Artificial consortia of *Bacillus amyloliquefaciens* HM618 and *Bacillus subtilis* for utilizing food waste to synthetize iturin A

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Abstract

Food waste is a cheap and abundant organic resource that can be used as a substrate for the production of the broad-spectrum antifungal compound iturin A. To increase the efficiency of food waste biotransformation, different artificial consortia incorporating the iturin A producer *Bacillus amyloliquefaciens* HM618 together with engineered *Bacillus subtilis* WB800N producing lipase or amylase were constructed. The results showed that recombinant *B. subtilis* WB-A13 had the highest amylase activity of 23406.4 U/mL, and that the lipase activity of recombinant *B. subtilis* WB-L01 was 57.5 U/mL. When strain HM618 was co-cultured with strain WB-A14, the higher yield of iturin A reached to 7.66 mg/L, representing a 32.9% increase compared to the pure culture of strain HM618. In the three-strain consortium comprising strains HM618, WB-L02, and WB-A14 with initial OD600 values of 0.2, 0.15, and 0.15, respectively, the yield of iturin A reached 8.12 mg/L, which was 38.6% higher than the control. Taken together, artificial consortia of *B. amyloliquefaciens* and recombinant *B. subtilis* can produce an increased yield of iturin A, which provides a new strategy for the valorization of food waste.

Keywords Antifungal lipopeptide · Waste substrate · Biotransformation · *Bacillus* species · Extracellular enzyme · Co-culture

Introduction

Food waste (FW) is generally discharged from restaurants, family kitchens, and public canteens (Zhang et al. 2014). In China, about 60 million tons of FW were generated annually in 2011, and this value was expected to reach 140 million in 2020 (Zhang et al. 2016; Ye et al. 2018). As such, a large amount of FW is a significant sanitary and ecological problem; there is an urgent need for rational and effective methods to valorize FW. Biochemically, FW is a mixture of organic substances, including starch, proteins, lipids, and cellulose (Dinesh et al. 2018; Kim et al. 2011). Accordingly, FW is rich in nutrients that can be used by microorganisms and converted into value-added products (Banu et al. 2020). In recent studies, FW has been biotransformed into pullulan (Rishi et al. 2020), Bt biopesticide (Zhang et al. 2015), bioflocculants (Liu et al. 2019), lipids and proteins (Zeng et al. 2017), biosurfactants (Chen et al. 2018), xanthan (Li et al. 2017), surfactin (Pan et al. 2021), ethanol (Wang et al. 2008), and L-lactic acid (Tashiro et al. 2016) through microorganism aerobic fermentation, as well as methane (Park et al. 2019; Yu et al. 2018), hydrogen (Kuang et al. 2020), and volatile fatty acid (Cheah et al. 2019) through anaerobic digestion. Compared with traditional disposal methods such as incineration or landflling (Tang et al. 2017), biotransformation of FW is much more environmentally friendly and sustainable.

Iturin A, an antifungal cyclic lipopeptide mainly produced by *Bacillus* species, is composed of a cyclic heptapeptide

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and a β-amino fatty acid chain with 13–17 carbon atoms (Zhao et al. 2019b; Kawagoe et al. 2015). Iturin A can be embedded in the fungal cell membrane to increase its permeability to K⁺ cations, which results in broad-spectrum antifungal activity (Cochrane and Vederas 2016). Therefore, iturin A possesses bright application prospects in the biological control of plant fungal pathogens (Wang et al. 2020; Calvo et al. 2019; Ambrico and Trupo 2017). Despite its high potential value, the high production cost coupled with low yield limits the further industrial application of iturin A (Dang et al. 2019; Shi et al. 2018). It is therefore imperative to reduce the production costs, and the use of cheaper raw materials is a promising strategy to achieve better process economics. In recent studies, various agro-industrial wastes or by-products such as rapeseed meal (Chen et al. 2021), rapeseed cake (Chen et al. 2019b), soybean curd residue (Mizumoto et al. 2006), soybean meal (Xu et al. 2020), and sunflower oil cake (Kumar et al. 2017) have been investigated as substrates for iturin A production. These studies have greatly stimulated the interest in exploring the utilization of FW to produce iturin A.

