Downscale fermentation for xylooligosaccharides production by recombinant Bacillus subtilis 3610

Cláudia Amorim¹, Sara C. Silvério¹, Raquel F.S. Gonçalves¹, Ana C. Pinheiro¹,², Soraia Silva³, Elisabete Coelho³, Manuel A. Coimbra³, Kristala L.J. Prather⁴, Lígia R. Rodrigues³,⁴,⁎

¹ CEB-Centre of Biological Engineering, Universidade do Minho, Campus de Gualtar, 4710-057 Braga, Portugal
² Instituto de Biologia Experimental e Tecnológica, Avenida da República, Quinta do Marquês, Estação Agronómica Nacional, Apartado 12, 2781-901 Oeiras, Portugal
³ QOPNA, Departamento de Química, Universidade de Aveiro, 3810-193 Aveiro, Portugal
⁴ Department of Chemical Engineering, Massachusetts Institute of Technology, 77 Massachusetts Ave, Cambridge, MA 02139, United States

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Abstract

The global demand of prebiotics such as xylooligosaccharides (XOS) has been growing over the years, motivating the search for different production processes with increased efficiency. In this study, a cloned Bacillus subtilis 3610, containing the xylanase gene xyn2 of Trichoderma reesei coupled with an endogenous secretion tag, was selected for XOS production through direct fermentation of beechwood xylan. A mixture of XOS with a degree of polymerization ranging from 4 to 6 was obtained, presenting high stability after a static in vitro digestion (98.5 ± 0.2%). The maximum production yield expressed as total XOS per amount of xylan (306 ± 4 mg/g) was achieved after 8 h of fermentation operating under one-time impulse fed-batch. The optimal conditions found were pH 6.0 and 42.5 °C, using 2.5 g/L of initial concentration of xylan increased up to 5.0 g/L at 3 h. Xylopentaose was the major oligosaccharide produced, representing 47% of the total production yield.

1. Introduction

In the light of current health consciousness, the demand of prebiotic food ingredients has been growing, as consumers prefer natural therapeutic agents to prevent widespread diseases, such as from the cardiovascular system, gastrointestinal tract or oncology (Kaprelyants, Zhurlova, Shpyrko, & Pozhitkova, 2017). GVR - Grand View Research Inc. (2016) estimates that global prebiotic market will worth 7.11 billion $ by 2024. According to the updated definition, a prebiotic can be defined as “a substance that is selectively utilized by host microorganisms conferring a health benefit” (Gibson et al., 2017). Xylooligosaccharides (XOS) are emerging prebiotics, presenting a remarkable potential as food ingredients due to their minimal recommended daily intake (1.4–2.8 g) (Finegold et al., 2014), heat and pH stability (Courtin, Swennen, Verjans, & Delcour, 2009), organoleptic properties and multidimensional effects on human health and livestock (Aachary, Gobinath, Srinivasan, & Prapulla, 2015). XOS were reported as able to stimulate the growth of bifidobacteria (Hsu, Liao, Chung, Hsieh, & Chan, 2004) and to produce lactate in a higher extent than other oligosaccharides (Rycroft, Jones, Gibson, & Rastall, 2001). In addition, these compounds are the only nutraceuticals that can be sourced from lignocellulosic biomass, such as agro-residues, which are an abundant and renewable resource (Cruz et al., 2018).

Chemically, XOS are oligosaccharides composed by 2–10 residues of (β1→4)-linked xylene. The main chain of xylose can comprise several substituents, namely acetyl groups, glucuronic acid, methylated glucuronic acid, arabinose, and hexose residues, depending on the biomass origin and the production process (Coelho, Rocha, Moreira, Domingues, & Coimbra, 2016). XOS are produced though the hydrolysis of xylan, which is the main constituent of hemicelluloses and the second most abundant carbohydrate in the lignocellulosic biomass (Bian et al., 2013). Particularly, they can be obtained by different hydrolytic methods including chemical, auto- hydrolysis, enzymatic hydrolysis, or a combination thereof (Carvalho, Neto, Silva, & Pastore, 2013). Chemical and auto-hydrolytic methods may originate undesired by-products, including toxic compounds such as hydroxymethylfurfural and furfural (Bian et al., 2014).

