Versatile microscale screening platform for improving recombinant protein productivity in Chinese hamster ovary cells

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Chinese hamster ovary (CHO) cells are widely used as cell factories for the production of biopharmaceuticals. In contrast to the highly optimized production processes for monoclonal antibody (mAb)-based biopharmaceuticals, improving productivity of non-mAb therapeutic glycoproteins is more likely to reduce production costs significantly. The aim of this study was to establish a versatile target gene screening platform for improving productivity for primarily non-mAb glycoproteins with complete interchangeability of model proteins and target genes using transient expression. The platform consists of four techniques compatible with 96-well microplates: lipid-based transient transfection, cell cultivation in microplates, cell counting and antibody-independent product titer determination based on split-GFP complementation. We were able to demonstrate growth profiles and volumetric productivity of CHO cells in 96-half-deepwell microplates comparable with those obtained in shake flasks. In addition, we demonstrate that split-GFP complementation can be used to accurately measure relative titers of therapeutic glycoproteins. Using this platform, we were able to detect target gene-specific increase in titer and specific productivity of two non-mAb glycoproteins. In conclusion, the platform provides a novel miniaturized and parallelisable solution for screening target genes and holds the potential to unravel genes that can enhance the secretory capacity of CHO cells.

The production of recombinant therapeutic proteins is a rapidly growing industry with a global sales value of approximately $140 billion in 20131. The main production host of biopharmaceuticals is Chinese hamster ovary (CHO) cells2, the productivity of which has improved substantially since the mid-1980s with focus on media composition and bioprocess design3,4. Production of therapeutic monoclonal antibodies (mAbs) in CHO cells in optimized bioprocesses typically reaches 5 g/L titers2 and specific productivity ($q_p$) of 50–90 pg per cell per day (pcd)3 resulting in production costs accounting for only 1 to 5% of the sales price5. However, significantly lower titers and $q_p$ are more common for glycosylated non-mAb biopharmaceuticals5,6. Thus, increasing titer and $q_p$ for non-mAb glycoproteins holds the potential to reduce production costs significantly.

Genetic engineering of CHO host cells is a well-established method for improving productivity by enhancing cell survival and $q_p$, or increasing cell growth and $q_p$. Several studies have demonstrated positive effects of overexpressing genes on productivity of recombinant glycoproteins in CHO cells (for reviews see7,8). However, contradictory effects on $q_p$ have been reported for some genes8,9. These discrepancies are likely a combined effect of multiple factors. For example, when working with stable clones, observed effects may be a consequence of clonal variation6,10. Moreover, the effects of ectopic expression of recombinant genes likely depend on physicochemical characteristics of the therapeutic glycoprotein9. A transient transfection-based overexpression screening platform has the potential to address these issues, because it enables complete interchangeability of which genes (hereafter referred to as target genes) and model therapeutic glycoproteins (hereafter referred to as model proteins) to express. Furthermore, clonal
Generic antibody-independent product titer determination methods would facilitate complete interchangeability of model proteins because specific antibodies for each individual model protein would not be needed. Split-GFP complementation-based product titer assays provide such an antibody-independent solution.

Recent advances in the split-GFP complementation technique have enabled a relatively small part (16 amino acid residues) of GFP to be tagged to the protein of interest; which is the 11th \( \beta \)-strand of GFP’s \( \beta \)-barrel (S11). The remaining part of GFP (GFP1–10; 214 amino acid residues) has been optimized for complementation and solubility (GFP1–10OPT). Separately, S11 and GFP1–10OPT are principally non-fluorescent; however, when the two fragments are mixed, the chromophore of GFP is formed and fluorescence complementation takes place. Fluorescence intensity upon complementation is proportional to the concentration of the S11-tagged protein when GFP1–10OPT is in excess, making this technique useful for product titer determination. A variant of the S11 peptide (M3 variant; S11_M3) was selected based on good complementation characteristics and minimal perturbation on solubility of tagged proteins in *E. coli*. The S11_M3-tag is therefore less likely to perturb folding processes and solubility than previous, larger versions of the split-GFP tag.

In the present study, we chose two non-mAbs as model proteins: human \( \alpha \)-1-antitrypsin (h\( \alpha \)1AT) and human C1 esterase inhibitor (hC1INH). h\( \alpha \)1AT is a protein of 394 amino acid residues with three N-glycans; hC1INH (478 amino acid residues) contains six N-glycans and at least seven O-glycans. Currently, h\( \alpha \)1AT and hC1INH are not produced commercially in cultured mammalian cells but are derived from human plasma for therapeutic use. In addition, recombinant hC1INH (Rucosent®) produced in milk of transgenic rabbits has recently been approved as a biopharmaceutical product.

