Precision Engineering of the Transcription Factor Cre1 in Hypocrea jecorina (Trichoderma reesei) for Efficient Cellulase Production in the Presence of Glucose

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In Trichoderma reesei, carbon catabolite repression (CCR) significantly downregulates the transcription of cellulolytic enzymes, which is usually mediated by the zinc finger protein Cre1. It was found that there is a conserved region at the C-terminus of Cre1/CreA in several cellulase-producing fungi that contains up to three continuous S/T phosphorylation sites. Here, S387, S388, T389, and T390 at the C-terminus of Cre1 in T. reesei were mutated to valine for mimicking an unphosphorylated state, thereby generating the transformants Tr_Cre1S387V, Tr_Cre1S388V, Tr_Cre1T389V, and Tr_Cre1T390V, respectively. Transcription of cel7a in Tr_Cre1S388V was markedly higher than that of the parent strain when grown in glucose-containing media. Under these conditions, both filter paperase (FPase) and p-nitrophenyl-β-D-celllobioside (pNPCase) activities, as well as soluble proteins from Tr_Cre1S388V were significantly increased by up to 2- to 3-fold compared with that of other transformants and the parent strain. The results suggested that S388 is critical site of phosphorylation for triggering CCR at the terminus of Cre1. To our knowledge, this is the first report demonstrating an improvement of cellulase production in T. reesei under CCR by mimicking dephosphorylation at the C-terminus of Cre1. Taken together, we developed a precision engineering strategy based on the modification of phosphorylation sites of Cre1 transcription factor to enhance the production of cellulase in T. reesei under CCR.

Keywords: carbon catabolite repression, transcription factor, Cre1, cellulase, phosphorylation, Trichoderma reesei
INTRODUCTION

The filamentous ascomycete *Trichoderma reesei*, a clonal derivative of the ascomycete *Hypocrea jecorina* (Kuhls et al., 1996), is widely used in industrial production of cellulases and xylanases (Dernl et al., 2019). There is a broad range of industrial applications of these cellulolytic enzymes in the food and feed, textile, and pulp and paper industries, as well as in the production of lignocellulosic bioethanol (Wilson, 2009).

Carbon catabolite repression (CCR) is a global regulatory system that is found in nearly all heterotrophic hosts (Görke and Stülke, 2008). Notably, the presence of monosaccharides, like D-glucose, during cellulolytic enzyme production triggers CCR, which is regulated mainly by the cellulase transcriptional repressor Cre1 and activators Xyr1, Clr1, Clr2, etc., and significantly downregulates the transcription of cellulolytic enzymes in *T. reesei* (Wang et al., 2019). Endogenous Cre1, which belongs to the group of C2H2-type zinc finger proteins, binds to the promoters of target genes, such as *cel7a*, and inhibits the transcription of encoded cellulase genes (Mello-de-Sousa et al., 2014).

Many reports have demonstrated that the knockout of cre1 in filamentous fungi alleviated CCR (Ilmén et al., 1996; Long et al., 2018). However, it was reported that Cre1 not only plays an essential role in correct nucleosome positioning, but also participates in several morphological functions, such as hyphal development and sporulation (Ries et al., 2014). Moreover, it was shown that the disruption of *cre1* in *T. reesei*, or its replacement with a *cre1* mutant, inhibits hyphal growth (Portnoy et al., 2011), affects biomass accumulation, and leads to a decrease in cellulase production (Nakari-Setälä et al., 2009; Rassinger et al., 2018; Wang et al., 2019).