The degradation of complex substrates, such as FW, is generally considered the rate-limiting step for microbial fermentation (Choi et al. 2018). Usually, it is preferred to add commercial enzymes to degrade FW into energy-rich small molecules that can be directly used by microorganisms (Rishi et al. 2020; Al-Dhabi et al. 2020; Prasoulas et al. 2020). Although the effect of this approach is obvious, the cost is relatively high. Presently, the co-cultivation of degradation and production strains has been favored for the transformation of complex substrates into value-added products, by virtue of its advantages in the minimal operational cost and synergistic metabolism of two or more organisms (Mohapatra et al. 2020; Izmirlioglu and Demirci 2017; Hashem et al. 2021). Recently, a two-strain consortium composed of an enzyme-producing fungus and *B. amyloliquefaciens* HM618 is constructed to produce surfactin using FW as the substrate (Pan et al. 2021). In an earlier study, distillers’ grains are directly used to produce surfactin by *B. amyloliquefaciens* MT45 and *B. amyloliquefaciens* X82, wherein the maximum surfactin level reached 3.4 g/L (Zhi et al. 2017). In addition, the successful co-cultures of *Bacillus coagulans* and *Bacillus thermoamylovorans* to produce L-lactic acid, *Saccharomyces cerevisiae* and *Phaffia rhodozyma* to produce ethanol, or *Bacillus* and *Enterococcus* to produce poly-3-hydroxybutyrate have been also reported (Tashiro et al. 2016; Ntaikou et al. 2018; Sindhu et al. 2020). In these studies, a target-product synthesizing strain was co-cultivated with an enzyme-producing strain, providing ideas for the valorization of FW to produce iturin A.

*Bacillus subtilis* WB800N, which has the advantages of a clear genetic background and strong protein secretion ability, is an excellent host for the production of heterologous proteins (Liu et al. 2018; Zhang et al. 2005). Because it cannot synthesize lipopeptides (Wu et al. 2018), strain WB800N is also suitable for co-cultivation with *B. amyloliquefaciens* HM618 to produce iturin A without affecting product analysis. In this study, recombinant *B. subtilis* was engineered to heterologously express amylase and lipase for the degradation of FW. Furthermore, recombinant *B. subtilis* was co-cultivated with HM618 to transform FW into iturin A. Finally, the inoculation sizes and time of strain HM618 and the engineered strains were optimized. To our knowledge, this is the first study to demonstrate the suitability of an artificial bacterial consortium to valorize FW for iturin A production.

### Materials and methods

#### Strains, plasmids, and reagents

The strains used in this study are listed in Table S1. *B. amyloliquefaciens* HM618 (CGMCC 7097), which was used to produce iturin A, was isolated from the soil. *B. subtilis* WB800N and the plasmid pHT43, used for expressing the heterologous lipase and amylase, were purchased from Bao-sai Biological Co., Ltd. (Hangzhou, China). *Escherichia coli* DH5α was used to construct the plasmids listed in Table S2.

Phanta DNA Polymerase was purchased from Novazan Biotechnology Co., Ltd. (Nanjing, China). All restriction endonucleases and T4 DNA ligase were purchased from New England Biolabs (Beijing, China). The BM Seamless Cloning and Lethal Based Fast Cloning kits were purchased from Biomed (Beijing, China) and Tiangen (Beijing, China), respectively. Iturin A standard was purchased from Sigma (USA).