We set out to establish a miniaturized and parallelisable transient transfection-based screening platform to improve the productivity of therapeutic glycoproteins in CHO cells (Fig. 1). The platform combines four techniques suitable for the 96-well format: (1) Lipid-based transfection and transient overexpression (2) Cell cultivation in half-deepwell (HDW) microplates. (3) Fluorescence image cytometry-based cell counting. (4) Split-GFP-based product titer assay. We wanted to investigate the general applicability and versatility of the platform. To this end, we analysed the effect on productivity of h\( \alpha \)1AT and hC1INH of overexpressing previously published target genes in CHO cells. By using this screening platform, we were able to demonstrate gene-specific increase in titer and \( q_p \) for our two model proteins.

**Materials and Methods**

**Plasmids and primers.** All primers used in this study are listed in Supplementary Table S1. The elements inserted into all constructs were verified by sequencing, and purified plasmid was obtained using the Nucleobond® Xtra Midi Kit (Machery-Nagel, Düren, Germany) according to the manufacturer’s instructions.

Plasmids encoding Rituximab and codon-optimized human Erythropoietin (hEPO) have previously been described. The human genes *Serpina1* and *Serping1*, which encode h\( \alpha \)1AT and hC1INH, respectively, either untagged or tagged with S11_M3 or S11_H7 (H7 variant of S11), were subcloned into pcDNA3.1(+)/Zeo (for schematic overview see Supplementary Fig. S1a; for sequences, see Supplementary Table S2).

Solid-phase cloning (SPC) was used to fuse the S11_M3-tag to the C-terminus of hC1INH. pcDNA3.1(+)/Zeo encoding hC1INH (PL_hC1INH_notag; Supplementary Table S3) was PCR amplified using Phusion® Hot-Start Flex (New England Biolabs, Ipswich, Massachusetts, USA) and primers #131 and #132. Primer #133 was designed to include 30 bases for hybridization, an amino acid linker (GGGSGGGS), and the S11_M3 sequence. The oligo
(primer #133) was genetically coupled to the C-terminus of Serping1 by head-to-tail SPC on a Magnatrix 1200 workstation (NorDiag AS, Oslo, Norway).

The remaining hO IAT and hC1INH constructs (Supplementary Table S4) were cloned by seamless PCR-based uracil-specific excision reagent (USER) fusion cloning, essentially as previously described23, using primers designed by the AMUSER software23. In brief, PCR amplicons (Supplementary Table S5) were made using the proofreading polymerase PfuTurbo and assembled by treating with USER Enzyme (New England Biolabs) and transforming into E. coli One Shot® Mach1™ competent cells (Life Technologies, Thermo Scientific, Rockford, IL).

All target genes were cloned into a vector harbouring an ‘Nt.BbvCI/PacI A USER cassette (PL_TGExpr) (Supplementary Fig. S1b). PL_TGExpr was made using Flexible Assembly Sequence Tag-USER-based cloning (FAST-USER) as previously described24 (Supplementary Table S6). Coding sequences of target genes were PCR-amplified using PfuTurbo polymerase and primers designed by AMUSER software (Supplementary Table S5). Coding sequences were inserted into the ‘Nt.BbvCI/PacI A USER cassette of PL_TGExpr as previously described25 (Supplementary Table S6).

**Cell cultivation.** CHO-S suspension cells (Life Technologies) were grown in CD CHO medium supplemented with 8 mM L-glutamine and 2 μL/mL anti-clumping agent (Life Technologies). Cells were expanded in Corning vent cap shake flasks (Sigma-Aldrich, St. Louis, MO) in a humidified incubator at 120 rpm (25 mm orbit), 37 °C, and 5% CO₂.

**Transfection.** Transfection in shake flasks, 6-well plates and 96-well microplates is described in Supplementary Information.