Furthermore, it was reported that CCR is achieved with the phosphotransferase system (Wang et al., 2017; Liu et al., 2020). Post-translational modifications, especially phosphorylation, of the proteins involved, including Cre1, play an essential role in signal transduction to achieve CCR (Jiang et al., 2018; Horta et al., 2019). Horta et al. (2019) revealed that there is large-scale phosphorylation and dephosphorylation of substrate-specific proteins, including Cre1, in *Neurospora crassa*, according to MS/MS-based peptide analysis. Nguyen et al. (2016) suggested that phosphorylation of Cre1 plays a role in the onset of CCR induction and sensing of the carbon source. Cziferszky et al. (2002) reported that phosphorylation of Cre1 at S241 in *T. reesei* has a significant effect on the repression of cellulolytic enzyme genes at the transcriptional level under CCR. Thus, the modification of specific phosphorylation sites in Cre1/CreA may be a rational strategy for releasing or attenuating CCR to achieve improvement of cellulase production.
In this study, the modification of phosphorylation sites at the C-terminus of Cre1 from *T. reesei* was performed for mimicking a dephosphorylated state, generating four transformants. Growth behavior, transcriptional profiles, and enzymatic activities of these transformants were investigated. Obtained data proved that the mutation of S388 to V388 at the C-terminus of Cre1 results in great improvement of cellulase production in the presence of glucose. Furthermore, it suggested that loss of phosphorylation at S388 at the continuous phosphorylation motif (SSTT) plays an important role in alleviating CCR.

**MATERIALS AND METHODS**

**Strains and Reagents**

The *T. reesei* strain (CCTCC M2015804) that was deposited in the China Center for Type Culture Collection (CCTCC) was used as the parent strain (Zhang et al., 2019) in this study, and pNPC was purchased from Aladdin (Shandong, China). All polymerase chain reaction amplifications were performed using DNA polymerase (Vazyme Co., Ltd., Nanking, China). RNaIso™ reagent and PrimeScript® RT reagent Kit With gDNA Eraser (Perfect Real Time) were purchased from Takara Bio Inc. (Shiga, Japan). FastStart Essential DNA Green Master was purchased from Roche (Basel, Switzerland). All other chemicals were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). ABclonal MultiF Seamless Assembly Mix was purchased from ABclonal (Wuhan, China). Primers were synthesized by Personalbio Biotech Co., Ltd. (Shanghai, China).

**Construction of cre1 Mutation Replacement Cassettes and Propagation of Transformant Strains**

All plasmids were constructed based on homologous recombination, which was mediated by the ABclonal MultiF Seamless Assembly Mix. The primers used for the construction of *cre1* mutation replacement cassettes are listed in Supplementary Table S1. All the plasmids and linearized expression cassettes used or constructed in this study were listed in Supplementary Table S2. The transformation of replacement cassettes into *T. reesei* was performed using previously described methods (Wang et al., 2019). The transformants were selected on plates containing minimal medium (MM) supplemented with 2% glucose and 200 µg/mL hygromycin. Propagation of transformant strains was performed based on a protocol published previously (Wang et al., 2019).

**Analysis of Growth Phenotype and Transcriptional Profile**

The method used for culturing fungi for growth phenotype, transcript analysis, and RT-qPCR assays was already described previously (Wang et al., 2019). The primers used for RT-qPCR are listed in Supplementary Table S1. Microscopic observation of hyphal morphology was performed based on a previously described method (Wang et al., 2019). Approximately, $1 \times 10^3$ spores were inoculated on slides with solidified medium containing PDA, wheat bran, MM with 2% glucose, or MM with 2% glycerol at 30°C for 48 h. Microscopic images of hyphae with 400 × magnification (Nikon Eclipse E100, Tokyo, Japan) were then captured using a Nikon D5000 camera.

**Cellulase Production and Determination of Enzymatic Activities, Soluble Protein, and Biomass**

The parent strain or mutants (approximately $1 \times 10^7$ spores) was inoculated in 300 ml flask including MM solution containing 1% glucose, 1% trytome, 1% wheat bran, and 0.5% CaCO₃ at 30°C and 200 rpm. After 48 h of cultivation, the 10 ml media as the seed was added into 500 ml flask including 100 ml liquid medium. In this study, batch cultures and pulse fed-batch cultures were cultivated at 30°C and 200 rpm for 7 days. The initial pH value was adjusted at pH 5.0 with 3 M NaOH. Batch culture was performed in the medium with MM solution containing 2% glucose based on a protocol published previously (Wang et al., 2019). Pulse fed-batch culture: For MM solution containing 2% avicel, 0.6% glucose was pulse-fed on the second and third days. Cultures were timely collected to measure pNPCase activity, as well as extracellular and intracellular protein concentrations. Protein and biomass concentrations were estimated according to a previously described method (Wang et al., 2019).

