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High-Throughput Selection for Cellulase Catalysts Using Chemical Complementation

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Abstract: Efficient enzymatic hydrolysis of lignocellulosic material remains one of the major bottlenecks to cost-effective conversion of biomass to ethanol. Improvement of glycosylhydrolases, however, is limited by existing medium-throughput screening technologies. Here, we report the first high-throughput selection for cellulase catalysts. This selection was developed by adapting chemical complementation to provide a growth assay for bond cleavage reactions. First, a *URA3* counter selection was adapted to link chemical dimerizer activated gene transcription to cell death. Next, the *URA3* counter selection was shown to detect cellulase activity based on cleavage of a tetrasaccharide chemical dimerizer substrate and decrease in expression of the toxic *URA3* reporter. Finally, the utility of the cellulase selection was assessed by isolating cellulases with improved activity from a cellulase library created by family DNA shuffling. This application provides further evidence that chemical complementation can be readily adapted to detect different enzymatic activities for important chemical transformations for which no natural selection exists. Because of the large number of enzyme variants that selections can now test as compared to existing medium-throughput screens for cellulases, this assay has the potential to impact the discovery of improved cellulases and other glycosylhydrolases for biomass conversion from libraries of cellulases created by mutagenesis or obtained from natural biodiversity.

Introduction

Directed evolution and biodiversity mining of environmental DNA are two powerful approaches in enzyme discovery. Directed evolution is an established method for generating enzymes with improved specific activities for industrial, research, and therapeutic applications.^{1–5} It consists of generating large numbers of protein variants and then assaying those variants en masse for the desired function. Biodiversity mining exploits the diversity found in unculturable microorganisms, which represent >99% of microorganisms,⁶ to find novel enzymes. It consists of amplifying environmental DNA in vitro, expressing the genes in a model organism, and testing them en masse for the desired function. Biodiversity mining has also

been successfully applied for the discovery of industrial, research and therapeutic enzymes.⁷ Directed evolution and biodiversity mining are limited to functional enzyme reactions that are inherently screenable or selectable, such as reactions where the product is fluorescent or an essential metabolite. To address this bottleneck, we developed chemical complementation, a general high-throughput assay for enzyme catalysts that relies on the yeast three-hybrid assay to link enzyme catalysis to reporter gene transcription in vivo. This assay has been previously applied to screen for bond cleavage reactions using a betalactamase enzyme and *lacZ* as the reporter gene.⁸ More recently, chemical complementation was adapted to select for bondformation reactions using a glycosynthase enzyme and a *LEU2* reporter gene essential for cell survival.⁹

One of the major bottlenecks to cost-competitive conversion of biomass to fermentable sugars for biofuel production is the high enzyme cost for cellulose degradation.¹⁰ Technological improvements in the breakdown of cellulose into sugars is often cited as one of the solutions to reducing cost.¹⁰ Cellulose, a β -1,4-glucose polymer, represents 38–50% of the total cellulosic

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Figure 1. Chemical complementation provides a high-throughput selection for cellulase catalysts. The discovery of improved cellulases, and other hydrolase enzymes important for biomass conversion, is limited by existing medium-throughput screening technologies. Chemical complementation has been adapted to provide a high-throughput selection for bond cleavage reactions and specifically cellulase catalysts. Chemical complementation detects enzyme catalysis of bond formation or cleavage reactions based on covalent coupling of two small molecule ligands. The heterodimeric small molecule reconstitutes a transcriptional activator, turning on the transcription of a downstream reporter gene. Here, a methotrexate (Mtx)-dexamethasone (Dex) yeast three-hybrid system is used. Endoglucanase activity is detected as cleavage of a glycosidic linkage between Mtx and Dex. A cellulase selection could be used as an initia high-throughput assay to evaluate very large libraries of cellulase variants generated either by mutagenesis or isolated from biodiversity sources. Active cellulases emerging from the high-throughput assay could be then tested using low-throughput assays using plant material that more closely mimic industrial conditions.

biomass composition.¹¹ Cellulose is hydrolyzed into glucose units by an ensemble of three enzymes: endoglucanases, also called cellulases [E.C. 3.2.1.4], exoglucanases [E.C. 3.2.1.91], and glucohydrolases [E.C. 3.2.1.21]. Glucose can then be fermented to ethanol by microorganisms. Because the ratelimiting step in enzymatic cellulose depolymerization is performed by exoglucanases and endoglucanases,¹² discovery of improved cellulases likely is a straightforward solution to decreasing costs.

