Identification of novel inhibitory metabolites and impact verification on growth and protein synthesis in mammalian cells

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1. Introduction

Biotherapeutics have emerged as one of the most effective treatment options for many diseases including cancers and autoimmune disorders. For more than three decades, mammalian production hosts have been used to produce these biotherapeutic molecules (Dhara et al., 2018). Of these hosts, Chinese hamster ovary (CHO) cells are used extensively at production scale to produce a majority of therapeutic proteins, such as monoclonal antibodies. The market value of CHO-expressed therapeutic proteins is expected to reach $217 billion by 2023 (Grilo and Mantalaris, 2019). This coupled with their biocompatibility through human-like post-translational modifications by the CHO cellular production machinery, make them the preferred hosts in the biopharmaceutical industry. Maximizing the protein producing capacity of the cells has been pursued in several ways such as supplementing cell growth medium with growth and productivity enhancers (Takagi et al., 2017), changing the operating mode of cell culture to fed-batch and perfusion (Hiller et al., 2019).
Solved oxygen, $pCO_2$, pH, etc.) to increase its longevity and productivity (Mohan et al., 2008).

Such process improvements have enabled the industry to reach very high viable cell densities in the order of $10^6$ cells$^{-1}$mL$^{-1}$, with production yields of several grams per liter (Kunert and Reinhart, 2016). Despite these improvements, limitations still exist in the productive capacity of the cells. One possible avenue to increase cellular productivity is addressing inefficient metabolism of available substrates (such as glucose and glutamine) in the cells, leading to the production and accumulation of waste metabolites over the duration of the cell culture (Pereira et al., 2018). CHO cells exhibit inefficient and poorly regulated metabolism if they consume more nutrients than required to support proliferation and production and thereby generate waste metabolites. Such inefficient metabolism can lead to the accumulation of potentially toxic metabolites which can inhibit overall cellular performance, especially at later stages of the culture process (Pereira et al., 2018). The impacts of undesired end-products such as lactate or ammonia have been widely reported (Lao and Toth, 1997; Chen and Harcum, 2005; Sun and Zhang, 2001; Freund and Coughlan, 2018; Buchsteiner et al., 2018). These by-products can become major rate-limiting factors, hindering further constructive utilization of the substrates or a decline in cellular performance with respect to cell proliferation and/or antibody production. To better regulate cell metabolism throughout the process and enhance cellular productivity, it is crucial to identify all inhibitory waste metabolites, study their corresponding biochemical precursors and evaluate their pathways of generation.

Liquid chromatography with tandem mass spectrometry (LC-MS/MS) is a powerful tool for analyzing potentially thousands of metabolites from a single sample. Owing to this, LC-MS/MS has been used in various metabolomic studies to identify markers in CHO cells which enhance growth and/or productivity. Recent study has shown LC-MS/MS can be used for quantitative assessments of medium additives and nutrients (Hoang et al., 2021). Additionally, LC-MS/MS has also been demonstrated to be capable of providing insights into cellular metabolism by profiling the extracellular metabolites generated during growth of CHO cells (Mohmad-Saberi et al., 2013). LC-MS/MS-based metabolomics can also probe the metabolic states of CHO cells by identifying growth limiting factors during different stages of culturing, including but not limited to oxidative stress and lipid metabolism (Chong et al., 2012). Other studies applied quantitative metabolomics to evaluate perfusion cell culture, which established a relationship between the multiple steady states in the process (Karst et al., 2017). Identification of metabolic by-products in CHO cell cultures that accumulate to toxic levels in CHO cell fed-batch processes and impact the culture performance has also been enabled through LC-MS/MS (Chong et al., 2011). Additionally, it has been shown that the extracellular environment is critical for cell proliferation and product titer quality in mammalian bioprocesses (Arden and Betenbaugh, 2004; Polanco et al., 2020). A recent study has demonstrated that accumulation of by-products such as isovalerate, indolelactate, indole-3-carboxylate, and others in high-density CHO cell fed-batch process was emerged from degradation of various essential amino acids such as leucine, tyrosine, and tryptophan (Mulukutla et al., 2017). Buildup of such waste inhibitory products inhibits cell culture performance and hinders productivity and thus, preventing or reducing the accumulation of these by-products will improve process performance. Therefore, it is of paramount importance to identify the inhibitory metabolites accumulated in mammalian bioprocess, and therefore unfolding the correlation between downstream metabolites to upstream feeding source.

In this study, LC-MS/MS based untargeted metabolomics platform is used to investigate the extracellular environment of a fed-batch CHO–K1 cell culture process to identify metabolites accumulating over the culture duration. Seventeen novel accumulating metabolites were tested for cell culture toxicity. A targeted metabolomic analysis was then used to confirm these results and eight were observed to quantitatively accumulate and negatively impact growth and antibody productivity. Sensitivity and pathway analysis on CHO-GS, CHO–K1, and HEK293 cell lines confirmed these inhibitory metabolites are not exclusive to CHO–K1 but can also be found across different mammalian host systems. The methodology developed here (see Fig. 1), as well as the metabolites identified which can be used to overcome growth and production bottlenecks in CHO and other mammalian production hosts can provide the foundation for development of robust future biomanufacturing platforms.