**96-well based cell counting assay.** Dye master mix was made daily: CD CHO + 8 mM L-glutamine + 5 μg/mL Hoechst-33342 (Life Technologies) + 0.4 μg/mL propidium iodide (Life Technologies). Master mix (200 μL) and cell suspension (3 μL) were mixed in a 96-well optical-bottom microplate (Greiner Bio-One, Frickenhausen, Germany) and cells were incubated for 40 min at RT. Image cytometry analysis was performed on a Celigo Imaging Cell Cytometer (NexCels Bioscience, MA, USA). Cells were identified using the blue channel (Hoechst-positive cells), and the red fluorescence channel (propidium iodide) was used to detect dead cells. Total cell density was computed from the total number of identified cells in the well. Cell viability was defined as the percentage of propidium iodide-negative cells. Based on total cell density and viability, viable cell density (VCD) was determined.

For comparison, VCD measurements were performed on a NucleoCounter NC-200 Cell Counter (ChemoMetec, Allerod, Denmark) using Via1-Cassettes™ using a Viability and Cell Count Assay (NucleoView software ver. 1.1.18.1) according to the manufacturer’s instructions.

**Deglycosylation of proteins.** N-glycans on proteins in supernatant samples were removed by PNGase F treatment (New England Biolab) under denaturing conditions according to the manufacturer’s instructions. N-glycans and O-glycans on proteins in supernatant samples were removed using Protein Deglycosylation Mix Kit (New England Biolab) under denaturing conditions according to the manufacturer’s instructions.

**Western blot analysis.** All samples for Western blotting were subjected to reducing SDS-PAGE on NuPAGE® Novex® 4–12% Bis-Tris Protein Gels (Life Technologies) in MOPS running buffer before transfer onto nitrocellulose membrane using the iBlot® 2 Dry Blotting System (Life Technologies). Primary antibodies: polyclonal goat hO1AT antibody (#ab7633, Abcam, Cambridge, UK) and rabbit polyclonal hC1INH antibody (#ab97348, Abcam). Secondary antibodies: horseradish peroxidase (HRP)-conjugated polyclonal donkey anti-goat IgG secondary antibody (#31458, Pierce, Thermo Scientific) and HRP-conjugated polyclonal goat anti-rabbit IgG secondary antibody (#ab6741, Abcam). Bound antibody was detected with ECL Prime Western Blotting Detection Reagent (GE Healthcare, Little Chalfont, United Kingdom).

**Split-GFP product titer assay.** Split-GFP product titer assay was performed essentially as described elsewhere, but with minor changes to the protocol (see Supplementary Information).

**Protein quantification by ELISA and bio-layer interferometry.** The hO1AT titer in supernatant samples was determined using a hO1AT ELISA Kit (#OKIA00048; Aviva Systems Biology, San Diego, CA, USA) according to the manufacturer’s instructions. The hO1AT calibrator from the kit (lyophilized serum) was used as standard. The titer of hC1INH in supernatant samples was determined using a hC1INH ELISA pair set (#SEK10995; Sino Biological Inc., Beijing, China) according to the manufacturer’s instructions. Purified recombinant his-tagged hC1INH (#10995-H08H, Sino Biological) was used as standard. The titer of hEPO in supernatant samples was determined using a hEPO ELISA Kit (#ab119522, Abcam) according to the manufacturer’s instructions. Purified recombinant hEPO (#Z02975-50, GenScript, Piscataway, NJ, USA) was used as standard. The Rituximab titer in supernatant samples was determined by bio-layer interferometry using an Octet RED96 (ForteBio, Pall, Menlo Park, CA, USA) as previously described, but with an increased shaking speed of 1000 rpm.

**Calculation of productivity parameters.** Integral of viable cell density (IVCD) and daily specific production rate were calculated as described elsewhere, but with minor changes to the protocol (see Supplementary Information).

**Statistical analysis.** Data (fold-changes) were tested for normality by D’Augustino-Pearson omnibus test using GraphPad Prism software (version 6.05 for Windows, GraphPad Software, La Jolla, CA, USA). Data (non-transformed and log2-transformed) in some groups did not pass the normality test, so statistical significance (p < 0.05) was assessed by performing the Mann-Whitney nonparametric test (two-tailed) on non-transformed fold-changes using GraphPad Prism software.
Results

Establishment of a 96-well-based cell-counting assay. In order to monitor cell growth in 96-well microplates, we wanted to use a 96-well-based cell-counting assay compatible with small sample volumes. Based on a study by He and co-workers, we established an assay that combines viability stain and 96-well-based imaging cell cytometry using 3 μL cell culture per sample (Fig. 1). Hoechst stain was used to identify the total number of cells (dead and alive) and propidium iodide stain was used to identify dead cells.

