Multivariate profiling as a pivotal tool in identifying critical raw materials that rescue N-linked glycosylation profiles

Steven B. Richardson1, Omar Wahab2, Christopher Kornfeld1, James S. Ross1, and Kevin Kayser1

1 Cell Sciences and Development, SAFC, 2909 Laclede Avenue, Saint Louis, MO 63103, USA
2 SAFC Irvine, Second Avenue, Heathcote Industrial Estate, Irvine, Ayrshire, KA12 8NB

Introduction

Over the past decade, notable advances in the characterization and control of post translational modifications (PTMs) to enhance therapeutic properties have continued to drive the development of many next-generation biopharmaceuticals. It has been widely reported that the quality of secreted therapeutic proteins is dependent on the consistency of attached glycan moieties. Insufficient glycosylation potentially impairs efficacy and safety. Our particular interest lies in identifying critical raw materials that rescue N-linked glycosylation profiles of therapeutic proteins from selected high-producing clones. This study represents an important step towards establishing GMP ready chemically defined supplements that significantly and reproducibly adjust glycan moieties. In addition, the multivariate profiling process described here represents a pivotal tool in the upstream characterization of biopharmaceutical platforms.

Technical Approach

Design of Experiments

Materials & Methods

Fed-Batch Culture

Glutamine Synthetase (-/-) recombinant IgG producing CHOZN™ cell line. Duplicate 30 mL cultures in 125 mL vented shake flasks at 37°C and 140 rpm with 80% Rh and 5% CO2. Glucose supplementation as needed. Cultures maintained in Ex-CELL CHO CD Fusion with 5% additions of imMEDIAte ADVANTAGE™ manufactured chemically defined feeds on days 3, 5, 7, 9, and 11.

Intact Protein Accurate Mass Spectrometry

A high-throughput SEC-MS (Waters Acquity UPLC®/Q-Tof Premier™) workflow was developed. Protein-A purification of day 14 culture supernatants followed by intact mass analysis of antibody heavy chain constituents. Automated deconvolution and data processing performed using Waters BiopharmaLynx™ software.

Results

Proprietary Feed Screen

Ex Cell Glycosylation Adjust (Gal+)

Conclusion

Identified critical raw materials that demonstrate 2-4 fold increases in relative G1F and G2F distributions.

Optimized supplement component ranges that minimally affect volumetric productivity and viable cell density.

Established culture supplementation protocol at 0.2% (v/v) beginning on day 2 and then every other day.

Demonstrated scalable performance over a wide variety of bioproduction cell origins including CHO-GS, CHO-M, DuxB11, and NS0 lines (data not shown).

Established multivariate profiling process as an important tool in upstream biopharmaceutical development.