Expressing accessory proteins in cellulolytic *Yarrowia lipolytica* to improve the conversion yield of recalcitrant cellulose

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**Abstract**

**Background:** A recently constructed cellulolytic *Yarrowia lipolytica* is able to grow efficiently on an industrial organo-solv cellulose pulp, but shows limited ability to degrade crystalline cellulose. In this work, we have further engineered this strain, adding accessory proteins xylanase II (XYNII), lytic polysaccharide monooxygenase (LPMO), and swollenin (SWO) from *Trichoderma reesei* in order to enhance the degradation of recalcitrant substrate.

**Results:** The production of EG I was enhanced using a promoter engineering strategy. This provided a new cellulolytic *Y. lipolytica* strain, which compared to the parent strain, exhibited higher hydrolytic activity on different cellulosic substrates. Furthermore, three accessory proteins, *TrXYNII*, *TrLPMOA* and *TrSWO*, were individually expressed in cellulolytic and non-cellulolytic *Y. lipolytica*. The amount of rh*TrXYNII* and rh*TrLPMOA* secreted by non-cellulolytic *Y. lipolytica* in YTD medium during batch cultivation in flasks was approximately 62 and 52 mg/L, respectively. The purified rh*TrXYNII* showed a specific activity of 532 U/mg-protein on beechwood xylan, while rh*TrLPMOA* exhibited a specific activity of 14.4 U/g-protein when using the Amplex Red/horseradish peroxidase assay. Characterization of rh*TrLPMOA* revealed that this protein displays broad specificity against β-(1,4)-linked glucans, but is inactive on xylan. Further studies showed that the presence of *TrLPMOA* synergistically enhanced enzymatic hydrolysis of cellulose by cellulases, while *TrSWO1* boosted cellulose hydrolysis only when it was applied before the action of cellulases. The presence of r*rTrXYNII* enhanced enzymatic hydrolysis of an industrial cellulose pulp and of wheat straw. Co-expressing *TrXYNII* and *TrLPMOA* in cellulolytic *Y. lipolytica* with enhanced EG I production procured a novel engineered *Y. lipolytica* strain that displayed enhanced ability to degrade both amorphous (CIMV-cellulose) and recalcitrant crystalline cellulose in complex biomass (wheat straw) by 16 and 90%, respectively.

**Conclusions:** This study has provided a potent cellulose-degrading *Y. lipolytica* strain that co-expresses a core set of cellulolytic enzymes and some accessory proteins. Results reveal that the tuning of cellulase production and the production of accessory proteins leads to optimized performance. Accordingly, the beneficial effect of accessory proteins for cellulase-mediated degradation of cellulose is underlined, especially when crystalline cellulose and complex biomass are used as substrates. Findings specifically underline the benefits and specific properties of swollenin. Although in our study swollenin clearly promoted cellulase action, its use requires process redesign to accommodate its specific mode of action.

**Keywords:** *Yarrowia lipolytica*, Cellulolytic biocatalyst, Consolidated bioprocessing, Accessory proteins, Xylanase, Lytic polysaccharide monooxygenase, Swollenin
Background

The use of lignocellulosic biomass (LCB) as a manufacturing raw material is regarded as a key feature of the bioeconomy, because it will allow industry to transit to a sustainable, low carbon future [1, 2]. However, processing lignocellulosic biomass in an economically viable way remains a challenge, despite the considerable worldwide efforts.

Cellulose, the main component of lignocellulosic biomass, is built from linear β-glucan chains containing several hundreds of β-1,4-linked glucosyl units. In nature, cellulose exists in both ordered crystalline forms, in which multiple polysaccharide chains are densely packed into microfibrils, and to a lesser extent in disordered amorphous forms. One of the aims of biomass transformation processes is to increase the proportion of amorphous cellulose, since this form is then more amenable for further processing [3]. However, when processing plant cell walls, cellulose crystallinity is not the only hurdle to overcome, since rather like reinforced concrete, cellulose ‘rods’ (bundles of microfibrils) are embedded in a matrix, which in this case is composed of hemicellulose and lignin [4]. Together, these features render lignocellulosic biomass particularly recalcitrant to all but the most severe processing strategies [5].

Biochemical conversion of lignocellulose into target products usually occurs in three macro-operations. Biomass pretreatment often involves a combination of physical and chemical strategies that are deployed to modify the structure of lignocellulosic biomass, separating cellulose from lignin and hemicellulose, reducing complexity and increasing the accessible surface area [6]. Pretreatment is followed by enzymatic depolymerization of the polysaccharides, generating fermentable sugars using cellulases and hemicellulases. Finally, fermentation using a suitable microorganism is employed to transform sugars into the desired product(s) [7]. Among these steps, pretreatment and enzymatic hydrolysis are still significant cost drivers, with enzyme loadings and thus costs still exceeding those that have been targeted for many years [8].

One way to reduce cost in biomass processing is to integrate some of the steps and reduce or eliminate the need for external commercial enzymes. This is known as consolidated bioprocessing (CBP) and to achieve this, it is necessary to employ a cellulolytic microorganism (or a microbial consortium) that can both hydrolyse biomass polysaccharides and convert sugars into a target product [9]. However, despite the attractiveness of this concept, it is difficult to achieve because very few naturally occurring microorganisms can perform these two functions in an economically viable and industrially compatible manner [10]. Therefore, it is pertinent to look towards microbial strain engineering strategies in order to confer to an engineered microorganism the attributes required for CBP.

To engineer a microorganism for CBP purposes, it is most common to first select a microorganism that displays a suitable metabolism for target product manufacture and then confer it with cellulosytic capability, using heterologous expression of cellulosases of fungal or bacterial origin [9]. Using this approach, several engineered CBP microorganisms have been successfully produced (reviewed in [9, 11]). However, these have mostly been tested using model cellulose substrates that are excessively amenable to enzyme hydrolysis and/or produce low product titers [7]. As a result, when such engineered CBP strains are confronted with industrial cellulose pulps, the addition of exogenous enzymes is necessary to achieve complete hydrolysis [11].

The extensively studied cellulosytic secretome of the soft-rot fungus *T. reesei* (syn. *Hypocrea jecorina*) contains several cellulosases that synergistically act on complex substrates [12]. In the *T. reesei* cellulosytic secretome, EG I and EG II are the main endo-acting enzymes, representing approximately 15 and 10% (w/w) of the total protein content, respectively [13], while CBH I and CBH II (Cel7A and Cel6A) are the major exo-acting components, representing 50 and 20% of the total protein content, respectively [1]. The minimal requirement for a cellulase cocktail composed of free enzymes includes at least one endoglucanase (EG, EC 3.2.1.4), one cellobiohydrolase (CBH, EC 3.2.1.91) and a β-glucosidase (BGL, EC 3.2.1.21) [14], the latter being necessary to ensure the production of glucose from celloolodextrins. Nevertheless, most commercial cocktails are more complex and are completed with a range of accessory enzymes, including hemicellulases, cinnamic acid and acetyl esterases that together remove hemicellulose and thus increase cellulose accessibility and decrease the risk of cellulase inhibition by xyl-o-oligosaccharides [15, 16]. Until recently, assembling a sufficiently comprehensive array of polysaccharide hydrolases in a cocktail constituted the main strategy to achieve the hydrolysis of cellulose pulps. However, the recent discovery of lytic polysaccharide monooxygenases (LPMOs) has dramatically changed this view and opened up new options for the engineering of efficient cellulosytic microbial cell factories [8].