Additionally, the use of chemicals together with extreme operational conditions require more robust and expensive equipment. Thus, the enzymatic process is generally preferred for XOS used in food (Kaprelyants et al., 2017), presenting higher efficiency and specificity towards the product, allowing the control of the degree of polymerization (DP) and to reduce the costs associated with the downstream process (Bian et al., 2014). Moreover, this process represents a...
more environment-friendly approach, since it operates at milder conditions not requiring the use of noxious chemicals (Azeelee et al., 2016). However, xylan is generally comprised in a xylan-lignin matrix structure present in the lignocellulosic biomass (Samanta et al., 2012). Thus, XOS are mainly produced by a two-step combination of different methods, including a first step of fractionation of the lignocellulosic material to obtain soluble xylan and then, its hydrolysis by xylanolytic enzymes (Carvalho et al., 2013). The low yields associated to the first step of extraction in addition to the cost of producing or purchasing commercially available xylanases, may compromise the economic viability of the production process (Reddy & Krishnan, 2016a). Amorim et al. (2018) addressed this problem by showing the potential of Bacillus subtilis to produce XOS in a single-step fermentation for one specific residue application, namely using brewer spent grain (BSG) as raw material. In this work, it is proposed the study of XOS production by direct fermentation using a commercial xylan in order to have a more detailed insight on the process and microorganism metabolism, particularly in terms of biomass growth evaluation, which is not possible when a complex residue is used as raw material. This comprehensive study may allow designing different one-step strategies to overcome the main issues associated with XOS production from agro-residues, specifically mass transfer and inhibition by the substrate. Moreover, this study reports for the first time the downscale process applied in the context of XOS production by direct fermentation of beechwood xylan, a hardwood tree that generates highly abundant forest residues. This study is particularly focused on (a) finding the B. subtilis clone with the highest potential to produce XOS; (b) proving that the best fermentation time corresponds to the time at which the maximum amount of oligosaccharides is reached (with minimal concentration of monosaccharides); (c) determining the profile curves associated to xylan concentration, temperature, and pH, which are three important process variables, as well as determining the biomass profile; (d) testing batch and fed-batch modes of operation to produce XOS. The main challenge of this work is to test the hypothesis of direct fermentation of hardwood xylans using genetically modified microorganisms as an efficient alternative for XOS production.

2. Experimental

2.1. Materials

The chemical composition of the beechwood xylan used in this study was provided by the manufacturer (CAS: 9014-63-5, Lot 141202, Megazyme, Bray, Ireland), containing xyllose (81.3%), glucuronic acid (13.0%), and other sugars (5.7%), namely glucose, arabinose, galactose, rhamnose, and mannose. All chemicals, media and media components were of analytical grade obtained from Sigma-Aldrich Chemical Ltd., unless specified otherwise.

2.2. Bacterial strains, pre-inoculum, and fermentation media

Bacillus subtilis 3610 wild type (wt) and two clones, previously described (Amorim et al., 2018), were used: clone 1 containing the xylanase gene xyn2 (X69574.1 Genbank NCBI) from Trichoderma reesei and clone 2 containing a secretion tag endogenous to B. subtilis (> tr[A0A052II12] Uniprot) coupled to xylanase N-terminus site. To prepare the pre-inoculum, one colony of B. subtilis grown on LB agar plate overnight at 37 °C was picked into 2 mL of LB medium (Difco, New Jersey, USA) and spectinomycin (100 μg/mL) was added when required for clones selection. The cells were then cultivated at 37 °C and 250 rpm during approximately 2 h until reaching OD_{600} ~ 1.0. This starter culture was then diluted to OD_{600} ~ 0.020 into the fermentation media. Fermentation medium consisted in a mixture of commercial beechwood xylan (Megazyme) in 2% (v/v) of Vogel medium (Vogel, 1956), autoclaved at 121 °C during 15 min.

2.3. Downscale fermentation of xylan by B. subtilis: optimization of process variables and operation mode

The Biolector apparatus (m2p-labs, USA) was used for high-throughput fermentations in flower plates (MTP-48-B, m2p-labs, USA). For all fermentations, each plate well was filled up to a total of 1 mL of xylan solution in 2% (v/v) of Vogel medium and inoculated as described in Section 2.2. The cells were cultivated during 24 h and samples were collected over time, centrifuged (3000 rpm during 3 min) and further analyzed (Section 2.7).

The performance of B. subtilis wt, clone 1 and clone 2 regarding the production of XOS were compared in batch conditions, using 10 g/L of commercial beechwood xylan at pH 7.0, 45 °C and 950 rpm. The selected transformant was used for further optimization studies.

The individual effects of substrate concentration (1.25, 2.5, 5.0, 10, 15 and 20 g/L), pH (5.0, 6.0, 6.5, 7.0 and 8.0) and temperature (37.0, 40.0, 42.5, 45.0 and 47.0 °C) were optimized in sequence by selecting the most favorable value of one parameter at each time.

