Engineering selection stringency on expression vector for the production of recombinant human alpha1-antitrypsin using Chinese Hamster Ovary cells

Christine L Chin1†, Hing K Chin1†, Cara S Hui Chin1, Ethan T Lai1, Say Kong Ng1,2*

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Background
Currently, the titers of biopharmaceutical production from Chinese hamster ovary (CHO) cells have achieved gram per liter range and this can be attributed to advances in bioprocess development, media development and cell line development. To obtain high producing cell lines, extensive screening of producer clones during cell line development is often necessary. To improve the efficiency and efficacy of generating and isolating high producing clones, various expression vector engineering technologies can be applied, for example ubiquitous chromatin opening element (UCOE) [1], matrix attachment regions (MARs) [2](Mirkovitch et al. 1984), site specific recombination [3-9], to improve selection stringency [10-13], and to co-localize the GOI with the selection marker [14-16].

In our previous studies, we have similarly shown that specific productivities can be improved when we increased selection stringency by destabilizing the selection marker through the addition of AU-rich elements (ARE) to promote mRNA degradation and murine ornithine decarboxylase (MODC) PEST region to enhance protein degradation [17]. While coexpression of GOI and selection marker using multiple promoters on the same vector may help in the co-localization, we have previously demonstrated that gene fragmentation can occur at a high level of 14% during stable transfection of dual promoter dicistronic vector in CHO-DG44 cells [18]. Subsequently, an attenuated IRES element was used together with the PEST region to allow for high recombinant protein titer using stably amplified cell pools [19].

In this study, we evaluated the use of tandem PEST sequence, further attenuation of the IRES element, and codon-deoptimization of the dhfr selection marker, to further optimizing the strength of selection marker expression in CHO cells for the production of recombinant human Alpha1-antitrypsin (rhA1AT), a serum protease inhibitor currently purified from human blood plasma as replacement therapy for patients who developed chronic obstructive pulmonary disease due to deficiency in the protein. Such vector combinations to attenuate translation initiation, protein elongation and protein stability for optimizing selection stringency have not been previously investigated. To our knowledge, there is also no report on high-titer production of rhA1AT in CHO cells, which is necessary for its manufacturability due to its high dosage requirement.

Experimental approach
7 expression vectors expressing rhA1AT that can be classified into 3 sets (Figure 1) were designed. Using rhA1AT as the gene of interest, the first vector set consists of pAID, pAIDp and pAIDpp. Comparing data from the use of pAIDp against pAID will allow us to validate the use of PEST element in improving stable recombinant gene expression, as observed in our previous studies [17,19]. The application of 2 tandem PEST elements in pAIDpp then allowed us to determine whether an additional PEST can further improve stable recombinant gene expression, as this has not been demonstrated in literature to our knowledge. The second vector set consists of
pAI709Dp and pAI772Dp. These 2 vectors incorporated mutations described by Hoffman MA and Palmenberg AC [20] into the attenuated IRES [21,22]. This is to evaluate whether the further attenuation of selection marker expression with these additional impediment in translation initiation can improve stable recombinant gene expression. The third vector set comprised of pAID* and pAID*p. These 2 vectors incorporated a codon de-optimized dhfr selection marker to evaluate the use of codon deoptimization as a strategy to further reduce selection marker expression levels, since it will theoretically reduce translation efficiency, a different aspect of gene expression that is not affected by the attenuated IRES and PEST. The selection and amplification efficiency, recombinant protein productivity, relative transcript copy numbers and dhfr expression levels were then analyzed.

**Results and discussion**

pAIDpp and pAI772Dp vectors gave further improvements in rhA1AT production when compared to pAID and pAIDp vectors, indicating that further selection marker attenuation can improve recombinant protein production. Using the pAI772Dp vector, we generated a cell pool that gave a maximum titer of 1.05 g/l of rhA1AT in an un-optimized shake flask batch culture using a 2-step amplification till 50 nM MTX that took less than 3 months (Table 1). Using the pAI772Dp and pAIDpp vectors, we generated cell pools that gave a maximum titer of 1.11 and 1.15 g/l respectively in un-optimized shake flask batch cultures at 300 nM MTX (Table 1). To our knowledge, this is the highest reported recombinant protein titer from shake flask cultures of stable mammalian cell pools.

Relative transcript copy numbers demonstrated that the transcription of rhA1AT and dhfr genes were correlated due to the IRES linkage, although the results also suggested that the protein expression were not solely dependent on transcript levels. Protein level analysis of dhfr validated that the cell pools were indeed expressing the modified dhfr of the correct molecular weight. In addition, it showed that the strategies of further attenuating dhfr protein expression with the use of a mutated IRES and tandem PEST, but not codon deoptimization, were effective in reducing dhfr protein levels in these MTX amplified cell pools in suspension serum free culture. Our data suggests the codon usage of surviving...
cells with codon deoptimized selection marker may be changed in our culture conditions to enable better cell survivability. Hence, this result suggest that codon usage of the selection marker should be considered for applications that involve gene amplification and serum free suspension culture, since the general expression and regulation of host cell proteins may be affected due to a change in codon usage in the surviving cells.