Belonging to the so-called Auxiliary Activity family 9 (AA9, formerly GH61) of the CAZY classification [17], LPMOs have been shown to significantly boost the overall efficiency of cellulose hydrolysis when using canonical hydrolytic cellulases. Studies reveal that the LPMOs catalyze the oxidative cleavage of insoluble polysaccharides using molecular oxygen or peroxide and an electron donor [18, 19]. Although most characterized
LPMOs oxidize polysaccharides at the C1 position (type 1 PMOs), yielding lactone [19–21], some perform oxidation at the C4 position (type 2 PMOs) or at both the C1 and C4 positions (type 3 PMOs), or at C6. In all cases, the action of LPMOs leads to the formation of a ketoaldose [20, 22, 23]. Considering the clear benefits of LPMOs for cellulase-mediated hydrolysis of cellulose, enzyme manufacturers have incorporated these into the latest generation of commercial cellulase cocktails [24].

In addition to LPMOs, it has been known for some time that fungal cellulolytic secretomes and plant cell walls contain non-enzymatic proteins that possess the ability to disrupt the ordered hydrogen-bonding network in crystalline cellulose. Although the exact role of proteins such as swollenins (fungal origin) and expansins (plant origin) has not been fully elucidated, research reveals that they enhance cellulose-mediated hydrolysis of cellulose, probably by disrupting the cellulose surface, making it more amenable to attack by cellulases [25–27].

Recently, we conferred cellulolytic properties to *Yarrowia lipolytica*, an oleaginous yeast that is recognized for its industrial usefulness and safety (it is classified by the FDA as a Generally Recognized As Safe strain) [28, 29]. To achieve this, we expressed in a *Y. lipolytica* strain a range of enzymes including BGLs, EGs, and CBHs, taking care to control the relative proportions of each of these in order to obtain a combination that is quantitatively similar to that of the native cellulase system of *T. reesei*. The engineered cellulolytic *Y. lipolytica* was shown to grow efficiently on industrial cellulose pulp (CIMV-cellulose, mostly amorphous), but displayed some difficulty to degrade recalcitrant crystalline cellulose [30]. Therefore, in the present work, we describe how the cellulolytic *Y. lipolytica* strain has been further manipulated to increase the hydrolysis of crystalline cellulose and complex substrates. This has been achieved by altering the proportions of the expressed cellulases and adding accessory proteins XYNII, LPMO, and swollenin.

**Results and discussion**

Enhancing the production of *T. reesei* endoglucanase I in cellulolytic *Y. lipolytica*

Previous studies have revealed that the efficient hydrolysis of pretreated LCB requires the presence of 25–35% (w/w) EG I in the cellulase cocktail, and that EG I cannot be replaced by EG II [31–33]. Additionally, our results showed that recombinant EG I exhibits a two-fold higher specific activity on insoluble substrates, such as Avicel, β-1,3 and β-1,4 glucans, than EG II. Together, these findings underline the important role of EG I for the hydrolysis of recalcitrant biomass. Unfortunately, in the previously described engineered cellulolytic *Y. lipolytica* strain (YLC6), the secretion yield of *T. reesei* EG I was approximately 60% lower than that of EG II [30]. Therefore, to alter this ratio, a hybrid promoter strategy [34] was employed, modifying the core TEF promoter element controlling EG I expression. Following successful promoter modification, analyses revealed that all of the previously introduced genes were present (Additional file 1: Figure S1) and that the new strain, YLC7, was able to grow efficiently on industrial cellulose pulp (CIMV-cellulose, mostly amorphous), but displayed some difficulty to degrade recalcitrant crystalline cellulose [30]. Therefore, in the present work, we describe how the cellulolytic *Y. lipolytica* strain has been further manipulated to increase the hydrolysis of crystalline cellulose and complex substrates. This has been achieved by altering the proportions of the expressed cellulases and adding accessory proteins XYNII, LPMO, and swollenin.
to hydrolyze CMC, PASC, and Avicel cellulose (Fig. 1). Comparing the performance of YLC7 to the parental strain YLC6 revealed that HTEF promoter-controlled expression of rTrEG I led to 18, 14, 10, and 17% increases in hydrolytic activity on CMC, PASC, Avicel, and CIMV-Cellulose, respectively. Using anti-His Western blot analysis, it was possible to correlate these increases with the improved expression level of rhTrEG I in the secretome of YLC6 (Additional file 1: Figure S2). Since our previous results showed that His6-tagging did not influence rTrEG I production [30], it is reasonable to attribute the increase in hydrolytic activity of YLC7 to its higher rTrEG I expression, although the reported cellulolytic activity is the composite result of all secreted cellulase activity.

Expression and characterization of T. reesei xylanase II in Y. lipolytica

The results of previous studies focused on the design of optimal cellulase formulations for complex biomass hydrolysis have emphasized the requirement for xylanase, which acts synergistically with cellulases [31–33]. Accordingly, we expressed the T. reesei xylanase II (rTrXYNII) in Y. lipolytica JMY1212 under TEF promoter control and using the lipase 2 pre-pro region to facilitate protein secretion, characterized the resultant recombinant enzyme, and then investigated how rTrXYNII could be expressed in the genetic background of YLC7. Screening on solid medium containing AZCL-arabinoxylan of Y. lipolytica transformants readily revealed the presence of clones producing either rTrXYNII or its His-tagged variant rhTrXYNII (Fig. 2a). To further confirm successful expression of TrXYNII, positive clones were grown in liquid YTD medium and xylanase activity was assayed in the culture supernatant. Accordingly, it was possible to demonstrate that during cultivation xylanase activity steadily increased over a 96-h period, reaching a final activity (in the case of rhTrXYNII) of 32.0 U/mL (Fig. 2b). Furthermore, using SDS-PAGE the comparison of the
rhTrXYNII-containing culture supernatant to that of a control culture (Fig. 2c) revealed the presence of a discrete species migrating to a position corresponding to an approximate Mw of 22 kDa, which is consistent with the theoretical Mw of TrXYNII (22.0 kDa).

In order to further investigate the properties of rhTrXYNII, the protein was purified from the culture supernatant (Fig. 2d) and its activity was measured on beechnwood xylan in different conditions of pH and temperature. This revealed that rhTrXYNII reached optimal activity (532 U/mg-protein) at pH 6.0 and 60 °C (Additional file 1: Figure S3), and that higher temperatures led to rapid enzyme inactivation (data not shown). These findings are consistent with previous data obtained for rTrXYNII expressed in Pichia pastoris [35]. Knowing the specific activity of rhTrXYNII, it was also possible to calculate that 62 mg/L of rhTrXYNII was secreted by Y. lipolytica, when growing in batch mode in YTD medium contained in flasks.