Currently, the discovery of improved endoglucanases and other glycosylhydrolase enzymes important for biomass conversion is based on medium-throughput screening technologies that rely on surrogate substrates.^{12,13} The two most widely used medium-throughput assays for endoglucanase activity are a halo assay carried out on Petri dishes, which detects hydrolysis of carboxymethylcellulose (CMC) based on Congo red staining of carbohydrate reducing ends, and a UV assay, which detects hydrolysis of *p*-nitrophenyl cellobioside (pNPC).¹² Because each gene in a screen must be assayed individually, even with automatation techniques, only 10^4 to 10^6 genes can be tested, at best. A selection for cellulase catalysts would allow the search of much larger cellulase catalysts libraries (>10⁸) because only cells expressing active enzyme variants survive. The problem is that cellulase catalysis is not inherently selectable.

Here, using chemical complementation, we have developed a *URA3* counter selection for cellulase activity (see Figure 1). To the best of our knowledge, this is the first high-throughput selection for cellulase catalysts and should significantly increase the number of variants that can be tested over medium-

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throughput screens. The method was developed using the endoglucanase Cel7B from Humicola insolens that catalyzes the hydrolysis of beta-1,4-linked glucosidic bonds in cellulose. First, a URA3 counter selection was adapted to link chemical dimerizer activated gene transcription to cell death. The classic yeast URA3 5-fluoroorotic acid counter selection was chosen as the reporter gene because it is the state of the art counter selection in the two-hybrid literature. Although the URA3 counter selection has been previously applied in an *n*-hybrid system,^{14,15} re-engineering of a chemical complementation URA3 counter selection strain was necessary as previously published strains were either not available or did not meet the parameters required by the chemical complementation strategy. Next, the URA3 counter selection was shown to detect cellulase activity based on cleavage of a tetrasaccharide chemical dimerizer substrate and decrease expression of the toxic URA3 reporter. Finally, the utility of the cellulase selection was assessed by isolating cellulases with improved activity from a cellulase family DNA shuffled library.

Results

Selection Scheme. In chemical complementation, enzyme catalysis of bond formation or cleavage is linked to cell survival based on covalent coupling of two small molecule ligands to reconstitute a transcriptional activator in vivo⁸ (Figure 2A). Bond formation is detected as activation of an essential reporter gene; bond cleavage, as repression of a toxic reporter gene. The assay is high-throughput because it can be run as a growth selection in which only cells containing the functional enzyme survive. The assay can be readily extended to new chemistry by

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Figure 2. Chemical complementation detects bond cleavage reactions. (A) In the cellulase growth selection, the Methotrexate-Cellotetraose-Dexamethasone (Mtx-Cel-Dex) heterodimer links a DNA-binding domain fused to dihydrofolate reductase (DBD-DHFR) and an activation domain fused to the glucocorticoid receptor (AD-GR), activating transcription of the toxic *URA3* reporter gene, thus leading to cell death. Active cellulases cleave the β -1,4-glucosidic bond in the Mtx-Cel-Dex linker, decreasing transcription of the toxic URA3 reporter gene and leading to cell survival. (B) Structure of the Mtx-Cel-Dex chemical inducer of dimerization used for this assay. C. The *URA3* counter selection strain was constructed in three steps. First, the *URA3* gene was introduced at the ura3-52 locus of the chemical complementation. Finally, the LexA DNA binding site and the tightly regulated Spo13 promoter were introduced upstream of the *URA3* reporter gene via homologous recombination.

synthesizing small molecule heterodimers with different chemical linkers as enzyme substrates.

