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Compartmentalized Proteomic Profiling Outlines the Crucial Role of the Classical Secretory Pathway during Recombinant Protein **Production in Chinese Hamster Ovary Cells**

Saumel Pérez-Rodriguez, Tune Wulff, Bjørn G. Voldborg, Claudia Altamirano, Mauricio A. Trujillo-Roldán, and Norma A. Valdez-Cruz*



coverage of 59% of the Chinese hamster proteome. Data are available via ProteomeXchange with identifier PXD021014. By using SAM and ROTS algorithms, 493 proteins were classified as differentially expressed, of which about 80% was proposed as novel targets and one-third were assigned to the CSP. Endoplasmic reticulum (ER) stress, unfolded protein response, calcium homeostasis, vesicle traffic, glycosylation, autophagy, proteasomal activity, protein synthesis and translocation into ER lumen, and secretion of extracellular matrix components were some of the affected processes that occurred in the secretory pathway. Processes from other cellular compartments, such as DNA replication, transcription, cytoskeleton organization, signaling, and metabolism, were also modified. This study gives new insights into the molecular traits of higher producer cells and provides novel targets for development of new sub-lines with improved phenotypes for RP production.

1. INTRODUCTION

Chinese hamster ovary (CHO) cells have been widely employed for expression of recombinant proteins (RP), both in research and biopharmaceutical industries. This expression system has been used for production of 84% of approved antibodies in 2015-2018 period, which represent over half of all approvals during this time.¹ The success of this cell line relies on several advantages such as a safety viral profile,² human compatible glycosylation,³ and availability of specific culture media and supplements,^{4,5} sub-lines with different capabilities,⁶ and improved expression vectors and selection strategies.⁷ Therefore, given the importance of CHO cells, a deeper knowledge of their biology through genomic, transcriptomic, proteomic, and metabolomic studies has been gained in the last few years.^{8–11} Transcriptomic and proteomic profiles acquired under low temperature,¹² butyrate addition,¹³ hyperosmotic pressure,¹⁴ cell engineering efforts,¹⁵ and contrasting phenotypes of protein degradation,¹⁶ production stability,¹⁷ and RP productivity,^{18–24} have led to the identification of targets related to improving productivity.

with different specific productivities were used as cellular models,

from which 4952 protein groups were identified, which represent a

Seven whole cell proteomic studies of phenotypes with different specific productivity (q_p) of fusion proteins,^{18,21} monoclonal antibodies (mAb),^{19,20,22,23} and antibody fragments²⁴ have been previously reported.

ER

Golgi Apparatus

Liquid

chromatography

A myriad of cellular processes such as chromatin organization, cell cycle, metabolism of nucleic acids, proteins, fatty acids and carbon, cytoskeleton organization, response against reactive oxygen species (ROS), and vesicle-mediated transport have been linked to productivity in these studies.¹⁸⁻²⁴ However, because this information has been obtained from whole cell extractions, highlighted categories represent abundant proteins and have limited coverage of

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Figure 1. Data processing and identification of DEPs. Cleaning, normalization, and imputation were applied to proteomic data prior to the identification of targets with a statistical differential expression between CRL-12444 and CRL-12445 cells. Gene sets and proteins were identified, mapped to *M. musculus* proteome, classified according to GO terms, and compared with previous reports. Proteins from the secretory pathway were further assigned to several biological processes. ECM: extracellular matrix; ER: endoplasmic reticulum; and PTM: post-translational modifications.

proteome from some organelles such as those from the classical secretory pathway (CSP). Due to the central role of this pathway in metabolism of proteins and lipids, organelle biogenesis, cell cycle, apoptosis and proliferation, and to be recognized as a bottleneck for protein secretion in mammalian cells,²⁵⁻²⁷ its characterization through subcellular proteomics represents a promising strategy for the identification of key targets associated to protein expression.

One established experimental approach to study individual organelles and especially the secretory pathway is the subcellular fractionation. This technique coupled to proteomics has been named subcellular proteomics,^{28,29} and its use over classical proteomics has shown a 30% increase in proteome coverage of pancreatic duct cells.³⁰ Other advantages of this approach include protein tracking, quantification of low abundance of proteins, unraveling of organelle dynamics, and comprehension of the function and regulation of little-known proteins.^{31–33} Actually, subcellular proteomics has been extensively used to study the protein composition of organelles^{34,35} and protein dynamics³⁶ and to explore new proteins involved in the secretory pathway.^{37,38} This approach has also been applied to characterize the subcellular distribution of proteins in breast cancer cells by combining

isopycnic centrifugation, gel electrophoresis, and label-free MS/MS, leading to the assignment of several cancer-related proteins to multiple subcellular locations and to the identification of targets related to various cellular processes with critical roles in breast cancer development.³⁹

Thus, in order to identify novel proteins from the secretory pathway of CHO cells linked to changes in RP production, we applied subcellular proteomics to two CHO cell lines producing a mAb against human interleukin 8 (IL-8) at different q_{p} .

Here, we took advantage of a strategy that has been effective to enrich the proteomic fractions of subcellular organelles (Figure 1), which allowed the identification of 493 differentially expressed proteins (DEPs) with statistical significance between the higher (CRL-12445) and lower producer (CRL-12444) CHO cells. These DEPs, and especially those from the CSP, will expand the engineering strategies to increase the titer and probably quality of RP. This proteomic analysis is a powerful approach to identify potential biotechnological targets that help to understand and improve RP bioprocesses.



Figure 2. Growth kinetics and metabolite profiles. (A) Viable cell concentration (circles) and viability (squares) of CRL-12444 (filled) and CRL-12445 (empty) cells were determined over time by the trypan blue dye exclusion method in a Neubauer chamber. Cells were collected for proteomic analysis at time indicated by the arrow. Concentration of glucose (B), lactate (C), glutamine (D), glutamate (E), and ammonium (F) were measured for CRL-12444 (filled circles) and CRL-12445 (open circles) cells. Standard deviation was calculated from three biological replicates.

2. RESULTS

2.1. CRL-12445 and CRL-12444 Cells Differed in Their Growth, Metabolism and Secretion Capacity. CRL-12444 and CRL-12445 cells, which secrete a mAb against human IL-8, were chosen as CHO cell models for differential proteomic comparison of the CSP. Prior to the proteomic analysis, these cells were characterized in terms of viable cell concentration, viability, metabolites, and mAb production. The growth curves and metabolite profiles are presented in Figure 2, while pH and ions profiles are shown in Figure S1. The calculated kinetics parameters are presented in Table 1. The lower producer CRL-12444 cells displayed a significantly higher specific growth rate (μ) (p < 0.01), maximum cell concentration (X_{max}) (p < 0.05), and a lower doubling time $(t_{\rm D})$ (p < 0.01) (Figure 2A and Table 1). In congruence with this faster cell growth, this cell line showed a 16.5% (p < 0.05) and 38.3% (p < 0.01) increment in glucose consumption (q_{Glc}) and lactate production (q_{Lac}) , respectively, that translated to a 1.18 times higher lactate/glucose apparent yield ($Y_{\text{Lac/Glc}}'$) (p <0.001) (Table 1). For these same cells, production rates of glutamate $(q_{\rm Glu})$ and ammonium $(q_{\rm NH_4^+})$ were also increased by 3.3 (p < 0.001) and 1.8 (p < 0.001) times, respectively (Figure 2E-F and Table 1), and glutamine was almost depleted at day 6, while a 3.29 mM concentration remained in CRL-12445 cell supernatants at this time (Figure 2D). Concentration of free glutamine was the net result of cellular import of the dipeptide alanyl-glutamine, its cleavage by peptidases, the incorporation of glutamine into cellular reactions, and its secretion to the medium.⁴⁰ Thus, given that the biochemical analyzer only quantifies free glutamine,