Expression of T. reesei lytic polysaccharide monooxygenase A in Y. lipolytica
In the T. reesei secretome LPMOs form a very minor component [36]. However, recent in vitro studies have shown that the inclusion of these enzymes in cellulase cocktails can reduce overall enzyme loadings, while maintaining the efficiency of cellulose conversion [37, 38]. Therefore, considering that it would be useful to include TrLPMOA in the cellulase cocktail expressed by YLC7, we cloned and expressed a His-tagged rhTrLP-MOA in Y. lipolytica JMY1212 under TEF promoter control and using the lipase 2 pre-pro region to facilitate protein secretion. Successful secretion of rhTrLPMOA into the culture supernatant was first established by SDS-PAGE and anti-His Western blot analyses. These revealed a smear in the Mw range 60–200 kDa, suggesting that the expected 34.4 kDa protein species was glycosylated (Fig. 3), which is unsurprising since yeasts are well known to perform hyper N-mannosylation [39]. To investigate this, rhTrLPMOA was submitted to Endo H-mediated deglycosylation and further SDS-PAGE/Western blot analysis, which revealed a better defined protein population that migrated to a position correlating to a median Mw of 55 kDa (Fig. 3a). This result suggests that even after the removal of N-glycosyl moieties, rhTrLPMOA still bears additional post-translational modifications, possibly O-glycosylation of serine/threonine-rich linker region [40].

To confirm that the expressed recombinant protein was active, the ability of rhTrLPMOA to mediate the reduction of O2 to H2O2 in the presence of ascorbate was established. Moreover, using this test, it was shown that the expression level of rhTrLPMOA steadily increased over a 72-h period, reaching a final activity in the culture supernatant of 0.75 U/L (Fig. 3b).

Characterization of the rhTrLPMOA expressed in Y. lipolytica
To further assess the functionality of rhTrLPMOA, the protein was purified in a single step, with an overall yield > 60% (Fig. 4a) and characterized. The purified rhTrLPMOA exhibited a specific activity of 14.4 U/g. This value is similar to that of the Neurospora crassa LPMO9F (15 U/g) expressed in P. pastoris, which is the most active LPMO reported so far using the same activity assay [8]. The amount of rhTrLPMOA secreted by Y. lipolytica in YTD medium during batch cultivation in flasks was approximately 52 mg/L, a yield that is moderate compared to other examples of LPMO heterologous expression [8, 41].

To further investigate the substrate specificities of rhTrLPMOA, we measured the production of H2O2 in the presence of 1.0% (w/v) PASC, CMC, Avicel, barley glucan (β-1,3; 1,4), and beechwood xylan. The repression in H2O2 production was observed for rhTrLPMOA in the presence of the four glucose-polymers (Fig. 4b) while xylan had no effect on H2O2 production. These results suggest that rhTrLPMOA displays broad specificity on β-(1,4)-linked glucans, but is inactive on xylan.

To further investigate the regioselectivity of rhTrLPMOA, tests were performed using celloolixitrins (DP4-6). These revealed the production of β-glucuronic acid, confirming that oxidation occurred at C1 (Fig. 4c). Moreover, HPAEC analysis failed to reveal any C4-oxidized species (data not shown). Taken together, these results indicate that rhTrLPMOA displays type 1-like LPMO activity, a conclusion that contradicts previous sequence and phylogeny analyses that suggest it is a type 3 LPMO (i.e., one that oxidizes at both the C1 and C4 positions) [40]. However, these apparently conflictual findings can be resolved by attributing TrLPMOA to a recently described type 3 subgroup (PMO3*) that contains LPMOs such as the one from Myceliophthora thermophila AA9 LPMO (MYCTH 92668) that also only performs oxidation at C1 [40].
higher than the expected Mw of rhTrSWO1 (50 kDa) (Fig. 5a). However, Western blot analysis using anti-His antibody confirmed that this large protein species was His-tagged (Fig. 5b), indicating that it is a highly glycosylated variant of rhTrSWO1, a finding that concurs with data in the literature [26, 27]. Using NetNGlyc 1.0 (http://www.cbs.dtu.dk/services/NetNGlyc/) to analyze the amino acid sequence of rhTrSWO1 has predicted that SWO1 bears three potential N-glycosylation sites. Therefore, to explore this possibility the protein was treated with Endo H and analyzed by Western blot. This revealed a newly formed discrete protein species that migrates to position corresponding to a Mw of approximately 75 kDa (Fig. 5b), a result consistent with the analysis of native SWO1 isolated from the T. reesei secretome [27]. Presumably the discrepancy between the predicted Mw for the polypeptide sequence of SWO1 and that measured (75 KDa) is due to other post-translational modifications, such as O-glycosylation of serine/threonine-rich linker region [26]. Finally, to further investigate the functionality of rhTrSWO1, the protein was purified (Fig. 5c).

Combining the action of LPMOA or SWO1 with cellulases for Avicel hydrolysis
To investigate the synergy of rhTrLPMOA and rhTrSWO1 with cellulases, tests were performed on microcrystalline cellulose (Avicel) using a commercial cellulase cocktail (Celluclast) supplemented with each recombinant protein at different concentrations. Celluclast was used because, unlike Cellic™ CTEC2 [37], it is devoid of LPMOs and SWO. Adding rhTrLPMOA alone significantly accelerated cellulose hydrolysis (Fig. 6a),
with rate being dependant on the concentration of rhTrLPMOA. When using 15 mg rhTrLPMOA per g cellulose, the conversion of Avicel (20 g/L) to soluble reducing sugars reached 60% within 72 h, which constitutes a 40% increase compared with the control (Fig. 6a). In contrast, adding rhTrSWO1 alone failed to produce any measurable effect on cellulose hydrolysis, a result that is consistent with previous data and suggests that swollenin does not display synergy with EG and CBH [42]. Nevertheless, the incubation of Avicel with rhTrSWO1 (5–15 mg rhTrSWO1/g cellulose) for a 24-h period prior to the reaction enhanced Avicel hydrolysis, with up to 70% conversion to reducing sugars occurring after 72 h when using 15 mg rhTrSWO1/g cellulose (Fig. 6b), although higher loadings did not procure further improvements (data not shown). Overall, these data indicate that swollenin adsorption is a prerequisite for amorphogenesis and that this is governed by a time-dependant equilibrium. For optimal Avicel amorphogenesis, it is necessary to leave sufficient time for several cycles of adsorption/desorption to occur [26, 42]. Finally, it is noteworthy that when using a similar protein load, the effect of rhTrSWO1 on cellulase-mediated Avicel hydrolysis was more potent than that of rhTrLPMOA, indicating that the use of SWO instead of LPMO in a biorefinery process might be more cost-effective. However, for this to be feasible, it would be necessary to propose a redesigned process and account for the cost burden associated with the time needed for the action of SWO.