#### Cultivation conditions

When constructing the plasmids, *E. coli* DH5α and the recombinant *B. subtilis* were cultivated in Luria–Bertani (LB) medium (10.0 g/L NaCl, 10.0 g/L peptone, 5.0 g/L yeast extract). If necessary, 100.0 mg/L ampicillin or 6.0 mg/L chloramphenicol was added. Before fermentation for the production of iturin A, all bacterial glycerol stocks were activated by culturing twice in 250-mL flasks containing 50 mL seed medium (60.0 g/L glucose, 10.0 g/L peptone, 10.0 g/L NaCl, 5.0 g/L yeast extract, 10.0 g/L beef extract) at 37 °C and 220 rpm for 24 h. Subsequently, the activated *B. amyloliquefaciens* HM618 and recombinant *B. subtilis* were seeded to an initial OD₆₀₀ of 0.20 into 500-mL flasks containing 200 mL of FW medium (60.0 g/L FW, 14.0 g/L sodium L-glutamate, 5.0 g/L yeast extract, 0.5 g/L KCl, 1.0 g/L KH₂PO₄, 1.0 g/L MgSO₄•7H₂O, 0.15 mg/L.
FeSO₄·5.0 mg/L MnSO₄·0.16 mg/L CuSO₄), which was based on replacing glucose in Landy medium with FW and cultivated at 37 °C and 180 rpm for 168 h.

### Plasmid construction

The lipase and amylase genes are listed in Table S3. The coding sequences of lip1a, lip2, lipA, and amyMH were synthesized and directly cloned into pHT43 by Tsingke Biotechnology Co., Ltd. (Beijing, China). For replacing the promoter P₅, or signal peptide SP₃, of plasmid pHT43 in situ, the fragments P43 and SP₄ were PCR amplified from the genome of *B. subtilis* 168 using the primer pairs P1/P2 and P3/P4 (Table 1), respectively. Then, the fragment P43 was assembled into the linearized pHT43 vector digested with *SacI* and *NotI* using the BM Seamless Cloning Kit, generating the promoter P43 replacement vector pHHT-P43. To obtain the signal peptide SP₄ replacement vector pHHT-SP₄, the fragment SP₄ was ligated into the linearized vector pHHT43, which was digested with *NotI* and *BamHI*, using T4 DNA Ligase. The promoter and signal peptide replacement vector pHHT-P43SP₄ was generated analogously. Additionally, *amyEBA* was cloned from the genome of *B. amyloliquefaciens* HM618 using the primer pair P6/P7, and four pHHT43 derivatives carrying *amyEBA* were generated through restriction coupled with ligation. The fragment encoding *amyEBA* was ligated into the linearized vector pHHT-P43lip1a, which was digested with *BamHI* and *XbaI*, generating pHHT-P43amyEBA.

### Composition analysis of FW

The FW was collected from the Xueyi canteen of Tianjin University (Tianjin, China), and mainly included rice, vegetables, and small amounts of soup. After picking out the bones, paper towels, and other impurities, FW was ground using a mechanical homogenizer and stored at ~40°C. The starch content was determined according to the method described by Ben Taher et al. (2017). The lipid content was measured via Soxhlet extraction (Pan et al. 2021). The crude protein content was estimated using the Kjeldahl method as described by Chen et al. (2017). The composition of the FW used in this study is listed in Table 2.

### Fermentation of the recombinant strains

All constructed plasmids were introduced into *B. subtilis* WB800N via natural competence (Anagnostopoulos and Spizizen 1961). The verified recombinant *B. subtilis* strains were cultured in LB medium at 37 °C and 180 rpm for 72 h. If necessary, 100.0 mg/L ampicillin, 6.0 mg/L chloramphenicol, or 0.5 mM IPTG was added. The biomass was assessed by measuring the OD₆₀₀ using a spectrophotometer (TU-1810, Beijing, China). The enzyme activity in the fermentation supernatant was measured after removing the cells by centrifugation at 12000 rpm for 5 min.