**In silico Prediction of Protein Domains and Homologous Sequence Alignment**

Partial in silico domain prediction of Cre1 (*Trichoderma reesei* NCBI Accession ID: AAB01677.1) was performed as previously described (Rassinger et al., 2018). Cre1 (*Trichoderma reesei*, NCBI Accession ID: AAB01677.1) and its homologs in *Penicillium oxalicum* (NCBI Accession ID: EPS28222.1), *Aspergillus oryzae* (NCBI Accession ID: AAK11189.1), *Aspergillus nidulans* (NCBI Accession ID: XP_663799.1), *Aspergillus niger* (NCBI Accession ID: AAA32690.1), *Penicillium oxalicum* (NCBI Accession ID: XP_755510.1), *Thermoascus aurantiacus* (NCBI Accession ID: AAT34979.1), *Talaromyces emersonii* (NCBI Accession ID: AAL33631.4), *Thermothelomyces thermophillus* (NCBI Accession ID: XP_003665891.1), and *Neurospora crassa* (NCBI Accession ID: EAA32758.1) were analyzed using ClustalX2.

**RESULTS**

**Phosphorylation of Cre1**

As shown in Figure 1A, we found that there is a conserved region (red box) at the C-terminus of Cre1/Crea/CreT in *Trichoderma reesei*, *Penicillium oxalicum*, *Aspergillus oryzae*, *Aspergillus nidulans*, *Aspergillus fumigatus*, *Aspergillus niger*, *Thermoascus aurantiacus*, *Talaromyces emersonii*, *Thermothelomyces thermophilus*, and *Neurospora crassa* using homology alignment. Most of these conserved regions contain up to three continuous S/T phosphorylation sites. Moreover, phosphorylation of the peptide sequence extending from amino acids 387 to 402...
Comparison of Phenotypic Characterization and Transcriptional Levels

Amino acid residues S387, S388, T389, and T390 of Tr_Cre1 of the parent strain were mutated to valine to mimic their dephosphorylation, generating the transformants named Tr_Cre1S387V, Tr_Cre1S388V, Tr_Cre1T389V, and Tr_Cre1T390V, respectively. The mutation of transformants was verified using PCR and DNA sequencing (Supplementary Figure S1). Phenotypic analysis of each strain was performed after incubating the plates for 6 days at 30°C. Figures 2A,B shows that no morphological changes on the colonies and hyphae were observed among the transformants and parent strain. It was proved that the substitution of S387, S388, T389, or T390 in Tr_Cre1 for valine had no significant effect on the growth behavior of transformants.

Subsequently, the expression levels of cel7a, xyr1, clr1, and clr2 in the parent and transformant strains were investigated using a...
mixture of glucose and avicel as carbon source (Figures 2C,D). No significant changes in xyr1, clr1, and clr2 expression were observed at the transcriptional level among the transformants and the parent strain, each grown in a mixture of glucose and avicel. However, transcription of cel7a in Tr_Cre1^{S388V} was markedly higher than in the parent strain when using a mixture of glucose and avicel as carbon source.

**Determining Cellulase Activity in the Presence of Glucose**

Furthermore, we compared the transformants and the parent strain cultured in either glucose-containing initial media in the batch culture or the supplemental media in the pulse fed-batch culture containing a mixture of glucose and avicel as carbon source. FPase and pNPCase (CBHI) activities, biomass, and soluble protein present in the supernatant of transformant and parent strain cultures were determined over 7 days of cultivation under either condition (Figure 3). There was no obvious difference in biomass among the transformants and the parent strain grown with glucose or the mixture of glucose and avicel before day 4. However, the biomass accumulation of Tr_Cre1^{S388V} was somewhat decreased after 6 days culture. After 7 days, FPase and pNPCase activities, as well as soluble protein from Tr_Cre1^{S388V} were significantly increased by 2.25-, 2.45-, and 1.93-fold, respectively, compared with those of the other transformants or the parent strain grown in glucose as a sole carbon source (Figure 3). Accordingly, FPase and pNPCase activities, and also soluble protein from Tr_Cre1^{S388V}, were significantly increased by 3.50-, 2.19-, and 1.78-fold, respectively, upon 7 days of culture with the mixture of glucose and avicel, compared with those of the other transformants and the parent strain (Figure 3). Our findings proved that the mutation of S388 at the C-terminus of Cre1 results in an improvement of cellulase production in the presence of glucose. In line with this, amino acid residues 387–390 at the C-terminus of Cre1 were predicted by the NetPhos 3.1 Server and KinasePhos:

1. http://www.cbs.dtu.dk/services/NetPhos/
2. http://kinasephos.mbc.nctu.edu.tw/index.php
FIGURE 3 | Determination of cellulase activity in the presence of glucose. T. reesei transformants and the parent strain were cultivated in liquid medium supplemented with 2% (w/v) (x-1) glucose or (x-2) a mixture of glucose and avicel as carbon source. Activity of $p$-nitrophenyl-$\beta$-D-cellobioside (pNPGase) (a), soluble protein (b), filter paperase (FPase) activity (c), and biomass accumulation (d) of each culture were measured in biological and technical duplicates. Enzymatic activities are given as mean values, with error bars indicating the standard deviation. Symbols: blank, parent strain; light gray, $Tr_{_{Cre1}^{S387V}}$; dark gray, $Tr_{_{Cre1}^{S388V}}$; oblique line, $Tr_{_{Cre1}^{T389V}}$; gridline, $Tr_{_{Cre1}^{T390V}}$. 
software as potential targets for phosphorylation. Additionally, it was found that S388 may be phosphorylated by cyclic adenosine monophosphate (cAMP)-dependent protein kinase A (PKA), a conserved key factor of a nutrient-sensing pathway that acts in parallel to the MAP kinase pathway (Ortiz-Urquiza and Keyhani, 2015), whereas S387, T389, and T390 are not. Furthermore, we suggest that dephosphorylation of S388 at the continuous phosphorylation motif (SSTT) plays an important role in alleviating CCR in the presence of glucose.

**DISCUSSION**

Carbon catabolite repression is a global regulatory system found in nearly all heterotrophic organisms (Liu et al., 2020). However, the substrate pairs inducing its expression as well as its underlying regulatory mechanisms need to be analyzed on a case-by-case basis, making it difficult to develop universal strategies to overcome its undesirable effects. A better fundamental understanding is still required to enable smarter designs for facilitating multiple substrate utilization.

Thus, several attempts were made to alleviate CCR by introducing modifications in Crel for improving cellulase production. It was proved that replacement of the endogenous transcription factor Crel/Crea by an artificial minimal transcriptional activator, such as Crel-96 or other Crel/Crea mutants, leads to the release or attenuation of CCR, which in turn supports cellulase production in *T. reesei* and *A. nidulans* (Shroff et al., 1997; Mello-de-Sousa et al., 2014; Rassinger et al., 2018; Zhang et al., 2018). However, Liu et al. (2019) demonstrated that full-length Crel might be necessary for the rapid growth of *T. reesei*. Our results suggested that dephosphorylation at the C-terminus of Crel has no effect on growth behavior because of the retaining of full-length Crel (Figures 2A,B). Meanwhile, it was assumed that phosphorylation of Crel in *T. reesei* affects its repressing activity on cellulase genes under CCR (Cziferszky et al., 2002; Nguyen et al., 2016).

Among the potential phosphorylation target sites in the region covering amino acids 387–390 at the C-terminus of Crel, S388 maybe be phosphorylated by PKA, in contrast to S387, T389, and T390. As a conserved nutrient-sensing pathway, the cAMP-PKA pathway plays an important role in the response to environmental carbon availability to activate or repress the relevant downstream genetic pathways (Ortiz-Urquiza and Keyhani, 2015; Hinterdobler et al., 2019). In *Aspergillus fumigatus* and *Neurospora crassa*, PKA is activated in the presence of glucose, thereby inhibiting alternative carbon source usage due to CCR (Brown et al., 2014; Huberman et al., 2016). In accordance with a previous study (Ribeiro et al., 2019), the *pkaA* deletion strain showed a significant improvement in cellulase activity in *Aspergillus nidulans*.