2. Materials and methods

2.1. Cells, media, and supplements

A CHO–K1 cell line expressing IgG (VRC01) antibody obtained from NIH (National Institute of Health) was used for this study with two proprietary media. For inoculation, basal medium (medium A) obtained from Sigma Aldrich (St. Louis, MO) and glutamine at 8 mM obtained from Corning (Corning, NY) were supplemented to cell culture on Day 0. For fed-batch process, enriched nutrient feed medium (medium B) obtained from Lonza (Portsmouth, NH) was used as feed medium and fed to cells after Day 0 until cellular viability dropped below 80%. The initial feeding day is dependent on the development of feeding strategy and is clearly addressed throughout the discussion. Cells were cultivated in sterile 125 mL shake flask obtained from Fisher Scientific (Waltham, MA) with a working volume of 30 mL in a humidified, shaking incubator (Model AJ125) obtained from ATR Biotech (Laurel, MD) operated at 125 RPM, 36.5 °C, and 5.2% carbon dioxide. Cells were inoculated at low viable cell densities (VCD) (approximately $0.5 \times 10^6$ cells$^{-1}$mL$^{-1}$) in basal medium and harvested when cell viability decreased below 80%. Sub-culturing was done every two to three days when the cells reached
VCD greater than 3 × 10^6 cells mL⁻¹. Cell viability was maintained above 90% throughout the inoculation phase. Stock solution of each candidate metabolite was prepared by dissolving pure metabolite standards obtained from Sigma Aldrich (St. Louis, MO) into ultrapure water from a MilliQ water purifier obtained from Millipore Sigma (Burlington, MA), and were supplemented in the basal media on Day 0 for both batch and fed-batch processes. For process validation, a CHO-S cell line cultured in medium A, and HEK293 cell line (CRL-1573.3) obtained from American Type Culture Collection (Manassas, VA) cultured in Irvine BalanCD medium obtained from Irvine Scientific (Santa Ana, CA).

2.2. Assessment of growth and productivity

VCD and viability were measured daily with a cell counter device (CeDex HiRes) obtained from Roche (Branchburg, NJ). Glucose, lactate, and ammonia concentration data from each culture condition were measured using an analyzer (Bioprofile FLEX) obtained from Nova Biomedical (Waltham, MA). Titer analysis was performed using a high-pressure liquid chromatography system (Agilent 1100 series) obtained from Agilent Technologies (Santa Clara, CA) with a protein A column (Poros A, 2 μm, 2.1 × 30 mm) obtained from Thermo Scientific (Walnut, CA). Glycoclean™S cartridges were from Prozyme (Hayward, CA). An inline filter (0.2 μm) was purchased from Restek Corporation (Bellefonte, PA) and a BEH amide column (1.7 μm, 2.1 mm × 50 mm) was purchased from Waters Corporation (Milford, MA). For summary of a detail protocol, interested reader can refer to Supplement S2.

2.3. Glycan analysis

Human serum IgG reference standard, 2-aminobenzamide (2AB), 2-picoline borane complex (2 PB), ammonium formate and acetonitrile (HPLC-grade) were obtained from Sigma Aldrich (St. Louis, MO). Magnetic beads coated with protein A and protein G were purchased from Thermo Fisher Scientific (Waltham, MA). A peptide:N-glycosidase F (PNGase F) enzyme kit containing 10× denaturing buffer, NP-40 non-ion surfactant detergent and 10× reaction buffer was purchased from New England Biolabs (Ipswich, MA) and stored as instructed by the manufacturer. Diol solid phase extraction cartridges HyperSep™ Diol SPE were purchased from Thermo Fisher Scientific (Waltham, MA). Glycoclean™S cartridges were from Prozyme (Hayward, CA). An inline filter (0.2 μm) was purchased from Restek Corporation (Bellefonte, PA) and a BEH amide column (1.7 μm, 2.1 mm × 50 mm) was purchased from Waters Corporation (Milford, MA). For summary of a detail protocol, interested reader can refer to Supplement S2.

2.4. Protein purification

The method for glycan analysis was followed closely to a previously established protocol (Sha et al., 2020). Simply put, cell culture samples were collected to contain approximately 0.1–0.5 g L⁻¹ titer (about 100–500 μL supernatant) to a 1.5 mL microcentrifuge tube, after which samples were spun down for 5 min at 1000 rpm. The supernatant which contained IgG was collected to a microcentrifuge tube, into which an aliquot of 10 μL magnetic bead slurry was added. The beads were washed twice, each time with 250 μL buffer. A magnetic stand was
incorporated into each wash to separate the beads from the buffer, after which the washing eluent was discarded. A volume of culture supernatant was calculated to contain approximately 20–50 μg mAb is then added to 1.5 mL microcentrifuge tube, after which the tube was vortexed gently and incubated on a shaker for 15 min. The tube was placed on the magnetic stand to separate the beads-IgG complex from the supernatant, after which the supernatant was removed. The beads were washed by two consecutive washes, with the first wash using 250 μL of binding buffer following by a second wash of 250 μL water. The beads were further suspended into 25 μL of elution buffer and incubated for 10 min. The tube was then placed on the magnetic stand and the supernatant (free of magnetic beads) is transferred to a new microcentrifuge tube which was pre-loaded with 8 μL neutralization buffer.

