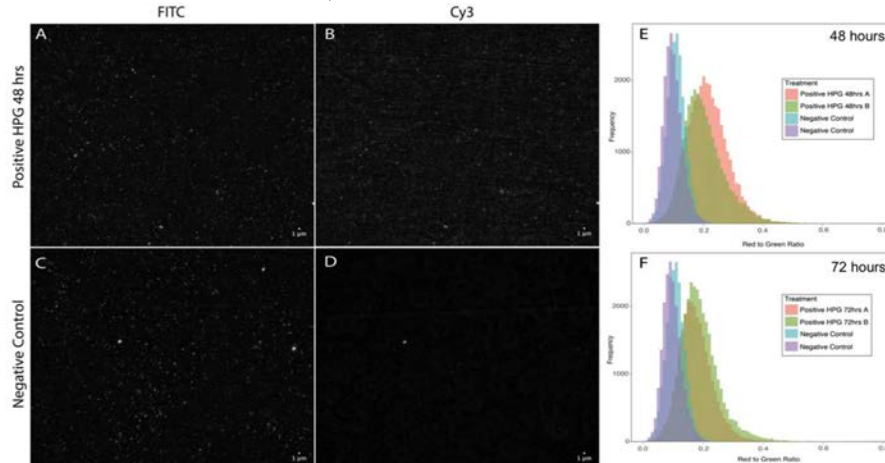


Watching cells resuscitate after long-term starvation

Sebastian et al., 2019

Studying virus turnover in bacterioplankton

Pasulka et al., 2018



Sorting and identifying the active fraction of cells in soil with BONCAT-FACS

Couradeau et al., 2019