We envisioned that chemical complementation could detect cellulase activity as cleavage of a β -1,4-glucosidic bond in a methotrexate-cellotetraose-dexamethasone (Mtx-Cel-Dex) substrate.^{9,16} Selection for bond cleavage required the construction of a chemical complementation counter selection where cleavage of the Mtx-Cel-Dex chemical dimerizer by a cellulase would relieve transcription of a toxic reporter gene. Thus, we adapted chemical complementation to incorporate the classic yeast counter selection marker *URA3*. In the presence of 5-fluoroorotic acid (5-FOA) the *URA3* gene product, orotidine-5'-phosphate decarboxylase, produces the toxic product 5-fluorouracil (5-FU), which is misincorporated into RNA and inhibits the nucleotide

synthetic enzyme thyamidylate synthase leading to cell death.¹⁷ Cell survival is achieved by cleavage of the heterodimeric small molecule, disrupting expression of the *URA3* gene.

Design of the Mtx-Cel-Dex Substrate. Because it is the chemical dimerizer substrate that imparts modularity to chemical complementation, we first synthesized a cellulase substrate carrying the Mtx and Dex handles. *E. carotovora* CelA, CelN, and CelV [E.C. 3.2.1.4] share 68% amino acid sequence identity with *Bacillus agaradharens* Cel5A. Therefore, we took advantage of the extensive crystallographic data on this cellulase to guide the design of the Mtx-Cel-Dex substrate.^{18,19} The high-resolution structure of *B. agaradharens* Cel5A predicts five subsites in the active site that accommodate five glucose units (-3, -2, -1, 1, 2). Given that in other cellulases, four subsites

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(-2, -1, 1, 2) contribute most to the binding energy, with the fifth contributing only slightly,²⁰ the cellulase substrate was designed to have four saccharide units. Because the tetrasaccharide is embedded between Mtx and Dex, it should recapitulate the cellulase oligomer substrates. The tetrasaccharide linker is β -1,4-linked Glu-Glu-Glu-Glu. The synthesis of Mtx-Cel-Dex was carried out chemoenzymatically. Mtx-Lac-F and Dex-Cel were synthesized using published strategies from our laboratory and the linkage between the two halves was synthesized by the previously reported Cel7B:E197A glycosynthase.²¹ The full assignment of Mtx-Cel-Dex can be found in Figure 2B and Figures S1 and S2 in the Supporting Information.^{9,16}

Construction of the Chemical Complementation URA3 Counter Selection Yeast Strain. Guided by the construction of a previously published reverse yeast two-hybrid strain, we reduced the basal transcription of the URA3 reporter gene by placing it under control of the tightly regulated Spo13 promoter.¹⁴ First, the FY251 ura3–52 chromosomal locus was replaced by a functional URA3 gene and selected for Ura⁺ phenotype. Next, the plasmids carrying the DNA binding domain-dihydrofolate reductase (DBD-DHFR) and the activation domain-glucocorticoid receptor (B42-GR) fusion proteins were introduced in the strain. Finally, a LexA(4op)-pSpo13 DNA fragment with four LexA binding sites was introduced upstream of the URA3 reporter gene and selected for a Ura⁻ phenotype. This selection provided numerous chemical complementation URA3 counter selection yeast strains (Figure 2C).

