Review

**Clostridium thermocellum** as a Promising Source of Genetic Material for Designer Cellulosomes: An Overview

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Abstract: Plant biomass-based biofuels have gradually substituted for conventional energy sources thanks to their obvious advantages, such as renewability, huge quantity, wide availability, economic feasibility, and sustainability. However, to make use of the large amount of carbon sources stored in the plant cell wall, robust cellulolytic microorganisms are highly demanded to efficiently disintegrate the recalcitrant intertwined cellulose fibers to release fermentable sugars for microbial conversion. The Gram-positive, thermophilic, cellulolytic bacterium *Clostridium thermocellum* possesses a cellulolytic multienzyme complex termed the cellulosome, which has been widely considered to be nature’s finest cellulolytic machinery, fascinating scientists as an auspicious source of saccharolytic enzymes for biomass-based biofuel production. Owing to the supra-modular characteristics of the *C. thermocellum* cellulosome architecture, the cellulosomal components, including cohesin, dockerin, scaffoldin protein, and the plentiful cellulolytic and hemicellulolytic enzymes have been widely used for constructing artificial cellulosomes for basic studies and industrial applications. In addition, as the well-known microbial workhorses are naïve to biomass deconstruction, several research groups have sought to transform them from non-cellulolytic microbes into consolidated bioprocessing-enabling microbes. This review aims to update and discuss the current progress in these mentioned issues, point out their limitations, and suggest some future directions.

Keywords: *Clostridium thermocellum*; cellulase booster; artificial cellulosomes; designer cellulosomes; *Kluyveromyces marxianus*; laccases; LPMOs

1. Introduction

During evolution, cellulolytic microbes developed an extracellular multienzyme complex called the cellulosomes to boost the cellulose degradation rate at maximum levels [1]. *Clostridium thermocellum*, a Gram-positive, thermophilic anaerobic bacterium, exhibits an outstanding capability for degrading cellulolytic biomass to release fermentable sugars of different lengths by means of its powerful cellulosomes. Since the polysaccharide compositions in plant cell walls vary substantially in both quality and quantity [2], *C. thermocellum* needs to mediate the composition of saccharolytic enzymes in the cellulosome complex to cope with the complexity and recalcitrance of a specific cell wall [3]. More than 70 cellulolytic enzymes-borne type-I dockerin (DocI) are noncovalently assembled onto the primary non-catalytic protein, termed scaffoldin (Scaf) CipA, through a calcium-dependent high-affinity interaction (i.e., dissociation constant \( K_D < 10^{-11} \) M) with the nine type-I cohesin (CohI) domains located on the CipA structure [4]. In addition, the CipA protein also contains a type-II dockerin (DocII) domain at its C-terminal to mediate the attachment
of the cellulosome complex to the bacterial cell surface [5]. Correspondingly, C. thermocellum possesses three types of surface-anchoring Scaf(s), namely SdbA, Orf2p, OlpB, which contain one, two, and seven type-II cohesins (CohII), respectively, responsible for the binding of the cellulosome complex to the cell surface through CohII–DocII interaction.

Inspired by the Lego-like architecture of the C. thermocellum cellulosome, various research groups have been seeking to construct designer cellulosomes (DCs) for basic and applied studies [6–8]. Although there are several conjugation techniques to design artificial cellulosomes [9], the present review mainly focuses on the CohI–DocI interaction-based artificial cellulosome construction. Furthermore, the related issues of this approach, such as enzymatic unit positions, types of enzymes, linkers between DocI and catalytic domain, and number of carbohydrate-binding modules (CBMs), are updated and discussed. Another strategy is to recombinantly produce a library of individual catalytic enzymes with diverse hydrolytic reaction modes and then formulate the specified enzymatic cocktails for specific substrates [10]. In addition, to convert non-cellulolytic biofuels-producing microbes into consolidated bioprocessing (CBP)-enabling microbes, which can conduct enzyme production, substrate saccharification, and fermentation of the released sugars into biofuels in a single step, various studies have been carried out to express heterologous cellulolytic enzymes in heterologous host cells to make them genetically engineered cellulolytic microbes [11]. These engineered microbes use a cell-surface display or secretion system to display their novel hydrolytic capability. These strategies, with strengths and weaknesses, are discussed in the present review.

2. Reservoir of Recombinant Enzymes for In Vitro Reconstitution of Artificial Cellulosomes

2.1. Impact of Processive Endoglucanase Component in Designer Cellulosome on Biomass Decomposition

In nature, an efficient decomposition of cellulosic biomass strongly depends on a fast adaptation of C. thermocellum to enable the regulation of its cellulosomal enzymes for a specific substrate composition. The prompt response of C. thermocellum is due to an elegant alternative sigma-anti sigma system [12] that quickly and flexibly regulates the expression of cellulosomal genes in its genome. As there are numerous cellulases with diverse functions, a comparative study on the hydrolytic activity profiles and product formation of 24 C. thermocellum cellulosomal cellulases was performed by Leis et al. (2017) [13]. The cellulases can be distinguished from each other based on the hydrolysis modes on Avicel and soluble sugars release. The first group comprising non-processive endoglucanases (EG), such as Cel8A, Cel5E, Cel5H, Cel9-44J, and Cel9T, released long-chain polysaccharides with degree of polymerization (DP) ≥ 5. This group was found to show the highest activity on CMC and barley β-glucan. The second group consists of exo-mode cellobiohydrolases (CBHs) (i.e., Cel48S, Cel9K, Cbh9A) with cellobiose as the only end-product during Avicel hydrolysis. The third group is comprised of all cellotetraose-type processive EGs (pEG4) family 9, namely Cel9F, Cel9N, Cel9P, Cel9Q, Cel9R, Cel9U, Cel9V, Cel9W, Lec9A, and Lec9B. All processive enzymes in this group, except for Cel9P that does not have a CBM3c, possess a GH9 catalytic module associated with a CBM3c. The last group includes processive cellulbiose-type EGs (pEG2) such as Cel5B, Cel5G, Cel5L, Cel5O, and Cel9D. The Cel5O is the unique EG that has a CBM3b module in this group, while the other members only possess catalytic modules. The next step of this study was to form artificial mini-cellulosomes to determine synergistic effects between cellulase components. By doing so, the selected cellulases were allowed to bind to the recombinant scaffoldin protein CipA8 carrying 8 CohI subunits on its backbone structure. Regarding enzymatic assays in this study, all the enzyme assays were performed in a total volume of 0.5 mL at 60 °C. The reaction buffer consisted of 0.1 M MOPS, pH 6.5, 50 mM NaCl, 10 mM CaCl₂, and 2 mM Tris(2-carboxyethyl)phosphine (TCEP) functioning as a reducing agent. For the single cellulase assay, the barley β-glucan, CMC, PASC (0.5%, w/v), or Avicel (1%, w/v) were used. Avicel (0.25%, w/v) was used for the determination of the complex cellulases activity with the standard enzyme load of 2 µg/mL.
while the 10-fold greater of Avicel concentration (2.5%, w/v) was employed for enzyme kinetics assay. To eliminate the potential inhibition of cellobiose against the complexed cellulases, 6 μg/mL of β-glucosidase from *Thermus thermophilus* was used. Reducing sugars released from the substrates were measured using the 3,5-dinitrosalicylic acid (DNS) method and one enzymatic unit was defined as 1 μmol of glucose equivalent released per minute. After 48 h of incubation on Avicel (0.25%, w/v) at 60 °C, all divalent complexes, namely Cip8A:Cel48S/Cel9K, Cip8A:Cel9K/Cbh9A, and Cip8A:Cel48S/Cbh9A, with only CBH components bound to Cip8A, exhibited less hydrolysis efficiency. The combination of Cip8A:Cel48S/Cel9K/Cel5L (SKL) with Cel5L is a processive pEG2 enzyme, which yielded the maximum reducing sugars released (737 μM/reaction) among all trivalent mini-cellulosomal complexes. Interestingly, without pEG2 enzyme in its complex, the tetravalent cellulose Cip8A:Cel48S/Cel9K/Cel8A/Cel9R (SKAR) showed lower activity compared to the SKL, which indicates the key role of the processive pEG2 enzyme in substrate decomposition. The nonavalent DC Cip8A:Cel48S/Cel9K/Cel8A/Cel9Q/Cel9T/Cel5G/Cel9R/Cel9-44J, which carried 9 enzymes of all 4 types of cellulases on its construct, reached approximately 52.6% of activity of the native Ct cellulosome (Figure 1).