**Construction of recombinant Y. lipolytica strains expressing multiple cellulases and accessory enzymes**

To further enhance the potency of cellulose hydrolysis by an engineered *Y. lipolytica*, the genes encoding *Tr*XY-NII, *Tr*LPMOA, and *Tr*SWO1 were singly expressed in the cellulase-producing strain YLC7 (Table 1). The presence of all of the exogenous genes was established using PCR (Additional file 1: Figure S1). In case of YLC8 (i.e.,
expressing rTrXYNII), subsequent screening on two different solid culture media containing AZCL-arabinoxylan and Azo-CMC respectively, revealed that clones displayed both xylanase and cellulase activity (Additional file 1: Figure S4; Fig. 2). Testing the ability of the supernatants of each of the daughter strains, YLC8 (rTrXYNII), YLC9 (rTrLPMOA), and YLC10 (rTrSWO1), respectively, to hydrolyze CMC, PASC, Avicel, and CIMV-cellulose (containing hemicellulose), revealed contrasting results (Fig. 1). A 12% increase in glucose production compared to YLC7 was procured by YLC8 on CIMV-cellulose hydrolysis, indicating that the presence of a xylanase is specifically beneficial to hydrolyze industrial cellulose pulp. This suggests that the xylanase works synergistically with the cellulases probably by removing hemicelluloses wrapped around glucan microfibrils and thus increasing the cellulose accessibility [43]. Indeed, the action of the xylanase was further confirmed by HPAEC analysis (data not shown), which revealed the presence of xylodextrins among the hydrolysis products. The presence of rTrLPMOA (YLC9) enhanced the hydrolysis of all of the substrates tested, which is consistent with its ability to oxidize cellulose (Fig. 1). In contrast, the co-expression of rTrSWO1 with the cellulases (YLC10) did not procure a significant improvement of hydrolysis, consistent with the observation that rTrSWO1 needs sufficient contact time and confirming that the mode of action of this protein is incompatible with a co-expression strategy. For this reason, and in view of the other results, only rTrXYNII and rTrLPMOA were retained for co-expression in the genetic

Fig. 5 Expression of TrSWO1 in Y. lipolytica a SDS-PAGE analysis of the culture supernatant of Y. lipolytica control strain expressing empty plasmid and the recombinant strain expressing rhTrSWO1; b Western blot analysis of the heterologous SWO1 protein produced by Y. lipolytica; lane 1, rhTrSWO1; lane 2, rhTrSWO1 treated by Endo H; c SDS-PAGE analysis of the purified rhTrSWO1
background of YLC7, thus producing YLC11. When using the latter strain for hydrolysis tests on the different substrates, it became clear that the highest enhancement was procured on CIMV-cellulose, while the hydrolysis of pure cellulose (CMC and Avicel) did not change when compared to YLC9 (Fig. 1). We further compared the hydrolytic efficiency of the YLC11 secretome with that of the commercial cocktail Cellic CTec2, using CIMV-cellulose, Avicel and wheat straw as substrates. The results showed that when using equivalent enzyme loadings (10 FPU/g-cellulose) the YLC11 secretome degraded all three substrates faster than Cellic CTec2 (Fig. 7). The CIMV-cellulose is the most amenable substrate, since the YLC11 secretome achieved 95% conversion of this substrate to glucose in 24 h, while Cellic CTec2 achieved 86% over the same time period (Fig. 7a). However, the superiority of the YLC11 secretome is better illustrated by its action on Avicel and wheat straw. On these substrates, the secretome achieved 75 and 30% conversion, respectively, whereas the action of Cellic CTec2 yielded 63% conversion of Avicel and 24% conversion of cellulose in wheat straw (Fig. 7b, c). When compared to the commercial cocktail, the higher potency of the YLC11 secretome can probably be attributed to several factors, the first being the presence of a more efficient CBH I [30]. Secondly, the ratio of the different enzymes is optimized in the YLC11 secretome and, thirdly, the amount of LPMO present is probably higher. Additionally, the presence of xylanase in the YLC11 secretome is not doubt useful for the hydrolysis of wheat straw. Therefore, we conclude that YLC11 is a particularly relevant strain for use with industrial cellulose pulp and recalcitrance biomass.

**Table 1 Microbial strains used in the present study**

| Strains     | Relevant genotype | Source of reference |
|-------------|-------------------|---------------------|
| T. reesei QM9414 | Wildtype         | DSMZ                |
| E. coli DH5   | Δθ0lacZΔm15, recA1, endA1, gyrA96, thi-1, hsdR17 (rk−, mk+), supE44, relA1, Δ(lacZYA-argF) U169 | Invitrogen          |
| Y. lipolytica JMY1212 (Zeta) | MATA, ura3-302, leu2-270-LEU2-zeta, xpr1-2-322 Δlip2, Δlip7, Δlip8 | [52]                |
| ylTXYNII     | Zeta, pTEF-XYNII-His6 | This investigation |
| ylLPMOA      | Zeta, pTEF-LPMOA-His6 | This investigation |
| ylTSWO1      | Zeta, pTEF-SWO1-His6 | This investigation |
| Y. lipolytica YLC6 | ΔpoxB12, pTEF-EGI, pTEF-EGII, pHTEF-NcCBHI, pHTEF-TrCBHIII | [30]                |
| YLC6b        | ΔpoxB12, pTEF-EGI-His6, pTEF-EGII, pHTEF-NcCBHI, pHTEF-TrCBHIII | This investigation |
| YLC7         | ΔpoxB12, pHTEF-EGI, pTEF-EGII, pHTEF-NcCBHI, pHTEF-TrCBHIII | This investigation |
| YLC7b        | ΔpoxB12, pHTEF-EGI-His6, pTEF-EGII, pHTEF-NcCBHI, pHTEF-TrCBHIII | This investigation |
| YLC8         | YLC7, pTEF-XYN II  | This investigation |
| YLC9         | YLC7, pTEF-LPMOA   | This investigation |
| YLC10        | YLC7, pTEF-SWO1    | This investigation |
| YLC11        | YLC7, pTEF-XYNII, pTEF-LPMOA | This investigation |
The growth of recombinant *Y. lipolytica* strains on cellulosic feedstocks

To investigate the suitability of the different strains described in this study for use in a consolidated bioprocessing concept, each of them (YLC6 to 11) was grown in liquid defined medium containing either CIMV-cellulose, Avicel, or milled wheat straw as the carbon source. Results revealed that all of the strains were able to use CIMV-cellulose, with conversion yields of more than 50% being achieved. In contrast, Avicel and wheat straw were less amenable to hydrolysis (Table 2). In terms of microbial biomass yields, these reached approximately 0.4 g-DCW/g-cellulose on CIMV-cellulose, but only (average) 0.3 and 0.1 g-DCW/g-cellulose on Avicel and wheat straw, respectively. It is noteworthy that the increased expression of r*TrEGI* (i.e., comparing YLC7 to YLC6) resulted in a slight increase in the conversion of all the cellulosic substrates, and predictably the expression of r*TrXYNII* (YLC8) enhanced the conversion of CIMV-cellulose (64.0 ± 0.4%), while not affecting Avicel conversion. Interestingly, the expression of r*TrXYNII* also enhanced the conversion of wheat straw (12.3 ± 0.5%), consistent with the fact that significant amounts of xylan are present in this substrate. Strain YLC9 expressing r*TrLPMOA* consumed 65% of CIMV-cellulose, 36% of Avicel, and 16% of cellulose in wheat straw, results that are comparable or better than those obtained with YLC7 and 8 (producing r*TrXYNII*). When both r*TrXYNII* and r*TrLPMOA* (YLC11) were present, the conversion of CIMV-cellulose and wheat straw were further enhanced (68 and 18%, respectively). Finally, while the treatment with r*TrSWO1* (15 mg-swollenin/g-cellulose) before strain cultivation failed to affect CIMV-cellulose conversion, it did prove to be beneficial for Avicel and wheat straw, with YLC11-mediated conversion rates being 42% (Avicel) and 22% (wheat straw), respectively, after 5 days of growth. However, it is also noteworthy that CIMV-cellulose is obtained using organosolv technology, which is known to produce relatively pure cellulose [44]. In further work, it will be interesting to confront YLC11 with cellulose pulps produced using less costly, but more ‘quick and dirty’ technologies [45].