### Amylase activity assay

The amylase activity was determined by measuring the release of reducing sugars using the dinitrosalicylic acid (DNS) method. The 1.5 mL reaction system consisted of 500 μL phosphate buffer (50 mM, pH 6.0), 900 μL of soluble starch (1.0%, w/v) as substrate, and 100 μL bacterial culture supernatant. The mixture was reacted at 37 °C for 10 min, followed by adding 1.5 mL of DNS mixture to end the reaction. After boiling for 10 min, the absorbance at 540 nm (A₅₄₀) was measured using a TU-1810 spectrophotometer. The control contained the same volume of fresh medium without inoculation. One unit of amylase activity was defined as the amount of enzyme required to produce 1.0 μmol of reducing sugars per minute.

### Lipase activity assay

Lipase activity was determined using the method described by Liu et al. (2017b). The 1.0 mL reaction system consisted of 750 μL of Tris–HCl buffer (50 mM, pH 8.0), 50 μL of para-nitrophenyl laurate (p-NPL, 10 mM) as the substrate, and 200 μL of the bacterial culture supernatant. The mixture was reacted at 37 °C for 10 min, followed by boiling for 3 min to end the reaction. Finally, the A₄₁₀ was measured.

### Table 1: Primer sequences. Arms of homology are marked as capital, and restriction sites are marked with italics and underline

| Primers | Sequences (5′-3′) |
|---------|------------------|
| P1      | ACGGCCAGTTGAAATTTCGAGTCCagtcctgcatgcag |
| P2      | TTCTCCTTATATTGCGGCCGCgtgtacctctctctctcataatag |
| P3      | GCGGCCGCGatgcggcaacctgcacag |
| P4      | GATTCAGcagcgctgacgcatgcttac |
| P5      | GTGCAGAATGTCGCATgegcgcgctgctactctctcttacataatag |
| P6      | gacgcatgttcagaaaaacgacatcacaacctc |
| P7      | tcacaattgccaagagataacccgtttaacce |

### Table 2: Components identification result of FW

| Component | Content (%) |
|-----------|-------------|
| Total solid (TS) | 26.79 ± 0.89 |
| Moisture | 73.21 ± 0.11 |
| Starch (based on TS) | 47.77 ± 5.33 |
| Crude lipid (based on TS) | 24.12 ± 0.90 |
| Crude protein (based on TS) | 6.07 ± 0.31 |
using the TU-1810 spectrophotometer. The control contained the same volume of fresh medium without inoculation. One unit of lipase activity is defined as the amount of enzyme required to produce 1.0 μmol of para-nitrophenol per minute.

**Iturin A isolation and high-performance liquid chromatography (HPLC) analysis**

A sample comprising 100 mL of the fermentation supernatant was collected by centrifugation at 11000 rpm and 4 °C for 10 min, after which the pH was adjusted to 2.0 with 6 M HCl and the acidified supernatant was stored overnight at 4 °C. The lipopeptide precipitate was collected by centrifugation at 8000 rpm and 4 °C for 10 min, and then resuspended with 100 mL methanol (AR grade) for iturin A extraction overnight. After centrifuging at 8000 rpm for 10 min, and evaporating until almost completely dry, the iturin A precipitate finally dissolved in 5 mL methanol (HPLC grade). The crude iturin A filtered through a 0.22-μm pore-size nylon membrane for quantitative analysis (Zhao et al. 2019a; Shi et al. 2018).

The resulting iturin A sample (10.0 µL) was subjected to reversed-phase HPLC on an LC-20A system (Shimadzu, Japan) equipped with a C18 column (Thermo Scientific, 250×4.6 mm, 5 μm) and a UV detector set to 220 nm. The mobile phase was a mixture of water containing 0.1% formic acid and acetonitrile (55:45, v/v) at a flow rate of 1.0 mL/min (Dang et al. 2019). The iturin A concentration was quantified by comparing the peak area with the calibration curve made using the authentic iturin A standard.

**Liquid chromatography–mass spectrometry (LC–MS) analysis**

The iturin A produced by *B. amyloliquefaciens* HM618 was determined by LC–MS using a Waters HPLC interfaced with an electrospray ionization mass spectrometry instrument (Thermo Electron, USA) and micrOTOF-Q software (Bruker Daltonics, USA). The ESI source was operated in positive ion mode, and the m/z scanning range was 800–2000. The mobile phase and column were the same as described above.