2.5. Enzymatic digestion of N-linked glycan from glycoprotein

Purified standard/mAbs solution is denatured by adding 4 μL of 10× denaturing buffer purchased from New England Biolabs (Ipswich, MA) and 6 μL of water. The mixture was then heated at 100 °C for 10 min. After cooling for 15 min, the mixture was added with 5 μL of 10× PNGase F reaction buffer and 5 μL of NP-40 in water. Water was added to the sample to reach a final 50 μL reaction volume. The PNGase F enzyme is added at 1 μL (500,000 units·mL−1), after which the digestion was carried out at 37 °C for 20 min.

2.6. Glycan labeling and cleanup procedure

The labeling solution was prepared to a final composition containing 0.35 M of 2AB, 1 M of 2 PB, 30% vol/vol (v/v) acetic acid in 70% DMSO. The labeling solution was added at 50 μL–50 μL of the digested protein solution, after which the mixture was covered with aluminum foil and incubated at 65 °C for 1 h.

For 2AB cleanup, the methodology presented in this paper utilized Diol SPE cartridge as an economical substitute to HILIC cartridge as previously reported has a similar and comparable final results (Sha et al., 2020). To use Diol SPE for sample cleanup, the cartridges were attached to a vacuum manifold equipped with a vacuum pump. The cartridges were first washed with 1 mL of water, followed by four consecutive washes of 1 mL ACN each wash. Labeled glycans were diluted in ACN at a ratio of 10/90% v/v. Sample were then loaded and pumped through the cartridge, after which two washes of 500 μL ACN each wash was added to remove unbound compounds including excess 2-AB. Finally, two consecutive aliquots of 100 μL water were added to elute the labeled glycans into fresh 1.5 mL microcentrifuge tube. The total volume of the eluent is 200 μL.

2.7. LC-MS/MS method for metabolites profiling

Metabolite standards were obtained from Sigma Aldrich (St. Louis, MO) and used for identification of inhibitor metabolites. Labeled amino acid standards set A (NSK-A) obtained from Cambridge Isotope Laboratories (Tewksbury, MA) was dissolved in water to prepare internal standards (IS) solution. IS solution of glycine (13C, 15N) was prepared at 5 M. The remaining labeled amino acid IS was prepared at 1 M. Samples were prepared by mixing 20 μL of sample with 5 μL IS and 100 μL Acetonitrile. Samples were vortexed for 30 s and centrifuged for 10 min at 16,000 g. The supernatant was then extracted for LC-MS/MS analysis.

For MS analysis, 3 μL of sample was analyzed through orbitrap LC-MS system (Thermo Scientific Accela) consisting of a LC column and a linear trap quadrupole velos mass spectrometer obtained from Thermo Scientific (Waltham, MA). Chromatographic separation was performed on a PEEK coated column (SeQuant ZIC-chICILC, 3 μm, 100 × 2.1 mm) with a guard kit (SeQuant ZIC-chICILC, 5 μm, 20 × 2.1 mm), both obtained from Millipore Sigma (Burlington, MA). For detail operating conditions regarding quantification method, interested reader can refer to Supplements S2.

2.8. Global metabolomics study

Hydrophilic metabolites (e.g., ACA derivatives, organic acids, sugars, and pyrimidines) were studied in the metabolome of HCD CHO fed-batch process using high resolution Orbitrap LC-MS system obtained from Thermo Scientific (Waltham, MA). Metabolomic data obtained through LC-MS were mapped to the human metabolome database (HMDB) to reveal additional metabolomic features. Overall, more than 30,000 hits were revealed using the Scaffold Elements software obtained from Proteome Software (Portland, OR). As an effort to acquire high-confidence metabolites identification in subsequent targeted metabolomic study and biological toxicity test, an MS2 scoring system was incorporated into the analysis, as used in previous studies (Huang et al., 2020). The MS2 score of each metabolite is determined by comparing MS2 precursor record from every MS2 scan through a dot product between the database recorded MS2 fragment scan and experimental analyte fragment MS2 scan.

For the untargeted global metabolomic study, since both types of media (medium A and medium B) were used simultaneously throughout different processes presented in this study, the MS signal intensity of both media are therefore normalized by constructing a blank consisted of medium A and medium B added to a final of volume with composition of medium A/medium B = 1/1.4 %v/v. Thus, the downstream ion intensity of each metabolite (Ii) is defined to be the detected ion intensity obtained from orbitrap MS subtracting the ion intensity of the blank. The standard deviation (σ) and average intensity value (I) of each metabolite i are calculated from triplicate measurements, from which the relative standard deviation (RSD) for each metabolite can be calculated (Eq (1)).

\[ RSD = \frac{\sigma}{I} \]