The results shown in Figures 2, 3 proved that the phosphorylation status at S388 of Crel in *T. reesei* plays an essential role in the regulation of transcription and synthesis of cellulolytic enzymes. Thus, we speculated that S388 of Crel in *T. reesei* was phosphorylated by one of the kinases acting in the PKA pathway in the presence of glucose, leading to CCR and, finally, to the CCR-mediated inhibition of the expression of cellulolytic enzymes. Our hypothesis is consistent with the report by Ribeiro et al. (2019), except for the phosphorylation site.

Here, we focused on the continuous phosphorylation of a particular motif (SSTT) at the C-terminus of Crel (Figure 1A) and analyzed mutants mimicking dephosphorylation of these sites. According to the results presented in Figure 3, an improvement of FPase and pNPCase activities contributed to a partial release from CCR. Cziferszky et al. (2002) reported that phosphorylation of another motif (HSNDEDD) was abolished when E244 was mutated to valine, following which CCR in *T. reesei* was depressed. Moreover, there is little change at the transcriptional levels of *xyl1, clr1*, and *clr2* expression in Tr_ CrelS388V, though FPase and pNPCase activities were greatly enhanced. Thus, we suggest that dephosphorylation of the motif (SSTT) has a great influence on the binding of target genes rather than the co-regulated gene, that is, *xyl1, clr1*, or *clr2*.

Therefore, we suggest that dephosphorylation modification of specific sites of Crel is a good strategy to reduce the CCR effect, and to enhance enzyme activity without affecting the growth of microorganisms. However, in order for this strategy to be effective it is crucial to identify the important phosphorylation sites, which requires a clear understanding of the regulatory pathway. Thus, this strategy will be hampered by the fact that the regulatory system is complex and currently not well understood (Ortiz-Urquiza and Keyhani, 2015; Adnan et al., 2017; Kunitake et al., 2019). Therefore, it is necessary to start from the downstream target, in this case Crel, to correctly identify the pivotal sites, providing a simple way to enhance the production of cellulase in fungal species exhibiting CCR.

**CONCLUSION**

This study precisely identifies the dephosphorylation/phosphorylation sites of Crel and the key kinase participating in CCR in *T. reesei*. Our work provides a potential workflow for creating *T. reesei* strains resistant to CCR when full-length Crel was retained. Above all, we developed a precision engineering strategy to improve the production of cellulases in fungal species under CCR, highlighting that the application of this technology could accelerate the industrial adoption of lignocellulosic biorefinery methods.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in NCBI database (https://www.ncbi.nlm.nih.gov/) under the accession numbers: AAB01677.1, EPS28222.1, AAK11189.1, XP_663799.1, AAA32690.1, XP_755510.1, AAT34979.1, AAL33631.4, and EAA32758.1.
AUTHOR CONTRIBUTIONS

LH, YT, WM, and KN performed the experiments. LH and XF wrote the manuscript and conceived the study. LH, YT, WM, SH, YL, and XF were involved in analysis and interpretation of experimental data. XF coordinated the project. All authors contributed to the article and approved the submitted version.

FUNDING

This work was supported by the National Key R&D Program of China (No. 2018YFB1501701), the National Natural Science Foundation of China (Nos. 31570040 and 31870785), the Key Technologies R&D Program of Shandong Province (No. 2018GSF121021), and the State Key Laboratory of Microbial Technology Open Projects Fund.

ACKNOWLEDGMENTS

The authors gratefully acknowledge Prof. Luying Xun’s help in revising this manuscript. The part contents in this article have been released as a pre-print at BioRxiv (Han et al., 2020).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fbioe.2020.00852/full#supplementary-material
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Conflict of Interest: SH was employed by the company Shandong Henglu Biological Technology Co., Ltd., Part of data in this manuscript have been issued in China patent application (CN 201911135400.2).

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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