Table 1. Kinetic and Stoichiometric Parameters of CRL-12444 and CRL-12445 Cells

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Parameter	CRL-12444	CRL-12445
$\mu^{a} \ (\mathrm{h^{-1}})^{**}$	0.031 ± 0.002^{g}	0.024 ± 0.001
$t_{\rm D}^{\ \ b}$ (h)**	22.6 ± 1.1	28.5 ± 0.7
$X_{\rm max} (10^6 \text{ cells/mL})^{c*}$	5.73 ± 0.57	4.65 ± 0.10
$q_{\rm Glc}^{\ \ \ \ }(\mu{ m mol}/10^6\ { m cells}^*{ m day})^*$	-3.17 ± 0.26	-2.72 ± 0.05
$q_{ m Lac} \; (\mu { m mol}/10^6 \; { m cells}^* { m day})^{**}$	4.15 ± 0.32	3.00 ± 0.15
$Y_{\rm Lac/Glc}'({\rm mol/mol})^{e_{***}}$	1.31 ± 0.01	1.11 ± 0.04
$q_{ m Glu} \; (\mu { m mol}/10^6 \; { m cells}^* { m day})^{***}$	0.20 ± 0.01	0.06 ± 0.01
$q_{\rm NH_4^+} (\mu { m mol}/10^6 { m cells*day})^{***}$	0.92 ± 0.06	0.50 ± 0.03
$q_{\mathrm{Ca}^{^{2+}}} \ (\mathrm{nmol}/10^6 \ \mathrm{cells}^*\mathrm{day})^*$	-2.57 ± 0.18	-4.27 ± 0.72
$q_{\mathrm{p}}^{f_{***}}$	1.0 ± 0.11	25.6 ± 1.5

^aSpecific growth rate. The asterisks indicate parameters that were significantly different (*: p < 0.05, **: p < 0.01, and ***: p < 0.001; t-test). ^bDoubling time. ^cMaximum cell concentration. ^dSpecific consumption (-) or production (+) rate. ^eLactate/glucose apparent yield. ^fRelative specific productivity. ^gStandard deviation from three biological replicates.

specific alanyl-glutamine consumption rate could not be determined and other techniques such as chemical derivatization coupled to liquid chromatography should be used for this purpose.⁴¹ Consumption or production rates of metabolites were in full agreement with values reported for adherent and suspension CHO cells, cultured in T-flasks, Erlenmeyer flasks, and bioreactors, under batch, fed-batch, or chemostat conditions.^{4,42–49} The 95% confidence interval of mean of $q_{\rm Glc}$ [-(2.02–4.33) μ mol/10⁶ cells*day)], $q_{\rm Lac}$ (2.54–7.18 μ mol/10⁶ cells*day), $Y'_{\rm Lac/Glc}$ (1.01–1.70 mol/mol), and $q_{\rm NH_4^+}$



Figure 3. Proteins with differential expression by cellular compartment. (A) Subcellular compartments (C1-C10) obtained by differential and isopycnic centrifugation from cell homogenates. (B) Number of DEPs identified for each compartment by SAM and ROTS algorithms, which were upregulated or downregulated in the higher producer CRL-12445 cells.

(0.17–1.88 μ mol/10⁶ cells*day) of all those reports comprises the values obtained in this study. Glutamate can be consumed or secreted in dependence on the medium and additives employed.^{4,42,44,47,49} In our case, glutamate was exported to the medium and its production rate ($q_{\rm Glu}$) was in accordance with other CHO cell clones.^{4,44}

No differences were observed in profiles of pH, sodium and potassium, at any time point, while calcium was consumed by both cells (Figure S1). Calcium consumption rate (q_{Ca}^{2+}) of CRL-12445 cells was superior to that of CRL-12444 cells (p < 0.05) (Table 1). MAb q_p of CRL-12445 (higher producer) cells was 26 times higher in average than that of CRL-12444 (lower producer) cells (p < 0.001) (Table 1), allowing these two cell populations to be used for differential subcellular proteomics analysis in order to identify proteins and suggest cellular processes linked to changes in RP productivity.

2.2. MAb from Both Cell Lines Showed a Similar Biological Activity. To analyze whether a 26-fold difference in q_p between both cell lines would affect the biological activity of secreted mAb, its binding to human IL-8 was assessed by Western blot (WB). For this purpose, antibody was affinitypurified to homogeneity, as confirmed by the presence of only light and heavy chains on reducing SDS-PAGE (Figure S2A). Non-reducing SDS-PAGE revealed four protein bands (Figure S2B), probably corresponding to different assemblies of light and heavy chains. Antibody from both cell lines strongly recognized monomers (10-15 kDa), dimers (15-25 kDa), and trimers (≈35 kDa) of recombinant IL-8 obtained in Escherichia coli lysates (Figure S2C-E), as expected. These results suggested that in spite of q_p changes, the folding state and antigen recognition of secreted mAb were not noticeably different between both cell lines, which indicates that in this case, productivity differences did not alter those structural properties of mAb directly linked to antigen recognition.

2.3. Around 10% of All Identified Proteins Were Differentially Expressed between Both Cell Lines. In order to capture the CSP subproteome, subcellular organelles from both cell lines were isolated by mechanical disruption followed by their separation in sucrose gradients, from which proteins were precipitated and subjected to shot-gun proteomics (Tables S1–S2). After data processing, 4952 protein groups were identified covering about 59% of all proteins reported from two CHO cell lines and seven tissues of Chinese hamster.⁵⁰ In addition to CHO cell proteins, the concentration of mAb light chain in subcellular compartments containing endoplasmic reticulum (ER) (C1, C3, and C9) was

also measured and used as an internal control of protein expression levels between both cell lines. In agreement with measured $q_{\rm p}$ intracellular mAb was higher in CRL-12445 cells than in CRL-12444 cells in all cases, showing statistical significance for C1 and C3 compartments (Figure S3, p < 0.05). Before proceeding with differential expression analysis, a correlation test was done to assess the relationship between each pair of biological replicates. As a result of analysis, all replicates were significantly associated (p < 0.001), and the Pearson's coefficient (R^2) was equal or superior to 0.79, indicating a positive correlation and that the replicates behaved very similarly. 80 and 70% of samples from CRL-12444 and CRL-12445 cells, respectively, showed a $R^2 \ge 0.90$ (Figures S4–S7).

In the higher producer CRL-12445 cells, SAM algorithm identified 125 upregulated and 285 downregulated proteins in comparison with the lower producer CRL-12444 cells, whereas ROTS reported 66 upregulated and 71 downregulated ones, accounting for a total of 493 DEPs that represent 10% of all identified proteins (Table S3). A group of 21 upregulated and 25 downregulated proteins were simultaneously detected by both algorithms (hereinafter shared upregulated or downregulated proteins). Besides the whole analysis of all DEPs, the contribution of each subcellular compartment (C1-C10, Figure 3A) to their identification was also examined (Figure 3B). Most DEPs came from compartments C3-C5 and C7, which are mainly enriched in Golgi Apparatus, nuclei, mitochondria, peroxisomes, and ER,⁵¹ showing an overrepresentation of proteins from the CSP. Upregulated proteins were positioned principally in an unidentifiable compartment (C4) and *trans*-Golgi, while downregulated ones were highly enriched in cis-Golgi.