Among the different accessory proteins that have been used in this study, LPMOA stands out, because it requires an electron donor and oxygen to function. Therefore, when tests were performed, ascorbic acid was included in the reactions. However, for CIMV-cellulose and wheat straw, this was unnecessary as similar yields of cellulose conversion and biomass production were achieved with YLC9 and YLC11 without the supplementation of ascorbic acid (Additional file 1: Table S2), probably because lignin present in the substrates acts as an electron donor [46]. Additionally, it has been shown that when using conventional cellulase cocktails supplemented with LPMOs, enhancement of cellulose hydrolysis will only occur under aerobic conditions [47]. In this respect, it is noteworthy that *Y. lipolytica* is strictly aerobic, thus the production of LPMOA in CBP-mode should...
be quite advantageous [29]. Finally, it is also important to note that *Y. lipolytica* can assimilate gluconic acid, an important attribute when LPMOA is present (Additional file 1: Figure S5).

Regarding swollenins, our study clearly confirms the great potential for these proteins, but underlines their incompatibility with a straightforward CBP concept. To draw benefit from swollenins, it will be necessary to devise a hybrid process in which exogenously produced swollenins are used at some point during biomass pretreatment, taking into account their specific operating requirements (i.e., contact time, temperature, and protein loading).

Finally, this study confirmed the usefulness of xylanases in the context of cellulose hydrolysis. Based on our findings and those of previous studies [48], we postulate that further engineering of *Y. lipolytica* to confer additional β-xylosidase activity and the ability to metabolize xylene efficiently [49] will procure a quite potent CBP strain. If successful, this could pave the way for the use of *Y. lipolytica* in the biorefinery industry.

### Conclusion

In the present study, a *Y. lipolytica* strain co-expressing core cellulolytic enzymes and accessory proteins has been successfully developed. The results reveal that accessory proteins can significantly enhance cellulose hydrolysis and confer cellulolytic *Y. lipolytica* with the ability on one hand to better handle industrial pulps containing xylan and on the other to hydrolyze more recalcitrant crystalline substrates. Overall, the data presented here confirm the feasibility of a *Y. lipolytica*-based consolidated bioprocess concept and reveal further ways to improve performance. In particular, the results obtained with swollenin provide insight into how this type of protein could be used in an advanced biorefinery concept involving a swollenin-mediated biomass pretreatment step and a CBP unit operation. Further strain engineering combined with appropriate process design will undoubtedly lead to much better performances in the future.

### Methods

#### Strains and media

The genotypes of the microbial strains used in the present study are summarized in Table 1. *E. coli* DH5 was purchased from Invitrogen (Paisley, UK) and used for plasmid construction. The *Y. lipolytica* strains were routinely cultivated in a medium composed of 1% w/v yeast extract, 1% w/v Bacto peptone, and 1% w/v glucose (YPD), solid media contained 1.5% agar. Transformants were selected on solid YNB medium (0.17% w/v YNB, 1% glucose or cellobiose w/v, 0.5% w/v ammonium chloride, with (for Ura<sup>+</sup>) or without (for Leu<sup>+</sup>) 0.2% w/v casamino acids and 50 mM sodium-potassium phosphate buffer, pH 6.8), supplemented with uracil (440 mg/L) or leucine (440 mg/L) depending on the auxotrophic requirements. The detection of xylanase activity in solid YNB medium was achieved by incorporating 2.0% w/v AZCL-arabinoxylan (Megazyme). For cellulase characterization, enzymes were produced in YTD medium (1% w/v yeast extract, 2% w/v tryptone, 5% w/v glucose and 100 M phosphate buffer, pH 6.8). To evaluate the growth of engineered cellulolytic *Y. lipolytica* on cellulose, transformants were aerobically cultivated in defined medium containing vitamins, trace elements [50], and salts, including 3.5 g/L (NH₄)₂SO₄, 3.0 g/L KH₂PO₄, 3.0 g/L NaH₂PO₄, and 1.0 g/L MgSO₄·7H₂O with 27.5 g/L CIMV-cellulose (91% w/v cellulose, provided by CIMV S.A.) [44], or 25 g/L Avicel PH-101 (Sigma) or 46 g/L wheat straw (particle size ~ 0.5 mm, 27% w/w arabinoxylan, and 44% w/w cellulose) [51]. In addition, ascorbate (1 mM) was added into the cultures as the reducing agent to reactivate LPMOA.

### Table 2 Comparison of cellulose utilization and biomass yield of cellulolytic *Y. lipolytica* grown on different cellulosic substrates for 120 h in aerobic cultivation

| Strains | CIMV-cellulose consumed % | Biomass yield % | Avicel consumed % | Biomass yield % | Cellulose consumed % (wheat straw) | Biomass yield % |
|---------|---------------------------|----------------|-------------------|----------------|------------------------------------|----------------|
| YLC6    | 58.6 ± 0.5 | 0.41 ± 0.03 | 30.2 ± 0.3 | 0.32 ± 0.03 | 9.7 ± 0.6 | 0.12 ± 0.01 |
| YLC7    | 61.5 ± 0.6 | 0.40 ± 0.05 | 32.0 ± 0.5 | 0.31 ± 0.02 | 10.5 ± 0.7 | 0.11 ± 0.01 |
| YLC8    | 64.0 ± 0.4 | 0.40 ± 0.02 | 31.8 ± 1.0 | 0.31 ± 0.02 | 12.3 ± 0.5 | 0.12 ± 0.01 |
| YLC9    | 65.4 ± 0.7 | 0.39 ± 0.03 | 36.3 ± 0.9 | 0.29 ± 0.03 | 15.8 ± 0.9 | 0.15 ± 0.01 |
| YLC10   | 59.0 ± 1.5 | 0.40 ± 0.04 | 31.0 ± 1.2 | 0.31 ± 0.04 | 10.3 ± 0.9 | 0.11 ± 0.02 |
| YLC11   | 67.8 ± 1.0 | 0.39 ± 0.02 | 36.5 ± 0.7 | 0.29 ± 0.02 | 18.4 ± 0.8 | 0.15 ± 0.02 |
| YLC11 + S<sup>e</sup> | 68.3 ± 1.3 | 0.38 ± 0.01 | 42.0 ± 1.1 | 0.30 ± 0.04 | 21.6 ± 0.5 | 0.17 ± 0.01 |

The results were calculated from at least three biological replicates, and are given as the mean value ± standard deviation. The initial cellulose content was 25 g/L for all the substrates.