Overall, VCD measurements of the 96-well-based assay were consistent with results obtained with the commercially available acridine orange- and DAPI-based NucleoCounter NC-200 Cell Counter (Fig. 2a,b). For analysis by the NucleoCounter NC-200 Cell Counter, dilution of cell cultures in which VCD exceeds 5 × 10^6 cells/mL is not recommended. Figure 2. Wide linear range and low variation in 96-well-based cell counting assay. Comparison of NucleoCounter® NC-200 and 96-well-based (combined Hoechst and propidium iodide stain) cell-counting assay (eight technical replicates). (a–d) A dilution series of exponentially growing healthy CHO-S cells was analyzed. (a) Viable cell density (VCD) is plotted against relative cell concentration. Black rectangles and red circles denote data points from 96-well-based cell-counting and NucleoCounter assay, respectively. Mean and standard deviation are depicted. The dashed black line and solid red line show trend lines from linear regression of 96-well-based cell-counting (R² = 0.999) and the NucleoCounter assay (R² = 0.998), respectively. The encircled data point was found to be an outlier in the linear regression analysis. (b) Zoomed view of the graph shown in panel (a). (c) Coefficient of variation (CV) of VCD measurements at different cell concentrations. The x-axis denotes relative cell concentration and VCD values in brackets (×10^5 cells/mL). White and grey bars denote the NucleoCounter and 96-well-based cell-counting assay, respectively. (d) Viability at different cell concentrations. The x-axis is as described in panel (c). Mean and standard deviation are depicted. (e–f) VCD and viability measurements of cells treated either with solvent (0.1% v/v DMSO) or with tunicamycin (2.5 or 10 μg/mL) for 24 h. White and grey bars denote the NucleoCounter and 96-well-based cell-counting assay, respectively. Mean and standard deviation are depicted.
Having all required techniques established, we wanted to test the applicability of our screening platform. As proof, we chose two model proteins, α-1AT and hC1INH, which are ideally suited for this purpose. We tagged all four variants N- and C-terminally with the S11-tag, which is approximately 2 kDa in size. However, the N-terminally tagged hC1INH variants gave rise to decreased titer of hC1INH indicating that the effects of S11-fusion are protein-specific. Moreover, viability measurements were consistent between the two assays (Fig. 2d). Next, we wanted to compare the performance of the two cell-counting assays with stressed cells. To this end, we treated CHO-S cells with the endoplasmic reticulum (ER)-stress inducer tunicamycin at two different concentrations for 24 h, because induction of ER stress affects the viability of mammalian cells. Although lower viability values of the tunicamycin-treated cells were observed in the 96-well-based assay compared with the NucleoCounter assay (Fig. 2e), VCD measurements of the two assays were consistent (Fig. 2f). Overall, the performance of the 96-well-based cell-counting assay was satisfactory and we concluded that it was suitable for monitoring cell growth in the 96-well format.

**Similar cell growth profiles and volumetric productivity in half-deepwells and shake flasks.** Having established a 96-well-based cell-counting assay, we were able to analyse cell growth in all wells in a 96-HDW-microplate and compare cell growth with standard Erlenmeyer shake flasks. Cells were seeded at 3 x 10^5 viable cells/mL and VCD and viability were measured daily for four days. Cell growth profiles in shake flasks and in wells in a 96-HDW-microplate were comparable (Fig. 3a). Viability during the four days of culture was in general high (>95%) and was consistent between shake flasks and HDWs (Fig. 3a).

Variation of VCD measurements was in general low (coefficient of variation [CV] = 5–8%) during the four-day period (Fig. 3b). Therefore, the observed variation seems to originate from small differences in seeding density (Fig. 3b) and variation of the 96-well-based cell-counting assay (Fig. 2c). More importantly, the CV of doubling time was 3.7% from day 0–6 (Fig. 3c) demonstrating low well-to-well variation. Moreover, well position had no observed effect on cell growth rate (Supplementary Tables S7 and S8).

Next, we wanted to compare volumetric productivity of different therapeutic glycoproteins in HDWs and shake flasks. To this end, we chose our two model proteins hα-1AT and hC1INH as well as hEPO and Rituximab. Two days after transfection, supernatant samples were obtained and product titer was determined. In general, volumetric productivity of each of the four proteins was comparable in HDWs and shake flasks (Fig. 2c). Moreover, well position had no observed effect on cell growth rate (Supplementary Tables S7 and S8).