2.4. Gene Ontology Analysis, Modified Pathways, and Enrichment of Gene Groups from DEPs. DEPs were classified according to gene ontology (GO) enrichment analysis through PANTHER, DAVID, and KEGG tools to determine which categories were over- or under-represented in the higher producer CRL-12445 cell line. By using PANTHER gene list analysis, shared upregulated (Figure S8A,C,E,G,I) and downregulated (Figure S8B,D,F,H,J) proteins pointed out to differences in metabolism, organization, location, or biogenesis of cellular components, and other cellular processes (Figure S8A,B), that were tracked to cells and protein-containing complexes and organelles (Figure S8C,D). Membrane proteins were upregulated while those from supramolecular complexes and extracellular regions were downregulated. Although central



Figure 4. Mapping of all DEPs to the CSP. Upregulated (red) and downregulated (green) proteins were mapped to organelles by using GO terms from Uniprot and DAVID and by matching with mouse database from COMPARTMENT. COPI/II: coat complex protein I/II vesicle, ER: endoplasmic reticulum, ERGIC: ER–Golgi intermediate compartment, SV: secretion vesicle, and PM: plasma membrane.

carbon pathways (glycolysis and pyruvate metabolism) were disturbed in both cells, the enrichment of tricarboxylic cycle indicated an improved oxidative metabolism in the higher producer cells⁵² (Figure S8G). Other reduced categories in CRL-12445 cells were proline biosynthesis, signaling molecules, and calcium-binding proteins (Figure S8H,J). Vesicle transport (Figure S9E,F), protein quality control by proteasome, and synthesis of DNA precursors (Figure S9G) were other functions positively associated to the higher producer cells, while cytoskeleton regulation by Rho GTPase and formyltetrahydroformate biosynthesis were negatively linked to this phenotype (Figure S9H). In addition to the gene list analysis of PANTHER, the over-representation test indicated that CRL-12445 cells increased protein translation and ER-to-i anterograde transport, while protein representation of ribonucleoprotein-associated processes and fatty acid oxidation was reduced (Tables S4–S13).

In agreement with an active oxidative metabolism and cytoskeleton rearrangement, the NAD metabolism (Figure S10A) and Rac GTPase-binding (Figure S10E) were new categories found, as upregulated in CRL-12445 cells by DAVID analysis. Conforming to the DAVID functional annotation tool, transport, secretion, endocytosis, lipid storage, actin polymerization, synthesis of ribose phosphate and ascorbic acid, proteasome, autophagosome, lysosomes, and endosomes were also upregulated (Table S14), while calcium transport, RNA splicing, and peroxisomes were downregulated (Table S15).

Besides GO terms, the differences in cellular pathways were explored through KEGG (Figure S11). A closer inspection to these reconstructed pathways exposed that pentose phosphate pathway, aminoacyl-tRNAs, RNA degradation, ABC transporter, actin cytoskeleton, and cholesterol metabolism were positively associated to higher producer cells, while oxidative phosphorylation, fatty acid degradation, nucleotide metabolism, transcription, ribosomes, ER folding, and cell adhesion were repressed (Table S16).

GSEA analysis revealed that gene sets corresponding to metabolism and transport of amino acids, cell growth, DNA repair, endocytosis, extracellular matrix (ECM) organization, glycosylation, vesicle-mediated transport, purine metabolism, and response to ROS were highlighted in the higher producer cells. On the contrary, transcription, processing, and splicing of RNAs, ribosome biogenesis, and response to ER stress were diminished (Tables S17–S19).

In line with our results, earlier omics studies suggest that during RP synthesis, CHO cells activate vesicle transport and DNA protection and modify their metabolism of carbohydrates, lipids, and amino acids, to increase protein secretion and energy availability and to avoid ROS-induced damage.^{15,18–24,53,54} The increment of productivity appears to be mediated by a higher transcription, chromatin remodeling, translation, and protein catabolism.^{15,19,24,53} In general, higher producer cells tend to diminish proteins participating in cell proliferation, and their cytoskeleton undergoes a rearrangement. Frequently, the abundance of cytoskeleton regulatory proteins changes, causing the restructuring of the filaments and microtubules that could promote vesicle-mediated transport.^{15,22,24} The regulation of intracellular calcium and calcium-dependent responses,^{20,24,54} annexin-dependent responses,²¹ and activity of MAP kinases,^{20,53} Ras,⁵³ insulin,⁵⁵ G proteins, Rho GTPases, phosphatases, and nuclear receptors²⁴ has been associated with an enhancement of RP production, as discussed in the Supporting Information.

2.5. Relevant New DEPs Involved in RP Production Were Identified by the Subcellular Fractionation Strategy Compared to Previous Whole CHO Cell **Proteomics.** The DEPs were compared with those reported in previous studies where q_p was used as a differentiation criterion between various cell lines (Tables S20–S21),^{18–24} in order to identify common targets relevant for RP production and to compare the detection capabilities between the subcellular fractionation strategy and the classical whole cell proteomics approaches. Only 33% of upregulated proteins were present in previous reports, whereas this percentage was even lower (17%) for the downregulated ones. The large number of unmatched proteins with preceding reports draws attention to the high percentage of new targets provided in this study, which are possibly related to the improvement of RP productivity in CHO cells. In this sense, subcellular proteomics strategy can be considered as an alternative to classical proteomics to explore the molecular traits of CHO cells, which are associated to a higher production phenotype.

2.6. One-Third of DEPs Belonged to the CSP. Given that the CSP has been recognized as a bottleneck for protein production in mammalian cells,^{56–58} all DEPs were mapped to organelles of this cellular route, to elucidate which molecular processes taking place along the secretory pathway could contribute to differences in q_p between both cell lines (Figure 4). Mapping of mouse orthologues was based on manual search in the literature, Uniprot, DAVID, and GeneCards resources and by matching with the mouse database from COMPARTMENT (Tables S22-S25).59 Interestingly, on average one-third of all DEPs were assigned to the secretory pathway, of which 21% were upregulated and 79% were downregulated, and most of them have not been identified in previous whole cell proteomic studies¹⁸⁻²⁴ (68% from upregulated and 87% from downregulated targets). These proteins were mainly located to ER (74% upregulated and 67% downregulated) and Golgi apparatus (59% upregulated and 54% downregulated). About 23% of all DEPs from the microsomal gradient belonged to the secretory pathway, and around 27% of all DEPs mapped to this pathway came from the microsomes. The changes in cellular levels of this subset of DEPs could positively impact translation, folding, traffic, and modifications of RP along the secretory pathway and will shed light on processes from this pathway related to RP expression.

3. DISCUSSION

The present study used subcellular proteomics to analyze the differential expression of proteins from the CSP of two cell lines with a 26-fold difference in their mAb $q_{\rm p}$, in order to identify possible bottlenecks for protein production in CHO cells, with special attention to the secretory pathway, which has been identified as one of the principal barriers during RP production in mammalian cells.^{25–27,56–58} Despite differences in $q_{\rm p}$, the lower (CRL-12444) and higher (CRL-12445) producer cells showed a comparable biological activity of mAb, assessed by antigen recognition in a WB assay (Figure S2), indicating that the differences in $q_{\rm p}$ did not alter mAb-binding capacity in these CHO cell clones. Unfortunately, experiments that analyze the influence of $q_{\rm p}$ on mAb binding are scarce or non-existent.