**Glucose analysis**

The content of glucose in the fermentation broth is an important indicator of the growth status of the bacteria. The fermentation broth was centrifuged at 12000 rpm for 10 min and filtered through a 0.22-μm pore-size aqueous membrane. Then, a sample comprising 10.0 mL was subjected to reversed-phase HPLC using an Aminex HPX-87H Ion Exclusion particles column (7.8 mm×300 mm, BIORAD) and a differential refractive index detector. The mobile phase was 5.0 mM sulfuric acid in water at a flow rate of 0.6 mL/min and the column temperature was maintained at 65°C (Liu et al. 2017a). The retention time of glucose was 9.3 min.

**Statistical analysis**

All experiments were performed in triplicate. Statistical analysis was performed using Microsoft Excel 2019 (Microsoft, USA). Differences with p values <0.05 (*), <0.01 (**), and <0.001 (***') were labeled with one, two, and three asterisks, respectively.

**Results and discussion**

**Identification of iturin A produced by *B. amyloliquefaciens* HM618**

Iturin A is a mixture of several homologues (Chen et al. 2019a). As reported, the hydrogen adduct ion m/z values of the four isoforms of iturin A with a C14 to C17 chain length were 1043.55, 1057.55, 1071.56, and 1085.58, respectively (Chen et al. 2019a; Dang et al. 2019). Consistent with the reported results, the LC–MS identification results showed that iturin A produced by *B. amyloliquefaciens* HM618 was composed of the four homologues of C14, C15, C16, and C17 (Fig. S1b). However, the tested C15 iturin A showed two liquid phase peaks, which may indicate two different sub-types. In addition, the C14 iturin A accounted for the largest proportion, while the amount of C17 iturin A was very small.

**Construction of engineered *B. subtilis* expressing amylase and lipase**

Since food waste is nearly half starch and about a quarter lipid, artificial consortia that use food waste as a substrate must produce amylases and lipases. As presented in Fig. 1, two amylase genes and three lipase genes were introduced into *B. subtilis* WB800N. During plasmid construction, the promoter and the signal peptide in plasmid pHT43 were also replaced. After screening, the recombinant WB800N-AmyE (engineered *B. subtilis* producing amylase) with higher amylase activity and the recombinant WB800N-Lip (engineered *B. subtilis* producing lipase) with higher lipase activity were designed to co-culture with *B. amyloliquefaciens* HM618. In the co-culture fermentation, the engineered *B. subtilis* are defined as the functional strains, secreting hydrolyses to convert the FW into glucose and fatty acids as precursors for the synthesis of iturin A.

A total of twenty pHT43 expression plasmids were constructed (Figs. S2–S8), confirmed by sequencing, and
introduced into \textit{B. subtilis} WB800N for lipase and amylase production. Three controls, \textit{B. subtilis} WB800N, WB + IPTG with added inducer during fermentation, and strain HM618, were also included. As shown in Figs. S9 and S10, the extracellular activities of the five enzymes consistently increased from 16 h, and reached a maximum at 48 h. According to the biomass curve (OD$_{600}$), the increase of enzyme activity was largely caused by the accumulation of biomass. Further analysis found that the three lipase genes were better expressed under the control of the IPTG-inducible promoter \( P_{garc} \). With the assistance of the signal peptide SPamyQ, the extracellular lipase activity of the recombinant strains was further increased. In addition, the results showed that the signal peptide SPamyQ greatly promoted the secretion of amylase AmyEBa, and the amylase activity of the WB-A13 and WB-A14 strains with the \( SPamyQ \) gene was significantly higher than that of other strains. Although an earlier study reported that methyl parathion hydrolase activity was increased more than 100-fold in recombinant strain WB800 under the control of the P43 promoter and \( SP_{nprB} \) signal peptide (Zhang et al. 2005), such a large increase could not be reproduced in this study.