To determine the dynamic linear range of the split-GFP assay, we subjected a dilution series of a synthetic α-1AT and C-terminally tagged hC1INH for all further experiments. To this end, we chose our two model proteins hα-1AT and hC1INH as well as hEPO and Rituximab. Two days after transfection, supernatant samples were obtained and product titer was determined. In general, volumetric productivity of each of the four proteins was comparable in HDWs and shake flasks (Fig. 3a). Viability during the four days of culture was in general high (>95%) and was consistent between shake flasks and HDWs (Fig. 3a).

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**Comparison of S11 variants fused N- and C-terminally to model proteins.** We wanted to investigate the applicability of using split-GFP methodology to determine product titer because this is a potentially versatile alternative to antibody-based product titer assays for non-mAbs (see Discussion). To this end, we tagged our two non-mAb model proteins hα-1AT and hC1INH N- and C-terminally with two different variants of the split-GFP S11-tag: M3 and H7 (Supplementary Fig. S1a). The H7 variant has been shown to improve expression levels of membrane proteins in E. coli compared with the M3 variant. The H7 variant (17 residues) is one amino acid residue longer than the M3 variant and contains seven histidine residues in contrast to the two histidine residues of M3 (Supplementary Fig. S2).

All hα-1AT and hC1INH variants were expressed transiently for two days. The H7 variant seemed to increase the titer of hα-1AT, whereas productivity of hα-1AT appeared unaffected by the M3 variant (Fig. 4a). In contrast, both variants gave rise to decreased titer of hC1INH (Fig. 4b) indicating that the effects of S11-fusion are protein-specific. C-terminally tagged hC1INH variants migrated slightly slower than untagged hC1INH consistent with fusion of the S11-tag, which is approximately 2 kDa in size (Fig. 4b). However, the N-terminally tagged hC1INH variants reproducibly migrated even slower than the C-terminally tagged hC1INH variants (Fig. 4b). This gel migration pattern was retained upon removal of N-glycans by PNGase F (Fig. 4c). However, removal of N- and O-glycans gave rise to co-migration of the N-terminally tagged M3 variant with the C-terminally tagged variants (Fig. 4d). Removal of N- and O-glycans also gave rise to a notable migration shift for the N-terminally tagged H7 variant but this variant reproducibly migrated slightly slower than the other three S11-tagged variants. The glycosidase mix used in this analysis is not able to remove all possible O-glycan structures. Thus, the N-terminally tagged H7 variant might contain such resistant O-glycan structure(s). We concluded that O-glycosylation was affected when the S11 peptide was N-terminally fused to hC1INH. Apart from this, all S11-tagged variants of hα-1AT and the C-terminally tagged variants of hC1INH appeared as expected; migrating slightly slower than the untagged variants consistent with fusion of the S11-tag (Fig. 4a,b).

Preliminary results indicated that the H7 variant gave rise to lower fluorescence complementation intensity than the M3 variant upon split-GFP complementation (data not shown). In order to elucidate this, we compared signal-to-noise ratios of all S11 constructs. All M3 variants gave rise to higher signal-to-noise ratio than the H7 variants (Fig. 4e,f), despite their titers being similar or lower (Fig. 4a,b). This suggests that the M3 variants have higher complementation capacity; thus, we chose this S11-tag variant for further experiments. We chose N-terminally tagged hα-1AT and C-terminally tagged hC1INH for all further experiments.

To determine the dynamic linear range of the split-GFP assay, we subjected a dilution series of a synthetic S11_M3 peptide to split-GFP product titer assay. Consistent with previously reported data, we observed a clear linear relationship between split-GFP complementation intensity and peptide concentration (Fig. 4g). The dynamic linear range spanned two orders of magnitude and showed intra-assay variation below 10% (Fig. 4h). This suggests that substantial changes in titer can be quantified without additional dilution steps.

**Split-GFP-based overexpression platform reveals target gene-specific effects on productivity.** Having all required techniques established, we wanted to test the applicability of our screening platform. As proof...
of concept, we tested seven target genes encoding proteins with previously reported effects on the productivity of recombinant glycoproteins in CHO cells (for reviews see7,9): five genes with positive effects (Xbp1s, Pdia3, Ppp1r15a, Ero1l, and ATF4) and two genes with mixed effects (Hspa5 and P4hb). Although not previously tested, Hsp90b1 was also included because its expression has been found to positively correlate with heterologous protein production33.

The functions of the proteins encoded by these eight genes are all related to the ER, as they are either involved in the unfolded protein response or are chaperones and/or oxidoreductases localized in the ER. As negative controls, two pro-apoptotic genes were included: Bak1 and Bax34.

