Changing oxidoreduction potential to improve water-soluble yellow pigment production with *Monascus ruber* CGMCC 10910

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Abstract

**Background:** *Monascus* pigments are widely used in the food and pharmaceutical industries due to their safety to human health. Our previous study found that glucose concentration induced extracellular oxidoreduction potential (ORP) changes could influence extracellular water-soluble yellow pigment production by *Monascus ruber* CGMCC 10910 in submerged fermentation. In this study, H\(_2\)O\(_2\) and dithiothreitol (DTT) were used to change the oxidoreduction potential for investigating the effects of oxidative or reductive substances on *Monascus* yellow pigment production by *Monascus ruber* CGMCC 10910.

**Results:** The extracellular ORP could be controlled by H\(_2\)O\(_2\) and DTT. Both cell growth and extracellular water-soluble yellow pigment production were enhanced under H\(_2\)O\(_2\)-induced oxidative (HIO) conditions and were inhibited under dithiothreitol-induced reductive conditions. By optimizing the amount of H\(_2\)O\(_2\) added and the timing of the addition, the yield of extracellular water-soluble yellow pigments significantly increased and reached a maximum of 209 AU, when 10 mM H\(_2\)O\(_2\) was added on the 3rd day of fermentation with *M. ruber* CGMCC 10910. Under HIO conditions, the ratio of NADH/NAD\(^+\) was much lower than that in the control group, and the expression levels of relative pigment biosynthesis genes were up-regulated; moreover, the activity of glucose-6-phosphate dehydrogenase (G6PDH) was increased while 6-phosphofructokinase (PFK) activity was inhibited.

**Conclusions:** Oxidative conditions induced by H\(_2\)O\(_2\) increased water-soluble yellow pigment accumulation via up-regulation of the expression levels of relative genes and by increasing the precursors of pigment biosynthesis through redirection of metabolic flux. In contrast, reductive conditions induced by dithiothreitol inhibited yellow pigment accumulation. This experiment provides a potential strategy for improving the production of *Monascus* yellow pigments.

**Keywords:** Water-soluble *Monascus* yellow pigments, Oxidoreduction potential, NADH/NAD\(^+\), Enzyme activity, Pigment biosynthesis genes
Water-soluble yellow pigments can be synthesized via chemical modification of intracellular alcohol-soluble pigments [12], but this strategy is a potential risk in food applications. In recent decades, considerable effort has been focused on enhancing water-soluble yellow pigment production, such as strain screening [13, 14], strain mutation [15], gene modification [16], and medium optimization [17, 18]. Chen has shown that both extracellular and intracellular yellow pigments are the main pigments produced during long periods of high cell density culturing by Monascus anka [19]. In addition, our previous studies found that extracellular water-soluble yellow pigments can be produced by M. ruber CGMCC 10910 using high glucose concentrations with low oxidation reduction potential (ORP) [20, 21]. However, the pigment yield and glucose utilization was still low.

Oxidoreduction potential was identified as a control parameter for fermentation processes [22–24]. Intracellular ORP is primarily determined by the ratio of NADH/NAD+ [25]. High NADH/NAD+ ratios can regulate intracellular metabolites to generate unusable byproducts, such as alcohol and lactate [26]. However, intracellular ORP can be influenced by changing extracellular ORP [24], while extracellular ORP can be changed by adding oxidative or reductive substances, such as H2O2, dithiothreitol (DTT), and potassium ferricyanide [24, 27]. It has been reported that spinosad and pseudoaglycone yields increased 3.11-fold in Saccharopolyspora spinosa fermentation under HIO conditions, and high citric acid productivity could be achieved in Aspergillus niger cultures with defined ORP profiles [22, 28]. Therefore, extracellular ORP provides an alternative parameter for the optimization of metabolic production. Monascus pigment biosynthesis occurs via a polyketide pathway and requires many primary metabolites, such as acetyl-CoA, malonyl-CoA, and NADPH [1, 31, 32]. The pentose phosphate pathway (PPP), a branch pathway of glycometabolism, was shown to be the primary source of NADPH for polyketide and lipid biosynthesis [29]. Recently, the biosynthetic gene cluster of Monascus pigments in the M. purpureus and M. ruber genome and the functions of some critical genes involved in the pigment biosynthetic pathway were reported [30, 31]. Both the targeted inactivation of MppKS5 (the homolog of MpigA in M. ruber) in M. purpureus and the targeted-deletion of MpigA in M. ruber resulted in the abolishment of pigment production, confirming that polyketide synthase is involved in pigment biosynthesis [31, 32]. The genes MpfasA2 (the homolog of Mpigl in M. ruber) and MpfasB2 (the homolog of MpigK in M. ruber) respectively encode fatty acid synthase α-subunit and fatty acid synthase β-subunit and supply the medium-chain (C8 and C10) fatty acyl moieties for Monascus pigments biosynthetic [31–34].