![Figure 1](image-url): Full synthetic cellulosome complex containing 25% of Cel48S, 12.5% each of Cel9K, Cbh9A, Cel8A, Cel9Q, Cel9T, and 4.2% each of Cel5G, Cel9R, and Cel9-44J (stoichiometric assembly ratio, % mol), whereas the stoichiometric binding capacity equals 1:8 (CipA:single enzyme). Abbreviations in the Figure 1: Ct, *C. thermocellum*.

### 2.2. Preparation of Recombinant *C. thermocellum* Cellulosomal Enzyme Library

In terms of industrial cellulose degradation, a highly versatile pool of free single saccharolytic enzymes enables a faster adaptation to a given substrate while reducing the addition of unnecessary enzyme components. Based on this argument, Leis and coworkers [10] aimed to prepare a full recombinant cellulosomal enzyme reservoir in which all predicted 73 DocI-containing enzymes-encoding genes in the *C. thermocellum* DSM 1313 genome and the *cip8A* gene were cloned and expressed in *E. coli* DH10B and DH5α cells. In total, 57 of the 73 DocI-bearing enzymes were successfully expressed and purified in a soluble form. An enzyme assay showed that the combination of two CBHs such as Cel48S and Cel9K and one EG Cel5L with the Scaf protein CipA8, namely CipA8:Cel48S/Cel9K/Cel5L complex (SKL), appeared to be sufficient for the hydrolysis of softwood pulp. However, the addition of a mannanase (Man26A) and a xylanase (Xyn10Y) to the SKL complex to form the SKLMY complex (i.e., CipA8:Cel48S/Cel9K/Cel5L/Man26A/Xyn10Y) dramatically increased the biomass degradation efficiency relative to the SKL. Enzyme assays were carried out at 60 °C in 0.5 μL of reaction buffer containing 0.1 M MOPS, pH 6.5, 50 mM NaCl, 10 mM CaCl$_2$. The activity of synthetic cellulosomes was measured using 0.25–0.5% (w/v) Avicel and 0.25% (w/v) micronized softwood pulp (UPM-Kymmen, Helsinki, Finland) as the substrates. The commercial Cellic Ctec2 (Novozymes, Sigma-Aldrich, St. Louis, MO, USA) was incubated at 50 °C in 0.1 M MES and 50 mM CaCl$_2$, pH 5.0. The enzyme β-glucosidase from *T. thermophilus* or *Thermoanaerobacter brockii* was supplemented to a final concentration of 6 μg/mL to avoid the inhibition of synthetic cellulosomes by cellobiose. After 48 h of incubation at 60–65 °C, the pentavalent SKLMY complex exhibited a higher thermostability, and displayed approximately 60% of enzyme activity relative to the commercial fungal enzyme mixture Cellic Ctec2. Moreover, the supplement of the 47 additional
recombinant cellulosomal enzymes from the above-mentioned enzyme reservoir or of the SM901 native free cellulosomal enzyme with varying amounts into the SKLMY complex increased its activity to a level comparable with that of the commercial enzyme blend Cellic Ctec2 (Figure 2) (Note: The mutant *C. thermocellum* strain SM901 was devoid of *cipA* gene). As far as we know, this is the first report in which 78% of all putative *C. thermocellum* cellulosomal enzymes, in full length, were recombinantly produced in *E. coli*. In addition, a core enzyme complex with only five enzymes enabling an effective hydrolysis of Kraft pulp was determined.

Figure 2. Construction of artificial cellulosome from a pool of recombinant cellulosomal enzymes from *C. thermocellum*. (A) The SKLMY complex with only 5 recombinant cellulosomal proteins exhibits 60% of catalytic efficiency as compared to the commercial Cellic Ctec2 cocktail. (B) The mixture of SKLMY complex and 47 recombinant enzymes at different ratios expresses a comparative catalytic efficiency with the commercial Cellic Ctec2.
Regarding the CohI–DocI binding flexibility, since there is no binding specificity between CohI and DocI originating from the same bacterium species *C. thermocellum*, an individual DocI-bearing catalytic component can theoretically randomly bind to any available CohI on the Scaf backbone architecture [14,15]. However, in the DCs constructed with divergent Coh–Doc pairs from different species, where the catalytic subunits were fused with divergent DocI with different specificities, the chimeric DocI is strictly required to bind to their corresponding CohI positioned on the chimeric Scaf [16]. The preparation of recombinant enzymes library from the same cellulolytic microbe, in our view, is an interesting and feasible strategy to quickly respond to the increasing interest in the search for a potent enzyme source in the cellulosic biofuel industry.

### 3. Incorporation of Novel Enzymes into Artificial Cellulosomes
#### 3.1. Integration of Laccase into Designer Cellulosomes

Although *C. thermocellum* cellulosome is among the fastest and most efficient biocatalysts known for decomposing lignocellulosic biomass, the low yield of its fermentation products limits the application of this bacterium in biofuel production systems. A DC can be constructed by artificially self-assembling divergent chimeric protein complexes including CohI-bearing Scaf, DocI-bearing saccharolytic enzymes, and CBMs into a whole complex. By virtue of their supra-modular nature, the DCs enable the control of the number of enzymes, the composition, and position of these enzymes on a Scaf construct [17]. Previous studies showed that the synergy and proximity effects among enzymes in DCs promote an enhanced solubilization of inherent recalcitrant cellulosic biomass in comparison with their free enzyme counterparts [7].

Lignocellulose, the most abundant polymers complex on Earth, is composed of cellulose, hemicellulose, and lignin. The Cel48S enzyme is the most abundant enzymatic subunit in the isolated cellulosome of *C. thermocellum* [3,18,19], and its important role in cellulose hydrolysis was obviously evidenced by Olson et al. [20] and Liu et al. [21]. Regarding catalytic components for DC construction, it is therefore not surprising that this cellobiohydrolase (CBH) family 48 was widely selected for constructing artificial cellulosomes and/or for preparing cellulase cocktails [10,13,22–24]. Similarly, the xylanase family 10 Xyn10Z responsible for xylan solubilization was frequently chosen for artificial cellulosomes [6,22]. The cellulase-xylanase synergy that facilitates the decomposition of complex substrates was noticed in previous studies [16,25–29]. In addition, the saccharolytic enzymes originated from the aerobic thermophilic bacterium *Thermobifida fusca*, with thermostability and potent catalytic efficiency [30], are identified as important starting materials for constructing DCs in several studies [16,29,31–33].