The fed-batch operation mode by one-time impulse was also studied in order to test the hypothesis of substrate inhibition and also to further optimize the process. Two initial concentrations of xylan were tested (2.5 and 5.0 g/L) which were increased up to 5.0 and 7.5 g/L, respectively, by one impulse at different times (3, 5, 7 and 9 h).

All fermentation assays were performed in duplicate and samples from the supernatant were collected, centrifuged and further analyzed in triplicate.

2.4. Quantification of biomass growth

The biomass growth obtained under optimal conditions in batch and fed-batch modes was accessed by cell counting using a Neubauer improved chamber. The samples were diluted in Vogel 2% (v/v) medium and observed under an inverted microscope (Eclipse TS100, Nikon) coupled with a SPOT Insight fire camera that allowed to capture the cells using a total magnification of 400 x . The cells pictures were processed in Spot software 5.0 setting manual exposure at 25 ms, 8 gain and 2 x 2 bin.

2.5. Sugar analysis and enzymatic activity

For all the fermentation assays, samples from the supernatant were collected, centrifuged and further analyzed. The DNS (3,5-dinitrosalicylic acid) method (Miller, 1959) was used as a screening qualitative technique to assess the total reducing sugars, RS (mg). The sugar production yield, Y_{RS} (mg/g), was defined as the ratio between RS (mg) and the mass of xylan (g) used in the fermentation.

The enzymatic activity of endo-(β1→4)-xylanase was accessed using a soluble chromogenic substrate according to the supplier guidelines for azo-xylan from birchwood (S-AXBP, Megazyme, Bray, Ireland).

2.6. XOS quantification

The XOS produced under optimum conditions were directly quantified by HPLC (Section 2.9), after previous purification of the fermentative broth with activated charcoal (Section 2.8). During the HPLC analysis the produced XOS were separated and individually collected according to their DP for chemical characterization analysis (Section 2.9). The DP of the XOS was estimated by TLC, as described by Amorim, Silvério, and Rodrigues, (2019), and confirmed by HPLC (Section 2.9) and GC–MS (Section 2.10). The XOS yield, Y_{XOS} (mg/g) was defined as the ratio between the total concentration of XOS (mg) and the mass of xylan (g) used in the fermentation.

2.7. Mass balance in xylose equivalents

For accurate comparison purposes, the process mass balance was...
performed in terms of xylose equivalents, Xeq (mg), per gram of xylan used in the fermentation.

The composition of XOS, YXOS, Xeq (mg/g), and the initial amount of xylan, Yxylan (mg/g), in Xeq, was determined by HPLC (Section 2.9) after acid hydrolysis according to (National Renewable Energy Laboratory) protocols (NREL/TP-510-42618-42622-4218).

The microorganism uptake of xylose, Yxup (mg/g), was quantified by the difference between the Yxylan and the yield in xylose equivalents of the supernatant obtained after the fermentation, YXS. The free xylose yield, Yxyl (mg/g) was defined as the ratio between the concentration of free xylose (mg) (Section 2.9) and the mass of xylan (g).

The residual xylan, YXR (mg/g), was determined as follow: YXR (mg/g) = Yxylan – (YXS + Yxup + Yxyl).

2.8. Partial purification of XOS produced by direct fermentation

To minimize the salt interference in the HPLC analysis, XOS produced by fermentation of xylan under optimal conditions were treated with activated charcoal as described in Amorim et al. (2019). The supernatant was collected (20 mL), centrifuged and loaded onto 13 g of activated charcoal. After the adsorption step, the charcoal was washed with MilliQ water, followed by sugar desorption using a mixture of butanol, acetic acid and water (2:1:1 v/v/v) as eluent. The collected eluent with sugars was evaporated to dryness and the samples were resuspended in water for further analysis (Sections 2.9 and 2.10).

2.9. Sugars analysis by HPLC

Two distinct HPLC methods were used, namely to quantify monosaccharides or oligosaccharides partially purified (Section 2.8). For monosaccharides quantification, an HPLC (Agilent Technologies, USA) fitted with RI and DAD detectors (Agilent Technologies, USA) and an Aminex HX 87H column (300 mm × 7.8; Biorad, USA) were used for quantification. A volume of 10 µL of sugar sample was eluted using 5 mM H2SO4 as the mobile phase at a flow rate of 0.7 mL/min and a temperature of 60 °C. Furfural and HMF concentrations were also determined by HPLC, using the DAD detector. For the oligosaccharides quantification, an HPLC (Jasco, Japan) fitted with RI detector (K-2300 Knauer, Germany) and a Prevail Carbohydrate column (250 mm × 4.6 mm; Grace, USA) were used. A mixture of acetonitrile and water 68:32 (v/v) was used as mobile phase at a flow rate of 0.9 mL/min, temperature of 30 °C and 20 µL injection volume. Pure XOS (X2-X6) were used as standards for the identification of the produced oligosaccharides DP.