The product profile of mppA (the homolog of MpigC in M. ruber), mppC (the homolog of MpigE in M. ruber) and mppE (the homolog of MpigG in M. ruber) mutants of M. purpureus substantiate that MppA-mediated ω-2 ketoreduction is a prerequisite for the synthesis of the pyranoquinone bicyclic core of the Monascus pigments and MppC activity determines the regioselectivity of the spontaneous Knoevenagel condensation. The mppB (the homolog of MpigD in M. ruber) in M. purpureus gene encodes a trichothecene 3-O-acetyltransferase (AT), which can transfer the medium-chain (C8 and C10) fatty acyl group into the polyketide chromophore to complete pigment biosynthesis. The mppD (the homolog of MpigF in M. ruber) gene in M. purpureus encodes an amine oxidase/esterase. The mppR1 (the homolog of MpigB in M. ruber) and mppR2 (the homolog of MpigI in M. ruber) genes in M. purpureus are regulatory genes encode transcription factors for pigment biosynthesis [30, 31, 34–36]. Those reports provide a frame of reference from which to investigate external factors on pigment production at the molecular level.

In this study, H2O2 was added to the fermentation medium of M. ruber CGMCC 10910. Its concentration and addition time were also optimized to improve extracellular water-soluble yellow pigment production. In addition, the effect of the oxidative environment induced by H2O2 on the extracellular and intracellular ORP, the expression levels of pigment biosynthesis genes, and the activities of key enzymes (PFK and G6PDH) in the glycolysis and pentose phosphate pathway (PPP) were also investigated.

Methods

Microorganism

Monascus ruber CGMCC 10910, which was deposited at the China General Microbiological Culture Collection Center (CGMCC), was used in this study, and it was cultivated on a potato dextrose agar (PDA) medium at 30 °C for 7 days and then stored at 4 °C.

Fermentation conditions

The seed culture medium contained 2% glucose, 0.3% yeast extract, 1% peptone, 0.4% KH2PO4, 0.05% KCl, and 0.001% FeSO4·7H2O. The fermentation culture medium contained 15% glucose, 0.5% (NH4)2SO4, 0.5% KH2PO4, 0.05% MgSO4·7H2O, 0.05% KCl, 0.003% MnSO4·H2O, 0.001% ZnSO4·7H2O, and 0.001% FeSO4·7H2O. For the seed culture, 5–6 loopfuls of single colonies (approximately 10 mm diameter) were scraped off an agar plate and inoculated into a 250 mL Erlenmeyer flask containing 50 mL of seed culture medium and agitated at 180 rpm at 30 °C. After 25 h, 2 mL of seed culture was used to inoculate a 250 mL Erlenmeyer flask containing 25 mL of
fermentation culture medium, which was then fermented for 12 days with an agitation speed of 180 rpm at 30 °C. Three duplicates were carried out for each condition.

**Addition method of H$_2$O$_2$ and dithiothreitol (DTT)**

H$_2$O$_2$ and DTT were filter-sterilized before being added to the medium. One, 5, 10, 15 or 20 mM H$_2$O$_2$ was added to the medium on day 2 of fermentation to study the effect of H$_2$O$_2$ concentration on cell growth and yellow pigment production. Ten millimolar H$_2$O$_2$ was added to the medium on day 0, 1, 3, and 5 of fermentation to study the effect of H$_2$O$_2$ addition time on cell growth and yellow pigment production. Three gram per liter DTT was added to the medium on day 3 to study the extracellular reducing conditions on cell growth and yellow pigment production.

**Determination of glucose concentration, extracellular oxidoreduction potential (ORP), and DCW**

After being cultured, the fermentation broth was vacuum-filtered through a 0.8 mm mixed cellulose esters membrane. The filtrate (extracellular broth) was appropriately diluted to determine the residual glucose concentration using the standard 3,5-dinitrosalicylic acid (DNS) method. Extracellular ORP was detected via an oxidation-reduction electrode (Leici, Shanghai) [20]. After filtration, the mycelia were washed 3 times with distilled water and dried at 60 °C in an oven for 12 h, until a constant mycelia weight was reached to determine dry cell weight (DCW) by gravity.