After comparing the extracellular enzyme activity of each strain at 48 h, the recombinant strains with higher amylase and lipase activity were selected. As shown in Fig. 2, the recombinant strain WB-A13 had the highest amylase activity, reaching 23460.4 U/mL, which is 8.25 times higher than that of the control strain WB + IPTG. Accordingly, the recombinant strains WB-A13 and WB-L01 were selected for co-culture with strain HM618. Similarly, the recombinant strains WB-A14 with outstanding amylase activity and WB-L09 with higher lipase activity were also selected to construct artificial consortia with strain HM618. In addition, recombinant strain WB-L02 which had the highest lipase activity of all strains without adding IPTG was also selected for co-culture experiments.

**Production of iturin A by artificial consortia consisting of strains HM618 and recombinant \textit{B. subtilis}**

Co-culture of strain HM618 with producing amylase recombinant \textit{B. subtilis}

Considering the fact that starch accounts for nearly half of the organic components of FW, improving the degradation efficiency of starch is conducive to obtaining large amounts of reducing sugars from the substrate. Hence, artificial consortia of strain HM618 and recombinant \textit{B. subtilis} expressing high levels of amylase were constructed for the production of iturin A. A pure culture of strain HM618 was included as a control. The iturin A production of the consortium comprising strains HM618 and WB800N-AmyE is shown in Fig. 3a. The iturin A level reached 7.66 mg/L in the co-culture of strains HM618 and WB-A14, representing a 32.9% increase over the pure culture of strain HM618 with the yield of 5.76 mg/L \((p < 0.05)\). Compared with the
co-culture of strains HM618 and WB-A14, the iturin A production was slightly decreased in the co-culture of strains HM618 and WB-A13. Thus, more iturin A was produced by the consortia than the pure culture, as anticipated. Accordingly, the choice of the co-cultured strain should be based on the composition of the complex substrate. In a previous
study, the food waste contained more crude lipids than starch, and *B. amyloliquefaciens* produced more surfactin when co-cultured with lipase-producing fungi than with amylase-producing fungi (Pan et al. 2021). Furthermore, some studies investigated the biotransformation of cellulose-rich biomass, such as rice straw (Mohapatra et al. 2020) and corn stover (Minty et al. 2013), into biofuels.

The amylase activity in the co-culture was detected every 24 h. As shown in Fig. 3b, the changes of amylase activity in the co-culture fermentation were almost the same as those of strain HM618 in the pure culture, but they were obviously different from those of recombinant *B. subtilis* under the pure culture conditions. Specifically, the amylase activity in the pure culture fermentation of the recombinant strains reached about 6000 U/mL at 24 h, and remained high until the late fermentation stage. However, the amylase activity in the co-culture fermentation was lower, less than 3000 U/mL, which was almost equivalent to the amylase activity of strain HM618 under the pure culture conditions. According to the glucose consumption curve shown in Fig. S11, strain HM618 could utilize glucose at a faster rate than strain WB800N, and also reached the stationary phase earlier during fermentation. Hence, strain HM618 theoretically prioritized the use of energy to gain an advantage in terms of biomass, reducing the growth of recombinant *B. subtilis* in the co-culture fermentation. Thus, the amylase activity of the co-culture was significantly lower than that of the pure culture of engineered *B. subtilis*.

Iturin A belonged is a secondary metabolite, and its synthesis depends on the accumulation of biomass (Xu et al. 2020). The amylase activity is related to the amount of energy spent and biomass accumulated in the fermentation, which in turn might influence iturin A production. In the first 48 h of fermentation, the amylase activity of the consortium with strains HM618 and WB-A14 increased to 2216.35 U/mg, which was 62% higher than in the pure culture of strain HM618 (1369.68 U/mg) at the same stage of fermentation (Fig. 3b). High amylase activity provided more carbon sources for the growth and metabolism of strain HM618, which can explain why the consortium with strains HM618 and WB-A14 yielded more iturin A.