While cellulose and hemicellulose are useful components to produce biofuels and other valuable biochemicals, lignin is often considered a barrier that needs to be removed during cellulosic substrate processing step. To this end, Davidi et al. [8] constructed a DocI-fused laccase Tfu1114 and incorporated it into a DC. Tfu1114 is a free monocupper polyphenol oxidase originated from the aerobic bacterium *T. fusca*, and it can remove some C–C and/or C–O bonds near to aromatic rings of lignin [34]. To overcome a challenge in the conversion of the free laccase Tfu1114 to the cellulosomal mode, a highly expressed GH10 xylanase (XynT6) from Geobacillus steaothermophilus was used as a solubility tag and fused at the N-terminus of the Doc-containing laccase to generate XynT6–c–Lac. The XynT6 itself was fused with the *Clostridium cellulolyticum* Doc to form XynT6–c to promote xylan degradation. For cellulose conversion, the EG Cel5A and CBH Cel48A from *T. fusca* were fused with the Doc(s) from *Ruminococcus flavefaciens* (f) and *C. thermocellum* (t), respectively, to form the f–Cel5A and t–Cel48A, respectively. In addition, the xylanase Xyn11V (Cthe_2972) from *C. thermocellum* ATCC 27405 was fused with an *Acetivibrio cellulolyticus* Doc (a) to form Xyn11V–a. To construct a backbone protein for the DC, a tetravalent ScafCATF with four diverse CohI(s) from *C. cellulolyticum* (C), *A. cellulolyticus* (A), *C. thermocellum* (T), and *R. flavefaciens* (F), and a CBM3a from *C. thermocellum* was produced. The individual enzymes and the scaffoldin were incubated for 2 h at 37 °C with 20 mM CaCl$_2$ to facilitate
cellulosome assembly. The resulting cellulosome complex (0.5 µM) was examined for wheat straw decomposition in a 200-µL reaction buffer containing 50 mM acetate buffer (pH 5.0), 10 mM CaCl₂, 2 mM EDTA, and 7 g/L wheat straw. The reaction mixtures were incubated at 50 °C for 72 h. As a result, the complexation of the tetravalent ScafCATF with all the four forms of Docl-bearing GHs, termed herein ScafCATF:XynT6–c–Lac/Xyn11V–a/t–Cel48A/f–Cel5A, displayed the highest saccharolytic activity, followed by the blend of four unbound enzymes t–Cel48A, f–Cel5A, Xyn11V–a, XynT6–c, and the mixture of t–[Cel48A, f–Cel5A, Xyn11V–a, and XynT6–c–Lac. Without laccase, the complexations of ScafCATF:t–Cel48A/f–Cel5A/Xyn11V–a/XynT6–c and ScafCATF:t–Cel48A/f–Cel5A/Xyn11V–a could not maintain their robust activities toward wheat straw, thus indicating the key role of the laccase Tfu1114 in biomass hydrolysis.

3.2. Incorporation of Lytic Polysaccharide Monooxygenases (LPMOs) into Designer Cellulosomes

LPMOs, relatively new enzymes discovered one decade ago, are copper-dependent and play pivotal roles in the depolymerization of sugar-based biopolymers [35]. The low-molecular-weight lignin-derived compounds (LMWLDCs) released from biomass solubilization was found to be able to donate electrons to LPMOs for the oxidation of cell-wall polysaccharides [36]. The activities of LPMOs are copper-dependent and can only be induced in the presence of oxygen molecules and an electron donor, which can be delivered from enzyme GMC oxidoreductases, from lignin, phenolic compounds and from light-activated photosynthetic pigments. Insoluble high-molecular-weight lignin functions as a source of electrons, while the LMWLDCs transfer electrons to LPMOs and being oxidized. The LMWLDCs will be reverted to reduced state by receiving electrons from the bulk lignin, thus enabling a new electron delivery [37]. In the study by Arfi et al. [33], the two free LPMOs, such as TfLPMO10A (hereafter termed E7) and TfLPMO10B (hereafter termed E8) from T. fusca, were converted to the cellulosomal mode by fusing to a DocA (a) from A. cellulolyticus (Figure 3). Two versions of E7 were constructed: the first version E7–a was formed with the DocA fused directly at the C-terminus of the catalytic domain AA10, and the second version E7–sk–a was generated with a 12-aa short linker (sk) between AA10 and C-terminal. Three versions of E8 included E8-a with the DocA placed at its C-terminus, E8Δ–a where the native CBM domain was altered by the DocA and E8ΔΔ–a with the FnIII and the CBM were replaced by the DocA. The cellulolytic activity of E8–a was similar to that of the WT counterpart, whereas E7–a without linker exhibited a dramatically reduced activity. A trivalent ScafABT was constructed with three Coh(s) from A. cellulolyticus, B. cellulosolvens, and C. thermocellum, and one CBM3a from C. thermocellum. Additionally, the endoglucanase TfCel5A and exoglucanase TfCel48A were converted to the cellulosomal mode t–Cel5A and b–Cel48A by replacing their CBMs with Doc(s) from C. thermocellum and B. cellulosolvens, respectively. Cellulase assays were performed by incubating the enzyme with 1% (w/v) Avicel at 50 °C for 72 h in 50 mM acetate buffer (pH 5.9) and 2 mM ascorbic acid. Enzyme concentrations were 1 µM for the WT and chimeric LPMOs enzymes or 0.5 µM for the DCs. The three LPMO(s)-bearing DC(s), namely ScafABT:E8Δ–a/b–Cel48A/t–Cel5A, ScafABT:E8ΔΔ–a/b–Cel48A/t–Cel5A, and ScafABT:E7–sk–a/b–Cel48A/t–Cel5A showed the highest activity towards Avicel, suggesting the superiority of cellulosomal architecture over the free enzymes (i.e., enzymes proximity synergy, robust substrate targeting, and enzyme-enzyme synergistic effects). Based on the hydrolysis capability of DCs, it should be noted that the LPMOs function as a booster when incorporated with other cellulases in increasing saccharolytic efficiency rather than being the main cellulose-degrading enzymes.
Figure 3. Hydrolysis of cellulose by LPMO-bearing DCs. (A) The WT form of Cel5A, Cel48A, E7, E8 and the converted enzyme variants t–Cel5A, b–Cel48A, E7–a, E7–sk–a, E8–a, E8Δ–a, and E8ΔΔ–a. (B) The trivalent DCs with E7 and E8 variants exhibit the highest enzymatic activities. (C) The mixtures of chimeric individual enzymes also possess potent hydrolytic activities. (D) The mixtures of WT E8, E7, Cel5A, and Cel48A showed average activities. (E) Without LPMO in the enzyme mixtures, the capability of cellulose conversion was dramatically reduced.

3.3. Incorporation of β-glucosidase into Designer Cellulosomes

Previous studies noticed the inhibitory effect of glucose [38] and cellobiose [39,40] on cellulosomes. Therefore, eliminating this feedback inhibition enables an enhanced biomass conversion [41]. The activity of β-glucosidase BglA was found to be too low in WT C. thermocellum strains [42], and due to the adsorption of cellulosomes in cellulose
matrix, only a small fraction of BglA is available to cellulosomes [43]. To address this problem, Gefen et al. (2012) [43] fused a β-glucosidase BglA protein with an unoccupied CohII and integrated this chimeric BglA–CohII complex into the purified *C. thermocellum* cellulosome. On Avicel and pretreated switchgrass, the cellulosomes, in combination with the resultant chimeric BglA–CohII complex, showed the highest saccharolytic efficiency, followed by the cellulosomes plus free WT BglA, and the *C. thermocellum* cellulosomes alone (Figure 4A). The enzyme activity was assayed in 50 mM citrate buffer (pH 6.0) and 15 mM CaCl$_2$ with 0.475 mg/mL cellulosome and 12.5% (w/v) Avicel or 4.2% (w/v) pretreated switchgrass as the substrate. The β-glucosidase concentration for the assay was 0.75 μM for WT BglA or BglA–CohII in an equivalent specific activity. After the first reaction, the resultant supernatants were transferred into a new reaction mixture with 0.5 μM WT BglA in 50 mM citrate buffer (pH 6.0) and 15 mM CaCl$_2$. The reaction was incubated at 60 °C for 3 h to convert all cellobiose into glucose. Overall, the study of Gefen and coworkers [43] suggested that spatial proximity of β-glucosidase to cellulosomes is a crucial parameter that influences saccharification efficiency. To corroborate the finding of Gefen et al., Li et al. [44] employed a thermotolerant BGL from *Caldicellulosiruptor* sp. F32 (BglA) to fuse with the CohII or with the CBM3a from *C. thermocellum* to generate CaBglA–CohII and CaBglA–CBM3a, respectively. On Avicel, the complexations of C1 cellulosomes and CaBglA–CohII via CohII–DocII interaction exhibited the highest activity, then the mixed proteins of cellulosomes with CaBglA–CBM3a, the mixture of cellulosomes with free CaBglA, and the cellulosomes alone. In addition, the glucose production also followed this order, indicating the greatest conversion rate of CaBglA in the closeness to cellulosomes. In contrast, the accumulation of cellobiose was the greatest in the sole cellulosomes treatment, suggesting an important role of Bgl enzyme in cellulose degradation.