Authors' details
1. Bioprocessing Technology Institute, Agency for Science, Technology and Research (A*STAR), Singapore. 2. Department of Pharmacy, Faculty of Science, National University of Singapore, Singapore.

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References
1. Benton T, Chen T, McIntee M, Fox B, King D, Crombie R, Thomas TC, Bebbington C. The use of UCOE vectors in combination with a preadapted serum free, suspension cell line allows for rapid production of large quantities of protein. Cytotherapy 2002, 3(1-3):43-46.
2. Mirkovitch J, Mirault ME, Laemmli UK. Organization of the higher-order chromatin loop: specific DNA attachment sites on nuclear scaffold. Cell 1984, 39(1):223-232.
3. Branda CS, Dymecke SM. Talking about a revolution: The impact of site-specific recombinases on genetic analyses in mice. Dev Cell 2004, 6(1):7-29.
4. Golic MM, Rong YS, Petersen RB, Lindquist SL, Golic KG. FLP-mediated DNA mobilization to specific target sites in Drosophila chromosomes. Nucleic Acids Res 1997, 25(18):3665-3671.
5. Groth AC, Fish M, Nusse R, Calos MP. Construction of transgenic Drosophila by using the site-specific integrase from phage phiC31. Genetics 2004, 166(4):1775-1782.
6. O’Gorman S, Fox DT, Wahl GM. Recombinase-mediated gene activation and site-specific integration in mammalian cells. Science 1991, 251(4999):1351-1355.
7. Voziyanov Y, Pathania S, Jayaram M. A general model for site-specific recombination by the integrase family recombinases. Nucleic Acids Res 1999, 27(4):930-941.
8. Voziyanov Y, Konieczenia JH, Stewart AF, Jayaram M. Stepwise manipulation of DNA specificity in Flp recombinase: progressively adapting Flp to individual and combinatorial mutations in its target site. J Mol Biol 2003, 326(1):65-76.
9. Wirth D, Gama-Norton L, Riemer P, Sandhui U, Schucht R, Hauser H. Road to precision: recombinase-based targeting technologies for genome engineering. Curr Opin Biotechnol 2007, 18(5):411-419.
10. Chen L, Xie Z, Teng Y, Wang M, Shi M, Qian L, Hu M, Feng J, Yang X, Shen B, et al. Highly efficient selection of the stable clones expressing antibody-IL-2 fusion protein by a dicistronic expression vector containing a mutant neo gene. J Immunol Methods 2004, 295(1-2):49-56.
11. Sautter K, Enenkel B. Selection of high-producing CHO cells using NPT selection marker with reduced enzyme activity. Biotechnol Bioeng 2005, 89(3):530-538.
12. Nava H, Yamamura K, Miyazaki J. Efficient selection for high-expression transfectants with a novel eukaryotic vector. Gene 1991, 108(2):193-199.
13. Westwood AD, Rowe DA, Clarke HR. Improved recombinant protein yield using a codon deoptimized DHFR selectable marker in a CHEF1 expression plasmid. Biotechnol Prog 2010, 26(6):1558-1566.
14. Kaufman RJ, Sharp PA. Amplification and expression of sequences cotransfected with a modular dihydrofolate reductase complementary dna gene. J Mol Biol 1982, 159(4):601-621.
15. Kaufman RJ, Davies MV, Wasley LC, Michnick D. Improved vectors for stable expression of foreign genes in mammalian cells by use of the untranslated leader sequence from EMC virus. Nucleic Acids Res 1991, 19(16):4485-4490.
16. Milbrandt JD, Hentz NH, White WC, Rothman SM, Hamlin JL. Methotrexate-resistant Chinese hamster ovary cells have amplified a 135-kilobase-pair region that includes the dihydrofolate reductase gene. Proc Natl Acad Sci USA 1981, 78(10):6043-6047.
17. Ng SK, Wang DJ, Yap MG. Application of destabilizing sequences on selection marker for improved recombinant protein productivity in CHO-DG44. Metab Eng 2007, 9(3):304-316.
18. Ng SK, Lin W, Sachdeva R, Wang DJ, Yap MG. Vector fragmentation: characterizing vector integrity in transfected clones by Southern blotting. Biotechnol Prog 2010, 26(1):11-20.
19. Ng SK, Tan TR, Wang Y, Ng D, Goh LT, Bardor M, Wong VW, Lam KP. Production of Functional Soluble Dectin-1 Glycoprotein Using an IRES-Linked Destabilized-Dihydrofolate Reductase Expression Vector. PLoS One 2012, 7(12):e52785.
20. Hoffman MA, Palmberg AC. Revertant analysis of J-K mutations in the encephalomyocarditis virus internal ribosomal entry site detects an altered leader protein. J Virol 1996, 70(9):6425-6430.
21. Bochkov YA, Palmenberg AC. Translational efficiency of EMCV IRES in bicistronic vectors is dependent upon IRES sequence and gene location. Biotechniques 2006, 41(3):283-284, 286, 288 passim.
22. Gurtu V, Yan G, Zhang G. IRES bicistronic expression vectors for efficient creation of stable mammalian cell lines. Biochem Biophys Res Commun 1996, 229(1):295-298.

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