Three of these strains were used to optimize the counter selection conditions by varying the level of galactose (0.5-2%), which affects the concentration of DBD-DHFR and AD-GR in the system, and 5-FOA (0.1-0.5%), which controls the toxicity level of the reporter gene (see Figure S3 in the Supporting Information). Using the optimized conditions, the counter selection strain showed on-off behavior between $1-10 \ \mu M$ Mtx-Dex, thus all future counter selection experiments were carried out at 1 μ M small molecule concentration. Next, the URA3 counter selection strain was tested with 1 μ M Mtx-Cel-Dex, the small molecule to be used in the cellulase selection, to confirm that the counter selection would also function properly with the cellulase substrate (Figure 3A). Finally, the chemical complementation URA3 counter selection yeast strain proved to be very stable after eight days under counter selection conditions. The cell growth of 93 strains under counter selection conditions was lower than the growth of the same 93 strains under noncounter selection conditions. This difference was statistically significant when analyzed with a paired one-tailed Student *t* test (p-value < 0.0001) (Figure 3B).

Chemical Complementation URA3 Counter Selection Links Cellulase Activity to Cell Survival. The chemical complementation URA3 counter selection strain was adapted to detect cellulase activity by introducing the Cel7B cellulase from *Humicola insolens*. As Figure 3C shows, expression of *H. insolens* Cel7B in the presence of Mtx-Cel-Dex conferred a growth advantage to the URA3 counter selection strain. Inside the cell, the Cel7B presumably cleaves the Mtx-Cel-Dex substrate, disrupting dimerization of the transcriptional activator, thus decreasing expression of the toxic URA3 reporter gene and leading to cell survival. As a negative control, we used the inactive cellulase variant Cel7B:E197A, obtained by mutation of the active-site nucleophile glutamic acid to alanine.^{9,21} The inactive variant lacks hydrolytic activity,^{21,22} thus it is unable to cleave the Mtx-Cel-Dex substrate, leading to transcription activation of the toxic *URA3* reporter gene and cell death.

Assessing the Utility of the Cellulase Selection. The utility of the cellulase selection was evaluated by isolating cellulases with improved activity from a cellulase library prepared by family DNA shuffling. First, the chemical complementation URA3 counter selection strain was confirmed to detect the cellulase activity of CelN, CelA and CelV (see Figure S4 in the Supporting Information). Together with *H. insolens* Cel7B the chemical complementation URA3 counter selection has successfully read out the four endoglucanases tested to date. The library was generated by shuffling the catalytic domains (signal peptides removed) of E. carotovora CelN, CelA, and CelV cellulases [E.C. 3.2.1.4], which share 90% sequence identity at the amino acid level and above 78-88% at the nucleotide level (see Figure S5 in the Supporting Information). To reach a large library size, we introduced the cellulase chimeras into the expression vector via S. cerevisiae in vivo homologous recombination, ^23, 24 yielding a library size of 1 \times 10 8 unique transformed cells. The quality of the library was determined by sequencing 10 chimeras before selection. All chimeras contained sections from the three starting cellulases and an average of 10 crossovers (see Figure S6 in the Supporting Information).

The cellulase selection can select for cellulase variants with increased activity from a family DNA shuffled cellulase library after just 5 days of selection. Using a colorimetric screen based on the quantification of aldehyde formation using CMC as the substrate,²⁵ we tested if variants isolated after selection had increased cellulase activity. As Figure 3D shows, the mean cellulase activity of the variants isolated after 5 days of selection is higher than that of variants isolated before selection. This difference is statistically significant (p-value < 0.005) when analyzed using a paired two-tailed Student *t* test.

In Vitro Characterization of Evolved Cellulase Variants. After 8 days of selection, four cellulase variants were isolated and characterized in vitro for hydrolysis of pNPC, the standard substrate for determining endoglucanase Michaelis—Menten constants. Four variants picked at random from the liquid selection after plating were overexpressed with a 6-His-tag in *E. coli*. Two variants had very low expression levels and no in vitro kinetics were measured. The other two variants, Cel_3.7 and Cel_5.7 showed excellent expression levels (see Figure S7 in the Supporting Information) and 3.7-fold and 5.7-fold increases in catalytic efficiency, respectively, over the best starting cellulase, CelN (Table 1 and Figure S8 in the Supporting Information).