* The cellulose was treated by SWO1 at the dosage of 15 mg/g cellulose for 24 h before enzymatic hydrolysis.
Plasmid constructions

The plasmids constructed in the present study are summarized in Table 3 and all primers are listed in Table 4. Briefly, the total RNA from 5-day cultured T. reesei QM9414 was isolated using RNase Plus Mini Kit (QIAGEN) and reverse transcription was performed with iScript™ cDNA Synthesis Kit (BIO-RAD) according to the manufacturer’s instructions. LPMOA gene (formerly known as GH61A, GenBank accession code: Y11113.1) was amplified from the cDNA of T. reesei by PCR using F (1) as forward primer and R (1) as reverse primer. A 15-base pair homologous sequence of the target plasmid was introduced into the upstream and downstream of each gene during PCR amplification. After that, the gene encoding LPMOA was fused with the PCR fragment (primers JMP1F/JMP1R) of secretion vector JMP62UraTFE under the control of TEF promoter and the pre-sequence of lipase 2 of Y. lipolytica by In-Fusion Cloning (Clontech). XYN II gene (GenBank accession code: XM_006968885.1), encoding T. reesei xylanase II, was amplified from the plasmid JMP63UraXYNII (kindly provided by Dr. Cédric Montanier, INSA-Toulouse) by PCR using primers F (2) and R (2) and fused with secretion vector JMP62UraTFE as described above. The SWO1 gene (GenBank accession code: A245918.1), encoding T. reesei swollenin, was synthesized by GenScript (USA) after codon optimization based on the codon bias of Y. lipolytica (Additional file 2), and cloned into the plasmid JMP62UraTFE.

For the expression of His-tagged proteins, XYNIH, LPMOA, and SWO1 were cloned by PCR with F3 as forward primer and R (3–5) as reverse primers using the expression vectors constructed in last step as template, and fused with PCR fragment (primers JMP2F/JMP2R) of the vector JMP62UraTB1his [30]. This enables the addition of a sequence encoding 6-histidine at the C-terminus of the proteins.

To improve the expression level of EG I, the plasmid JMP62UraEG1 was digested using BamHI/AvrII, and the EG I gene was recovered and then inserted into the corresponding sites of the previously constructed plasmid JMP62UraTFE under the control of TEF promoter with an enhancer comprising 4 tandem copies of upstream activation sequences (4UASs) [30].

After construction, all expression vectors were verified by DNA sequencing (GATC Biotech, Konstanz, Germany). For Y. lipolytica transformation, vectors were digested using NotI, thus generating a linear DNA with Zeta sequences at both extremities. Then the gel purified expression cassettes were introduced into the Zeta docking platform of Y. lipolytica JMY1212 Zeta for the expression of single recombinant protein, or randomly into the genome of ΔpoxB12 strain, for co-expression of multiple cellulases, using the lithium acetate method [52]. For the latter case, the LoxP-Cre recombination system was used for marker rescue and to ensure the multistep

Table 3 Plasmids used or created in the present study

| Plasmids          | Description                  | Source of reference |
|-------------------|------------------------------|---------------------|
| JMP62UraTFE       | URA3, pTEF                   | [52]                |
| JMP62LeuTF        | LEU2, pTEF                   | [52]                |
| JMP62UraTB1his    | URA3, pTEF-BGL1-His6         | [57]                |
| PUB4-CRE          | hph, hpAd-CRE                | [53]                |
| JMP62UraHETF      | URA3, pHETF                  | [30]                |
| JMP62LeuHETF      | LEU2, pHETF                  | This investigation  |
| JMP62hphHETF      | Hph, pHETF                   | This investigation  |
| JMP62LeuHETF-EG1  | LEU2, pHETF-EG1              | This investigation  |
| JMP62UraXYNIIhis  | URA3, pTEF-XYNII-His6        | This investigation  |
| JMP62UraLPMOAhis  | URA3, pTEF-LPMOA-His6        | This investigation  |
| JMP62UraSWO1his   | URA3, pTEF-SWO1-His6         | This investigation  |
| JMP62UraXYNII     | URA3, pTEF-XYNII             | This investigation  |
| JMP62LeuXYNII     | LEU2, pTEF-XYNII             | This investigation  |
| JMP62hphXYNII     | hph, pTEF-XYNII              | This investigation  |
| JMP62UraLPMOA     | URA3, pTEF-LPMOA             | This investigation  |
| JMP62LeuLPMOA     | LEU2, pTEF-LPMOA             | This investigation  |
| JMP62hphLPMOA     | hph, pTEF-LPMOA              | This investigation  |

Table 4 Sequences of the oligonucleotide primers used in this study

| Primer names | Sequence (5′–3′), 15-bp homologous sequence for fusion is underlined |
|-------------|---------------------------------------------------------------------|
| F1          | GTCCTCCAGAACAGGCAGCAGATGGCATATTAGTCCATGTCATCAACG                     |
| R1          | TACACAGACACCCTAGGGCTATAGTTAAGGCCAGCTGGGTAGTGGG                       |
| F2          | GTCCTCCAGAACAGGCAGCAGATGGCATATTAGTCCATGTCATCAACG                     |
| R2          | TACACAGACACCCTAGGGCTATAGTTAAGGCCAGCTGGGTAGTGGG                       |
| JMP1F       | CTTAGGGGTCTCTGGGTATCACTAAGCTATT                                      |
| JMP1R       | TCCTCTTCTGGAGAAGACTCGG                                              |
| F3          | ACCACCCGAGAGTCCCAATGAAAGCTTCCACATCCC                                |
| R3          | ATGGTAATGATGGTGGTGGCTACCAGGTAGAGGAGGAG                               |
| F4          | ATGGTAATGATGGTGGTGGCTACCAGGTAGAGGAGGAG                               |
| R5          | ATGGTAATGATGGTGGTGGCTACCAGGTAGAGGAGGAG                               |
| JMP2F       | CACCATCATCACCATCATATTAAAAT                                          |
| JMP2R       | GGCTCTTCTGGAGAAGACTCGG                                              |
| HphF        | CCCACACATCATCCACATGAAAGGCTTCCACATCCC                                |
| HphR        | TAGACGGGACAGGGCTTATCTTCTTGGCTTGAGGAGGAG                               |
| LHF         | GGGCTCTTCTGGAGAAGACTCGG                                              |
| LHFR        | TCGCTCTTCTGGAGAAGACTCGG                                              |
| XLF         | GCCTCTTCTGGAGAAGACTCGG                                              |
| XLR         | GCCTCTTCTGGAGAAGACTCGG                                              |