2.10. XOS characterization

Partially methylated alditol acetates were obtained from XOS samples using CH3I as methylating reagent (Coelho et al., 2016) and analyzed by GC-MS, as described by Amorim et al. (2018). XOS DP was calculated as the relative amount of total xylose divided by the amount of terminally linked xylose, as xylose (Xyl) should not occur as branching residues. The ratio between the branching points in substituted Xyl residues (Xylsubt) and the total amount of Xyl (Xyltotal): [Xylsubt/Xyltotal] was used to calculate the branching degree, where Xylsubt is the sum of the amount of monosubstituted residues (2,4-Xyl + 3,4-Xyl) + twice the amount of dissubstituted residues (2,3,4-Xyl).

2.11. In vitro gastrointestinal digestion

The supernatant containing the mixture of XOS produced under optimal conditions was lyophilized and used to test its stability under a harmonized static in vitro digestion method as described by Minekus et al. (2014). Briefly, 5 mL of XOS samples were prepared and subsequently exposed to conditions simulating the mouth, stomach and small intestine. At the oral phase, the sample was mixed with simulated salivary fluid solution (SSF), CaCl2(H2O)2 (to obtain 1.5 mmol/L in the fluid) and purified water (at volume necessary to achieve 1x of concentration of SSF). The mixture was incubated at 37 °C in a shaking bath (B. BRAUN BIOTECH model CERTOMAT WR, Melsungen, Germany) under agitation at 120 rpm during 2 min. The gastric phase consisted in the addition of simulated gastric fluid (SGF), porcine pepsin (2000 U/mL in the final mixture) and CaCl2(H2O)2 (to obtain 0.15 mmol/L in the fluid). The mixture was adjusted to pH 3.0 through the addition of HCl 1 M. Purified water was added in order to achieve 1x concentration of SGF. The final mixture was incubated over 2 h at 37 °C and under agitation at 120 rpm. The intestinal phase was simulated by the addition of simulated intestinal fluid (SIF), porcine pancreatin (100 U/mL in the final mixture), bile solution (10 mmol/L in the final mixture) and CaCl2(H2O)2 (to obtain 0.6 mmol/L in the fluid). The mixture pH was adjusted to 7.0 by adding the necessary volume of NaOH 1 M or HCl 1 M. Purified water was added in order to achieve 1x concentration of SIF. The final mixture was incubated during 2 h at 37 °C under agitation at 120 rpm. Samples were collected after the in vitro digestion and the reaction was stopped by adding the enzyme inhibitor pefabloc (1 mmol/L) (10 µL per 1 mL of sample). All samples were tested at least in triplicate.

3. Results and discussion

3.1. Optimization of the production of XOS by direct fermentation of xylan using the B. subtilis

3.1.1. Evaluation of the transformants performance for the production of XOS

The performance of the B. subtilis wild type (wt) as XOS producer by direct fermentation of xylan was compared with the clone 1 and clone 2 cultured in microbioreactor, using 10 g/L of xylan at pH 6.0, 45 °C and 950 rpm (Fig. 1). These operational conditions were defined based on previous literature reports (Amorim et al., 2018; Helianti et al., 2016; Irfan, Ashgar, Nadeem, Nelofer, & Syed, 2016).

All tested strains achieved the maximum value of YRS, YXOSmax, after 8 h of fermentation. Nonetheless, clone 2 (127 ± 8 mg/g) presented higher YXOSmax values when compared to the wild type (93 ± 7 mg/g), showing a statistically significant increase of 37% (t-student test, α = 0.05). Moreover, all the production curves obtained present similar profiles, which may indicate that the expression of the heterologous
xylanase did not impose a metabolic burden to the cells. Although being recognized as a high protein-secreting microorganism (Van Zyl, den Haan, & la Grange, 2013), B. subtilis wt and clone 1 achieved similar YRSmax and enzymatic activity values, suggesting that the secretion tag present in clone 2 possibly plays a significant role in the secretion of the heterologous xylanase, increasing its performance in terms of sugars production from xylan.