2.9. Impact determination and metabolite characterization

CHO–K1 and CHO–S cultures were cultivated in their respective media (CHO–K1 in medium A and CHO–GS in modified medium A). Both media were supplemented with 6–8 mM of glutamine (Corning INC, Corning, NY). Cells were cultured in 125 mL Erlenmeyer flask obtained from Fisher Scientific (Waltham, MA) with a working volume of 30 mL. Sub-culturing of cells was performed every two to three days when cells reached VCD of more than 3 × 10⁶ cells·mL⁻¹. All the metabolites screened for the spike-in studies and confirmation batch runs were obtained in pure forms from Millipore Sigma (Burlington, MA) and are listed in Table S2. To examine the impact of metabolites for the screening studies, the metabolites were spiked into the CHO cell cultures at two different concentrations (1 mM and 5 mM). Whereas, during metabolite characterization studies, an additional level was tested taken from the measured accumulation levels of the inhibitory metabolite from targeted metabolomics performed on CHO–K1 fed-batch cultures. Stock solutions of metabolites (except for trans-cinnamic acid and hydroxyphenyllactic acid due to their low solubility) were dissolved with water or DMSO depending on compound solubility. The solutions were added to the culture on Day 0. The pH of the medium was adjusted to minimize the effect of pH changes resulting from the metabolite additions. The inhibitor-spiked media were evaluated for osmolality, verifying a minimal change of less than 10% in osmolality (less than 4% on average), which remains well within the acceptable range of cell culture media osmolality variation. Cells were inoculated at a seeding cell density of 0.5 × 10⁶ cells·mL⁻¹. Samples were collected for cell count and metabolite measurement every 24 h. Cell count was performed using a hemocytometer until the viability dropped below 70%. Spent medium analysis for glucose and lactate measurements was performed using a biochemistry analyzer obtained from YSI (Yellow Springs, OH). All the conditions were tested with biological duplicates and the flasks were maintained at 37 °C in a humidified shaker incubator.
obtained from Infors HT (Weymouth, MA) at 5% CO₂ levels.

3. Results and discussion

3.1. High cell density process development

A high cell density (HCD) fed-batch process was conducted with nine different feeding conditions on a CHO-K1 cell line using medium A, as illustrated in Table 1. When studied the effect of different feeding volumes on cell culture performance, the study found that medium feeding condition (condition 5, feeding volume 3 mL) resulted in the highest IVCD (see Fig. 2 B), which was 30.8% higher than the high feeding condition (condition 3, feeding volume 4.5 mL) and 25.9% higher than the low feeding condition (condition 7, feeding volume 1.5 mL). High feeding volume was found to be directly correlated to higher glucose consumption and lactate accumulation rate when compared against the medium and low feeding condition (see Fig. 2 C and D). However, high feeding volume did not translate to a better growth profile, as the medium feeding condition (feeding volume 3 mL) showed highest VCD and growth rate (Fig. 2 A). Similarly, when studied the effect of different initial feeding day, the study found that feeding on Day 3 (condition 5) resulted in the highest attainable cumulative IVCD (see Fig. 2 F), which was 11.8% higher than the early feeding condition (condition 2, Day 2) and 10.9% higher than the late feeding condition (condition 8, Day 4) with all three conditions maintaining the same 3 mL feeding amount. Feeding too early (condition 2, feeding Day 2) resulted in cells reaching production phase at lower peak VCD. On the other hand, feeding too late (condition 8, feeding Day 4) resulted in cells rapidly reaching highest peak VCD (Day 8) but rapidly approaching death phase and consequently translated to the shortest culture duration (12 days culture duration, Fig. 2 E). A similar metabolism profile was also observed when the effect of different initial feeding days on cellular performance was studied, as the earliest feeding day (condition 2, feeding Day 2) with the most abundant nutrients available in the spent medium allowed cells to up-regulate their glycolysis pathway (higher glucose uptake) and higher lactate production when compared against the mid and late feeding conditions (see Fig. 2 G and H). Interestingly, the lactate accumulation rate of Day 3 feeding was found to be the lowest when compared against Day 2 feeding and Day 4 feeding. Altogether, the results obtained from employing different feeding volumes (Fig. 2 A) and different initial feeding days (Fig. 2 E) suggest that cells exhibit more rapid substrate utilization and metabolic activity rates when nutrients are supplemented at excess levels. The data further confirm that waste metabolites generated through upregulated pathways can inhibit cellular growth due to lactate production (Chen et al., 2001; Sha et al., 2020), reactive oxygen species generation (Sha et al., 2020; Ha and Lee, 2000), and endoplasmic reticulum stress (Zhong et al., 2012). On the other hand, the study showed that lower feeding volume (condition 4) or later feeding day (condition 8) both result in higher peak cell density but a shorter culture period due to a shortage of nutrients. For the growth profile and consumption profiles of all other conditions studied in this HCD process development, interested readers can refer to Figure 5 1.

Table 1

| Condition | Initial Feeding Day | Feeding Volume (% Seeding Volume) |
|-----------|---------------------|-----------------------------------|
| 1         | 2                   | Low – 5%                          |
| 2         | 2                   | Medium – 10%                      |
| 3         | 2                   | High – 15%                        |
| 4         | 3                   | Low – 5%                          |
| 5         | 3                   | Medium – 10%                      |
| 6         | 3                   | High – 15%                        |
| 7         | 4                   | Low – 5%                          |
| 8         | 4                   | Medium – 10%                      |
| 9         | 4                   | High – 15%                        |

Ultimately, the feeding strategy obtained from condition 5 was selected for future study due to its highest attainable IVCD across all the conditions tested.

3.2. Inhibitory metabolites identification: global metabolomics study

Hydrophilic metabolites were explored in the global metabolomic study. A rigorous ranking system was used to screen metabolites generated in a fed-batch process for their inhibitory impact. First, a MS2 score was calculated and assigned to each global metabolome feature, from which only features with MS2 score greater than 0.5 or RSD less than 0.1 were further considered for targeted metabolomics study and biological verification impact analysis. Since a metabolite can have multiple retention times due to different liquid chromatography conditions, only one metabolite was selected for data-mapping to the human metabolome database. The remaining metabolite candidates were then selected for toxicity tests and targeted metabolomics verification to obtain high-confidence identification of waste inhibitory metabolites. For further detail regarding metabolomics information, interested reader can refer to Table 5 3. The final list of all fifteen high-confidence metabolites is shown in Table 1. The global metabolomics analysis study was also repeated for CHO batch process.