This report was supported by a grant awarded to Dr. Guo (Guo T. et al. Biotechnol Biofuels (2017) 10:298) by the German Research Foundation (DFG) (GRK 1701:3/1-3) and the European Commission, FP7 Project (FP7/2007-2013) (project no. 249023).
insertion of the target genes [53]. However, after the integration of six cellulase genes (BGL1, BGL2, EGI, EGH, CBHI, and CBHII), the removal of the selection markers (URA3 and LEU2) from cellulolytic Y. lipolytica caused loss of the previously introduced genes, probably due to the presence of multiple LoxP sites. In order to further introduce accessory proteins into cellulolytic Y. lipolytica, the hygromycin phosphotransferase coding gene hph, conferring hygromycin resistance to Y. lipolytica, was amplified from the vector pUB4-Cre by PCR using primers HphF and HphR. Then, the hph gene was fused with DNA fragment amplified from the vectors JMP62Leu-TEFLPMOA, JMP62LeuTEFXYNII, and JMP62LeuTEFSWO1 using primers LHFF and LHFR, respectively. The plasmid for co-expression of TrLPMOA and TrXYNII was constructed by insertion of the expressing cassette containing TrLPMOA under the control of TEF promoter, obtained by PCR amplification from the vector JMP62UraTEFLPMOA using primers XLF and XLR, into the XbaI site of the vector JMP62hphTEFXYNII. The successful multiple integration of the heterologous genes into the genome of Y. lipolytica was verified by PCR using gene-specific primers. In addition, transformants expressing multiple enzymes were tested for growth on cellobiose, and for degradation of Azo-CM-Cellulose and AZCL-arabinoxylan. Clones displaying both activities were retained for further analysis. Table 3 summarizes the expressed cellulase genes and their corresponding Y. lipolytica transformants.

Protein production and hydrolytic activity assay
Recombinant protein production by Y. lipolytica was carried out in liquid YTD medium in shake flask at 28 °C and 120 rpm for 5 days. PASC was prepared from Avicel PH-101 as previously described [54]. The overall cellulolytic activities were measured on CMC, PASC, CIMV-cellulose, and Avicel PH-101 using a previously described method with slight modifications [55]. Briefly, the reaction mixture contained 1% (w/v) cellulosic substrate, 50 mM citrate buffer (pH 4.8), and proper volume of diluted enzyme solution.

Xylanase activity was measured on 2% (w/v) beechwood xylan (Megazyme) in 50 mM citrate buffer (pH 6.0). The reaction was conducted at 50 °C for 10 min. The reducing sugars were quantified using the dinitro salicylic acid (DNS) reagent [30]. One unit of activity (U) was defined as the amount of enzyme or culture supernatant required to release 1 μmol of reducing sugars per min for xylanase activity, or 1 mg of reducing sugars per min for cellulosytic activity.

In addition, glucose and xylose were measured using an Aminex HPX87-H column (Bio-Rad Laboratories, Germany), operating at 50 °C using a mobile phase (5 mM H2SO4) flowing at a rate of 0.5 mL/min. Glucose and xylose were detected using a Shodex RI-101 refractive index detector (Showa Denko, New York, NY, USA).

All protein concentrations were measured using the Bradford method and bovine serum albumin as a standard [56]. All enzymatic activity measurements were performed in triplicate unless otherwise stated.

SDS-PAGE and Western blot analysis
SDS-PAGE was conducted using Mini-PROTEAN TGX Stain-Free precast gels (Biorad) according to the manufacturer’s instructions. Fifteen microlitre of culture supernatant or enzyme solution was loaded into each well. Western blotting of proteins was performed as described previously [57]. Crude supernatant of Y. lipolytica JMY1212 expressing LPMOA and SWO1 fused with the His6 tag were concentrated tenfold by ultrafiltration on an Omega™ membrane disc filter with a 10 kDa cut off (Pall, France). Blots were sequentially treated with mouse non-position-specific His-Tag antibody 1:2500 (THE™ from Genscript, Piscataway, NJ, US) and the alkaline phosphatase-conjugated goat anti-mouse IgG. Phosphatase activity was detected by NBT/BCIP as substrate.

Purification and deglycosylation of recombinant proteins
Yarrowia lipolytica JMY1212 expressing His6-tagged XYNII, LPMOA and SWO1, respectively, was grown in 200 mL YTD medium at 120 rpm, 28 °C for 48 h. After centrifugation (8000×g for 5 min at 4 °C), the supernatant was concentrated tenfold by ultrafiltration with an Omega™ membrane disc filter at 10 kDa cut off (Pall, France), and applied to 2 mL of TALON Metal Affinity Resin (Clontech, Takara-Bio, Kyoto, Japan). Subsequently, protein was eluted using buffer containing imidazole according to the manufacturer’s instructions. Deglycosylation was carried out by treating the purified proteins with endoglycosidase H (New England Biolabs, Beverly, MA, USA) to remove N-linked carbohydrates at 37 °C for 1 h. Protein samples were analyzed by SDS-PAGE and visualized with colloidal Coomassie blue staining.

Amplex Red/horseradish peroxidase assay
The oxygen reactivity of TrLPMOA was measured in a quantitative time resolved assay of H2O2 using Amplex Red and horseradish peroxidase as described previously [8, 22]. The reaction mixture (100 μL) contained 100 mM sodium acetate buffer (pH 6.0), 50 μM Amplex Red reagent (Invitrogen, France), 7.14 U/mL horseradish peroxidase, 0.1–0.5 μM LPMOA and 30 μM ascorbate as reductant, and was assembled in the well of a 96-well microtiter plate and incubated at 30 °C. The fluorescence (excitation and emission wavelengths of 560 and 595 nm, respectively) was measured using a Tecan
Enzymatic hydrolysis of cellulose feedstocks

Enzymatic hydrolysis of different cellulose feedstocks was carried out in 50 mM citrate buffer (pH 4.8). CIMV-cellulose, Avicel and wheat straw, respectively, was added into the reaction to yield a final cellulose concentration of 2.0% (w/v). Celfic® CTec2 (a gift from Novozyme) or concentrated secretome of Y. lipolytica YLC11 was added at 10.0 FPU/g-cellulose. The reaction mixture was incubated at 50 °C under shaking for 120 h. Samples were taken at regular intervals to determine the concentration of glucose using HPLC.

Growth of yeast expressing multiple cellulases on cellulose feedstocks

Yeast growth on different cellulose feedstock was performed in 50 mL defined medium containing CIMV-cellulose, Avicel cellulose or wheat straw in 250-mL Erlemeyer flasks. Yeasts were pre-cultivated in defined media on glucose until mid-exponential phase and the cells were collected by centrifugation. After washing with deionized water, the cells were used to inoculate the defined medium containing cellulose to yield an initial biomass concentration of 1.0 g-DCW/L. In addition, CIMV-cellulose, Avicel cellulose, and wheat straw were incubated with swollenin at 50 °C under shaking for 24 h prior to the inoculation of strain YLC11. All the cultivations were conducted at 28 °C and under shaking at 120 rpm. Samples were taken at the end of 5 days to determine concentrations of biomass and residual cellulose (see below). The data are presented as mean values of at least three biological replicates ± standard deviation.

Analysis of cellulose residues and determination of dry cell weight

The quantification of residual cellulose and dry cell matter was conducted as previously described with slight modifications [30]. Briefly, the cell pellets mixed with cellulose from the above cultures were harvested by centrifugation at 8000 × g for 10 min at 4 °C. After two washes with distilled water, the collected cellulose-cell pellet was freeze-dried and weighed. The amount of remaining cellulose and hemicellulose was calculated from the total glucose and xylose released from diluted acid hydrolysis of the residues with 2.5% sulfuric acid at 121 °C for 1 h. Dry cell weight was deduced by subtracting the amount of cellulose or lignocellulose from the weight of cellulose or lignocellulose-cell pellet. Glucose and xylose were measured by HPLC as described before [57]. The biomass yield was calculated as the ratio of the amount of biomass obtained divided by the amount of carbon source consumed.