The free xylose yield, Yxyl, was determined by HPLC (Fig. 1) for clone 2. Interestingly, when YRSmax was achieved (8h), only low amounts of free xylose were present in the liquid media (Yxyl = 3.5 ± 0.6 mg/g), suggesting that the YRSmax value is mainly represented by oligosaccharides. Thus, it seems that there is an optimal time around 8h, tmax, which corresponds to the time phase at which the oligosaccharides are accumulated, before starting to be degraded into xylose (maximum between 10 and 14h) (Fig. 1). The TLC analysis of the fermentation supernatant after 8h of fermentation shows defined bands in the region of 4-, 5- and 6-DP XOS and a vestigial band is visible in the xylose region (Appendix A, Fig. A1 in Supplementary material).

The improvement of the clone 2 performance was also confirmed by xylanase activity analysis, presenting an increase of approximately 50% in the maximal activity at 12h when compared to the wild type (data not shown). Based on these results, B. subtilis clone 2 was selected for further optimization studies (Section 3.1.2).

### 3.1.2. Optimization of the initial pH, xylan concentration and temperature for XOS production using the B. subtilis clone 2

Extracellular pH, substrate concentration and temperature are process variables affecting significantly the growth of bacteria and enzyme production. These variables were separately studied in order to evaluate their individual effect on the XOS production by direct fermentation of xylan using clone 2 (Fig. 2).

The effect of the initial pH on the YRS was studied at 45°C, 10.0 g/L of xylan and 950 rpm, using different pH, from pH 5.0 up to 8.0 (Fig. 2A). The highest YRSmax (128 ± 8 mg/g) was achieved after 8h of fermentation at pH 6.0, when as expected, the Yxyl was maintained at low levels (3.3 ± 0.8) mg/g. In this sense, pH 6.0 was selected for further studies, corresponding to the pH previously used to compare the transformants performance (Section 3.1.1).

The optimal pH for this process was possibly influenced by several phenomena, i.e. it was established as a function of the pH conditions more favorable for Bacillus growth, xylanase production and xylan solubility. In particular, B. subtilis growth in minimal medium is usually performed around pH 7.0 (Jaacks, Healy, Losick, & Grossman, 1989), while the optimal pH for B. subtilis xylanase activity has been reported around pH 5.0–5.5 (Irfan et al., 2016; Milessi et al., 2015; Reddy & Krishnan, 2016b), as well as for T. reesei xyn2 (Tenkanen, Puls, & Potanen, 1992). Interestingly, the optimal pH for production of AXOS by direct fermentation of BSG with B. subtilis clone 2 was pH 7.0 (Amorim et al., 2018). However, in that process the exposure of the carbohydrates present in the lignocellulosic biomass to the enzymes is improved by alkaline conditions (Maurelli, Ionata, La Cara, & Morana, 2013).

The effect of the xylan concentration on YRS was studied using different concentrations of xylan in 2% (v/v) Vogel medium at pH 6.0, 45°C and 950 rpm (Fig. 2B). The YRSmax was achieved after 8h of fermentation for all the concentrations tested.

For 2.5 and 5.0 g/L of xylan, similar YRSmax values were obtained (133 ± 10 and 133 ± 8 mg/g, respectively). However, for 2.5 and 1.25 g/L, the measured concentrations of sugars are close to the lowest sensitivity limit of the DNS method, which may also explain the higher error associated to the YRS values, in particular for the YRS values at 0h. On the other hand, concentrations of xylan higher than 5.0 g/L had a negative impact on the YRSmax reducing significantly (t-student test, α = 0.05) its value (YRSmax of 97 ± 8 mg/g for 20.0 g/L of xylan). This effect could be caused by different possible phenomena, such as promoter regulation effect; inhibition by the end-product and/
or substrate; mass transfer and aeration issues (Akpinar, Ak, Kavas, Bakir, & Yilmaz, 2007; Helianti et al., 2016). The latter is generally associated to the increase of the viscosity and density of the reaction mixture, which occurs when higher concentrations of substrate are present (Akpinar et al., 2007).

A possible effect on promoter regulation caused by the gene manipulation of the microorganism was dismissed as the same negative effect of increasing concentrations of xylan was observed under the same conditions for B. subtilis wt (Appendix A, Fig. A2 in Supplementary material). The hypothesis of inhibition by the end product was tested with the addition of xylose to the original medium containing xylan. As expected, the xylanase activity was residual in this scenario, due to the preferential consumption of free xylose in the media (data not shown). Lastly, the hypothesis of inhibition by the substrate was assessed through the fed-batch production strategy (more detailed information on this is presented in Section 3.2).

In summary, attending the sensitivity limit of both DNS and HPLC techniques, the optimal concentration of xylan selected was 5.0 g/L, which was used for the subsequent studies.