Sequencing of the improved cellulase variants revealed nucleotide segments from all three starting cellulases CelV, CelA and CelN (see Figure S9 in the Supporting Information). Cel_3.7 is most similar to the parent gene CelA, while Cel_5.7 is most

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Figure 3. Chemical complementation *URA3* counter selection strain. (A) *URA3* reverse yeast three-hybrid system. Absence of small molecule leads to cell survival and an increase in absorption at 600 nm. Presence of either 1, 5, 10 μ M Mtx-Dex, or 1 μ M Mtx-Cel-Dex leads to toxic *URA3* reporter gene transcription, cell death, and a decrease in absorption at 600 nm. The experiments were carried out in triplicate. Shown is the mean growth and the error bars represent the standard deviation from the mean. (B) Reversion rate of the chemical complementation *URA3* counter selection strains carrying Cel_5.7 were tested for growth under counter selection conditions (1 μ M Mtx-Dex) and under noncounter selection conditions (No Mtx-Dex) as a control. Shown is the mean growth and the error bars represent the standard error form the mean (p-value < 0.0001). (C) The chemical complementation *URA3* counter selection strain detects cellulase activity. Presence of 1 μ M Mtx-Cel-Dex and active cellulase Cel7B leads to cell survival and an increase in absorption at 600 nm. Presence of 1 μ M Mtx-Cel-Dex and inactive cellulase Cel7B:E197A leads to cell death and a decrease in absorption at 600nm. As a control, the *URA3* counter selection strain was also grown in the absence of Mtx-Cel-Dex and either the presence of Cel7B or Cel7B:E197A. All experiments were carried out in triplicate. Shown is the mean growth and the error bars represent the standard deviation from the mean. (D) The chemical complementation *URA3* counter selection erriches for cellulase activity after five days of selection. Twenty-two cellulase variants isolated before selection and 22 variants isolated after 5 days of selection were tested for cellulase activity using carboxymethylcellulose (CMC) in crude cell extracts. Increase in aldehyde formation was detected as an increase in absorption at 660 nm. Shown is the mean (p-value < 0.005).

Table 1. Michaelis–Menten Parameters of the Starting Cellulas	е
Genes CelN, CelA, CelV and the Isolated Variants Using	
p-Nitrophenyl Cellobioside ^a	

cellulases	$\textit{K}_{m}(\textit{mM};\textit{mean} \pm \textit{s.d})$	$\textit{k}_{\rm cat}~(\rm s^{-1};~mean~\pm~s.d.)$	$k_{cat}\!/K_{M}$ (M $^{-1}$ s $^{-1};$ mean \pm s.d)
Cel V	4.6 ± 0.3	$1.10 \times 10^3 \pm 50$	$2.4 \times 10^{5} \pm 1.8 \times 10^{4}$
Cel A	7.5 ± 0.7	$1.45 \times 10^3 \pm 65$	$1.9 \times 10^5 \pm 2 \times 10^4$
Cel N	3.4 ± 0.2	$1.27 \times 10^{3} \pm 42$	$3.7 \times 10^{5} \pm 2 \times 10^{4}$
Cel_3.7	3.4 ± 0.1	$4.5 \times 10^{3} \pm 132$	$1.35 \times 10^{6} \pm 7 \times 10^{4}$
Cel_5.7	2.51 ± 0.08	$5.3 \times 10^{3} \pm 124$	$2.09 \times 10^{6} \pm 2 \times 10^{4}$

^{*a*} The standard deviations were calculated from at least three independent measurements.

similar to the parent gene CelN. Cel_3.7 has nine crossovers. When compared to CelA, Cel_3.7 differs by five point mutations: V86A, Q186L, L193M, R208C, and V254I. Cel_5.7 has eight crossovers. When compared to CelN, Cel_5.7 differs by eight point mutations: A86V, S126N, E154D, T172S, C208R, T262S, A267T, and A272T. To map the mutated residues onto the cellulase scaffold, homology models of Cel_3.7 and Cel_5.7 were made using the *B. agaradherans* Cel5A structure with a cellopentoside inhibitor.²⁶ The majority of the isolated mutations were remote from the enzyme active site.