Infinite M200Pro plate reader (Tecan, Männedorf, Switzerland). The activity of TrLPMAO was derived from the data using a standard H2O2 calibration curve. The inhibition of H2O2 production was used to test the activity of TrLPMAO on various β-glucan and xylan substrates as described previously [41]. A final concentration of 1.0% (w/v) PASC, CMC, Avicel PH-101, barley glucan (β-1,3; 1,4), and xylan was added into the above reaction mixture. All measurements were performed in triplicates.

**Cello-oligosaccharide cleavage assay**

The cleavage of cello-oligosaccharides by LPMOA was assayed by incubating 16 mg of TrLPMAO, 1 mM of ascorbate, and 5 mM cello-oligosaccharides (DP4-DP6) in 50 mM sodium acetate buffer (pH 4.8) at 50 °C for 24 h under shaking (1000 rpm). To stop the reaction, samples were boiled (100 °C for 10 min) and then cooled before adding an aliquot of Novozyme 188 β-glucosidase (gift from Novozyme, Denmark) (12.0 IU/g-cello-oligosaccharides, 810 IU/mL). After that, the reaction was incubated for a further 12 h and then, the oxidized mono-saccharides in the supernatant were measured using a d-gluconic acid/d-glucono-δ-Lactone assay kit (Megazyme) according to manufacturer’s instructions. All assays were carried out in triplicate.

**Study of the synergy of LPMOA and SWO1 on enzymatic hydrolysis of Avicel**

Enzymatic hydrolysis of cellulose was carried out in 50 mM citrate buffer (pH 4.8) containing 2.0% (w/v) microcrystalline cellulose (Avicel PH-101, Sigma). Cellic® CTec2 (a gift from Novozyme) or concentrated secretome of Y. lipolytica YLC11 was added at 5.0 FPU/g-cellulose and 12.0 IU/g-cellulose, respectively. The reaction (1 mL) was conducted in Eppendorf tubes (2 mL) using a thermomixer at 50 °C and 1000 rpm. To study the synergistic effect of TrLPMAO on the enzymatic hydrolysis of cellulose, different amounts of this protein (5, 10, and 15 mg-protein/g-cellulose, respectively) were added to the reaction mixtures containing Avicel (2.0% w/v). In addition, ascorbate (1 mM) was added into the reactions as the reducing agent to reactivate LPMOA. To study the synergistic effect of swollenin on the enzymatic hydrolysis of cellulose, this protein was either added into reaction mixture together with cellulases at different concentrations (5, 10, and 15 mg-swollenin/g-cellulose respectively) or, incubated with the Avicel at 50 °C under shaking for 24 h prior to the addition of cellulases. Control experiments were conducted under the same conditions, with or without, BSA (at 5, 10, and 15 mg-protein/g-cellulose). Samples were taken at regular intervals to determine the concentration of reducing sugars using DNS.
Additional files

**Additional file 1: Figure S1.** PCR verification of *Y. lipolytica* transformants expressing multiple cellulases and accessory proteins (A) YLC7, Lane 1 to 6: BGL1, BGL2, 4UASTrEGI, TEGII, 4UASNCBH1, 4UASTrCBHII, (B) YLC8, Lane 1 to 6: BGL1, BGL2, 4UASTrEGI, TEGII, 4UASNCBH1, 4UASTrCBHII, YC10, Lane 1 to 7: BGL1, BGL2, 4UASTrEGI, TEGII, 4UASNCBH1, 4UASTrCBHII, TelFMOA, (C) YLC10, Lane 1 to 7: BGL1, BGL2, 4UASTrEGI, TEGII, 4UASNCBH1, 4UASTrCBHII, TelFMOA, (D) YLC10, Lane 1 to 7: BGL1, BGL2, 4UASTrEGI, TEGII, 4UASNCBH1, 4UASTrCBHII, TelFMOA. **Figure S2.** Western blot analysis of the heterologous rhTrEGI protein secreted by the engineered *Y. lipolytica* strains: lane 1, Endo H-treated secretome of YLC6b (20 μL), lane 2, Endo H-treated secretome of YLC7b (20 μL). **Figure S3.** Characterization of the recombinant XYNI expressed in *Y. lipolytica*. (a) Effect of pH on the activity of rhXYNI. (b) Effect of temperature on the activity of rhXYNI.

**Figure S4.** Screening of *Y. lipolytica* expressing cellulases and accessory enzymes on YNB indicator plate containing supplemented with 0.2% w/vAv-CM Cellulose. Lane 1, Y. lipolytica control; Lane 2 to 4, YLC8, YLC9 and YLC10. **Figure S5.** The growth of *Y. lipolytica* in defined medium containing 10 g/L gluconic acid or 10 g/L glucose. **Table S1.** The sequences of the oligonucleotide primers used for PCR verification of *Y. lipolytica*-transformants. **Table S2.** Comparison of cellulase utilization and biomass yield of cellulolytic *Y. lipolytica* grown on different cellulogenic substrates for 120 h in aerobic cultivation without the addition of acetic acid.

**Additional file 2.** Nucleotide sequence of codon-optimized SWO1.

**Abbreviations**

BCP: 5-bromo-4-chloro-3-indolyl phosphate; BGL: β-glucosidase; CBH: cellobiohydrolase; CBP: consolidated bioprocessing; CMC: carboxymethyl cellulose; DCW: dry cell weight; EG: endoglucanase; Endo H: endoglucosidase; LCB: lignocellulosic biomass; LPMO: lytic polysaccharide monooxygenase; MW: molecular weight; NBT: nitroblue tetrazolium; Nc: Neurospora crassa; PAGE: polyacrylamide gel electrophoresis; PASC: phosphoric acid swollen cellulose; r: recombinant proteins; rh: His6-tagged recombinant proteins; HPACE: high performance anion exchange chromatography; SWO: swollenin; Tr: Trichoderma reesei; yl: Y. lipolytica.

**Authors’ contributions**

ZPG, SB, SD, AM, and MIOD conceived the study and participated in its design. ZPG designed the constructs, carried out all the experiments, and drafted the manuscript. SB, SD, JMN, AM, and MIOD revised the manuscript. All authors read and approved the final manuscript.

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**Acknowledgements**

The authors would like to express their gratitude to Nelly Monties for her help with chromatographic analyses. We thank Dr. Cédric Montanier for kindly providing us with CIMV-cellulose. We thank the ICEO facility dedicated to enzyme screening and discovery, and part of the Integrated Screening Platform of Toulouse (PICT, IBiSA) for providing access to its screening, protein purification and analytical facilities.

**Competing interests**

The authors declare that they have no competing interests.

**Availability of supporting data**

All data generated or analyzed in this present study are included in this published article and a supporting material “Additional file 1.”

**Consent for publication**

All authors consent for publication.
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