In the study of Moraïs and colleagues [24], a trivalent ScafGTV composed of 3 CohI(s) from thermophilic species *Archaeoglobus fulgidus* (G), *C. thermocellum* (T), and *Clostridium clari flavum* (V), respectively, and the *C. thermocellum* CBM3a for substrate targeting was constructed for artificial cellulosome formation (Figure 4B). The native DocI of EG Cel8A and its thermostable mutant Cel8A* were replaced by the DocI from *A. fulgidus* (g) to generate Cel8A–g and Cel8A*–g, respectively, and the free *C. thermocellum* BglA as well as its thermotolerant version BglA* were converted into cellulosomal mode BglA–v using the DocI (v) from *C. clari flavum* to create BglA–v and BglA*–v, respectively. The mesophilic tetravalent ScafBTFA bearing 4 divergent CohI(s) originated from *Bacteroides cellulosolvens* (B), *C. thermocellum* (T), *R. flavefaciens* (F), and *A. cellulolyticus* (A), respectively, was constructed and used as a control to compare thermostability with that of ScafGTV. Moreover, two matching DocI(s) from *B. cellulosolvens* (b), and *R. flavefaciens* (f) were used to generate Cel8A*–b, and BglA*–f. Enzymatic activity was measured with 0.5 μM free enzyme each or 0.5 μM optimized DCs in 50 mM acetate buffer (pH 5.0) using 1% (w/v) Avicel as a substrate at different incubation times and temperatures. After 96 h of incubation at 60 °C on 1% (w/v) Avicel, reducing sugar released from the complexation of ScafGTV:Cel8A*–g/Cell48S*–t/BglA*–v was the highest, followed by those from the ScafGTV:Cel8A–g/Cell48S–t/BglA–v, the thermostable free enzymes Cel8A*–g, Cell48S*–t, BglA*–v, and WT free enzymes Cel8A–g, Cel48S–t, and BglA–v, which indicates the superiority of the thermostable Scaf and catalytic modules in substrate saccharification at elevated temperatures (Figure 4B). As expected, the complex ScafBTFA:Cel8A*–b/Cell48S*–t/BglA*–f only sustained thermostability at 50 °C for 48 h of incubation. Recently, the *Thermanaerobacter brokii* β-glucosidase CgIT was fused with a DocI originated from enzyme *C. Cel48S* to promote its incorporation into artificial cellulosomes. This is one of the most efficient thermophilic β-glucosidases (BGL) identified thus far [45]. A high ratio (i.e., four to five) of BGL to cellulases (i.e., Cel48S, Cel9K, Cel8A) on the CipA exhibited the highest activity on crystalline cellulose, however, an increased or decreased ratio of CgIT to cellulases (i.e., seven CgIT to two cellulases or two CgIT to seven cellulases, respectively) also led to a decreased hydrolytic efficiency, which indicates that an appropriate enzymatic ratio of enzyme components in artificial cellulosomes is crucial for a robust saccharolytic activity (Figure 4C). This is in line with conclusions presented in...
the studies of Tang et al. [46] and Qi et al. [47] as an imbalance ratio of BGL to cellulase or an excessive BGL supplementation to enzyme assay buffer, respectively, did not benefit the saccharification process.

Figure 4. The incorporation of BGL into DCs aids to improve saccharification of cellulose. (A) The native non-cellulosomal BglA from C. thermocellum was transformed into the cellulosomal mode and integrated into Ct cellulosome via CohII–DocII interaction. (B) The thermostable mutants Cel8A, Cel48S, and BglA were used as cellulosomal enzyme components to construct trivalent artificial cellulose. (C) The impact of ratio of BglA-to-cellulases on hydrolytic activity. Note: The asterisk (*) indicates the thermostable mutant strain.

3.4. Impacts of Catalytic Enzyme Positions on Enzymatic Activity of Designer Cellulosomes

The position of catalytic subunits in the Scaf structure may govern the catalytic activity of DCs. In a study by Stern and coworkers [31], three recombinant T. fusca enzymes such as the processive EG Cel9A, EG Cel5A, and exoglucanase Cel48A were recombinantly generated and converted into the cellulosomal mode for DC constructions. Besides, ten chimeric Scaf(s), namely Scaf4, Scaf5, Scaf6, Scaf9, Scaf18, Scaf19, Scaf20, Scaf21, Scaf22, and Scaf23, with different designated orders of enzymes and CBMs were also produced. The enzymatic assay was carried out in 50 mM acetate buffer (pH 5.0), 12 mM CaCl₂, 2 mM EDTA, 10% (w/v) Avicel, incubated at 50 °C for 24–72 h with shaking. The two optimal trivalent Scaf configurations Scaf20 (i.e., ScafABT:a–Cel9A/b–Cel48A/Cel5A–t) with the processive EG Cel9A and the thermostable mutants Cel8A, Cel9R, Cel5G, and Cel48S, respectively (Figure 5A). Based on their hydrolytic efficiencies, the suggestion for future artificial cellulosomes design is that the processive EG should not be placed in the middle of the trivalent chimeric Scaf because the processive EG functions by cutting β-1,4-glycosidic bonds in the interior of the cellulose chain, followed by a processive cleavage of the even-numbered short...
oligosaccharides [48], its middle position within the trivalent Scaf likely restricted its movement along the substrate. In contrast, the processive EG should be located close to a CBM in the Scaf to properly fulfill its catalytic function. The third conclusion that can be drawn is that two enzymes with parallel characteristics (i.e., processive Cel9A and Cel5A from \textit{T. fusca} or processive Cel9R and Cel5G from \textit{C. thermocellum}) should not be placed in proximity with each other since this might lead to an unnecessary functional competition between them.

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure5.png}
\caption{Influences of enzyme positions on activity of chimeric trivalent cellulosomes. (A) Hydrolysis of Avicel by \textit{T. fusca}-based trivalent cellulosomes. (B) Avicel degradation by \textit{C. thermocellum}-based trivalent cellulosomes.}
\end{figure}

3.5. Impact of Mechanical Stability of CohI on Hydrolytic Activity of Designer Cellulosomes

The mechanical stability of CohI within a Scaf is an important parameter since it impacts the catalytic activity of cellulosomes. Although the nine CohI(s) display a high level of sequence identity [49], their robustness against mechanical stress appears to be dramatically different [50]. On the CipA protein, the region between the CBM3a and the DocII was called the connecting region, while the region that consists of CohI-1, and CohI-2 was called the external region. Previous studies revealed that CohI(s), which were located in the connecting region, demonstrated higher mechanical stability in comparison to those positioned in the external region [50, 51]. In a recent study of Galera-Prat et al. [52] the low mechanical stability CohI-1, with unfolding force \((F_U) = 124 \pm 25 \text{ pN} \) at \(400 \text{ nm/s} \) of pulling velocity, was used to produce C1 and E1, and the high mechanical stability CohI-7, with \( F_U = 480 \pm 77 \text{ pN} \) at \(400 \text{ nm/s} \) of pulling velocity, was employed to generate C7 and E7 (Figure 6). The resulting C1, E1, C7, and E7 were anchored on an artificial monovalent Scaf at predesignated positions, and then the monovalent Scaf was allowed to concurrently bind to a polystyrene microparticle, which mimicked bacterial cell surface, via a CohII of the anchoring protein SdbA. Two artificial monovalent ScafC1 and ScafC7 were designed to have one C1 and C7, respectively, placed between the CBMs and the X module/DocII (XDocII). The other ScafE1 and ScafE7 carrying one E1 and E7, respectively, positioned in the external region. Cel5A, a free EG from the thermophilic bacterium \textit{T. fusca}, which was transformed to cellulosomal mode by fusing a DocI of the Xyn10Z from \textit{C. thermocellum}, was employed as a DocI-bearing enzyme. The complexation of ScafC1:Cel5A, ScafC7:Cel5A, ScafE1:Cel5A, and ScafE7:Cel5A was confirmed by SDS-PAGE and these monovalent cellulosomes also showed similar hydrolytic activity when compared each other, but higher than that of the free Cel5A, maybe due to the targeting effect of the CBM3a in the monovalent cellulosomes. The monovalent cellulosomes were attached onto the CitS1-coated microparticle through a CohII of the anchoring Scaf SdbA to generate the bound ScafC1:Cel5A + CitS1-coated microparticle, ScafC7:Cel5A + CitS1-coated
microparticle, ScafE1:Cel5A + CtS1-coated microparticle, and ScafE7:Ce5A + CtS1-coated microparticle complexes, and then subjected to mechanical stress. Under high mechanical stress, the activities of monovalent cellulosomes in their bound and free state were measured.