Though lack of correlation between μ and q_p has been reported for a panel of recombinant mAb expressing CHO subclones,⁶⁰ other studies have documented an inverse relationship between these variables on batch and chemostat cultures,^{61–63} which is coincident with our work (Table 1). It could be suggested that at higher growth rates, there is an increasing demand for energy and biomolecule precursors for biomass synthesis, which compromises the cellular resources for RP production. In line with this explanation, a higher X_{max} (p < 0.05) was reached by the lower producer and faster growing CRL-12444 cells in comparison to CRL-12445 cells, similar to what has been reported for a ht-PA producing cell line.⁶³ On the contrary, at lower μ values, a greater proportion of resources can be used for mAb production in CRL-12445 cells, resulting in a significantly higher q_p (p < 0.001).

In order to capture the cellular processes from the CSP that could be limiting for protein production, a subcellular fractionation protocol was applied to cells collected during the exponential growth phase,⁵¹ from which 10 subcellular compartments were isolated and their proteins submitted to shot-gun proteomics. This subcellular proteomics strategy led to the identification of 493 DEPs, of which around 80% have not been described as relevant for protein production,



Figure 5. Homeostasis and stress of ER and UPR. Downregulated (green) and upregulated (red) proteins belonging to this category inhibited or activated various molecular processes that could lead to, suppress, or be triggered by ER stress. ER: endoplasmic reticulum, ERAD: ER-associated degradation, and TM: transmembrane.

providing hundreds of new targets that can be modified in CHO cells. This high number of new targets reflects the potential of subcellular proteomics over classical approaches to increase proteome coverage and the identification of low abundance cellular proteins.^{30,31,33} Actually, the 59% proteome coverage of this study exceeds that achieved in other differential proteomic studies $(5-31\%)^{18,20-22}$ which reinforces the importance of subcellular fractionation prior to proteomics to allow enrichment of low abundance of proteins, such as those from the CSP and mitochondria.^{30,51} The fact that one-third of all DEPs were assigned to the CSP (Figure 4) supports the use of this technique and highlights the large number of molecular processes from the secretory pathway that can be considered as a limiting factor for RP production in these cells. It is noteworthy that a quarter of all DEPs came from the microsomal gradient (Figure 3), demonstrating the potential of this novel sucrose gradient to study the CSP of mammalian cells.⁵¹

To comprehensively understand the biological processes associated to a higher producer phenotype, the DEPs from the CSP were grouped into eight different categories according to their known functions in protein production in this pathway (21 downregulated and 13 upregulated proteins). Additional discussion about metabolism of carbon, nitrogen, nucleic acids, proteins, cofactors, vitamins, cytoskeleton organization, and cell signaling is provided in the Supporting Information.

3.1. ER Stress and Unfolded Protein Response. ER stress and unfolded protein response (UPR) comprise eight (CLCC1, DNAJC3, EMC7, OS9, MINPP1, TMED4, UFC1, and PRKCD) and two (PITPNM1 and SURF4) proteins directly or indirectly involved in these processes, respectively (Figure 5). CLCC1 is a chloride permeable channel with low expression in CHO cells⁶⁴ whose loss increases sensitivity to ER stress and triggers an UPR.65 Its decrease in the higher producer cells could be a consequence of the absence of ER stress and UPR, and of a higher folding capacity of this cell line, explanation that could also be applied to the downregulation of DNAJC3, EMC7, OS9, and UFC1. The UPR increases the amount of DNAJC3 in ER lumen,^{66–68} where it functions as a co-chaperone with HSP40 and HSC70.⁶⁹ Another UPR-induced chaperone is EMC7, a subunit of the EMC complex that participates in ER-mitochondria tethering, folding of transmembrane proteins,⁷⁰⁻⁷² and disposal of misfolded proteins by interaction with ER-associated degradation (ERAD) machinery.⁷³ OS9 expression has also been increased after an UPR,^{74,75} which favors the degradation of glycosylated and retention of non-glycosylated substrates,^{74,76,77} stabilizes the ERAD SEL1/HRD1 complex⁷⁷ and functions in complex with BiP and GRP94 chaperones



Figure 6. Affected targets involved in protein synthesis, translocation into ER lumen, and vesicle-mediated traffic throughout the secretory pathway. Upregulated (red) and downregulated (green) targets were shown. Arrows indicated the direction of the transport. PM: plasma membrane, ER: endoplasmic reticulum, TM: transmembrane, mAb: monoclonal antibody, ERES: ER exit sites, and ERGIC: ER–Golgi intermediate compartment.

during protein folding.⁷⁶ UFC1 mediates ufmylation of various cellular targets and, although the biological function of this post-translational modification (PTM) is poorly understood,^{78–80} its known that ER stress increases the formation of ufmylation complex and, congruently, blocking ufmylation leads to ER stress.⁸¹

In the case of PRKCD, this protein kinase is recruited to ER membranes and stress fibers upon activation, where it is required for a full development of UPR and apoptosis.⁸²⁻⁸⁵ MINPP1 and TMED4 are involved in maintaining a functional apoptosome, and their cellular amounts increase as a result of ER stress.^{86,87} As a consequence of this same stress, PITPNM1 assembles into a phosphorylated PYK2/PITPNM1 complex that plays a role in calcium- and phosphoinositide-dependent signaling pathways,⁸⁸ and SURF4 initiates or maintains UPR by preventing calcium from entering the cell.⁸⁹ The downregulation of all proteins from this category supports the hypothesis that the higher producer cells have not develop an ER stress, UPR, or stress-dependent apoptosis, probably because their maximum folding capacity has not been reached yet. Among the preceding omics comparing CHO cell populations with different RP production, only one transcriptomic study suggested the upregulation of UPR and ERAD,⁵⁴ which was not subsequently confirmed by the proteomic study of these same cells,¹⁵ similar to our results. It should be noted that in particular cell lines, the combination

of intrinsic physicochemical properties of an IgG and a deficient ER export machinery can activate a full UPR.²⁵

3.2. Homeostasis of ER and Golgi Apparatus. Seven downregulated proteins were implicated in this group (CLCC1, EMC7, UFC1, PITPNM1, TMF1, GOLPH3L, and GOLGA5). The functions of CLCC1, EMC7, and UFC1 in organelle homeostasis are linked to their role in protein folding, ER stress, and UPR, while the changes in the remaining four proteins affect the Golgi structure (Figures 5 and 6). PITPNM1, a phosphatidylinositol (PI)-transfer protein, maintains the Golgi morphology by regulating diacylglycerol homeostasis,⁹⁰ and the depletion of golgin and Rab interacting protein TMF1^{91,92} leads to changes in stacking of Golgi cisternae.^{91,93} GOLPH3L overexpression induces Golgi compaction, while its depletion has opposite effects, demonstrating its function in the Golgi morphology.⁹⁴ The golgin GOLGA5 affects dramatically Golgi morphology, probably as a side effect of its functions in vesicle transport. Its downregulation leads to fragmentation⁹⁵ or dramatic loss⁹⁶ of Golgi membranes, while its overexpression could stabilize Golgi structure⁹⁷ or induce Golgi ribbon fragmentation.⁹⁶

On the other hand, five upregulated members (DDHD2, RHBDD1, SCFD1, STX17, and FKBP1A) were also classified in this category (Figures 5 and 6). SCFD1 and STX17 function in maintaining the Golgi apparatus structure by their role in protein traffic,^{98,99} with knockdown of STX17 causing

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disruption of the ER-Golgi intermediate compartment (ERGIC), fragmentation of Golgi apparatus, and a disturbed COPI vesicle localization.98 The role of DDHD2 in the structure and functions of the secretory pathway remains controversial, insomuch as its knockdown does not affect the Golgi structure,¹⁰⁰ but its overexpression can result in Golgi dispersion and aggregation of ER and ERGIC^{101,102} or no morphological changes at all.¹⁰³ RHBDD1 preserves ER homeostasis by aiding at disposal of misfolded transmembrane proteins,¹⁰⁴ and FKBP1A, a cytosolic isomerase, by stabilizing RyR and IP3R calcium receptors, ^{105,106} preventing in this way calcium leakage from ER lumen. The adjustment in the cellular concentration of proteins of this category suggests the presence of morphological changes in the CSP that are associated with differences in q_p between both cell lines, while maintaining at the same a proper CSP structure for synthesis, traffic, and modifications of recombinant and self-proteins.