3.3. Inhibitory metabolites identification: metabolites screening for impact

The first level of screening through spike-in study was performed in CHO–K1 batch cultures at two concentrations of metabolite (1 mM and 5 mM) in the initial growth medium. The rationale behind the selection of these initial concentrations was two-fold: i) high cell density CHO culture was observed to exhibit accumulation of metabolic by-products in the range of micromolar to a few millimolar amount in their extracellular environment based on previous study, and ii) previous study has verified certain metabolite candidate to show potential inhibitory effect on CHO growth 1 mM and 5 mM concentrations (Mulukutla et al., 2017).

The pH of the initial growth medium was initially measured (pH = 6.95), to which the pH of each experimental condition was adjusted to ensure no inhibitory effects due to pH shifts resulting from the respective metabolite addition. Cell culture growth parameters such as viable cell density (VCD) and viability (%) were monitored in which each of the cell cultures supplemented with the target metabolite at these two different concentrations were compared with the un-supplemented control cell cultures. A metabolite was therefore considered toxic to the cells if the VCD was lowered by more than or equal to 15% compared to the control in either the exponential or the stationary phase of cell culture.

For the metabolites impact screening study, ornithine and hypoxanthine were not included in the metabolite candidates spiking list due to their known physiological roles as substrates in cellular metabolism (Chen et al., 2012; Zajac et al., 2010). The remaining fifteen metabolites obtained from untargeted global metabolomics analysis were selected for biological verification to evaluate the toxicity effect on cellular performance in a standard culture bioprocess. The task was undertaken to provide a mean to mitigate the generation of metabolite that was observed to accumulate at high concentration during stationary phase of CHO fed-batch culture, of which their metabolic build-up does not contribute to cellular performance. Metabolites that show no inhibitory effect on CHO cultures, elucidated through spike-in study, would not be further considered in downstream metabolomics pipeline towards structural verification through targeted metabolomics and subsequent level determinations. The metabolites included in the spike-in study (and their abbreviations) are provided along with the rationale of identification confirmation or removal (see Table 2). The inhibitory impact on cellular performance due to spiking of metabolites presented as VCD impact heatmap is shown in Fig. 3 B. Of the fifteen metabolites considered for biological verification, hydroxyphenyllactic acid (HPL), deoxycytidine (DCI), l-allothreonine (LAT) were removed from the
metabolite candidates list since spiking of these metabolites into CHO cultures showed no inhibitory impact on cellular performance as shown in the VCD impact heatmap illustrated in Fig. 3B. Additionally, 6-aminocaproic acid (6CA) was also removed due to having high RSD value (inconsistent MS data intensity) across replicate runs. Finally, glutaric acid (GA), trans-cinnamic acid (TCA), and p-aminobenzoic acid (pAA) were removed from the metabolite candidates list through comparison of their MS3 spectrum and LC retention time obtained from the sample to the reference standard, after which the concentration at the end of culture was measured (see Table 2). For comparison of the MS3 spectrum between the candidate features, interested readers can refer to Table S3.
3.4. Inhibitory metabolites characterization in CHO-K1 and CHO-GS batch process

Verification on the identity of waste metabolites and quantification of their accumulated concentration in CHO batch culture was performed through targeted metabolomics study. Overall, careful study on the metabolism of cells allowed identification of eight metabolites accumulating at high level in the harvested samples. The impact of these metabolites on culture performance was further characterized in batch and fed-batch culture of CHO-K1, as well as batch culture of CHO-GS and HEK293 to determine whether the toxicity on cellular performance is universal across different mammalian cell lines. Overall, careful analysis of these metabolites of interest enabled a broader classification on the metabolic pathways responsible for their accumulation. The identities of the metabolites and their respective metabolic pathway are shown Fig. 4. The cellular level of metabolites accumulation as measured from samples obtained at the harvest day of CHO-K1 batch process is shown in Table 2.

The results obtained from the spiking studies of CHO-K1, and CHO-GS process are shown in Fig. 5 and Fig. 6. Here, metabolites CMP, GMP and ACA (aconitate) are known to accumulate from pathways emerging from the central carbon metabolism, such as the TCA cycle, purine, and pyrimidine metabolism. Of the three central carbon intermediates, the study showed that only CMP was shown to have a negative effect on cell growth at all tested concentrations while GMP and ACA were toxic at two concentration levels (1 mM and 5 mM) and one concentration level (1 mM) respectively out of the three concentrations that were tested. Surprisingly, the negative impact of ACA on cell growth was only found in cultures where ACA was supplemented in lower concentrations. At 5 mM spiking concentration to batch process, ACA showed a positive impact on cell growth. It is also important to note that, cultures spiked with ACA showed a decrease in specific productivity along with CMP and GMP when tested at “cell” concentration (see Fig. 5 J). Overall, at the end-of-culture (“cell”) concentration noted in Table 2, the three metabolites CMP, GMP and ACA had an impact on CHO cell cultures, although the toxic effect due to GMP to cells was only significant at millimolar levels (see Fig 5 A and Fig. 6 B). Thus, it was concluded from the study that CMP and GMP are growth inhibitors as well as productivity inhibitors. In the case of ACA (aconitate), this metabolite is generated as a TCA cycle intermediate which is formed during the conversion of citrate to iso-citrate catalyzed by mitochondrial aconitase (Aco2) in mammalian cells. Previous studies conducted on CHO process revealed that knockout of Aco2 negatively impacts cell growth (Dhami et al., 2018). This infers that supplementing ACA to cells drives the activity of TCA cycle in the forward direction to generate more ATP required for growth and protein production. Hence, ACA showed no growth toxicity and reduced productivity in CHO cells at “cell” concentration and in fact, at 5 mM enhanced growth in comparison to the control (see Fig. 5 C).