To form the monovalent cellulose complex, 0.5 μM enzyme was incubated with the monovalent Scaf in a 1:1 ratio. The reaction was assayed in the reaction buffer containing 50 mM Tris (pH 7.4), 300 mM NaCl, 1 mM CaCl$_2$, 1 mg/mL BSA, 5–20% (w/v) Avicel, incubated at 37 °C for 60 min. As a result, only the hydrolytic profile of ScafC1:Ce5A + CtS1-coated microparticle complex was negatively influenced while the others’ activities remained indistinguishable (Figure 6). According to the authors [52], this could be explained by the low mechanical stability of C1 that prevented the binding of the monovalent ScafC1:Ce5A to the CtS1-coated microparticle. However, in our opinion, the release of the monovalent ScafC1:Ce5A from the ScafC1:Ce5A + CtS1-coated microparticle complex is not as important as the release of the enzyme Ce5A from the ScafC1:Ce5A structure since it was the Ce5A that took responsibility for the hydrolysis of Avicel, but not the cell surface-mimicking microparticle. Moreover, the finding from a report of Hong et al. [53] showed that the deletion of the C-terminal DocII caused the detachment of cellulosomes from the cell surface, only negatively affecting 9% of the cellulose hydrolysis rate, which suggests a small contribution of cellulosome–cell synergy. Therefore, we preferably agree with the suggestion made in the previous study [51] that the higher unfolding rate of C1, when subjected to a force pulse, might trigger the release of enzyme-bearing DocI from its structure.

Figure 6. (From the top) The CohI located in the external and connecting regions on the CipA scaffoldin protein were used to construct mini-Scaf. Mechanically unstable Coh1 was positioned between CBM3a and DocII (ScafC1) or outside of the CBM3a (ScafE1), and mechanically stable Coh7 was placed between CBM3a and DocII (ScafC7) or outside of the CBM3a (ScafE7). The complexations of ScafC1:Ce5A, ScafE1:Ce5A, ScafC7:Ce5A, and ScafE7:Ce5A was bound to CtS1-coated microparticle for biomimetic cellulosome construction.
3.6. Incorporation of Hyper-Thermophilic Enzymes into Designer Cellulosomes

Artificial cellulosomes with hydrolytic enzyme components derived from mesophilic bacteria, cellulolytic fungi, and the mildly thermophilic C. thermocellum have long been developed [54]. However, for a broader spectrum of industrial applications, better thermostable DCs are highly demanded. At 78 °C, the hyper-thermophilic, anaerobic bacterium Caldicellulosiruptor bescii degrades the unpretreated switchgrass at high substrate loading (200 g/L) [55] and Kataeva et al. [56] found that this bacterium solubilized carbohydrate and lignin concurrently. The CbCel9A/Cel48A is the most abundant of the carbohydrate-active enzymes (CAZymes) in the C. bescii exoproteome (representing almost 50% of the C. bescii exoproteome), and the most efficient single catalytic enzyme ever tested for biomass depolymerization [57]. This bifunctional enzyme consists of one family 9 processive EG Cel9A–CBM3c, one family 48 cellobiohydrolase Cel48A, and two family 3b CBM3b modules [58]. These elements were linked together by proline/threonine-rich linker peptides (Cel9A–CBM3c–CBM3b–CBM3b–Cel48A). Recently, in a study by Kahn et al. [17], the potent Cel9A/Cel48A originated from C. bescii DSM 6725 was chosen for designing artificial cellulosomes. Technical challenge did not allow an adequate expression of a full-length DocI-bearing Cel9A/Cel48A in E. coli, thus separated catalytic modules Cel9A and Cel48A were used for the conversion into the cellulosomal mode (Figure 7A). In addition, a glycoside hydrolase family 5 (GH5) EG Cel5D with a CBM28, which acts in synergy with Cel9A/Cel48A, was selected for incorporating into the artificial cellulosomes. Since C. bescii only produces free enzymes, three pairs of CohI–DocI, namely T–t, V–v, and G–g from C. thermocellum ATCC 27405, C. clariflavum, and the hyper-thermophilic archaeon A. fulgidus, respectively, were used to transform the free enzymes C. bescii Cel9A/Cel48A and Cel5D into the cellulosomal modes, creating the resulting chimeric Cel5D–g, Cel5D–t, Cel9A–sl–v, Cel48A–sk–t, Cel9A–lk–v, and Cel48A–lk–t. (Note: sk is a short linker of 3–7 aa, and lk is a long linker of 73 aa). Accordingly, three monovalent Scaf each carrying a single CohI and a CBM3a from C. thermocellum (i.e., ScafG, ScafT, ScafV) and one trivalent Scaf bearing 3 CohI(s) (i.e., ScafGTV) were designed (Figure 7A). After inoculated at 75 °C for 24 h, all three complex structures ScafG:Cel5D–g, ScafT:Cel5D–t, and ScafV:Cel5D–v almost maintained their stability (92–100%), whereas 35%, 32%, and 42% reductions in the stability of the individual unbound enzymes Cel5D–g, Cel5D–t, and Cel5D–v were observed, respectively. Under the same incubation condition, the stabilities of individual chimeric ScafG, ScafT, and ScafV were reduced by 26%, 11%, and 40%, respectively. The trivalent ScafGTV alone showed 100% stability at 70 °C and 60% at 75 °C for 24 h. The complexation of ScafGTV with any single form of the chimeric Cel5D enzyme, termed ScafGTV:Cel5D–g, ScafGTV:Cel5D–t, ScafGTV:Cel5D–v, also exhibited an absolute stability at 70 °C and decreased to 70–85% at 75 °C after 24 h of incubation. Intriguingly, the complexation of ScafGTV with all 3 forms of Cel5D enzyme, termed ScafGTV:Cel5D–g/Cel5D–t/Cel5D–v, was much stable at 75 °C with 96% stability. The optimized 0.05 µM enzyme concentration was used to compare the activity of all the free enzyme and the single enzymes bound to their monovalent Scaf. For cellulosome complex formation, each Scaf was incubated with its corresponding enzymes at 37 °C for 2 h in the buffer containing TBS, 10 mM CaCl2, and 0.05% Tween. The enzymatic assay was performed on 4% (w/v) Avicel at 60 °C and 75 °C for 24–72 h in 50 mM acetate buffer, pH 5.5, 900 rpm. With regards to the effects of linker length on DC activities, the DC with long-linker Cel9A–lk–v appeared to have higher activity than other DCs with short-linker Cel9A–sk–v (Figure 7B). In a previous study, Caspi et al. [59] also found that length of the linker between catalytic module and the DocI had little effect of the activity of hydrolytic enzyme. Instead, the location of the DocI, at the N- or C-terminal of the enzyme, likely had more effect on enzyme activity. It is noteworthy that most cellulosomal enzymes bear DocI on the C-terminal sides of the catalytic modules, therefore, the fusion position of any heterologous DocI to catalytic entities should be thoroughly scrutinized. Tang et al. [46] noticed that natural linkers from T. reesei CBH1 and C. cellulovorans EngB improved the activity of Sf-dBGL1 by 2-fold, whereas the synthetic linker from a commercial plasmid pYD1 did not. In contrast, the synthetic linker
was found to increase the activity of enzyme Te-dCBH1, but the naturally derived linkers did not. Especially, for the Ct-CelA, none of the three linkers could enhance its activity. Based on these instances, we might assume that the impacts of linkers on enzyme activity may vary case by case, thus no consensus on this issue has been reached yet.

Figure 7. Development of E. coli-based DC. (A) Conversion of the wild-type (WT) C. bescii Cel9A, Cel48A into chimeric forms using E. coli as an expression host. (B) Influence of linker length in DC enzyme components on hydrolytic efficiency (24 h of incubation at 75 °C, 4% (w/v) Avicel). (C) Comparison of hydrolytic activities between C. bescii-based DC, chimeric C. bescii enzymes, WT C. bescii enzymes, and native Ct cellulose, (72 h of incubation at 75 °C, 4% (w/v) Avicel). (D) Effects of temperature on the activities of C. thermocellum-based DC and C. bescii-based DC (72 h of incubation at 75 °C, 4% (w/v) Avicel).

