Bioorthogonal non-canonical amino acid tagging - BONCAT -

Hatzenpichler lab www.environmental-microbiology.com doi: 10.13140/RG.2.1.3698.7040/1

First, some definitions

bioorthogonal

non-interacting with cellular functionalities

non-canonical

synthetic, not part of biological machinery

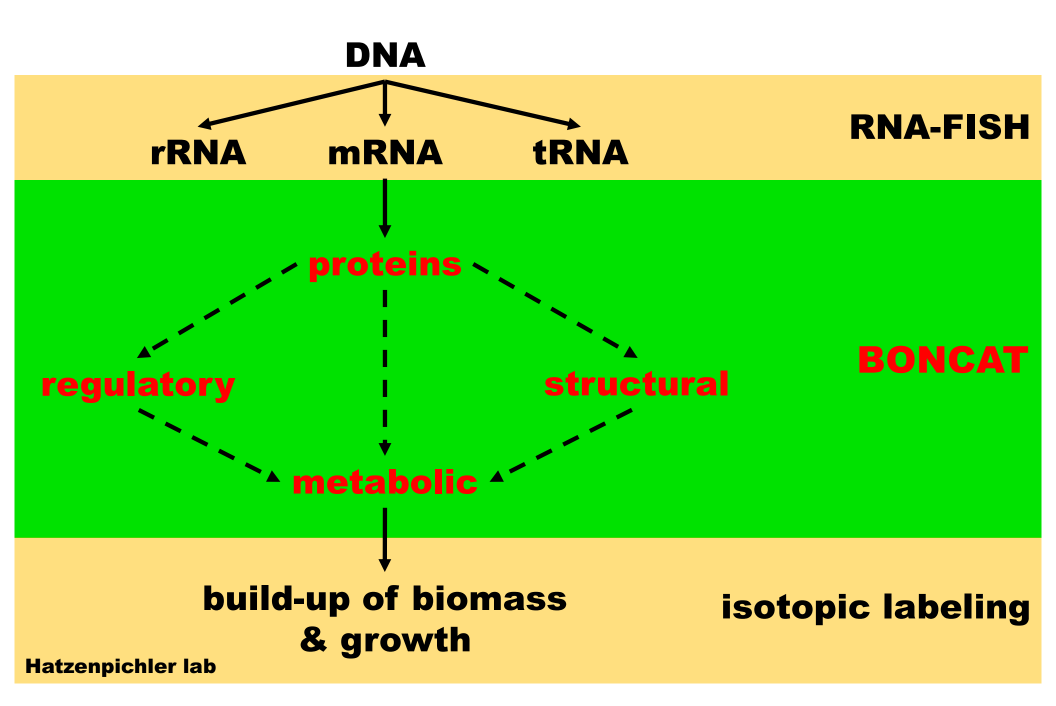
Click chemistry

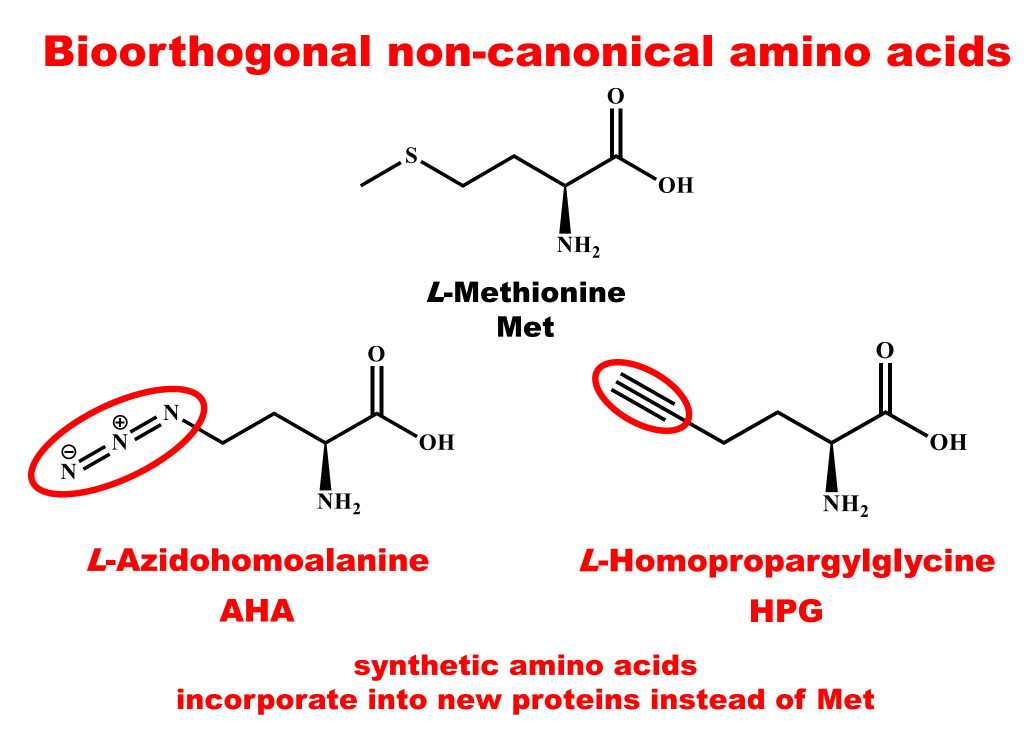
complete conversion of reagents to single product