3.3. Anterograde and Retrograde Transport. This category comprises seven downregulated proteins (PITPNM1, ZFPL1, GOLGB1, GOLGA5, PACS-1, TMF1, and COPG1) that participate in different transport routes (Figure 6). Downregulation of PITPNM1 inhibits export of plasma membrane (PM) proteins and glycosaminoglycans from the trans-Golgi network (TGN).⁹⁰ ZFPL1 mediates ERGIC to cis-Golgi transport of cell surface proteins, probably by acting as a tethering factor.¹⁰⁷ GOLGB1, with an analogous function to ZFPL1, mediates retrograde transport,¹⁰⁸ maintenance of Golgi resident proteins,^{109,110} and anterograde transport of ECM components¹¹¹ and PM proteins.¹¹² GOLGA5, another tethering factor^{95,113} and a RAB1 effector,^{96,97} participates in intra-Golgi retrograde transport^{95,113} and functions partially in anterograde transport of call surface proteins^{96,113} TME1 else anterograde transport of cell surface proteins.^{96,113} TMF1 also functions in intra-Golgi retrograde transport⁹² and has been involved in endosomes to TGN transport.⁹³ PACS-1 mediates transport from endosomes^{114,115} and PM^{115,116} to TGN, and probably works in other post-Golgi transport routes,¹¹⁵ through connection of cargoes and adaptor proteins (Figure 6). The downregulation of COPG1 is challenging to explain due to the wide range of transport processes in which COPI vesicles could participate.^{117–119} Since these vesicles are heterogeneous in their composition of γ and ζ subunits, coatamer populations that carry $\gamma 1$ or $\gamma 2$ subunits coexist at the same time in mammalian cells and, therefore, the deficiency of γ 1 subunit (COPG1) could be rectified by γ 2 in the higher producer cells.^{120,121} Additionally, only receptor tyrosine kinase nuclear signaling or specific transport routes in the higher producer clone could be affected by COPG1 down-regulation.^{117,118}

Regarding upregulated targets, eight members (ARF3, DDHD2, PDCD6, RAB32, RHBDD1, SCFD1, TRAPPC9, and STX17) were also included in this transport category (Figure 6). ARF3 is an ADP-ribosylation factor that supports COPI vesicle biogenesis,^{122,123} mediates transport from ER to endolysosomes,¹²⁴ endosome to PM, ERGIC to *cis*-Golgi, *cis*-Golgi to ER,^{125,126} and TGN to PM,¹²⁷ and is required for the integrity of recycling endosomes.¹²⁵ The role of DDHD2 in the early secretory pathway is controversial, being necessary for Golgi to PM¹⁰⁰ or Golgi to ER¹⁰³ transport. PDCD6 is a calcium-binding protein recruited to ER exit sites (ERES) that participates in formation and trafficking of COPII vesicles by enhancing outer coat recruitment,¹²⁸ loading of cargo and other required proteins, functioning as a cargo receptor for certain substrates,¹²⁹ and recruiting other proteins to

ERES.^{130,131} Its reported upregulation in recombinant CHO cells in comparison to non-producer cells²³ validates this protein as a relevant target for RP expression. RAB32 participates in protein traffic in the late secretory pathway, where it targets LRRK2 to transport vesicles and recycling endosomes,¹³² mediates traffic from endosomes to TGN,¹³ and maintains CI-MPR in endosomes.¹³³ RHBDD1 is an ER resident trans-membrane protease that negatively regulates exosome secretion¹³⁴ or facilitates protein secretion in microvesicles by mediating their ER to Golgi transport.¹³⁵ TRAPPC9 is a non-essential subunit of the TRAPP complex that allows shedding of the inner coat from COPII vesicles to facilitate tethering, docking, and fusion of these vesicles with their target membranes. Its binding to p150Glued links vesicle movement along microtubules with tethering, printing a suitable directionality to this movement.¹³⁶ The SNARE STX17 recycles between ER and ERGIC and has demonstrated to be essential for an adequate secretion of proteins¹³⁷ and maintenance of ERGIC and Golgi compartments.98 SCFD1 cooperates with SNARE complexes in membrane fusion events, 138-140 is required for correct targeting of PM and lysosomal proteins,^{139,141} and its overexpression has increased the q_p of RP in yeasts^{142,143} and mammalian cells.⁵⁸ It also functions in Golgi to ER traffic^{99,138} and certain intra-Golgi transport steps.⁹⁹

In summary, the transport between ER and Golgi apparatus is highly enhanced in the higher producer cells, while some intra-Golgi transport routes are downregulated for specific and non-mAb-related cargoes. Transport from and to the late secretory pathway seems to be profoundly remodeled to the detriment of PM and ECM proteins, cargoes that could be dispensable during a higher production of a mAb. Endosomesrelated transport and anterograde fusion events also appear to be highly stimulated during a higher RP production.

Omics data have been shown that an increase of q_p in different CHO cell lines leads to a rearrangement of intracellular traffic. An accumulation of adapter proteins (AP2 and AP3), molecular motors (kinesin and myosin), coat subunits (COPA, COPG2, and COPB1), small GTPases (RABs, SAR1A, and ARFs), and proteins related to vesicle formation (PDCD6) and membrane fusion (recognition and anchoring factors, NSF, SNAREs) has been observed.^{20,22–24,53,54} In common with our results, information suggests an increase in biogenesis, transport, recognition, and fusion of vesicles that could contribute to a higher productivity.

3.4. Production of ECM Components and Secretory Cargoes. Downregulated GOLGB1, SURF4, MINPP1, TMED4, FKBP14, GOLM1, PRKCD, PITPNM1, and MCFD2 (Figures 4 and 6) were classified in this category by their positive effect on ECM production. The knockout/ knockdown of GOLGB1 has shown a negative impact on secretion of proteoglycans and proteins involved in their synthesis, collagens, ECM proteins, and glycosaminoglycans.^{109,111,144} This golgin also co-localizes with dymeclin, another protein involved in collagen secretion.¹⁴⁵ PRKCD and FKBP14 are also directly linked to collagen secretion as the depletion of the former substantially decreases the transcription, transport, and secretion of collagens,^{146,147} while the latter binds to and participates in the refolding of hydroxylated states of collagens.^{148,149} PITPNM1 favors glycosaminoglycans transport from TGN to PM by controlling diacylglycerol accumulation in Golgi apparatus,⁹⁰ and MINPP1 is tightly linked to ECM production during the proliferation 150 or maturation 151,152 of chondrocytes.