After 72 h of incubation at optimal temperature 60 °C for C. thermocellum enzymes, with the synergy of Cel48S and Cel9R [60], the C. thermocellum-based DC (i.e., ScafTBA:Cel5G–
t/b–Cel48S/a–Cel9R) [31] produced higher reducing sugar release than the ScafGTV:Cel5D–g/Cel48A–lk–t/Cel9A–lk–v, indicating a predominant efficiency of the *C. thermocellum* enzymes at mild thermophilic conditions. However, after 72 h of incubation at 75 °C, an opposite situation was observed as the complexation of ScafGTV:Cel5D–g/Cel48A–lk–t/Cel9A–lk–v showed a far higher saccharolytic activity than that of the *C. thermocellum*-based DC and even higher than that of the native *C. thermocellum* cellulosome whose cellulose-degrading capacity dramatically lost at such a higher temperature (Figure 7C,D). Synergy between enzymes, however, was not observed in the *C. bescii*-based DCs. In fact, the catalytic efficiencies of the individual bound Cel48A–lk–t and Cel48A–sk–t and their unbound forms were dramatically low on Avicel due to the lack of an appropriate glycosylation system in the *E. coli* XL1 cells [61].

### 3.7. Effects of Glycosylation on Enzyme Activity

The lack of a proper glycosylation system in *E. coli* seems to prevent *C. bescii* enzymes from fully expressing their capability [17]. Therefore, to improve the cellulolytic activity of the DCs through glycosylation, in a recently published paper, Kahn and coworkers [61] used *C. bescii* as an expression host to produce glycosylated CbCel9A–lk–v, CbCel48A–lk–t, and chimeric CbScaf–GTV. The DocI-bearing Cel5D failed to express in *C. bescii* and the CbCel48A–lk–t could not bind to Scaf–T, thus the non-glycosylated DocI-bearing versions of Cel48A and Cel5D, with short linker, were expressed in *E. coli* to produce EcCel48A–sk–t and EcCel5D–g, respectively (Figure 8A). Additionally, the other DC components from their previous study [17] such as EcScaf–GTV and EcCel9A–lk–v were used to form *E. coli*-based DC (Figure 8A). As a result, the *C. bescii*-based DC (CbScafGTV:EcCel5D–g/EcCel48A–sk–t/CbCel9A–lk–v) showed the highest activity, followed by the mixture of unbound individual enzymes EcCel5D–g + EcCel48A–sk–t + CbCel9A–lk–v, *E. coli*-based DC (EcScafGTV:Eec5D–g/EcCel48A–sk–t/EcCel9A–lk–v), and the cocktail of unbound individual enzymes EcCel5D–g + EcCel48A–sk–t + EcCel9A–lk–v (Figure 8B,C). The result revealed that glycosylated enzymes, even in their unbound form, can express a comparable activity relative to the unglycosylated DCs. Moreover, the attachment of O-linked α-1,2-galactose-containing oligosaccharides on the linker of catalytic modules likely has more impact on the overall DC activity than the attachment of oligosaccharides on the Scaf protein. Notably, glycosylation also improves thermostability of the DC, suggesting that a posttranslational modification is an important part in DC engineering (Figure 8D). The findings from Kahn and coworkers [17,61] indicate that the hyper-thermotolerant cellulolytic enzymes from *C. bescii*, with a marvelous thermostolerant property, are potential catalysts for biomass decomposition. A better thermostolerance enzyme also makes enzyme recycling more feasible and cellulosic substrates-based biofuels more cost-effective relative to the fossil-derived fuels. However, the discrepancy in released reducing sugar data from these two reports remains to be explained because the authors used the same protocol for enzyme assay. For instance, with the same enzyme assay protocol with 4% (w/v) Avicel, 72 h of incubation at 75 °C, in their previous paper [17], the authors found approximately 18.5 mM sugar released when using *E. coli*-based DC (EcScafGTV:EcCel5D–g/EcCel48A–lk–t/EcCel9A–lk–v), whereas in their second paper [62], they only recorded about 7.3–7.8 mM reducing sugar released from the EcScafGTV:EcCel5D–g/EcCel48A–sk–t/EcCel9A–lk–v. Note that no statistically significant difference was found in the catalytic efficiency of the EcCel48A–lk–t and EcCel48A–sk–t [17]. The inconsistency makes their conclusion less pronounced because the glycosylated *C. bescii*-based DC (CbScafGTV:EcCel5D–g/EcCel48A–sk–t/CbCel9A–lk–v), which was stated as the best DC in this study, only produced roughly 11–12 mM reducing sugar [61].
Figure 8. Development of C. bescii-based DC. (A) Conversion of the WT C. bescii enzymes into chimeric forms using E. coli or C. bescii as an expression host. (B) Influence of linker length in DC enzyme components on hydrolytic efficiency (24-h incubation at 75 °C, 4% (w/v) Avicel). (C) Comparison of hydrolytic activities between C. bescii-based DC, chimeric C. bescii enzymes, WT C. bescii enzymes, and native C. thermocellum cellulosome, (72-h incubation at 75 °C, 4% (w/v) Avicel). (D) Effects of temperature on the activities of C. thermocellum-based DC and C. bescii-based DC (Heat treatment at 85 °C for 3 h, followed by 16-h incubation at 75 °C, 4% (w/v) Avicel).
4. Conversion of Non-Cellulolytic Biofuel Microbes into Consolidated Bioprocessing Microbes

4.1. Creation of Cellulolytic Bacillus Subtilis

Although well-known for the robust cellulose machinery, the slow growth rate of C. thermocellum with strictly anaerobic culturing conditions, the highly cost of cellulose production due to the low productivity, and its metabolic intermediates likely appear undesirable for industrial applications [62]. The Gram-positive bacterium Bacillus subtilis has been commonly used as a workhorse to produce numerous recombinant proteins [63]. As a super CBP microbe remains unfound, a co-culturing system that harmoniously combines one cellulolytic B. subtilis and another potent bioethanol-producing yeast is likely a feasible approach. In the study of Chang et al. [22], two polycistronic operons with the same gene collection but different orders, namely Type-I: cipA–cel9K–cel48S–cel9R–sdbA–cel8A–xyn10C–xyn10Z and Type-II: cipA–xyn10Z–xyn10C–cel8A–sdbA–cel9K–cel9R–cel48S, were constructed using the ordered gene assembly in B. subtilis (OGAB) method [64] to evaluate the influence of enzyme position on catalytic performance (Figure 9). Genes in these two operons were driven by a strong, thermo-inducible Pr promoter from phage lambda [65]. To measure the enzyme activity, 50 μL of the 25-fold condensed culture supernatant of engineered B. subtilis was mixed with 50 μL of 50 mM sodium acetate (pH 5.0), 2% (w/v) PASC or 2% (w/v) xylan. The type-I engineered B. subtilis strain released higher reducing sugars than the type-II strain when growing on Avicel and filter paper, whereas the type-II strain exhibited more robust capability of degrading Napier grass powder than the type-I strain, suggesting an important role of xylanases enzyme in the decomposition of native plant biomass. The finding was also in agreement with the finding of Stern et al. [31], in that the enzyme position within a given enzyme complex, e.g., DCs, is of importance for the optimum solubilization of a specific substrate.

Figure 9. Two cell-surface displayed artificial cellulosomes anchored on B. subtilis cells. The upper operon represents the order of enzyme-encoding genes where the CBH and EG genes were placed near the Pr promoter. The lower operon displays enzyme-encoding genes where the Xyn genes were positioned near the Pr promoter.