+ mild conditions

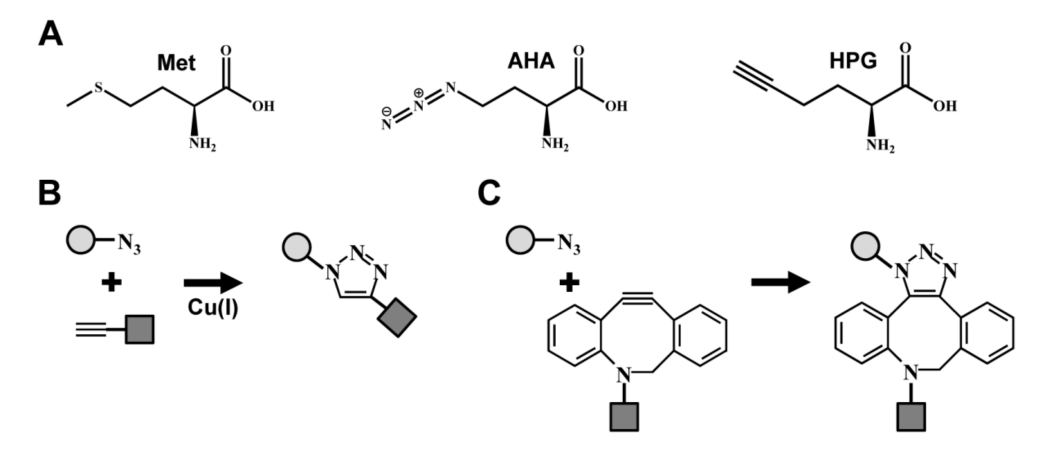
- + very fast
- + in water

Activity assays on individual cell level





Azide-alkyne click reactions



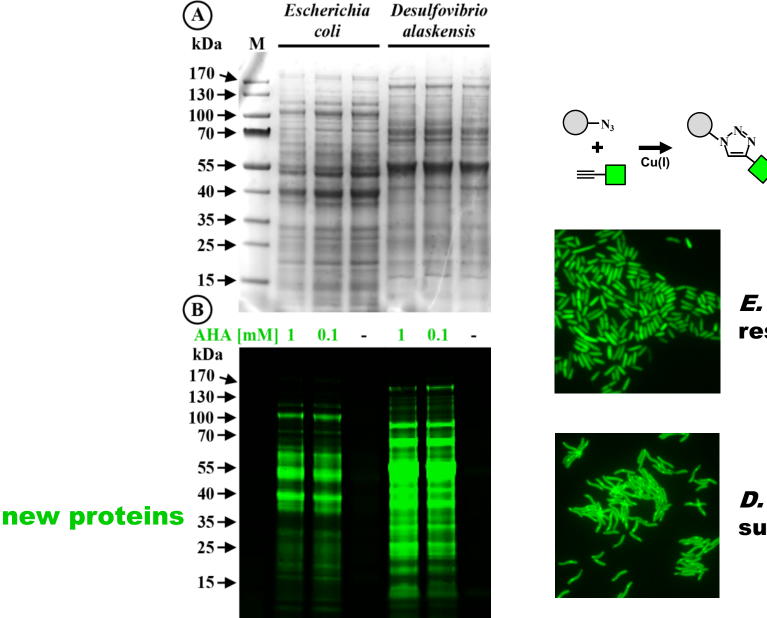
A. Structures of Met and its surrogates AHA and HPG, which compete with Met during translation.

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.

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Incorporation into newly made proteins