Of the eight identified toxic metabolites, HICA and MSA can be classified as emerging from the branched-chain amino acids (BCAA) which include isoleucine, leucine, and valine. HICA is the degradation product of leucine, and MSA is formed from the isomerization product of isoleucine. From our study, both compounds were observed to have a toxic effect on growth, with HICA showing a slightly higher toxicity as compared to MSA. Likewise, the negative effect on productivity of these two metabolites is less than five percent which makes them predominantly growth inhibiting metabolites (see Fig. 5 I and J). Historically, branched-chain amino acid metabolites (such as isovaleric acid) have been shown to be toxic to CHO cellular growth in fed-batch cultures (Li
et al., 2010). However, to the best of our knowledge, there was no literature evidence showing these novel metabolites secreted at toxic level to cell growth through BCAA degradation pathway during fed-batch culture, which highlights the current metabolomic study as an exhaustive effort to characterize CHO cellular metabolism in different culturing settings.

Other growth medium components supplemented to CHO cells (such as arginine, tryptophan, and vitamin B12) have also been linked to the identified toxic compounds (i.e., NAP, ICA and TRI). Of the three metabolites, ICA was previously reported to secret at high level into cell culture environment in CHO-GS cell line (Li et al., 2010). It was also shown in the same study that reduction of ICA through modulation of amino acid concentration or elimination of ICA production through cell line engineering could improve CHO cellular performance. Similar finding was observed in this current metabolomics study where ICA was found to be eminently toxic to cellular performance when supplemented at high level and moderately toxic at concentration up to 1 mM (see Fig. 5 B), although it has only been reported from literature that accumulation of ICA in the spent media was less than 10 μM concentration. At 5 mM spiking concentration, ICA was reported to completely inhibit cellular growth. This tryptophan degradation metabolic product also decreased cellular productivity (see Fig. 5 J) when supplemented at micromolar level into CHO cultures corroborating the toxicity of ICA to these cell lines.

Finally, TRI and NAP which are generated as by-products of the nicotinate and arginine metabolism, respectively, were observed to be toxic to cells at lower concentration. Particularly in the study, NAP was shown to inhibit cellular growth at the measured cellular concentration (0.6 μM). When supplemented back into cell culture at millimolar level, the toxic effect was not observed, which suggests that NAP was redirected into the arginine metabolism pathway. Specifically, NAP potentially can be converted back to putrescine which is a substrate from the polyamine pathway (see Fig. 4 and Figure S2), which generates growth factors to promote cell proliferation in mammalian systems (Hölttä and Pohjanpelto, 1982) and thereby neutralizing the inhibitory effect of NAP on cell growth, as shown in Fig. 5 B. Alternatively, excess NAP could also drive the metabolic flux in the forward direction downstream of its formation and be utilized by cells to support growth. In a similar fashion, TRI was found to be less toxic as the concentration increased from micromolar level to millimolar level. TRI is a metabolite by-product of nicotinate metabolism, which is a part of vitamin and cofactor metabolism. Since these chemical compounds are supplemented in lower amount in the growth media, trace amount (about a few μM) of their metabolic by-products was shown to negatively impact cellular performance in the current study. Thus, with the aid of spiking in inhibitor spiked-in growth medium at varied concentration, the conducted study was able to characterize the varied impact of the identified metabolites on CHO cultures. The impact of the eight identified metabolites in higher cell density culture often found in fed-batch process was further examined in subsequent study as detail below.

### 3.5. Inhibitory metabolites characterization in CHO-K1 fed-batch process

In a standard industrial culture bioprocess, cells are exposed to inhibitory metabolites for an extended culture period as cells are cultivated in fed-batch mode. Thus, as an effort to validate metabolite inhibitory impact in a long-term process at higher cell density, a fourteen days fed-batch process with metabolites spiked in on Day 0 was performed. Cultivation strategy of cells followed closely to the method previously described in Section 2.1. Feeding strategy was designed based on the best performing condition as studied in the HCD fed-batch process development, namely feeding of 3 mL enriched medium B (10% of inoculation volume) starting on Day 3. Metabolites were spiked into fed-batch cultures on Day 0 at the end-of-culture concentration previously measured from samples obtained on Day 14 from the HCD fed-batch process, as shown Table 2. In this study, ICA which was proven as a growth inhibitor in past study was used as a positive inhibitor control (Mulukutla et al., 2017).

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**Fig. 4.** [PRINT IN COLOR] Metabolites Generation Pathway and Metabolic Network Mapping in Mammalian Systems. Mapping of identified metabolites and their pathway of generation to CHO metabolic network. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
At the end of the growth phase (Day 4), the VCD across all culture conditions were experimentally measured (Fig. 7 A) and IVCD for each condition was calculated (Fig. 7 B). Overall, conditions spiked with inhibitory metabolites previously identified from the batch study showed a lower growth profile as compared against the control (without spiking of metabolites). Except for the ACA spiked condition, the remaining cultures showed more than 10% IVCD decrease when compared against the control (see Fig. 7 C). From the study, it was shown that of the seven metabolites, MSA, CMP and ICA appeared to pose a stronger inhibitory impact on cell density (IVCD decrease of 24.2%, 23.2%, and 22.0% in comparison to the control, respectively).

