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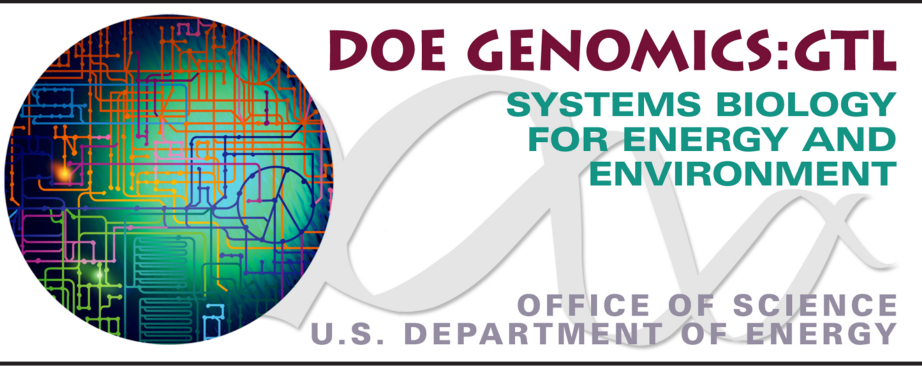
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# Energy Conservation Mechanisms for Syntrophic Growth of Desulfovibrio vulgaris and Methanococcus maripaludis

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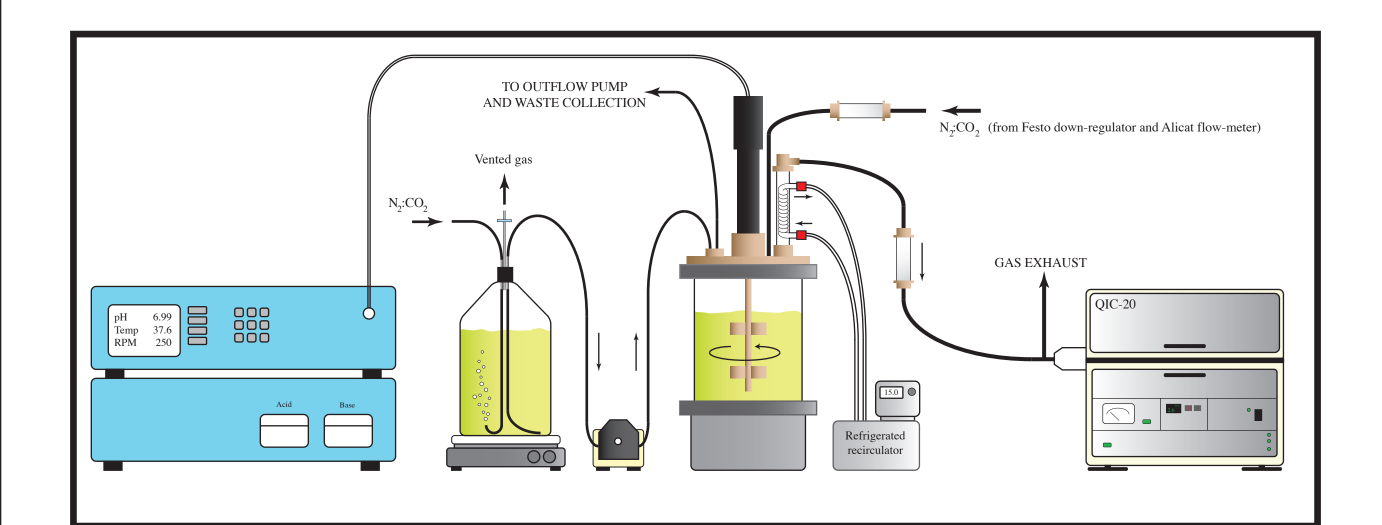
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## **ABSTRACT**

In the absence of electron acceptors, many *Desulfovibrio* species grow on non-fermentable substrates via syntrophic association with hydrogen consuming methanogens. We examined the physiology of *D. vulgaris* Hildenborough growing syntrophically with *Methanococcus maripaludis* LL using a combination of transcriptional and deletion mutant analyses. Syntrophic cocultures were established in chemostats on minimal media amended with lactate but lacking electron acceptor. Replicated whole genome transcriptional analyses identified 169 and 254 genes that were significantly up- or down-regulated, respectively, relative to sulfate-limited monocultures grown at the same generation time. The majority of up-regulated genes were associated with energy production/conservation, signal transduction mechanisms, and amino acid transport/metabolism. A number of the down-regulated genes were associated with signal transduction mechanisms, inorganic ion transport/metabolism and amino acid transport and metabolism. In order to elucidate possible roles of several highly up-regulated genes associated with electron transfer and energy conservation, we constructed mutants of *Desulfovibrio* deleted in a subset of these genes. Cocultures developed with these mutants displayed a range of growth yields, implicating a putative carbon-monoxide induced hydrogenase (Coo, DVU2286-93) and a high-molecular weight cytochrome (Hmc, DVU0531-6) in energy conservation during syntrophic growth. Mutant monocultures grew to the same density on lactate/sulfate as the wildtype. The *cooL* and *hmc* mutants grew significantly slower and to approximately 25% yields of wildtype cocultures. Together, these data suggest a role of these genes in energy conservation of *D. vulgaris* Hildenborough during syntrophic growth.