Discussion

Together these results establish that chemical complementation provides the first high-throughput assay for cellulase catalysts, a key target in the economical conversion of cellulosic biomass to ethanol. The chemical complementation *URA3*

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selection detected all known endoglucanases that we tested-four in total. After 5 days, the cellulase growth selection was shown to enrich for cellulase variants with increased activity from a family DNA shuffling library based on carboxymethylcellulose hydrolysis in cell culture. The enrichment was statistically significant, with a p-value < 0.005. After 8 days of selection, a cellulase variant with a 6-fold increase in k_{cat}/K_{M} over the best parent enzyme for the hydrolysis of *p*-nitrophenyl cellobioside was isolated. Given that the parent cellulases are already highly active with k_{cat}/K_{M} values on the order of 10⁵ M⁻¹ s⁻¹, it is quite challenging to generate variants with improved activity. By way of comparison, it is worth noting that the most significant increase in endoglucanase activity by directed evolution in the literature is a 2.2-fold increase in activity using CMC as the substrate.²⁷ The Department of Energy's stated goal for improvement of cellulases for biomass conversion is a 10fold increase in activity of industrial cellulases on crystalline cellulose.28

For the first time, chemical complementation has been used to detect bond cleavage as cell survival. Together with the previously reported selection for bond formation reactions,⁹ we now have functional selections for bond cleavage and bond formation reactions. The *URA3*/5-FOA counter selection was chosen as the platform for the chemical complementation bond cleavage selection strain because it is the state of the art counter selection platform in the yeast two-hybrid system. Currently we are working on the design of optimized counter selections with improved dynamic ranges for biodiversity mining and directed evolution.

While applied here to cellulases, the chemical complementation counter selection should be readily adapted to other glycosylhydrolases important for bioenergy production. Once the *URA3* counter selection yeast strain was developed, all that was required to detect cellulase activity was the addition of the Mtx-Cel-Dex substrate and a cellulase. With the same ease, chemical complementation could be adapted to select for other hydrolase catalysts important for biomass conversion, such as hydrolases able to degrade pectin, hemicellulose, and even lignin. The generality of the chemical complementation selection for bond cleavage reactions should also allow testing other glycosylhydrolases not involved in bioenergy.

Ultimately, we envision using the chemical complementation high-throughput selection followed by a low-throughput assay on crystalline cellulose that mimics industrial bioreactor conditions to discover new, industrially useful cellulases. The chemical complementation assay could be used to search environmental DNA libraries to discover new cellulases, or for directed evolution to increase the activity of known cellulases. A low-throughput assay could then evaluate which of these cellulases had improved activity under industrial conditions. Although the chemical complementation assay cannot mimic the natural polymer, crystalline cellulose substrate, notably the assay does successfully read out the four endoglucanases we have tested to date. With library sizes of ca. 10⁸ achievable in yeast, chemical complementation has the potential to increase by several orders of magnitude the number of variants that can

be initially tested, thus increasing the likelihood of discovering improved variants.

Experimental Section

URA3 Counter Selection Yeast Strain Construction. Standard methods for molecular biology in *S. cerevisiae* and *E. coli* were used.^{29,30} The URA3 gene from pMW112 was recombined at the ura3-52 locus of FY251 (MATa trp1 Δ 63 his3 Δ 200 ura3-52 leu2 Δ 1 Gal+) and selected on plates lacking uracil to give strain VC2169Y. Vectors pBC398 and pKB521 carrying the GR-B42 and LexA-DHFR constructs were transformed into VC2169Y to give strain VC2291Y. The LexA(4op)-Sp013 reporter construct was obtained by fusion PCR of the LexA(4op) from pMW112 and the Sp013-promoter from MaV95.¹⁴ The reporter construct was amplified with oligos carrying 30bp homology to the URA3 promoter and the URA3 gene, recombined at the URA3 locus of VC2291Y, and selected on 5-FOA plates to give the URA3 counter selection strain VC2240Y.