The effect of temperature on the YRS was evaluated at pH 6.0, 5.0 g/L xylan and 950 rpm, using a temperature range from 37 °C up to 47 °C (Fig. 2C). The highest YRSmax was obtained at 42.5 °C (135.6 ± 8.3 mg/g) after 8 h of fermentation with a correspondent Yxyl of 2.5 ± 0.4 mg/g, while the lowest YRSmax was observed at 37 °C (105 ± 8 mg/g). The optimal temperature possibly stands under the synergy between the optimal temperature for B. subtilis growth, between 30 and 37 °C (Korsten & Cook, 1996), the positive effect of high temperatures in reducing xylan viscosity and consequently increasing oxygen and mass transfer, and the optimal temperature for B. subtilis xylanase activity, between 50 and 56 °C (Banka, Guralp, & Gurali, 2014; Irfan et al., 2016; Milessi et al., 2015) and T. reesei xylanase (xyn2) activity, around 60 °C (La Grange, Pretorius, & van Zyl, 1996). However, this positive effect appears to be more significant for direct fermentation of lignocellulosic biomass, since it improves the accessibility of the enzymes to the hemicelluloses (Amorim et al., 2018).

3.2. Fed-batch production of XOS by B. subtilis clone 2

As previously demonstrated in Section 3.1.2, higher concentrations of xylan had a negative impact on YRS, possibly due to mass transfer issues and xylanase inhibition by the substrate and the product. In this sense, the fed-batch operation mode was seen as a potential production approach to overcome those issues. Two initial concentrations of xylan were tested, 2.5 and 5.0 g/L, which were increased up to 5.0 and 7.5 g/L of total substrate, by one impulse addition of a concentrated xylan solution at different fermentation times (3, 5, 7 and 9 h) (Fig. 3A and B). The pH and temperature conditions previously selected as optimal for batch production (Section 3.1.2) were also used in these experiments, namely 42.5 °C and pH 6.0.

For both initial concentrations of xylan, the impulse at 3 h lead to the highest YRSmax which was achieved after 8 h of fermentation, similarly to what occurs in the batch operation mode. Nonetheless, only for an initial xylan concentration of 2.5 g/L, the YRSmax increased significantly (t-student test, α = 0.05), around 12%, when compared to the optimized batch mode (136 ± 8 mg/g). The TLC analysis of the fermentation supernatant at 8 h shows strong bands in the region of 4-, 5- and 6-DP XOS and only a vestigial band can be observed in the xylose region (Appendix A, Fig. A1 in Supplementary material).

A two-time impulse fed-batch was also tested at 3 and 5 h, using an initial xylan concentration of 2.5 g/L (w/v) which was increased to a total amount of 5.0 g/L with the impulses. However, when compared to one impulse, this scenario affected negatively both YRSmax, decreasing 8%, and tmax, increasing 3 h (data not shown), when compared to the batch results, possibly due to substrate inhibition.

The improvement of the YRSmax using the one-time impulse fed-batch corroborates the hypothesis of substrate inhibition, however a
more detailed study should be performed to evaluate the mechanisms underlying this phenomenon.

Interestingly, the most considerable improvement of \( \text{Y}_{\text{XOSmax}} \) was due to the genetic engineering of the microorganism, 37% higher than the wild type, while the process optimization led to a less significant increase of the yield (around 16%). The optimization of the microorganism performance may represent a key factor to ensure the economic viability of XOS production by direct fermentation. Furthermore, the two-step metabolism of xylan, including the initial production of oligosaccharides followed by their degradation, is one of the main advantages of the direct fermentation approach, since it allows to minimize the amount of free xylose. This fact considerably simplifies the downstream process, which generally represents up to 80% of the total entire production costs (Urmann, Graals, Joehnck, Jacob, & Frech, 2010).

Fig. 3C shows the biomass growth quantified under the best conditions selected for the batch (Section 3.1.2) and fed-batch fermentation. Interestingly, the batch mode leads to a higher cell density at 8 h (207 × 10^6 cells/mL), although presenting a higher doubling time (98 min) when compared to fed-batch (171 × 10^6 cells/mL and 87 min). Higher number of cells may imply higher number of xylanases in the medium, as well as xylosidases, also responsible for XOS degradation. Thus, since the production of XOS is growth-associated, the direct fermentation of xylan requires a precise amount of biomass to allow a high production and a slow degradation of XOS, leading to their accumulation in a specific time.