4.2. CBP-Enabling Saccharomyces cerevisiae

CBP is a promising approach to reduce the cost of biofuel production from cellulosic biomass. However, the well-known biofuel workhorses such as S. cerevisiae and Kluyveromyces marxianus lack a hydrolytic enzyme system to efficiently degrade cellulose and hemicellulose. Advances in modern genetic engineering and synthetic biology enable biofuels scientists to introduce many saccharolytic genes derived from divergent species, both prokaryotes and eukaryotes, into the target host cells to make them engineered CBP microbes. The direct conversion of cellulose into bioethanol through yeast surface-
displayed synthetic cellulosomes is a promising approach for CBP [46]. In a study by Fan et al. [11], individual minicellulosomes, each carrying a primary ScafI, an EG (CelCCA), a cellobiohydrolase (CelCCE), and a BGL (Cel_2454) from \textit{C. cellulolyticum}, were engineered to display on cell surface of \textit{S. cerevisiae}. The enzymes BGL, EG, and CBH were fused with DocI-1 from \textit{C. cellulovorans}, DocI-2 from \textit{C. cellulolyticum}, and DocI-3 from \textit{C. thermocellum}, respectively. The primary ScafI contained a C-terminal DocII, a CBM, and three CohI-1, CohI-2, and CohI-3 that were able to interact with their matching DocI-1, DocI-2, and DocI-3, respectively. The ScafII carrying one to four CohII(s) that mediate the anchoring of minicellulosomes onto cell surface. The display of ScafII was mediated using a pYD1 Yeast Display Vector Kit containing a yeast-secretion signal. The supernatants containing cellulases and miniScaf were mixed in 100 mM Tris-HCl buffer (pH 5.5), 10 mM CaCl$_2$, 2 h of incubation at 4 °C for minicellulosome assembly. The reaction was conducted by incubating the minicellulosomes in 100 mM Tris-HCl (pH 5.5), 10 mM CaCl$_2$, at 30 °C for 16 h with 0.1% (w/v) Avicel, PASC or CMC. However, ethanol titer produced by the recombinant \textit{S. cerevisiae} EBY was extremely low with only 1.41 mg/L and 1.1 mg/L on 0.1% (w/v) Avicel and 0.1% (PASC), respectively. In a study by Liang et al. [66], five enzymes, including a GH61a (a LPMO) from \textit{Thermoanascus aurantiacus}, a cellobiose dehydrogenase (CDH) from \textit{Humicola insolens}, a CBH and an EG from \textit{Trichoderma reesei}, and a BGL from \textit{Aspergillus aculeatus} were assembled onto a five-CohI \textit{C. thermocellum} miniScaf [66]. This miniScaf was successfully displayed on the surface of \textit{S. cerevisiae} through an AGA1 and AGA2 yeast display system, and the engineered \textit{S. cerevisiae} produced 2.7 g/L ethanol on 0.1% PASC.

A double-layered cellulosome was synthesized in a study by Tang et al. [46] where an artificial ScafAGA3 bearing the repeated N-terminus of Aga1p (tAga1p) was displayed on the cell surface of \textit{S. cerevisiae} through the Aga1p C-terminal domain. The ScafAGA3 was used as an anchoring protein via Aga1p-Aga2p linkage and the ScafCipA3 functioned as the primary Scaf for cellulase assembly (Figure 10). The ScafCipA3, which contained three CohI(s) from \textit{C. thermocellum} on its structure, carried three cellulases from divergent microbes via CohI–DocI interaction. For the conversion of free cellulases into cellulosomal mode, a BGL from \textit{Saccharomycopsis fibuligera}, a CBH from \textit{Talaromyces emersonii}, and an EG CelA from \textit{C. thermocellum} were fused with the traditional \textit{C. thermocellum} DocI(s). The novel disulfide bonds showed higher display efficiency of the ScafAGA3 on the cell surface in comparison with that of the conventional CtScafCipA3. The result demonstrates that the covalent disulfide bonds of tAga1p–Aga2 appeared to outperform the non-covalent bonds of the conventional Coh–DocI pair in cellulase assembling. For enzyme activity measurement, the enzymes were mixed with 5 mM p-nitrophenyl-β-D-cellobiose (pNPC) or carboxymethylcellulose sodium salt (CMC-Na) as the substrates in 50 mM citrate buffer (pH 5.0) at 50 °C for 30 min for CBH and EG measurements, respectively. The resultant engineered \textit{S. cerevisiae} carrying the double-layered cellulosome ScafAGA5–ScafCipA3:CBH1/CelA/BGLA produced 1.52 g/L ethanol on 1% (w/v) PASC. In summary, despite many efforts to create the CBP \textit{S. cerevisiae}, the bioethanol produced by these engineered strains have been still modest due mainly to the lower efficiency of the surface-displayed cellulosomes. Some catalytic proteins assembled onto the DCs did not have sufficient ability to digest high substrate concentrations to supply an abundant fermentable sugar source for the CBP \textit{S. cerevisiae}. 
Yeast

Scaf A5

Scaf Cip A3

Figure 10. The attachment of trivalent Scaf on S. cerevisiae cell surface via covalent disulfide bonds of tAga1p–Aga2. This approach helps to improve surface display efficiency relative to the common CohII–DocII binding.

4.3. CBP-Enabling Pichia pastoris

Pichia pastoris has been widely used for the expression of diverse recombinant proteins. The limited production of endogenous secretory proteins in P. pastoris makes the purification of recombinant proteins easy [67]. Additionally, an appropriate posttranslational modification is an advantage in producing proteins with correct folding and proper biological activity. Recently, an indirect P. pastoris surface-display method was developed to create a CBP cell factory [68]. Instead of using the usual non-covalent interactions between Coh and DocI from C. thermocellum, the colicin E7 DNase (CE7) and its matching immunity protein 7 (Im7) was used as a CE7–Im7 protein pair. The CE7 was mutated to inactivate DNase activity but retained its full binding affinity to generate CL7 tag, and the Im7, a cognate inhibitor of CE7, was employed to form an ultra-high-affinity IM7–CL7 protein pair (K_D ~10^{-14}–10^{-17} M) [69]. The IM7–CL7 protein pair was used as an alternative to the conventional Coh-DocI pair for cellulosome assembly (Figure 11). A CBH from Yarrowia lipolytica, an EG Cel9D from C. thermocellum DSM1237, a BGL from Thermoanaerobacterium thermosaccharolyticum, and a CBM from T. fusca were fused with N-terminal CL7 tags and recombinantly expressed in E. coli. The surface anchoring protein SED from S. cerevisiae was fused to the IM7 scaffoldin protein to mediate the attachment of the enzyme complex onto cell surface. In turn, the IM7 proteins were engineered to display for two or three times to carry two to three catalytic modules, thus generating Y-IM2 and Y-IM3 yeasts, respectively. Subsequently, the engineered P. pastoris yeasts were incubated with the E. coli lysates containing cellulosomal-mode-cellulases and CBMs to promote the assembly of minicellulosomes on cell surface via IM7–CL7 interaction. On Avicel and PASC, the engineered Y-IM2 performed better than Y-IM3 with 2.5 g/L and 1.2 g/L ethanol, respectively, whereas the Y-IM3 surpassed Y-IM2 on CMC substrate with up to 5.1 g/L ethanol. Moreover, a synergistic effect on CMC hydrolysis was also observed in Y-IM3, indicating a positive relationship between the number of Scaf IM7 and the catalytic efficiency.
The novel ultra-high-affinity IM7–CL7 protein pair ($K_D \sim 10^{-14} - 10^{-17}$ M) was used for cellulase assembly onto $P. \text{pastoris}$ cell surface. (A) The engineered $P. \text{pastoris}$ with one BGL, and one EG enzyme on its surface. (B) The engineered $P. \text{pastoris}$ with one CBH and one CBM on its surface. (C) The engineered $P. \text{pastoris}$ with one CBH, one EG, one CBH, and one CBM on its surface.