Others targets involved in the secretion of soluble cargoes were also downregulated in the higher producer cells. GOLM1 has been positively correlated with secretion of matrix metalloproteinases MMP-1, 2, 9, and 13153-155 and participates directly in traffic and secretion of MMP-2 and the extracellular chaperone clusterin.¹⁵⁶ TMED4 possibly functions in transport of the hormone precursor proopiomelanocortin (POMC) in the intermediate pituitary melanotrope cells of Xenopus laevis.¹⁵⁷ MCFD2 forms a cargo receptor together with LMAN1 to transport coagulation factors V and VIII from ER to ERGIC and probably alpha1-antitrypsin.¹⁵⁸⁻¹⁶⁰ SURF4 is implicated in the secretion of a broad range of cargoes that include glycoproteins,¹⁶¹ LPP,¹⁶² and enamel ECM proteins, hormones, and proteases.¹⁶³ Since cargoes carried by SURF4 from ER to Golgi apparatus contain an ER-ESCAPE motif¹⁶³ that it is not present in the light or heavy chains of anti-IL8 mAb,¹⁶⁴ its lower abundance is unlikely to affect antibody secretion. The downregulation of all these nonessential proteins suggests that the release of cellular resources and their redirecting to the production of RP are common strategies of the higher producer cells, which could be applied during cell line development through the knockdown or knockout of these extracellular proteins and those related to their production. Indeed, the disruption of up to 14 genes coding for extracellular proteins has been shown to improve cell density, viability, and transient mAb productivity.¹⁶

3.5. Glycosylation. Glycosylation was represented by four downregulated members in CRL-12445 cells (GOLGB1, ZFPL1, GOLGA5, and TMF1). While GOLGA5 is required for a mature glycosylation pattern of lysosomal and PM proteins,95 its relevance for secreted cargoes remains unexplored. GOLGB1 knockdown decreases protein concentration and induces mislocalization of many glycosyltransferases, such as B4GALT1, MGAT1, and ST6GAL1,¹⁰⁹ and shifts N-glycans toward a high-mannose type in cancer cell lines.¹¹⁰ The Oglycosylation appears to be disturbed as well given that TMF1 maintains Golgi localization of GalNAc-T293 and knockdown of ZFPL1 decreases O-linked N-acetylglucosamine.¹⁶⁶ In this context, a carbohydrate analysis of mAb is necessary to confirm a disturbed glycosylation in detriment to O-glycan and complex N-glycan patterns in the higher producer cells. MAb N-glycosylation occurs mainly at Asn297 in the Fc region, modulating antibody effector functions through binding to $Fc\gamma$ receptors and complement activation, whereas O-glycosylation occurs at serine and threonine residues without a consensus sequence.¹⁶⁷ Therefore, since the Fab portion and not the Fc region of mAb binds to the antigen, these possible changes in glycosylation patterns of antibodies between both cell lines are not expected to impact IL-8 binding, as demonstrated by WB (Figure S2D,E). On the contrary, these plausible glycosylation changes are expected to alter mAb effector functions.

3.6. Autophagy. Autophagy could be activated in the higher producer CRL-12445 cells through downregulation of PRKCD and ZFPL1 and upregulation of RAB32, SCFD1, and STX17. Loss of PRKCD and ZFPL1 triggers autophagy in rat proximal tubular cells^{168,169} and human gastric carcinoma cell lines.¹⁶⁶ On the other hand, overexpression of RAB32 increases the number of autophagosomes, whereas its knockdown leads to autophagy blockade.^{170,171} SCFD1 is required for transport of lysosomal enzymes from ER to Golgi apparatus and, although its knockdown triggers induction of autophago-

somes as a consequence of ER stress and UPR, autophagy does not occur in its absence because of the lack of lysosomal enzymatic activities.¹⁴¹ The SNARE STX17 is recruited to autophagosomes upon autophagic stimuli,^{172,173} where it binds to other SNAREs,¹⁷⁴ mediates the recruitment of protein complexes required for maturation or fusion of autophagosomes,^{175,176} and participates in autophagosome–lysosome fusion.^{172,174,176} Furthermore, STX17 could increase the transcription of proteins involved in lysosomal functions and autophagy.¹⁷⁷

Autophagy has been described as a survival mechanism of eukaryotic cells to protect themselves from stressful conditions, provide the necessary energy and biomolecule precursors, and remove damaged organelles.¹⁷⁸ In the case of CHO cells, this process can be activated by nutrient depletion, hyperosmotic stress, and sodium butyrate addition,^{178,179} conditions that are absent from the exponentially growing cells sampled in this study. Besides a positive impact of autophagy on viability and longevity of cell cultures,^{180,181} its chemical induction by 3-MA^{179,182,183} or rapamycin¹⁸⁰ on CHO cells has been associated with an increase of q_p or product titer. After all, a profound study is becoming mandatory to elucidate the relationship between basal autophagy and q_p in recombinant CHO cells during the exponential growth phase.

3.7. Proteasomal Activity. The upregulation of two critical chaperones, PSMG1 and POMP, that participate coordinately in the assembly and maturation of proteasomes,184-187 proposes that the proteasomal activity could be enhanced in the higher producer cells. PSMG1 participates in α -ring assembly of proteasomes, prevents premature nuclear translocation of their intermediates, and favors an adequate folding for further interactions.^{184–186} Subsequently, POMP recruits this α -ring and the β subunits to ER membranes for formation and dimerization of hemi-proteasomes and mediates maturation of 20S proteasomes.^{185,186,188} In view of the fact that PSMG1 and POMP are continuously degraded during maturation of 20S proteasomes,^{185,187} increased levels of these proteins are required to sustain or increase the proteasomal activity. Their role in maintaining active proteasomes has been essential to sustain cell proliferation, signaling, and anti-oxidant defenses, avoid apoptosis, and support ER homeostasis.^{188–191}

3.8. Protein Synthesis and Translocation into ER Lumen. The two upregulated members of this group (SRP72 and SRPA) act coordinately during the synthesis of proteins and their translocation into ER lumen, which indicates a high upregulation of this pathway in the higher producer cells (Figure 6). SRP72 is a member of the signal recognition particle (SRP), a complex that mediates mRNA-ribosome targeting to Sec61 translocon and favors signal peptide removing.^{192–194} SRP72 is also required for nuclear export, stability, and function of SRP.^{192,194–196} SRPA, the α -subunit of SRP receptor (SRPR),^{197,198} is a peripheral membrane protein that targets SPR ribosomes to ER membranes through its binding to the β -subunit of SRPR.^{197,199} Immediately after targeting, GTP is incorporated into the complex, mRNAribosome cargo is transferred to Sec61 for the subsequent translocation of nascent polypeptide into ER lumen, and the SRP-SRPR complex is dissociated.¹⁹⁷⁻¹⁹⁹ SRPA also favors the SRP pathway for protein translocation by displacing Sec62 from Sec61-Sec62 complexes and making Sec61 available for SRP.²⁰⁰ In CHO cells, endogenous levels of SRPA have demonstrated to positively correlate with an increase in productivity of therapeutic mAbs.²⁰ Actually, the overexpression of most components from the SRP pathway, including SRPA, has substantially increased the q_p of CHO cell clones producing trastuzumab and infliximab,²⁰¹ which is in accordance with the upregulation of proteins from this pathway found in our study.