3.3. Quantification and chemical characterization of XOS produced by B. subtilis clone 2

The amount of XOS produced by B. subtilis clone 2 was quantified by HPLC after their partial purification (Sections 2.6 and 2.7). Table 1 shows the production yields in equivalents of xylose for XOS, \( \text{Y}_{\text{XOS}} \) free xylose, \( \text{Y}_{\text{Xyl}} \), uptake of xylan, \( \text{Y}_{\text{Xup}} \), and residual xylan, \( \text{Y}_{\text{xr}} \), obtained per gram of xylan after 8 h of fermentation under fed-batch mode at optinal conditions selected in Section 3.2.

The \( \text{Y}_{\text{XOS}} \) quantified by HPLC (306 ± 4 mg/g) deviates greatly from \( \text{Y}_{\text{XOS}} \) obtained by DNS (148 ± 10 mg/g). It is important to note that DNS was used previously as a qualitative technique to set the fermentative profile evolution in terms of total reducing sugars (Sections 3.1–3.2). However, it is important to highlight the limitations of this technique for evaluating a mixture of oligosaccharides, since DNS assay can only be accurate for the evaluation of a single reducing sugar (Miller, 1959). Indeed, this technique responds differently to XOS of different DPs, showing lower reactivity to lower-ACP XOS (Jeffries, Yang, & Davis, 1998), which may explain the differences observed between \( \text{Y}_{\text{XOS}} \) and \( \text{Y}_{\text{XOS}} \).

The relative amount of \( \text{Y}_{\text{XOSeq}} \) in the mixture at 8 h was higher than the \( \text{Y}_{\text{Xup}} \) and \( \text{Y}_{\text{Xup}} \), together, however the residual xylan represents the most considerable portion, suggesting that the microorganisms degrade the xylan in a stepwise process. In this sense, a continuous process could be a potential approach to overcome the dynamics of the microorganism metabolism. Nonetheless, it is important to highlight that \( \text{Y}_{\text{Xup}} \) could be overestimated, as it may be including the XOS with DP > 6 which cannot be detected by HPLC within the sensitivity limits. This is corroborated by the TLC analysis that presents a blurred band corresponding to the region of DP > 6 XOS (Appendix A, Fig. A1 in Supplementary material).

Table 2 shows a reduction of both \( \text{Y}_{\text{Xyl}} \) and \( \text{Y}_{\text{Xup}} \) values when compared to the wild type (Table 2). This performance improvement was mostly due to the genetic manipulation of B. subtilis, which allowed a reduction in 4- and 2-fold for \( \text{Y}_{\text{Xyl}} \) and \( \text{Y}_{\text{Xup}} \), respectively. Similar XOS production yields per gram of xylan using an enzymatic approach has been reported (Azeele et al., 2016; Biam et al., 2013; Kallel et al., 2014). To hydrolyze pretreated sugarcane bagasse using a xylanase of Pichia stipites during 12 h, 318.0 mg/g were obtained (Biam et al., 2013); a yield of 307.4 mg/g was obtained after an 8 h hydrolysis of garlic straw xylan with Bacillus majovensis UEB-FK xylanase (Kallel et al., 2014), and 351.5 mg/g of XOS were produced from pretreated kenaf stems using an enzymatic cocktail of Xyn2:A-nabfA during 48 h (Azeele et al., 2016). However, these processes require an extra step for the enzyme production (when it is not purchased), which is not being considered in the final yield, increasing significantly the cost of the production process. These constraints highlight the potential of using a single step approach to produce XOS.

Additionally, it is important to refer that the \( \text{Y}_{\text{XOS}} \) obtained in the present study is lower than the one obtained for BSG (542 mg/g) (Amorim et al., 2018) after 12 h of fermentation using the same modified microorganism. A possible explanation for this evidence could rely on the complexity of the lignocellulosic biomass in BSG, which may provide other compounds such as sugars and other elements stimulating biomass growth and xylanase activity. Goyal, Kalra, Sareen, and Soni, (2008) reported a higher activity of Trichoderma viride xylanase using lignocellulosic biomass as compared to that with xylan as carbon source. To confirm this evidence, a preliminary study was performed using T. reesei for direct fermentation of xylan, resulting in lower xylanase activity values when compared to the use of BSG as carbon source (data not shown). Therefore, the successful use of this single step strategy requires the combination of suitable lignocellulosic materials and effective xylanase producers.

The XOS produced were characterized by GC–MS (Section 2.10). For that purpose, the XOS were previously separated by HPLC (Section 2.9) and individually collected according their DP for neutral sugars analysis and glycosidic linkage analysis (Table 3).