4.4. Consolidated Bioprocessing-Enabling K. marxianus

In $C. \text{thermcellum}$, aside from bearing CohI(s) on its backbone, the CipA has a DocII subunit that interacts with the CohI on the bacterial cell surface to mediate the attachment of the entire Scaf and cellulolytic enzymes complex to the $C. \text{thermcellum}$ cell surface. The anchoring outer layer protein B (OlpB) has seven CohI(s) subunits, so it can theoretically concurrently carry up to 63 saccharolytic enzymes. Generally, the incorporation of more catalytic enzymes onto the primary Scaf results in higher synergy for substrate decomposition [70]. The thermostolerant $K. \text{marxianus}$ has several advantages (e.g., a wide range of sugars assimilation, thermo, salinity, and toxin tolerance, a wide range of pH, high ethanol yield at elevated temperature) for industrial applications [71,72], and therefore it could be used as an alternative to $S. \text{cerevisiae}$ in specific conditions. Recently, the anchoring protein OlpB was successfully introduced and expressed in $K. \text{marxianus}$ cells by Anandharaj and coworkers [73]. In this study, lytic polysaccharide monoxygenases (LPMO), the copper-dependent enzymes, which can efficiently solubilize crystalline cellulose by cleaving polysaccharides through an oxidative mechanism, was converted to the cellulosomal mode for constructing DC. These enzymes were brought to light in the last decade and some previous studies that integrated LPMO enzyme into cellulolose complex were performed by Arfi et al. (2014) [33] and Liang et al. [66]. Apart from the LPMO from $\text{Thermo} \sigma \text{auro} \text{ntiacus}$ (TaLPMO) and its electron donor cellubiose dehydrogenase from $\text{Myceliophthora} \text{thermophila}$ (MtCDH), an EG from $T. \text{reesei}$ (TrEgIII), a synthetic CBH (NpaBGS), and a BGL from $\text{Neocallimastix} \text{patriciarum}$ (NpaBGS) were fused with a DocI of $C. \text{thermcellum}$ to create DocI-fused enzyme subunits, namely TrEgIII–t, CBHII–t, NpaBGS–t, TaLPMO–t, and MtCDH–t, respectively (Figure 12). In addition, a cell-surface protein glycosylphosphatidylglycerol (GPI) from $S. \text{cerevisiae}$ (ScGPI) was used to facilitate the anchoring of the artificial cellulosome to the $K. \text{marxianus}$ cell surface because the anchoring domain of $C. \text{thermcellum}$ is not suitable for eukaryotic hosts. The supernatants of red fluorescent protein (RFP)-fused DocT (RFP–DocT), OlpB–ScGPI, and CipA1B9C were mixed to form the entire cellulolose complex, namely OlpB–ScGPI–CipA1B9C–RFP–DocT, and their assembly was confirmed using an epifluorescence microscopy. The hydrolytic activity of cellulolose on 1% (w/v) Avicel was quantified by mixing the concentrated supernatants of the YP culture broth with Avicel and incubated at 40 °C for 6 h. On the model micro-crystalline cellulose Avicel, the amounts of sugar release were significantly influenced by the number of Coh and the number of CBMs, whereas the effect of these components was less pronounced on amorphous cellulose phosphoric acid swollen cellulose (PASC) hydrolysis (Figure 12A,B). The results confirmed the decisive roles of the scaffoldin protein CipA in cellulose solubilization rate as elucidated in a previous study by Olson et al. [74]. To the best of our knowledge, this is the first time an assembly of 63 saccharolytic cellulosomal enzymes from different species was engineered and expressed on a heterologous host’s surface, enabling a greater biomass degradation rate for future consolidated bioprocessing (CBP) microorganisms (Figure 12C).
Figure 12. Engineered *K. marxianus* carrying the largest cellulosome complex on its cell surface. (A) (from the top) Full-length of CipA with 1 CBM3a (1B9C), full-length of CipA with 2 CBM3a(s) (2B9C), full-length of CipA without CBM3a (0B9C), truncated CipA with 6 CohI(s) + 1 CBM3a (1B6C), and truncated CipA with 3 CohI(s) + 1 CBM3a (1B3C), respectively. (B) Ethanol productivity of different engineered *K. marxianus* strains. On Avicel, the SH-2B9C strain yielded the highest ethanol titer, followed by the SH-1B9C strain, and the lowest ethanol concentration was found in SH-0B9C which does not have any CBM, indicating the decisive role of a CBM in crystalline cellulose hydrolysis. On PASC, however, the lowest ethanol titer was recorded in SH-1B3C strain, suggesting that the numbers of catalytic subunits, but not CBM3a(s), dictate the efficiency of amorphous cellulose solubilization. (C) The engineered *K. marxianus* with the whole cellulosome complex consisting of total 63 enzyme subunits from divergent species on its surface. Abbreviations in the Figure 12: B, Carbohydrate-binding module; C, Cohesin.

5. Promise, Limitations and Future Directions

The search more effective biomass-degrading DCs or cellulase blends is an onerous task. For further optimization of artificial cellulosomal compositions, more investigations are
required to support a better understanding of the hydrolysis modes, especially synergistic effects between divergent cellulolytic enzymes in a whole complex. Despite some difficulties in the conversion of free enzymes into cellulosomal modes due to the incompatibility between host cells and the expression of the transgenes, this approach has been a common strategy in recent years because it enables scientists to exploit novel enzyme sources with advantageous characteristics (i.e., hyperthermal stability, lignin degrader, cellulase boosters). Furthermore, the reasons underlying unsuccessful cases of many *C. thermocellum* cellulosomal enzymes that could not be recombinantly produced in *E. coli* cells remain unknown but may be also related to the suboptimal heterologous protein expression in the host cells. Although O-linked α-1,2-galactose-containing oligosaccharides were found on the linkers of scaffoldin proteins from *C. thermocellum* and *Bacteroides cellulosolvens* for decades, the impact of glycosylation on the performance of artificial cellulosomes was recently confirmed as the glycosylated DCs exhibited enhanced saccharolytic effectiveness and thermal stability. The glycosylation of DCs should therefore be taken into consideration as a new parameter for DCs construction in future studies. Besides, mechano-stability of the Coh(s) is also an important parameter for preserving net saccharification activity under great mechanical load. Furthermore, the finding of new protein pairs other than CohI-DocI pairs is a new trend noticed in some recently published papers to improve binding affinity between saccharolytic domains, scaffoldin protein, and/or cell-surface-display effectiveness.

The development of robust microbes that can ferment cellulose to biofuels or other valuable products is an interesting but challenging proposition for the biofuels industry. Despite the many efforts that have been deployed thus far to convert the non-cellulolytic ethanol producers into CBP microbes, it is likely that these engineered microbes, including *S. cerevisiae*, *P. pastoris*, and *K. marxianus*, could not perform well as we might expect. For instance, although well-armed with up to 63 cellulolytic enzymes on the surface, the engineered *K. marxianus* only produced 3.09 g/L and 8.61 g/L ethanol from microcrystalline cellulose Avicel and amorphous cellulose PASC, respectively, let alone the native recalcitrant biomass. These ethanol titers, though higher than any other engineered yeast cellulosome, are still quite low to make the artificial cellulolytic *K. marxianus* CBP ready.

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**Abbreviations**

A. cellulolyticus, *Acetivibrio cellulolyticus*; A. fulgidus, *Archaoglobus fulgidus*; A. aculeatus, *Aspergillus aculeatus*; B. cellulosolvens, *Bacteroides cellulosolvens*; BGL, Beta-glucosidase; CBM, Carbohydrate-binding module; CBP, Consolidated bioprocessing; *C. cellulolyticum*, *Clostridium cellulolyticum*; C. clariflavum, *Clostridium clarilavum*; CMC, Carboxymethylcellulose; CMC-Na, Carboxymethylcellulose sodium salt; C. bescii, *Caldicellulosiruptor bescii*; CBH, Cellobiohydrolase; CohI, Type-I cohesin; CohII, Type-II cohesion; C. thermocellum, *Clostridium thermocellum*; DC, Designer cellulosome; DocI, Type-I dockerin; DocII, Type-II dockerin; DP, Degree of polymerization; EG, Endoglucanase; GH, Glycoside hydrolase; H. insolens, *Humicola insolens*; HPAEC-PAD, High pressure anion exchange chromatography coupled with pulsed amperometric detector; *K. marxianus*, *Kluyveromyces marxianus*; lk, Long linker; PASC, Phosphoric acid swollen cellulose; pNPC, p-nitrophenyl-β-D-celllobioside; pNPG, p-nitrophenyl-β-D-glucopyranoside; R. flavifaciens, *Ruminococcus flavifaciens*; sk, Short linker; Scaf, Scaffoldin; SLH, Surface layer homology; T. brockii, *Thermoanaerobacter brockii*; T. fusca, *Thermobifida fusca*; T. reesei, *Trichoderma reesei*. 
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