4. CONCLUSIONS

In the present study, a subcellular proteomics strategy was applied to two CHO cell lines producing a mAb, with a 26-fold difference in their q_{p} , to identify targets from the secretory pathway associated with RP production. Metabolic analysis showed that an efficient consumption of glucose and glutamine, lower production of harmful metabolites, and an improved oxidative metabolism could be used as high productivity markers during clonal selection. Approximately, 80% of 493 DEPs were recognized as new targets, of which a third was assigned to the secretory pathway, demonstrating a greater capacity of subcellular proteomics to identify low abundance of proteins compared to classical proteomic strategies. Differential proteomic comparison indicated that an overexpression of proteins involved in protein synthesis and translocation into ER lumen, autophagy, proteasomal activity, and vesicular trafficking, and a downregulation of those related to the production of ECM and secretory cargoes, represent viable strategies to increase product titer during cell culture bioprocesses. On the other hand, increased ER stress, UPR, and ERAD were associated with a lower productivity, suggesting that cells showing these traits should be discarded during clone isolation. Restructuring of intra-Golgi transport, morphological changes of secretory pathway, and mAb PTMs need further studies for their validation and understanding. The approach applied in the present work allowed the identification of hundreds of new protein targets that will help to understand the biological processes associated with protein productivity and provided the basis to design new sublines with novel gene modifications and stronger capabilities for RP production. The cellular concentration of those proteins identified in this and previous classical proteomic studies as differentially expressed should be manipulated in future by genetic approaches in recombinant CHO cells producing mAbs or other RPs, in order to confirm which of them might impact productivity. Future subcellular proteomics will allow to increase the number of targets related to q_{p} , in comparison to the classical approaches performed until now. This strategy proved to be an effective tool to gain a deeper insight into the molecular processes related to protein production in CHO cells.

5. EXPERIMENTAL PROCEDURES

5.1. Cell Lines and Culture Conditions. Cell lines CHO DP-12 clone #1933 CRL-12444 and clone#1934 CRL-12445, which secrete a mAb against human IL-8, were acquired from American Type Culture Collection (ATCC) and adapted to growth in CDM4CHO medium. Cells were seeded at 0.50×10^6 cells/ml in 250 mL Erlenmeyer flasks with a filling volume of 20% in CDM4CHO medium (Hyclone, Logan, UT, USA) supplemented with 6 mM stable glutamine (L-alanyl-L-glutamine dipeptide, Biowest LLC, Kansas City, MO, USA), 0.002 mg/mL Humulin N (Eli Lilly, Indianapolis, IN, USA), and 200 nM methotrexate (Pfizer, New York, NY, USA), at 60 rpm, 37 °C in a 5% CO₂ atmosphere in a humidified incubator. Cell concentration and viability were recorded every 24 h by

cell counting in a Neubauer chamber using the trypan blue dye exclusion method.

Each biological replicate represents a different frozen vial from a working cell bank. Biological triplicates were used for the kinetic and metabolic characterization of cultures. Two biological replicates, collected during the exponential growth phase (72 h) from each cell line, were processed for the subcellular proteomic analysis, where each replicate was a pool of nine Erlenmeyer flasks, in order to collect enough cells.

5.2. Quantification of Metabolites, lons, and pH. Concentration of glucose, lactate, glutamine, glutamate, ammonium, sodium, potassium, and calcium, and pH were measured in supernatants every 24 h by using BioProfile FLEX2 Automated Cell Culture Analyzer (Nova Biomedical, Waltham, MA, USA). Specific consumption or production rates were calculated from the exponential growth phase as the ratio between the net concentration of analyte and integral viable cell concentration (IVCD), determined as area under curve by trapezium rule using GraphPad Prism Software v5.01 (GraphPad Software, San Diego, CA, USA).

5.3. Quantification of q_p . MAb concentration was measured in supernatants every 24 h by using Human IgG ELISA Quantitation Set (E80-104, Bethyl Laboratories, Inc., TX, USA), according to manufacturer's protocol. SigmaFast OPD substrate (Sigma-Aldrich, Merck KGaA, Darmstadt, Germany) was prepared according to manufacturer's recommendations and incubated at room temperature for 15 min. The reaction was stopped with 10% (v/v) HCl, and the absorbance was recorded at 490 nm. q_p was calculated from the exponential growth phase as the ratio between the net product titer and IVCD.

5.4. MAb Purification. MAb purification was carried out under native conditions to preserve its biological activity. The mAb was purified by Protein A-Agarose affinity chromatography (MabSelect SuRe, GE Healthcare Bio-Sciences, USA) from two culture replicates of each cell line. The supernatant was centrifuged, diluted two-fold in equilibrium buffer (150 mM NaCl, 20 mM sodium phosphate, pH 7.2), 0.2 μ m filtered, and loaded into a 5 mL column at a flow rate of 2 mL/min. After washing the column with equilibrium buffer until absorbance has reached the baseline, antibody was eluted with 0.1 M sodium citrate (pH 3.0), neutralized with 5% (v/v) 1 M Tris-HCl (pH 9.0), and extensively dialyzed against phosphate buffer (pH 7.4) (137 mM NaCl, 2.7 mM KCl, 8.1 mM Na₂HPO₄, 1.8 mM KH₂PO₄) at 4 °C.

5.5. Human IL-8 Expression in *E. coli.* BL21(DE3) cells were transformed with IL-8-pMCSG7 plasmid (DNASU Plasmid Repository, Arizona State University, AZ, USA)²⁰² and cultured in ampicillin-supplemented LB medium, at 37 °C and 150 rpm. Expression of human IL-8 was induced with 1 mM IPTG for 4 h. Next, cells were centrifuged at 8161g for 10 min and disrupted by sonication (Soniprep 150, MSE, Heathfield, East Sussex, UK) in lysis buffer (100 mM NaCl, 1 mM EDTA, 50 mM Tris–HCl pH 8.0). Homogenates were solubilized in isoelectric focusing (IEF) buffer [7 M urea, 2 M thiourea, 2% (w/v) CHAPS, 40 mM DTT], clarified by centrifugation, precipitated by acetone, and solubilized again in IEF buffer. Proteins were quantified and subjected to reducing SDS-PAGE.

5.6. Determination of Protein Concentration. Protein concentration was determined by the Bradford method in 96-well microplates using Dye Reagent Concentrate (Bio-Rad, Hercules, CA, USA), and bovine serum albumin (GE

Healthcare Bio-Sciences, USA) was used as standard, according to manufacturer's recommendations. The samples stored in IEF buffer were diluted 5 times in MilliQ water before quantification.

5.7. SDS-PAGE for Analysis of Affinity-Purified mAb and Human IL-8. Laemmli buffer was added to samples at a final composition of 60 mM Tris-HCl pH 6.6, supplemented with 10% (v/v) glycerol, 70 mM sodium dodecyl sulfate (SDS), and 0.2 mM bromophenol blue, with (reducing conditions) or without (non-reducing conditions) 2.5% (v/ v) 2-mercaptoethanol. Next, these samples were boiled at 95 °C for 5 min, with the exception of those stored in IEF buffer, centrifuged at 8161g for 5 min, and applied to 7.5, 12, or 15% SDS-polyacrylamide gels. Page Ruler Prestained Protein Ladder (ThermoFisher Scientific) was selected as the molecular weight (MW) marker. Samples were resolved in a SE260 Mighty Small II Deluxe Mini Vertical Protein Electrophoresis System (Hoefer, Holliston, USA) at 60 mA, using Tris-Glycine pH 8.3 [25 mM Tris, 192 mM Glycine, 0.1% (w/v) SDS] as running buffer. Gels were stained with Coomassie Brilliant Blue R250 (Sigma-Aldrich, Merck KGaA, Darmstadt, Germany) and destained in 5% (v/v) methanol and 7.5% (v/v) acetic acid.