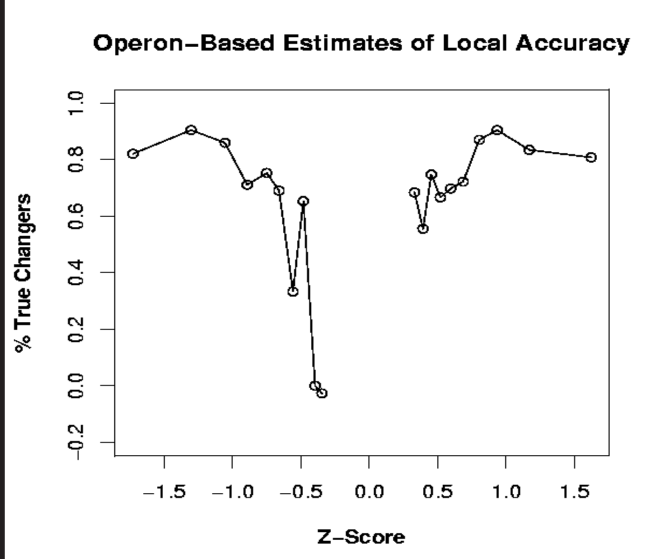
## CHEMOSTAT CONFIGURATION



Chemostats are run using a 24 hr retention time at 37 °C and a stirring speed of 250 rpm. The headspace of the chemostat is flushed with a mixture of  $N_2$ : $CO_2$  (90:10) at a rate of 0.20 - 0.50 ml/min. The headspace gas composition is sampled in 15 min. intervals using a Hiden QIC-20 mass spectrometer.