**Co-culture of strain HM618 with lipase-producing recombinant *B. subtilis***

Lipids are the second most abundant nutrient in FW, and the fatty acids released by the hydrolysis of neutral lipids are also an important precursor for the synthesis of iturin A, making it promising to construct artificial consortia with lipase-producing recombinant *B. subtilis*. However, the iturin A production of the co-culture of strains HM618 and WB800N-Lip was even less satisfactory (Fig. 3c). Although the yield of iturin A in the co-culture of strains HM618 and WB-L02 reached 9.99 mg/L, which was the highest yield of iturin A in this co-culture fermentation, it was still less than the yield of the control group (10.51 mg/L). Interestingly, the yield of iturin A in the pure culture of strain HM618 was 10.51 mg/L (Fig. 3c), compared to only 5.76 mg/L in the first fermentation (Fig. 3a). The difference of iturin A yields might be caused by the instability of FW components that were pulverized into solid–liquid mixture. In general, relatively more iturin A was produced by the co-culture of strains HM618 and WB800N-AmyE, which may be related to the high starch content in FW. The change trend of lipase activity in the consortia was similar to that of amylase (Fig. 3d). Notably, the lipase activity of the consortia was almost indistinguishable from that of strain HM618 in the pure culture. In particular, the lipase activity at 24 h was higher in the pure culture of strain HM618 than in any of the three consortia, suggesting that the growth of the two stains might be affected in the consortia. Furthermore, less active lipase appeared to be insufficient to hydrolyze enough lipids for iturin A synthesis. These might explain why the pure culture of strain HM618 produced more iturin A.

In order to make full use of FW, the recombinant strains WB-A14 and WB-L02 were selected for co-culture with strain HM618. However, complex interactions between microorganisms inevitably occur in co-culture fermentation, including competition for resources and sharing of metabolites (Mee and Wang 2012). Therefore, a reasonable distribution of substrates and metabolic intermediates in the mixed culture is of great significance for increasing the yield of iturin A. It was reported that optimizing the inoculation sizes of *Paenibacillus polymyxa*, which can synthesize acetoin, as well as optimizing the inoculation time of an *E. coli* strain that produces vitamin B2, can result in an artificial consortium that can produce more acetoin (Liu et al. 2017a). Similarly, optimizing the inoculum sizes of two *B. amyloliquefaciens* strains in a co-culture fermentation with distiller’s grains resulted in an increase in the surfactant yield to 2.54 g/L (Zhi et al. 2017). Therefore, the inoculation method should be optimized to balance the energy and biomass distribution between *B. amyloliquefaciens* HM618 and the substrate-degrading recombinant *B. subtilis* strains as much as possible.

**Effect of the inoculation method on iturin A production in the co-culture fermentation of strains HM618, WB-A14, and WB-L02**

The recombinant strains that degrade the food waste were added into the FW medium earlier, to obtain more sufficient small molecules when the strain HM618 was inoculated. Taking into account the higher biomass of recombinant *B. subtilis* strains after 24 h, different inoculum sizes of strain HM618 were tested. As shown in Fig. 4a, when the strain
HM618 was directly inoculated with strains WB-L02 and WB-A14, the yield of iturin A reached 8.12 mg/L, which was an increase of 38.6% compared with the control group \((p < 0.01)\). When strain HM618 inoculation was delayed for 24 h, the yield of iturin A only reached between 2.05 and 2.52 mg/L, which was significantly less than the control group with a yield of 5.86 mg/L \((p < 0.01)\), even if the initial \(OD_{600}\) of strain HM618 was increased to 0.6. This was not in line with our expectation that the yield of iturin A would be dramatically increased when strain HM618 inoculation was postponed for 24 h. This inconsistency might be explained by analyzing other results of the mixed fermentation. As shown in Fig. 4b, the amylase activity in consortia b–d at 24 h was higher than that of consortium a, and this advantage lasted until the later stage of fermentation. Accordingly, the glucose concentration in consortia b–d at 24 h was almost 4 times higher than in consortium a, and this gap gradually increased with the increase of fermentation time (Fig. 4d). These results indicated that the amount of \(B.\ subtilis\) was fully accumulated when strain HM618 was inoculated at 24 h. Consequently, the growth of strain HM618 was affected significantly, resulting in the decreased yield of iturin A in the mixed fermentation. In addition, the changes of lipase activity shown in Fig. 4c also indicated that strain HM618 no longer had a growth advantage in the three-strain consortium when its inoculation was postponed for 24 h. Although the three-strain consortium containing strains HM618, WB-L02, and WB-A14 showed certain advantages in the production of iturin A compared with other fermentation methods, the energy distribution between the three strains required additional optimization.