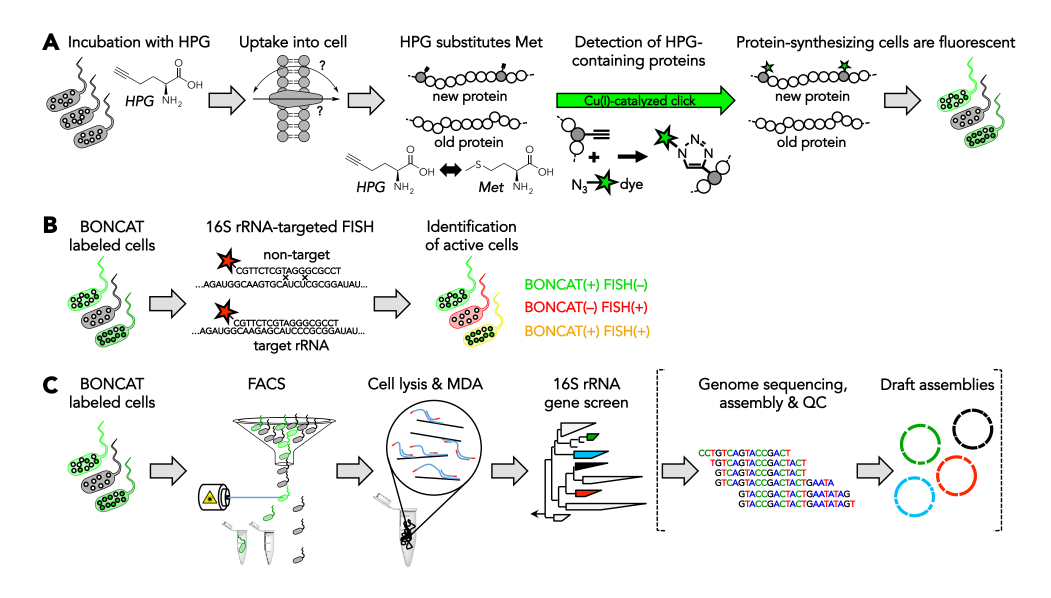


E. coli respiring glucose

D. alaskensis sulfate reducer

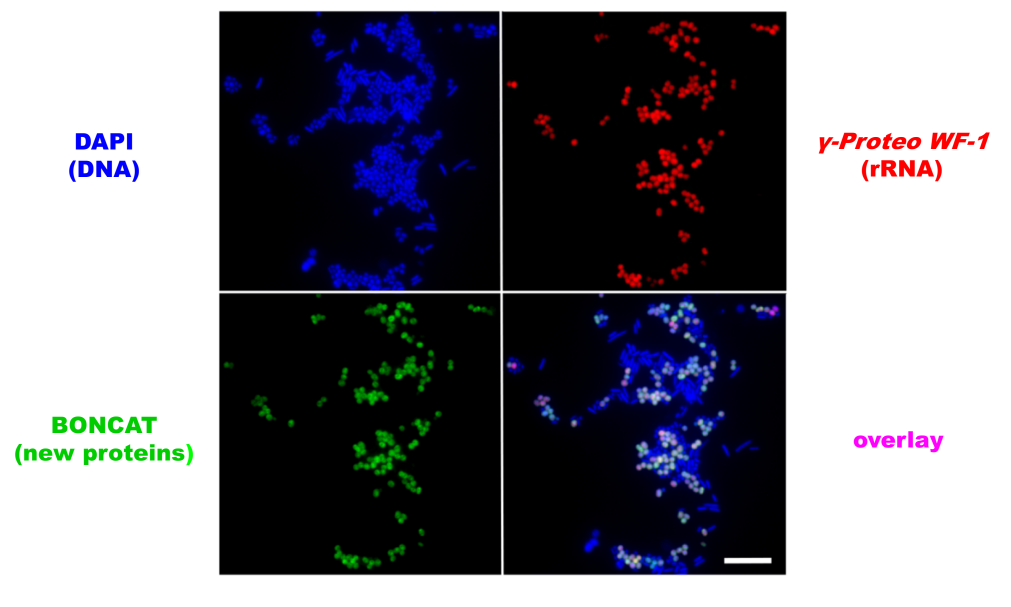
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Visualizing, identifying, and sorting translationally active microbes



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Identification of translationally active cells



Bar = 10 µm

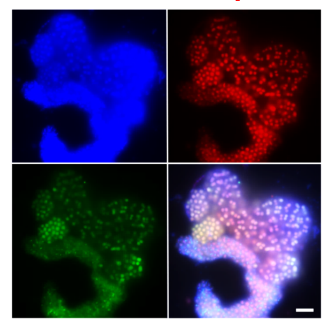
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BONCAT-FISH of uncultured microbes

Arch915

EUB338 I-III

Gam42a + competitor



Methane seep ANME-SRB consortium

Tongue biofilm and saliva

Freshwater from Lily pond on Caltech campus



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

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 ¹⁵NH₃ incorporation (nanoSIMS)

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

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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 available1 hazide-alkyne~\$500epi-scope

BONCAT in environmental microbiology (as of Mar 2018)

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 in the environment; 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 Mar Sci, doi: 10.3389/fmars.2014.00048 (2014)

> application of BONCAT to seawater; correlation of BONCAT with MAR

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, Vol. 7: Single-cell and single-molecule methods Springer Protocols Handbooks, doi 10.1007/8623_2015_61 (2015)

> description of how to design and 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)

application of BONCAT-FISH and BONCAT-FACS to ANME-SRB consortia from three methane seep sediments; development of activity-based cell-sorting via bioorthogonal labeling

Pasulka AL et al.

Interrogating marine virus-host interactions and elemental transfer with BONCAT and nanoSIMS-based methods

Environ Microbiol, DOI: 10.1111/1462-2920.13996 (accepted) 2017

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

Leizeaga et al.

Using Click-Chemistry for Visualizing *in Situ* Changes of Translational Activity in Planktonic Marine Bacteria Front Aquat Microbiol, doi: 10.3389/fmicb.2017.02360 (2017)