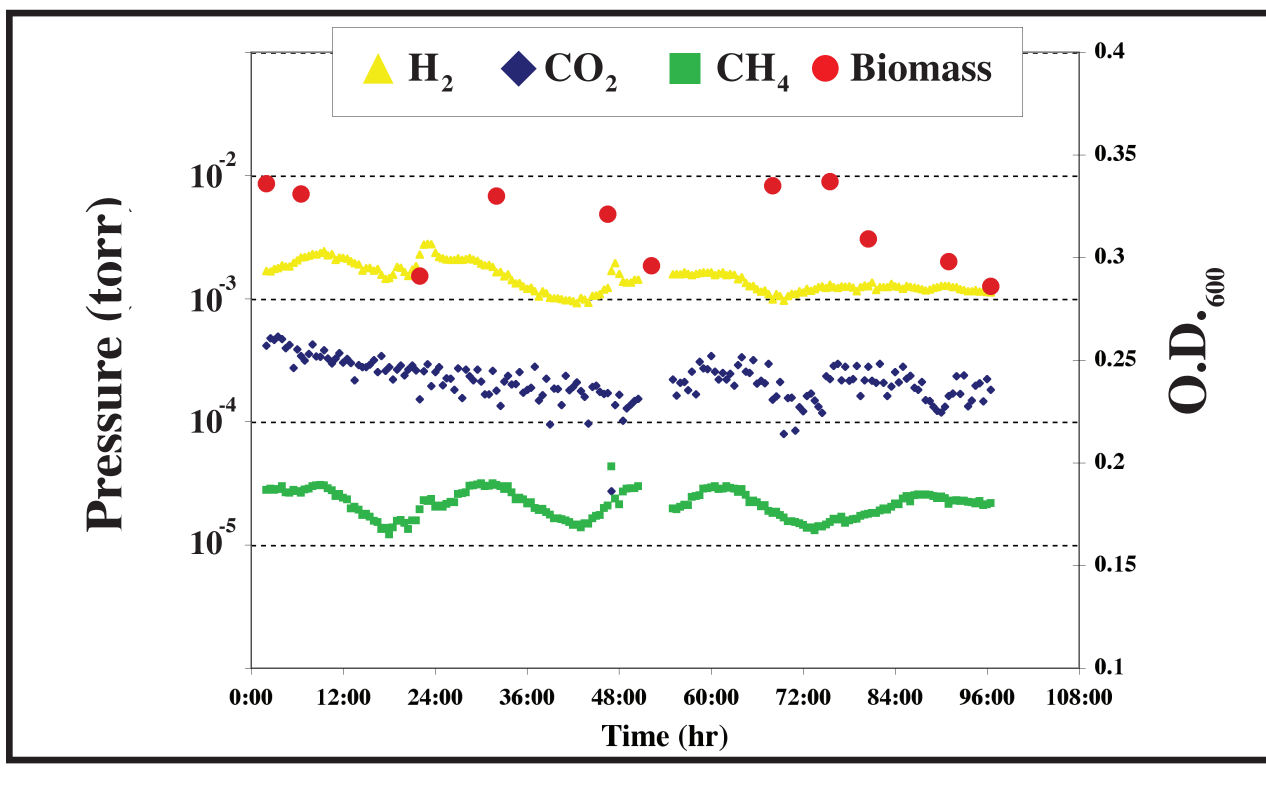
## TRANSCRIPTIONAL ANALYSIS

Triplicate biological replicates of cocultures and sulfate-limited *D. vulgaris* monocultures were analyzed by the ESPP Functional Genomics Core using custom-designed wholegenome microarrays. Microarray slides were designed with duplicate spots of each openreading frame (ORF) for both organisms. At least three slides were used for each biological replicate. The ESPP Computational Core calculated RNA/DNA expression ratios for each ORF and the log<sub>2</sub> ratio comparing the coculture versus sulfate-limited monoculture growth conditions was determined. Z-scores for each ORF were calculated to determine statistical



significance. Operon-based estimates of local accuracy indicate an absolute Z-score of 1.0 accurately predicts expression changes between the two conditions. Using this value, 169 ORFs displayed significant up-regulation. 254 ORFs were statistically down-regulated. Genes were assigned clusters of orthologous group (COG) functional codes based on previous genome annotations.

# BIOMASS & HEADSPACE GAS MEASUREMENTS



Steady-state was assumed when O.D.<sub>600</sub> measurements varied by less than 10% of initial value for 3 rentention times. *D. vulgaris:M. maripaludis* cell ratio was ~4:1 throughout steady-state as determined by DAPI-stained cell counts.

