Multi-omic profiling of EPO-producing CHO cell panel reveals metabolic adaptation to heterologous protein production

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Multi-omic profiling of EPO producing Chinese hamster ovary cell panel reveals metabolic adaptation to heterologous protein production

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Introduction

Heterologous protein production in CHO cells imposes a burden on the host cell metabolism and impact cellular physiology on a global scale. In this work, a multi-omics approach was applied to characterize the physiological impact of erythropoietin production and discover production bottlenecks in a panel of CHO-K1 cells in batch and chemostat culture.

Objectives

- Generate a panel of EPO producing clones with broad specific protein productivity.
- Identify the production bottleneck in the secretory protein production pathway.
- Characterize the burden of EPO production on the host cell metabolism and identify bottlenecks in energy metabolism.

Cell lines

The panel consist of 7 EPO-producing CHO K1 clones spanning a 25-fold productivity range.

Investigating the protein expression bottleneck

To identify the protein expression bottleneck, we cultured three clones (C1, C4 and C7) in chemostat culture for 31 days and determined EPO gene copy numbers, mRNA levels and secreted EPO. The post-transcriptional efficiency (ratio of EPO per mRNA) indicated a production bottleneck downstream of transcription in clone 1 and 4 relative to clone 7.

Common protein expression bottlenecks

Common bottlenecks found in production of secreted heterologous proteins in CHO cells, as reported in the literature.

Metabolic impact of EPO production

To discover bottlenecks in glucose metabolism, we quantified intracellular metabolites of glycolysis (data not shown) and associated energy metabolites across the EPO producing clones in mid-exponential phase in batch culture. The differences in concentration showed no correlation to specific EPO productivity suggesting EPO production is not limited by energy supply.

To investigate the impact of EPO production on metabolism, we reconstructed of glycolysis, TCA and amino acid catabolism in CHO cells (see link below) and integrated differential gene expression data from phase II of chemostat culture. The data integration revealed decreased transcription level of genes responsible for degradation of the amino acids most frequently found in EPO. This, indicating possible regulatory adaptation of gene expression towards decreased amino acid catabolism specific for the most abundant amino acids in EPO, in the high producer relative to the low producer.

Conclusion

We provide evidence that EPO production up to 5 pg/cell/day is not limited by metabolism (i.e. glycolysis and associated energy metabolism) or bottlenecks in gene dosage, transcription and post-translational processing of EPO. Furthermore, we demonstrate that the amino acid catabolism can adapt to different amino acid demand imposed by heterologous protein production within 9 generations during prolonged chemostat cultivation.

References