3.6. Productivity and product quality

Monoclonal antibody production was measured from samples collected at the harvest day from the fed-batch process. Overall, titer production for cultures spiked with HICA, CMP, TRI, ICA, NAP, and MSA showed more than 10% decrease in productivity when compared against the control (without spiking of metabolites). Except for the ACA spiked condition, the remaining cultures showed more than 10% IVCD decrease when compared against the control (see Fig. 7 C). From the study, it was shown that of the seven metabolites, MSA, CMP and ICA appeared to pose a stronger inhibitory impact on cell density (IVCD decrease of 24.2%, 23.2%, and 22.0% in comparison to the control, respectively).

3.7. Validation of inhibitory metabolites in multiple mammalian cell lines

Accumulation of inhibitory metabolites during growth and production phase was also studied in different mammalian cell lines capable of expressing therapeutic proteins. Here, six of the eight novel inhibitory metabolites that accumulated during CHO-K1 fed-batch process were also found to accumulate in CHO-GS and HEK293 bioprocess. The accumulation concentration measured at harvest day was shown in Table 3. The results suggest that metabolites identified using untargeted and targeted metabolomics strategy as shown here are not strictly exclusive to CHO-K1 but can also be found across multiple mammalian systems used to express therapeutic proteins.

Interestingly, the concentration of the inhibitory metabolite ACA that accumulated during the cultivation of CHO-GS and HEK293 system is lower compared to CHO-K1. An examination of the metabolism of the individual cell lines may help to explain the observed accumulation trend of different metabolites across these cell lines. For CHO-GS, glutamine synthetase expressed during transcription allows glutamine being resynthesized from glutamate which likely lowers the deamination activity of converting glutamate into $\alpha$-ketoglutarate, therefore decreasing the amount of $\alpha$-ketoglutarate available for the TCA cycle. Reduction of $\alpha$-ketoglutarate may require upregulation of other upstream enzymatic activity to provide adequate protons for the TCA cycle. In other mammalian system, this could be achieved through higher metabolic activity of proline which yields $\alpha$-ketoglutarate as the direct beneficial by-product. However, as CHO is known to be dysfunctional in metabolism of proline, other enzymatic activity pathways must be upregulated to account for the reduction in G2F glycosylation in cultures spiked with HICA.
α-ketoglutarate, one of which is aconitase activity (Valle et al., 1973). Aconitase is known to facilitate the consumption of aconitate to form isocitrate which is precursor to formation of α-ketoglutarate. Thus, upregulation of aconitase activity reduces the accumulation of aconitate, and hence explains the reduction in culture concentrations of the metabolite ACA observed in the case of CHO-GS.

Regarding the decrease in the accumulation of ACA in the case of HEK293, a further look into the metabolic system of HEK293, suggests a plausible explanation for the observed trend. For HEK293, the expression of the c-MYC regulator gene could account for increasing the intracellular concentration of pyrroline-5-carboxylate reductase (PYCR) (Tanner et al., 2018). As PYCR facilitates the breaking down of 1-pyrroline-5-carboxylate (P5C) in the proline cycle, additional availability of PYCR may drive the accumulation of P5C followed by a consequential reduction in glutamate (Le Chatelier’s principle), and ultimately, leading to a decrease in the amount of α-ketoglutarate. The rest of the pathway follows similar reasoning as made above for CHO-GS which ultimately accounts for the observed reduction of ACA available in the TCA cycle. Interestingly, while ACA was found to accumulate in both HEK293 and CHO-GS throughout the culture duration, the measured concentration of ACA at harvest day was found to be lower in HEK293 compared to CHO–K1. Difference in level of gene expression and enzymatic activity between CHO–K1 and HEK system may provide an explanation for the differences in levels between these two hosts. As mentioned above, CHO metabolic systems are known to be proline auxotrophic which prevent any production of α-ketoglutarate through this pathway. This, however, is not the case for HEK293, therefore excess amount of α-ketoglutarate yielded from the proline metabolism cycle found in HEK293 would drive the equilibrium of the consumption reaction of ACA to the right (Le Chatelier’s principle), therefore reducing the accumulation of ACA at throughout the culture duration. A more in-depth analysis is needed to be conducted in future study to verify the observed trend in metabolites accumulation in HEK293 cell line.

The concentration of CMP, TRI, and HICA were found to be more abundant in CHO-GS. Depletion in the synthesis of α-ketoglutarate in CHO-GS due to CHO being proline auxotroph promotes the activity of other relevant metabolic pathways, namely glycolysis and the metabolism of other amino acids, including leucine and tryptophan. This would account for the increase in accumulated concentration at end-of-culture observed in the case of CMP (direct by-product of glycolysis), HICA (direct by-product from the metabolism of leucine), and TRI (direct by-product from the metabolism of tryptophane) observed throughout the study. These pathways are therefore activated to stimulate the production of α-ketoglutarate in the CHO-GS cell lines that are lost from activation of the glutamine synthase gene.