Methylation analysis showed the presence of linear XOS composed of

\[ \text{XOS} \rightarrow \text{Xyl} \rightarrow \text{Xyl} \rightarrow \text{Xyl} \rightarrow \text{Xyl} \]

Table 1

<table>
<thead>
<tr>
<th>Product Yield</th>
<th>mg/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Y}_{\text{XOS}} )</td>
<td>306 ± 4</td>
</tr>
<tr>
<td>Xyloctotetraose</td>
<td>55.7 ± 0.9</td>
</tr>
<tr>
<td>Xylopanetetraose</td>
<td>145 ± 1</td>
</tr>
<tr>
<td>Xylohexaose</td>
<td>105 ± 2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Process mass balance</th>
<th>mg/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Y}_{\text{XOSeq}} )</td>
<td>348 ± 4</td>
</tr>
<tr>
<td>( \text{Y}_{\text{Xyl}} )</td>
<td>1.3 ± 0.2</td>
</tr>
<tr>
<td>( \text{Y}_{\text{Xup}} )</td>
<td>142 ± 7</td>
</tr>
<tr>
<td>( \text{Y}_{\text{xr}} )</td>
<td>507 ± 11</td>
</tr>
</tbody>
</table>

Table 2

<table>
<thead>
<tr>
<th>dp</th>
<th>( \text{Y}_{\text{XOS}} ) (mg/g)</th>
<th>( \text{Y}_{\text{Xyl}} ) (mg/g)</th>
<th>( \text{Y}_{\text{Xup}} ) (mg/g)</th>
<th>( \text{Y}_{\text{xr}} ) (mg/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>824 ± 5</td>
<td>944 ± 7</td>
<td>984 ± 7</td>
<td>997 ± 7</td>
</tr>
<tr>
<td>2</td>
<td>16 ± 1</td>
<td>3.5 ± 0.6</td>
<td>2.5 ± 0.4</td>
<td>13 ± 0.2</td>
</tr>
<tr>
<td>3</td>
<td>301 ± 6</td>
<td>193 ± 8</td>
<td>154 ± 7</td>
<td>142 ± 7</td>
</tr>
</tbody>
</table>

(a) Non-optimized batch (10.0 g/L of xylan, pH 6.0 and 45°C).
(b) Optimized batch (5.0 g/L of xylan, pH 6.0 and 42.5°C).
(c) One-time impulse fed-batch (2.5 g/L of initial xylan, impulse of 2.5 g/L xylan at 3h, pH6.0 and 42.5°C).
mainly of (1→4)-linked-xylpyranosyl residues with a small amount of branched residues. The fraction A contained XOS with an average DP of 4 with 5.3% of branching, were the branching points are mainly in disubstituted xylose residues with terminally linked arabinofuranose. The presence of glucose (22.7%), galactose (4.4%), and arabinose (2.7%) in fraction A may be explained by the composition of the commercial beechwood xylan (Section 2.1), as this fraction with the lowest DP level analysed (DP = 4) was a minor portion of the total amount of the produced XOS (18%, Table 1).

Fraction B was almost composed of xylopentaose linear oligosaccharides and fraction C presented an average degree of polymerization of 6 xylose residues with 2.2% of branching (Table 3).

Interestingly, the DP-pattern obtained with clone 2 (4–6) is more characteristic of T. reesei xyn2 activity (Tenkanen et al., 1992) than B. subtilis endo-xylanase, which is associated to the production of lower DP XOS (between 2 and 4) (Reddy & Krishnan, 2016b). These results suggest that there is a dominance of xyn2 activity over the endogenous xylanase system of B. subtilis.

To modulate the gut microbiota the prebiotic oligosaccharides must be resistant to digestion in upper gastrointestinal tract, i.e. reaching the large intestine intact (Gibson et al., 2017). The stability of the produced XOS to the digestive system (i.e. gastric acidity, digestive enzymes) was tested under a harmonized static in vitro digestion method. After digestion, 98.5 ± 0.2% of the oligosaccharides remained in its complex form, without being degraded. This means the produced XOS were resistant to hydrolysis along the simulated harsh conditions of the gastrointestinal system, suggesting that they could reach the large intestine intact.

4. Conclusions

The current study clearly highlights the potential of using genetic engineering tools to improve the performance of B. subtilis to produce XOS, a prebiotic functional food ingredient. Cloning of T. reesei xylanase gene coupled with a secretion tag into B. subtilis wt allowed increasing the production of XOS and reducing the production and uptake of free xylose obtained by direct fermentation of beechwood xylan. When comparing to the enzymatic process, direct fermentation is a more attractive and advantageous approach to produce XOS, presenting the potential of reducing costs associated with enzymes purchase and downstream process.

Acknowledgments

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.carbpol.2018.09.088.
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