5.8. Protein Precipitation. Proteins from *E. coli* homogenates and sucrose gradients were precipitated by acetone, as described by Crowell et al.²⁰³ and Pérez-Rodriguez et al.²⁰⁴ Briefly, NaCl was added at a final concentration of 100 mM, followed by addition of 4 volumes of cold 80% (v/v) acetone and overnight incubation at -20 °C. After centrifugation at 16,000g for 25 min, the protein precipitate was washed twice with 4 volumes of cold 80% (v/v) acetone and air-dried.

5.9. MAb Binding to Human IL-8 by WB. Activated polyvinylidene fluoride membranes and reduced 15% SDS-PAGE gels, with 30 μ g of proteins by lane from chemically induced *E. coli* homogenates, were soaked in transfer buffer [20] mM Tris, 154 mM glycine, 0.08% (w/v) SDS, 20% (v/v) methanol] for 5 min. Protein transfer was carried out in a Trans-Blot SD Semi-dry Transfer Cell (Bio-Rad, Hercules, CA, USA) at 20 V for 60 min and verified by Ponceau staining [0.5% (w/v) Ponceau S, 1% (v/v) acetic acid]. Membranes were blocked in 3% (w/v) skimmed milk and 0.05% (v/v) Tween-20 in phosphate buffer for 1 h and incubated with 5 μ g/mL of purified anti IL-8 antibodies for 2 h. HRPconjugated anti-human antibody was diluted 1200-fold and incubated with membranes for 1 h. Membranes were revealed with 0.05% (m/v) 3,3'-diaminobenzidine and 0.001% of 30% H₂O₂ solution in phosphate buffer.

5.10. Subcellular Fractionation. All protocols for subcellular fractionation comprising cellular disruption and differential and isopycnic centrifugation have been described previously⁵¹ and are available at dx.doi.org/10.17504/protocols.io.bf9sjr6e and dx.doi.org/10.17504/protocols.io.bf9sjr6e and dx.doi.org/10.17504/protocols.io.bgc4j-syw. In brief, cells were suspended in HEPES buffer (1 mM EDTA, 10 mM HEPES, pH 7.4), incubated on ice for 30 min, and broken up with 25 strokes in a Dounce homogenizer. Cold sucrose was added to restore osmolarity at 0.25 M, and nuclear, mitochondrial, and microsomal pellets were collected at 3000g for 10 min, 9000g for 15 min, and 100,000g for 1 h at 4 °C, respectively. Supernatant from the last centrifugation step was named cytosol. Pellets were diluted in 0.25 M sucrose and separated in sucrose gradients at 154,693g for 3 h at 4 °C (Beckman Coulter, IN, USA). 30–60 and 10–60% sucrose

gradients were employed for nuclear and mitochondrial suspensions and for microsomal preparations, respectively. Proteins from all isolated subcellular fractions were acetoneprecipitated and used for MS/MS analysis.

5.11. Mass Spectrometry Analysis. 200 μ g of precipitated proteins were solubilized in 50 μ L of guanidine hydrochloride (GuHCl) buffer [6 M GuHCl, 5 mM Tris (2-carboxyethyl) phosphine, 10 mM chloracetamide, 100 mM Tris–HCl, pH 8.5] and incubated at 99 °C for 10 min. Of these, 100 μ g was digested with trypsin for 12 h, after which trifluoroacetic acid was added at 0.5% (v/v), and 20 μ g of digested peptides were stage tipped according to Rappsilber et al.²⁰⁵

Liquid chromatography was developed on a capillary-flow UltiMate 3000 RSLCnano system with a capLC system (Thermo Fisher Scientific, Waltham, MA, USA) coupled to a 15 cm C18 easy spray column 50 μ m \times 150 mm, 2 μ m Acclaim PepMap C18 column at 1.2 μ L/min (Thermo Fisher Scientific, Waltham, MA, USA). Using a stepped 3-45% acetonitrile gradient for 120 min, the samples were sprayed into a Q-Exactive HF-X mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) operated in the Top 12 Datadependent acquisition mode. First of a full scan was collected at 60,000 resolution, maximum injection time 50 ms, AGC Target 3.0×10^6 , followed by up to 12 MS2 scans at 15,000 resolution with maximum injection time 30 ms, dynamic exclusion set to 25 s, and higher-energy collisional dissociation (HCD) collision energy at 28%. Data were analyzed using MaxQuant software, carbamidomethyl and oxidation of methionine residues were established as fixed and variable modifications, respectively, one missed cleavage was allowed, and false discovery rate (FDR) was fixed at 1%. Data were searched against the CHO proteome from Uniprot (UP000001075) while including a list of known contaminants. The concentration of mAb light chain in intracellular compartments containing ER was measured and used as an internal control of cellular protein expression levels. MAb light chain sequence was added to the search database.¹⁶⁴ Tolerance of 20 ppm was set for the first search mass and MS/MS. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE²⁰⁶ partner repository with the data set identifier PXD021014.

5.12. Processing, Comparison, and Classification of **Proteomic Data.** A flow diagram that represents data processing, identification of DEPs and gene groups, and their classification into GO categories is depicted in Figure 1. Contaminants and groups identified in reverse database or by PTM were discarded, and protein groups were reduced to one member. All subsequent analyses for each subcellular compartment were carried out in R language.²⁰⁷ The two best-performing normalization methods from Normalyzer v1.1.1²⁰⁸ were selected for all further analyses. Missing values from each group were imputed by using the quantile regression imputation of left-censored data (QRILC) method.²⁰⁹

Protein expression was compared by using SAM²¹⁰ and ROTS²¹¹ algorithms. A cut-off of two-fold change and FDR ≤ 0.05 were used in SAM package, while B = 1000, K = 2000, and FDR ≤ 0.05 were used for ROTS. DEPs were aligned against *Mus musculus* proteome (UP000000589) using BLAST + v2.2.24 and mapped and classified by BlastKOALA from KEGG,²¹² PANTHER v14.0,²¹³ and DAVID v6.8 (EASE < 0.05).²¹⁴

Defined sets of genes were statistically compared using the javaGSEA desktop application.²¹⁵ *M. musculus* GSEA GO terms were obtained from prebuilt gene sets of GO2MSIG.²¹⁶ Collapse data set to gene symbols and metric for ranking genes were set to "false" and "diff of classes," respectively. FDR <0.25 and p < 0.05 were selected as the cut-off value for statistically significant gene sets.²¹⁵

5.13. Statistical Analysis. Pearson's correlation test was carried out between each pair of biological replicates by using *ggpubr* R package,²¹⁷ to evaluate their relationship. The concentration of mAb light chain in subcellular compartments containing ER and growth and metabolic parameters were compared between both cell lines by using Student's *t*-test in GraphPad Prism Software v5.01 (GraphPad Software, San Diego, CA, USA).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.0c06030.

Profiles of pH and cations in cell supernatants; purification and biological activity of mAb; intracellular concentration of mAb light chain; correlation analyses between replicates; classification of DEPs by PAN-THER, DAVID, and KEGG; differential proteomic analysis of glycolysis, lipid synthesis, pentose phosphate pathway, TCA cycle, synthesis of cholesterol and proline, fatty acid degradation, and one carbon metabolism; sample identity by each experiment and cell line; classification of altered proteins by KEGG; and differentially enriched gene sets by GSEA (PDF)

Unprocessed MaxQuant output; DEPs by ROTS and SAM algorithms; classification of altered proteins by PANTHER and DAVID; comparison of DEPs with previous proteomics studies; and mapping of altered proteins to the CSP (XLSX)

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Notes

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