Cellulase Mutagenesis. The expression vector pPPY2148 was constructed by digestion of p425Met25 (ATCC 87323) with *Hind*III and *Pst*I and introduction of an 800bp stuffer flanked by SfiI sites. A C-terminal 6-Histag was introduced to the catalytic domains, signal peptides removed, of the cellulases CelN,³¹ CelA,³² and CelV,³³ and subcloned into pPPY2148 to generate vectors pP-PY2230, pPPY2234, and pPPY2236, respectively. The cellulase genes were shuffled and amplified with primers incorporating 30bp homology to the promoter and terminator of pPPY2148 and subcloned into pPPY2148 via *S. cerevisiae* in vivo homologous recombination.

Yeast Transformation and Cellulase Selection. VC2240Y was transformed with the 12.5 μ g of cellulase chimeras and 3.6 μ g of linear pPPY2148 via electroporation using high transformation efficiency procedures with slight variations to give a library size of 1 × 10⁸ unique transformants. After recovery, 10 μ L were used to determine the library size. The rest was washed, resuspended in water and placed at 4 °C for 3 days to eliminate any residual uracil inside the cell. Once the library was determined to be >1 × 10⁷, it was resuspended in counter selection media (SC(HTL⁻) 2% gal, 2% raf, 0.2% 5-FOA, and 1 μ M Mtx-Cel-Dex, at pH 5) to a final volume of 2 mL. The selection was run for 8 days, and samples were taken on days 5 and 8. To isolate selected variants, samples of the selection were diluted and plated under nonselective conditions.

CMC Cellulase Secondary Screen. Cellulase activity was detected using a carboxymethylcellulose colorimetric secondary assay. This secondary screen for aldehyde formation using CMC was adapted to be carried out in a 96-well plate format.²⁵ The 22 colonies isolated before selection and the 22 colonies isolated after 5 days of selection were grown in 10 mL synthetic media lacking leucine for two days. The same number of cells (2×10^7) from each cell culture were arrayed on a 96-well plate and lysed using 100 μ L YPER (Novagen). The cell extract (80 μ L) was transferred onto a second 96-well plate containing 120 µL 2% CMC (low viscosity, Sigma) in 0.1 M NaOAc buffer pH 5, and the reaction was incubated at 37 °C for 24hrs. The next day, 10 µL of the reaction mixture were incubated with 90 μ L of 0.1% tetrazolium blue (0.5 M potassium sodium tartrate, 0.05 M NaOH) and heated at 98 °C for 10 min. The reaction was cooled to room temperature and absorption at 660 nm was taken immediately.

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In Vitro Characterization of Cellulases on pNPC. The three starting cellulases and the improved cellulase variants were subcloned into the *E. coli* T7 vector, pAED4, between NdeI and *Hin*dIII sites to give vectors pPPY2256 (CelN), pPPY2258 (CelA), pPPY2257 (CelV), pPPY2291 (Cel_3.7), and pPPY2292 (Cel_5.7). The cellulases were purified using Quick spin Ni-NTA affinity column (Qiagen). Protein concentration was determined using U.V. absorption ($\varepsilon = 76\,890, A_{280}$). To determine the kinetic constants for the hydrolysis of pNPC, the cellulase chimeras were incubated at room temperature in phosphate buffer (25 mM K₂HPO₄, 100 mM NaCl, pH 7) containing seven different concentrations of pNPC ranging from 12 mM to 0.185 mM. The release of *p*-nitrophenol ($\varepsilon_{pH7} = 4000 \text{ M}^{-1} \text{ cm}^{-1}, A_{420}$) was recorded continuously in a SpectraMax Plus 384 spectrophotometer at 420 nm.

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Supporting Information Available: A more complete description of the general methods for molecular biology, strain and vector construction, DNA family shuffling, yeast transformation, secondary screening, and enzyme purification. This material is available free of charge via the Internet at http://pubs.acs.org.

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