In order to evaluate if the inhibitors identified in the current study also exhibited an effect on cell growth for cell lines beyond CHO–K1, negative impact study on culture performance of CHO-GS process was conducted on the six metabolites already seen to accumulate in CHO-GS cultures (see Table 3) along with ICA and GMP. Similar to the previously conducted batch screening studies on CHO–K1 cell line, the metabolites were spiked in the CHO-GS cultures at two different concentrations (1 mM and 5 mM). Here, CHO-GS cells were cultivated in the second generation of medium A, hereby addressed as medium A 1.1 (medium A 1.1 includes similar amino acids as medium A 1.0 but eliminates or reduces other non-essential inputs). The results of the impact study of the metabolites on CHO-GS cell line study are shown in Fig. 6. The VCD profile of the condition with inhibitory metabolites spiked in on Day 0 was compared against the control group. It was observed that all experimental conditions spiked with metabolites showed inhibition on cellular growth at higher concentration (5 mM) during both exponential and stationary phase of the culture process. A similar trend was also observed for most of the metabolites at lower (1 mM) concentration, except for GMP, NAP and TRI which only showed inhibition in cell growth during the early exponential phase at 1 mM concentration. The study demonstrates the universal nature of the identified metabolites as they were shown to generate during cellular metabolism and have an impact on growth across multiple mammalian cell lines when cultivated in different growth media.
Overall, this inhibitory impact study showed that accumulated inhibitors were found predominantly in fed-batch process, which suggests that reducing certain nutrient inputs might help to mitigate the accumulation of wasted inhibitory metabolites. Perhaps, the most crucial initial step is to identify and regulate inefficient metabolic fluxes occurring throughout glycolysis and amino acids consumption. Generation of inhibitory metabolites can then be best controlled during the stage when they are generated and accumulated into the cytoplasm, prior to their release into the surrounding growth medium. Overall, the study indicates that accumulation of growth inhibitors secreted in a culture process can be controlled via up- or down-regulation of upstream metabolic fluxes where the end-of-culture metabolites was either consumed or generated. Further study regarding genetic intervention of the corresponding genes responsible for these metabolic pathways can...
perhaps offer a robust set of tools to control and manipulate the generation of growth inhibitory metabolites in a culture process of mammalian cells.

4. Conclusions

In conclusion, eight metabolites derived as by-products from the metabolism of nutrients in mammalian cells were verified to show inhibitory impact on CHO cells cultivated in batch and fed-batch process. Experimental study was conducted to verify six of the identified eight metabolites to be novel growth inhibitors: aconitic acid (ACA), 2-hydroxyisocaproic acid (HICA), methylsulfinic acid (MSA), cytidine monophosphate (CMP), trigonelline (TRI), and n-acetyl putrescine (NAP). Metabolites spiking study on CHO batch process was conducted at 1 mM and 5 mM spiking concentration. At both tested concentrations, seven out of eight metabolites (ACA, CMP, GMP, HICA, ICA, MSA and NAP) were toxic to cellular growth, whereas four out of eight (NAP, TRI, CMP and GMP) showed negative impact on titer productivity. Spiking study was also conducted on CHO fed-batch cultures at cellular accumulated level in a fed-batch process. Except for ACA, all metabolites were shown to have inhibitory impact on the growth profile, with three metabolites (MSA, CMP and ICA) showing significant degree of growth suppression. Regarding protein production, six metabolites (HICA, CMP, TRI, ICA, NAP, MSA) showed inhibitory impact on specific titer productivity and IgG glycosylation profile. Consequently, spression in glycosylated activity also suppressed the galactosylation profile, which were shown in the study to inhibit the formation of mono-galactosylated biantennary (G1F) and biantennary galactosylated (G2F) N-glycans, with a more profound impact on the latter as expected due to the additional requirement of a galactose subunit. Spiking study was also repeated on CHO-GS cell line. Of the eight metabolites included in the study, only four (CMP, GMP, NAP and MSA) showed negative impact on cellular growth at 5 mM spiking concentration. The remaining metabolites (ACA, TRI, ICA, HICA) showed growth suppression at both 1 mM and 5 mM spiking concentrations. Overall, the study suggests that accumulation of growth inhibitors secreted in cell culture process can be controlled via up- or down-regulation of upstream metabolic fluxes where the metabolites accumulated at the end-of-culture was either consumed or generated. Future studies, such as genetic intervention of the corresponding genes responsible for these metabolic pathways can perhaps offer a robust set of tools to control and manipulate the generation of growth inhibitory metabolites in a culture process of mammalian cells.

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://doi.org/10.1016/j.mec.2021.e00182.

Conflict of interests

All the authors declare no conflict of interest.

Declaration of interest

None.

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Table 3

Metabolite accumulation in CHO-K1, CHO-GS, and HEK293 production host at harvest day in fed-batch process measured through LC-MS. Here, the reported molar concentration of each metabolite was normalized against the measured viable cell density.

| Metabolite | CHO-K1 | CHO-GS | HEK293 |
|-----------|--------|--------|--------|
| ACA       | 4.5574 | 1.3474 | 17.4934 |
| HICA      | 0.0750 | 0.7713 | 0.0387 |
| MSA       | 0.0280 | 0.0632 | 0.0067 |
| CMP       | 0.0826 | 1.9144 | 0.0607 |
| TRI       | 0.0006 | 0.0198 | 0.0005 |
| NAP       | 0.0011 | 0.0049 | 0.0042 |

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