Eight individual transfection pre-mixes were made per target gene because we observed some transfection-based variation (Supplementary Fig. S3). Although not separated in time, data from individual transfections were processed statistically as individual data points in order to cover as much variation as possible.

Co-expression of ATF4 gave rise to increased qP and titer of hα1AT-S11 three days post-transfection (Fig. 5a,b). As for hC11NH-S11, co-expression of Xbp1s gave rise to increased qP and titer (Fig. 5c,d). Although less prominent, effects of ATF4 and Xbp1s were also observed two days post-transfection (Supplementary Fig. S4). Interestingly, three of the five genes previously reported to have a positive effect on specific productivity had a negative effect on both model proteins (Fig. 5a–d): Pdia3, Ppp1r15a, and Ero1l. Apart from reduced viability for Ppp1r15a, only

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Figure 3. Shake-flask-like cell growth profile and volumetric productivity in a 96-HDW-microplate. (a–c) Comparison of cell growth in shake flasks (three flasks) and HDW-microplate (96 wells). CHO-S cells were seeded at 3 × 10⁵ cells/mL in shake flasks (25 mL in 125 mL flasks) or in HDWs (250 μL) and VCD and viability were measured daily (day 0–4). (a) VCD (solid lines) and viability (dashed lines) versus culture time. Red and black lines show data from shake flasks and HDWs, respectively. VCD values are plotted on a logarithmic scale. (b) Coefficient of variation (CV) of VCD measurements in HDWs during the four days of culture. (c) Doubling time from day 0–3 in 96 wells calculated by exponential growth equation (least squares fit) in GraphPad Prism (GraphPad Software). Median and interquartile range are depicted and CV is shown above the data points. (d–g) Comparison of volumetric productivity in shake flasks and HDWs of hα1AT, hEPO, Rituximab and hC11NH. CHO-S cells were transiently transfected with plasmids encoding the aforementioned proteins and supernatants two days post-transfection were obtained and titers were determined by ELISA (hα1AT, hEPO and hC11NH) and bio-layer interferometry (Rituximab). 16 and 4 separate transfections were performed in HDWs and flasks, respectively. Mean, standard deviation, and percentage difference are shown.
Figure 4. Analysis of split-GFP S11-tag variants. (a–f) CHO-S cells in 6-well plates were transfected with plasmids encoding hα1AT or hC1INH either S11_M3 or S11_H7-tagged at the N- or C-terminus or untagged. Supernatants two days post-transfection were analyzed. (a) α-hα1AT and (b–d) α-hC1INH Western blots under reducing conditions. The Western blots shown are representative of three experiments. (c,d) Proteins in supernatants were subjected to PNGase F and O-glycosidase (Protein Deglycosylation Mix) treatment under denaturing conditions where indicated. (e,f) Supernatants were analyzed by split-GFP product titer assay. Signal-to-noise (S/N) ratio was determined by defining noise as split-GFP complementation signal from mock transfected cells. Mean and standard deviation are depicted (n = 3). (g) A dilution series of a synthetic S11_M3 peptide was analyzed by split-GFP product titer assay. The trend line from linear regression and correlation coefficient (R²) are shown. The encircled data point was found to be an outlier in the linear regression analysis. Mean and standard deviation from eight technical replicates are depicted. (h) Intra-assay variation of the split-GFP product titer assay was analyzed from the data shown in panel (e). The coefficient of variation (CV) at different concentrations of the S11_M3 peptide are shown. The two-headed arrow denotes the defined dynamic linear range with low intra-assay variation (CV < 10%).
slight differences in viability (Supplementary Fig. S5) and IVCD (Supplementary Fig. S6) were observed. Minute amounts of hα1AT-S11 and hC1INH-S11 were produced upon co-expression of Bax and Bak1 (Fig. 5a–d). As expected from these pro-apoptotic genes, a significant drop in viability was observed (Supplementary Fig. S5) resulting in a drop in IVCD (Supplementary Fig. S6).

In order to validate the split-GFP product titer assay for our two model proteins, we wanted to compare this assay with the widely used protein quantification method; ELISA. To this end, product titer was determined in selected hα1AT and hC1INH supernatant samples by ELISA (Fig. 5e,f). Relative titer determined by split-GFP complementation for hα1AT-S11 and hC1INH-S11 was within ±4% compared to ELISA. Thus, we concluded that the split-GFP-based transient expression platform is able to accurately quantify relative titers of therapeutic glycoproteins in complex samples (spent media).