al Up-regulated

50

Down-regulated

10

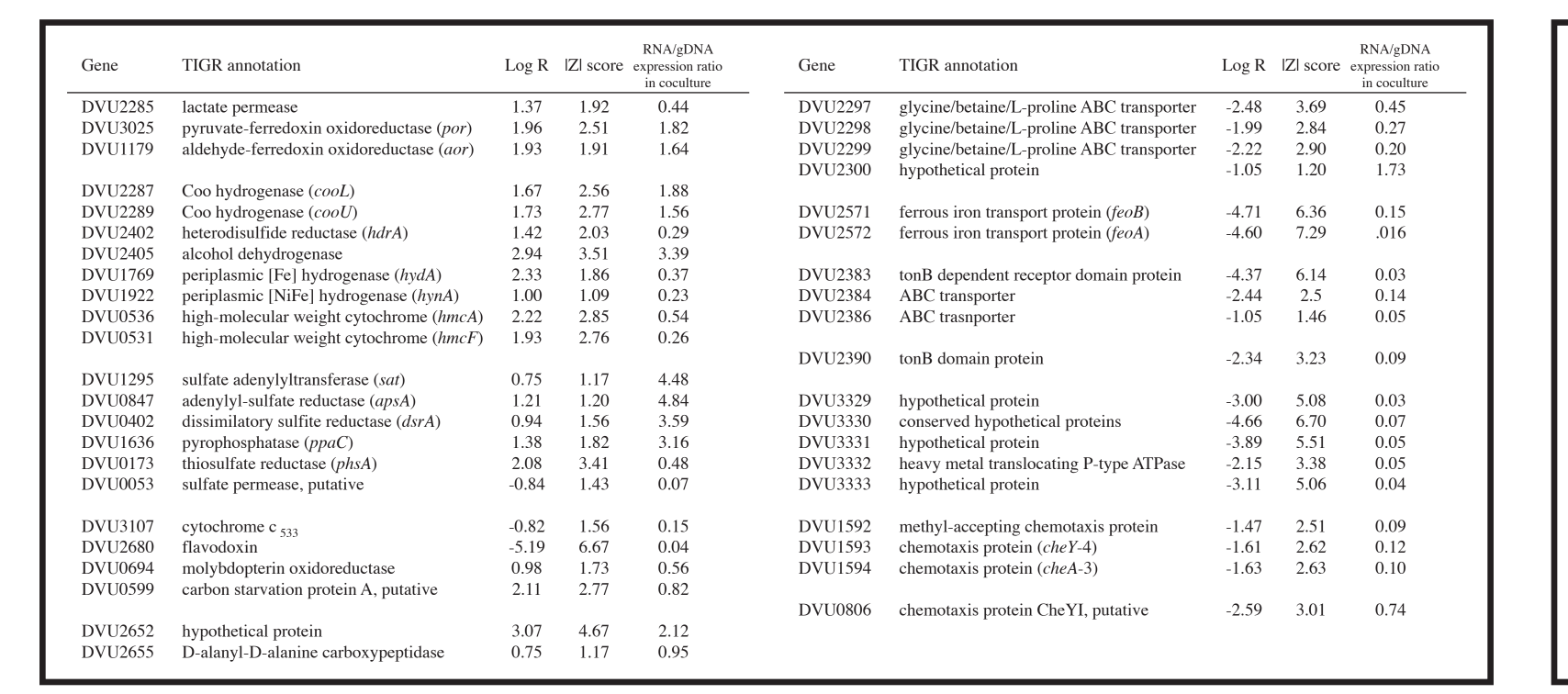
10

Down-regulated

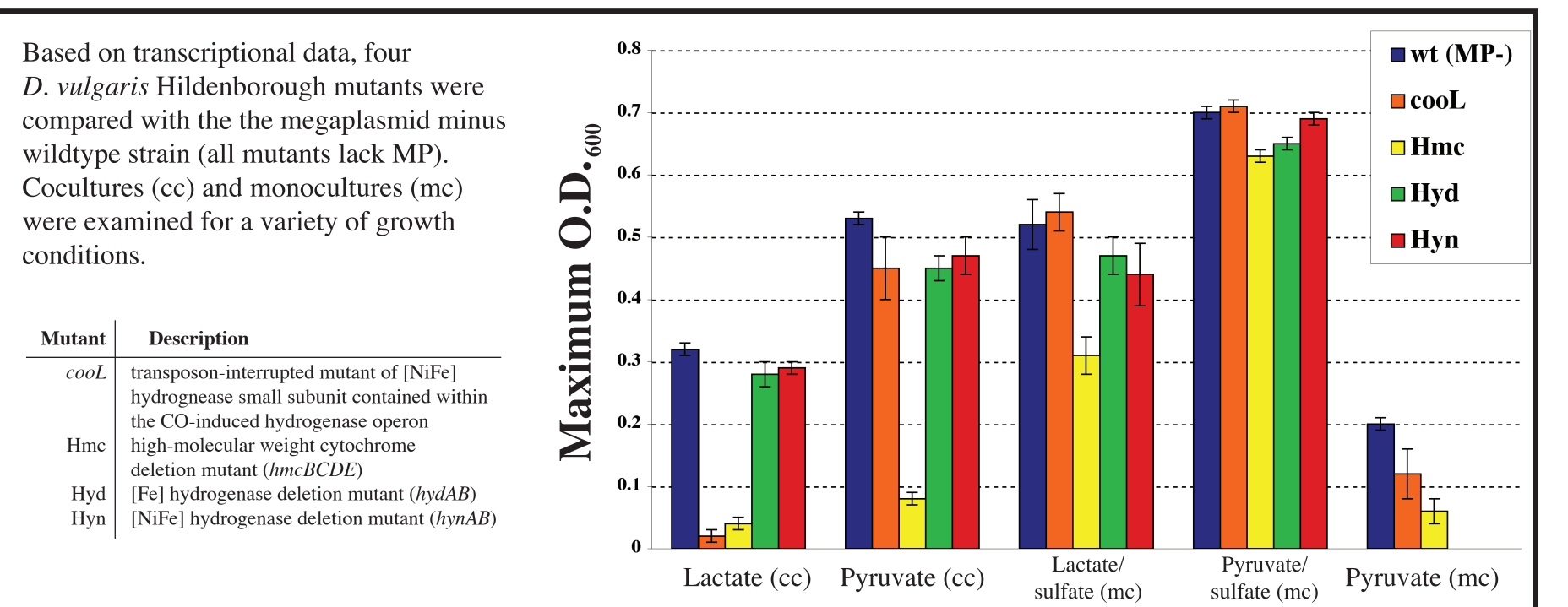
Cluster of orthologous groups up- and down-regulated during syntrophic growth. Categories are amino acid transport (E), carbohydrate transport and metabolism (G), cell division and chromosome partitioning (D), cell envelope biogenesis (M), cell motility and secretion (N), chromatin structure and dynamics (B), coenzyme metabolism (H), DNA replication, recombination and repair (L), defense mechanisms (V), energy production and conservation (C), function unknown (S), general function prediction only (R), inorganic ion transport and metabolism (P), intracellular trafficking and secretion (U), lipid metabolism (I), nucleotide transport and metabolism (F), post-translational modification, protein turnover, chaperones (O), secondary metabolites biosynthesis, transport and catabolism (Q), signal transduction mechanisms (T), transcription (K) and translation,