The broader application of iturin A is still limited by its high production cost (Dang et al. 2019; Shi et al. 2018). Although the production of iturin A from FW in the pure culture of strain HM618 was unstable (Fig. 5), the co-culture of enzyme-producing \(B.\ subtilis\) and strain HM618 could generally improve the yield. Moreover, the feasibility of

**Fig. 4** Effects of inoculation time in mixed culture fermentation of three-strain consortia. Iturin A production (a), amylase activity (b), lipase activity (c), and glucose content (d) under mixed culture of three-strain consortia. HM618 was solely cultured with initial \(OD_{600}\) of 0.2 as the control. Consortium a, \(L02_{0.15} + A14_{0.15} + HM_{0} + 0.2\); b, \(L02_{0.15} + A14_{0.15} + HM_{24h} + 0.2\) (\(L02\) and \(A14\) were inoculated with \(OD_{600}\) 0.15 and 0.15 respectively; after 24 h, HM618 was inoculated with \(OD_{600}\) of 0.2); c, \(L02_{0.15} + A14_{0.15} + HM_{24h} + 0.4\); d, \(L02_{0.15} + A14_{0.15} + HM_{24h} + 0.6\)

**Fig. 5** Comparison of iturin A production. Iturin A was synthesized from strain HM618 pure culture using Landy medium and FW medium, while from consortium comprising strains HM618 and WB-A14, consortium comprising strains HM618 and WB-L02, and the three-strain consortium comprising strains HM618, WB-L02, and WB-A14 respectively.
reducing the synthesis cost of iturin A by using FW was analyzed. As presented in Fig. 5, the yield of iturin A obtained by cultivating strain HM618 with Landy medium was 9.69 mg/L, compared with 5.76 mg/L, 10.51 mg/L, and 5.86 mg/L obtained using different consortia in FW medium. After replacing glucose with food waste, the production of iturin A did not change significantly. Therefore, this research verified the application value of FW in reducing the production cost of iturin A.

The ability of the artificial consortia to synthetize iturin A was increased to some extent by adjusting the inoculation time. In agreement with previous studies (Chen et al. 2020; Ntaikou et al. 2018; Minty et al. 2013), our results confirm that the consortia contained different functional strains could convert the waste substrates into target product. Based on this research, it may be possible to develop consortia that encompass more species (Zhi et al. 2017) or are more purposefully modified via synthetic genetic circuits (Minty et al. 2013; Dang et al. 2019) to achieve the cost-effective biotransformation of food waste into iturin A.

Conclusion

Food waste is a suitable raw material for iturin A production. Consortia that incorporate the engineered *B. subtilis* strains with excellent amylase and lipase activity constructed via genetic methods resulted in the increase of iturin A production by *B. amyloliquefaciens* HM618 compared with the pure culture. Co-culture of strains HM618 and WB-A14 produced the higher iturin A yield among the tested two-strain consortia. Furthermore, the iturin A yield of the three-strain consortium comprising strains HM618, WB-A14, and WB-L02, respectively, inoculated at initial OD<sub>600</sub> values of 0.2, 0.15, and 0.15, was 38.6% higher than that of strain HM618 pure culture. These results confirm that food waste can be converted into value-added chemicals using artificial consortia.

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Data availability The datasets used and analyzed during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate Not applicable.

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