Positive effects of Atf4 and Xbp1s validated on untagged model proteins. Lastly, we wanted to test whether the positive effects of Atf4 and Xbp1s could be detected on untagged hα1AT and hC1INH. To this end, untagged hα1AT and hC1INH were expressed in CHO-S cells in a 96-HDW-microplate for three days with and without co-expression of Atf4 and Xbp1s. Titer and qₚ for untagged hα1AT and hC1INH co-expressed with a mock plasmid were 72 mg/L and 13 pcd, and 3.2 mg/L and 0.5 pcd, respectively. Increased titer and qₚ were observed for hα1AT and hC1INH upon co-expression of Atf4 and Xbp1s, respectively (Fig. 6a,b). Although the effects were 8–16% higher with S11-tagged than untagged model proteins (Supplementary Table S9), we concluded that our split-GFP-based transient expression platform is able to identify effects on productivity of non-mAb glycoproteins.

Discussion

Compared with conventional shake flask cultivation of CHO cells, cultivation in 96-well microplates significantly increases throughput. Originally, growth conditions in 96-well microplates were suboptimal due, for example, to excessive evaporation and insufficient oxygen transfer rate. However, these issues have been solved by Duetz covers, which reduce evaporation while allowing sufficient oxygen transfer. The dramatic effect of the Duetz cover on evaporation was recently demonstrated by showing a 5.5-fold decrease in evaporation compared with gas-permeable seals. In the present study, evaporation of 1.4% v/v per well per day was observed in 96-HDW-microplates with Duetz covers (data not shown). We were able to demonstrate cell-growth profiles in
from Fig. 5). The variation of deviation, and statistically significant (p ≤ 0.05) fold-changes are depicted. Eight separate transfections were performed in each experiment and experiments were performed twice (n = 2; 16 wells). Mean, standard deviation, and statistically significant (p ≤ 0.05) fold-changes are shown. Eight separate transfections were performed in each experiment and experiments were performed twice (n = 2; 16 wells). Mean, standard deviation, and statistically significant (p ≤ 0.05) fold-changes are depicted.

This range likely explains why unexpectedly low titers and observed variation (CV: 9.2%) in transfection efficiency, which ranged from 55% to 80% (Supplementary Fig. S3).

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encoding untagged hC1INH was co-transfected with mock plasmid or plasmid encoding

Mean, standard deviation, and statistically significant (p ≤ 0.05) fold-changes are depicted. Eight separate transfections were performed in each experiment and experiments were performed twice (n = 2; 16 wells). Mean, standard deviation, and statistically significant (p ≤ 0.05) fold-changes are depicted.

6-well microplates that were comparable to those in shake flasks (Fig. 3a). Similarly, shake-flask-like cell-growth profiles have been reported in 24-well plates using Duetz covers38. The well-to-well variation of VCD on day 4 in 24-well plates was reported to be 9.0%38, which is comparable with the value of 7.5% observed in 96-well microplates in this study (Fig. 3b). No effect of well position on cell growth rate was observed in 24-well plates, which is consistent with our results in 96-well microplates (Supplementary Table S7).

Similarly, we observed relatively low variation of titer and qg (Figs 5 and 6). Mean variation of titer and qg was 9.7% and 14.9%, respectively, for both model proteins and all target genes except Bak1 and Bax (data from Fig. 5). The variation of qg is consistent with the previously reported variation of qgantibody (16%) from polyethylenimine-mediated transfection of CHO cells in flasks10. The variation of titer is comparable with the observed variation (CV: 9.2%) in transfection efficiency, which ranged from 35% to 80% (Supplementary Fig. S3). This range likely explains why unexpectedly low titers and qg were observed for some wells (e.g. Errol and Hsp90b1 in Fig. 5a,b and Atf4 in Fig. 6a). To our knowledge, this is the first published example of transient protein production in CHO cells cultivated in 96-well microplates. Recently, a study was published on transient expression of IgG1 in HEK293 cells by polyethyleneimine-mediated transfection in 96-deepwell microplates using Duetz covers37. Unfortunately, comparison of well-to-well variation from that study is not possible, because only plate-to-plate variation was reported. Importantly, we were able to show that volumetric productivity of four different therapeutic glycoproteins in shake flasks was comparable in HDWs (Fig. 3d–g). However, some differences were observed ranging from −16% to +14% between shake flasks and HDWs. Moreover, variation in titer was higher in HDWs compared to shake flasks: average CV of all four proteins HDWs and shake flasks was 7% and 4%, respectively. Overall, it is likely that these differences partly arise from differences in the transfection procedure. Complexation of the transfection reagent and plasmids is a time-dependent process39 and the complex incubation time likely varies more when transfecting cells in HDWs compared to shake flasks. Automation of the transfection process by a liquid handling robot could potentially minimize this variation to some extent.