V C S R P U I F O Q T K

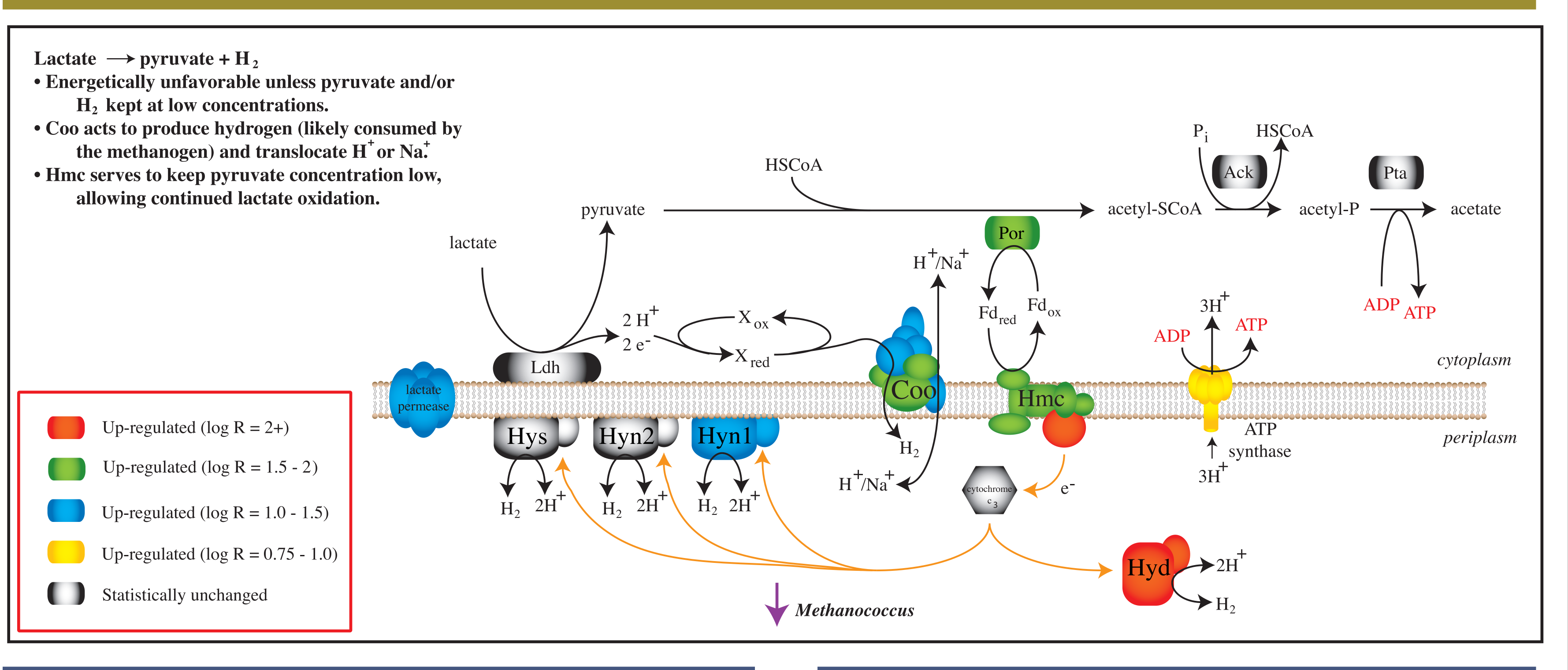
## TRANSCRIPTIONAL RESPONSE



## PHENOTYPIC DIFFERENCES BETWEEN D. VULGARIS MUTANTS



# ENERGETIC OVERVIEW



# **FUTURE WORK**

- Develop syntrophic growth pathway on other compounds, such as ethanol.
- Refine metabolite interaction model between *D. vulgaris* and *M. maripaludis* LL using transcriptional, proteome and phenotypic analyses.
- Explore salt, nitrate, pH, etc., stress response of coculture.
- Compare responses with other organisms, especially other *Desulfovibrio* strains.

## ACKNOWLEDGMENT

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