Sixteen wells per gene were used in the screening platform. The sample size can be reduced to three and six wells to detect fold-changes of 1.5 to 1.3, respectively, with power of 0.80 and type I error of 0.05 based on the observed variation of 15%. Moreover, several plates can be analysed in parallel. Finally, the screening platform is suitable for automation by a liquid handling robot because the majority of the experimental work involves liquid transfer between 96-well microplates. These options have the potential to increase the throughput and number of identified positive effects significantly of the screening platform. Importantly, positive effects on productivity of target genes must eventually be further characterized in an industry-like context as well as product quality attributes such as glycosylation and bioactivity.

Effects on productivity were less pronounced on untagged model proteins than S11-tagged model proteins (Supplementary Table S9). This discrepancy is likely a consequence of fusing the S11-tag to the model proteins. Although the S11-tag is relatively small (~2 kDa), fusion of this tag to proteins could change the overall conformation as well as post-translational modifications. Indeed, the H7 S11-variants of hα1AT led to increased titers and all S11-tagged variants of hC1INH gave rise to decreased titers compared to untagged protein (Fig. 4a,b). Moreover, N-terminally fusion of the S11 tag to hC1INH altered O-glycosylation (Fig. 4c,d). These results clearly show that fusion of the S11 tag to the protein of interest is a possible drawback of this platform. However, further optimization of the S11-tag sequence might be able to address this issue to some extent — for example by altering the glycine-serine linker sequence40. Despite the effects of the S11-tag on titers and post-translational modifications,
we could reproduce that Atf4 and Xbp1s overexpression significantly improved productivity of untagged ho 1AT and hC1INH, respectively (Fig. 6a,b). Moreover, relative titers obtained by ELISA and split-GFP complementation were consistent (Fig. 5e,f). Thus, split-GFP complementation seems to be a versatile antibody-independent method for determining relative productive titers of recombinant glycoproteins produced in CHO cells. For mAbs, several well-established high-throughput and automatable product titer assays exist. However, this is not the case for the majority of non-mAbs; split-GFP-based product titer determination therefore seems particularly valuable for non-mAb recombinant proteins. Besides the application as a discovery tool in the platform presented here, the split-GFP methodology could be useful in initial trials of producing protein variants by transient expression in the biopharmaceutical industry. However, the split-GFP system is not suitable for larger scale production in stable cell lines, because the S11-tag eventually becomes an undesired trait in for example detailed characterization of the biopharmaceutical industry.

For non-mAb recombinant proteins. Besides the application as a discovery tool in the platform presented here, the split-GFP-based product titer determination therefore seems particularly valuable for non-mAbs. Furthermore, both pro-apoptotic genes (Bak1 and Bax) clearly affected cell viability (Supplementary Fig. S5). Alternatively, mouse genes can be replaced by homologous hamster genes owing to the recent drafts of the Chinese hamster and CHO cell line genomes48–51. However, not all hamster genes are at present annotated correctly when compared to mouse homologues (our unpublished observations). Consequently, using hamster genes in our screening platform could lead to erroneous conclusions, until sequencing and annotation of the hamster genome has been further refined.

Bacterial cell line engineering seems to be a constructive way of improving secretory capacity and several examples of synergistic effects on productivity of co-expressing two target genes have been reported52–54. More than 20,000 genes are present in the Chinese hamster genome1 and combinatorial approaches dramatically increase the number of conditions possible to be tested. Compared with current lower throughput setups, our platform facilitates cost-efficient combinatorial screening of genes. In conclusion, we believe that our target gene-screening platform holds the potential to unravel novel genes and combinations of genes that can enhance the secretory pathway machinery in CHO cells.

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Author Contributions
H.G.H. designed the experiments and wrote the manuscript. A.M.L. provided the list of target genes and the cloning design of the target genes. H.G.H., C.N.N., S.K., L.M.G. and M.L. performed the experiments. H.G.H., C.N.N., J.R., G.M.L., M.R.A. and H.F.K. analysed the data. All authors reviewed the manuscript.

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