**Pigment analysis using UV–visible spectrophotometer**

The filtrate (extracellular broth) was appropriately diluted to determine the extracellular water-soluble yellow pigment concentration. The absorbance spectrum of the water-soluble yellow pigments was recorded by a UV–visible spectrophotometer (Unico, USA) from 300 to 550 nm at 1 nm intervals, and the absorbance units (AU) at the peak wavelength of 350 nm was multiplied by the dilution ratio, which was used as an index for the intracellular yellow pigment concentration.

**Analyses of pigment compositions by HPLC**

Analyses of pigment compositions were performed using an Alliance e2695 HPLC system (Waters, Milford, CT, USA) equipped with a 2998 Photodiode Array (PDA) detector (Waters, Milford, CT, USA) and a Zorbax Eclipse Plus C18 column (250 × 4.6 mm, 5 μm, Agilent, Palo Alto, CA, USA). The temperature of the column oven was set at 30 °C. A mixture of H$_3$PO$_4$ solution (pH 2.5, phase A) and acetonitrile (phase B) were used as the mobile phase using the following gradient program: 0 min, 80% A; 20% B; 25 min, 20% A, 80% B; 35 min, 20% A, 80% B; 36 min, 80% A, 20% B; 41 min, 80% A, 20% B. The PDA was set at 200–600 nm, and the flow rate of the mobile phase was 0.8 mL/min [21].

**Determination of the pigment composition using LC–MS**

LC–MS (liquid chromatography–mass spectrometry) consisted of a HP1100 HPLC system (Agilent, Palo Alto, CA, USA) and a microTOF-QII mass spectrometer (Bruker, Rheinstetten, Germany). The C18 column and chromatographic conditions were the same as mentioned above, except for mobile phase A (water, 0.1% formic acid).

**NADH and NAD+ determination**

The intracellular NADH and NAD+ concentrations were determined using procedures described in previous studies [25, 37]. Mycelia were rapidly ground into a powder using liquid nitrogen; 0.1 g of mycelia powder was quickly transferred into a sample tube containing 1 mL of acidic extracting solution (0.2 M HCl for NAD+) or alkaline extracting solution (0.4 M KOH for NADH). Next, the samples were heated at 50 °C for 10 min and then centrifuged at 12,000g, 4 °C for 10 min. The supernatant transformed into a tube and was neutralized using alkaline extracting solution or acidic extracting solution. After neutralization, the samples were centrifuged at 12,000g, 4 °C for 10 min. The supernatant was collected for NAD+ and NADH determination using the Coenzyme I NAD(H) content test kit (Nanjing Jiancheng Bioengineering Institute, Nanjing, China) according to the manufacturer’s instructions. The kit is based on an enzymatic cycling assay method.

**Determination of PFK and G6PDH enzyme activities**

Mycelia were rapidly ground into a powder using liquid nitrogen; 0.1 g of mycelia powder was quickly transferred into a sample tube containing 1 mL of crude enzyme extracting solution (25 mM Tris–HCl (pH = 7.6), 10 mM MgCl$_2$, 20 mM NH$_4$Cl, 0.5 mM DTT) and then...
fully shocked [38, 39]. The protein concentration was quantified using the standard Bradford method. Activity of 6-phosphofructokinase (PFK) was determined using a PFK test kit (Nanjing Jiancheng Bioengineering Institute, Nanjing, China) according to the manufacturer’s instructions. Activity of glucose-6-phosphate dehydrogenase (G6PDH) was analysed as reported with minor modifications [39]. A total of 3 mL of the reaction mixture containing 600 µL of 0.2 M Tris–HCl (pH = 8.0), 150 µL of 5 mM glucose-6-phosphate, 50 µL of 5 mM NADP++, 2 µL of β-mercaptoethanol, and 20 µL of crude enzyme solution was incubated at 30 °C for 20 min. The G6PDH activity of each sample was then determined spectrophotometrically by measuring the formation of NADPH at 340 nm.

Gene expression analysis

The relative expression levels of key genes involved in pigment biosynthesis were analysed using real-time quantitative PCR as described previously [21]. Mycelia were collected for total RNA extraction using the Plant RNA Extraction Kit (TaKaRa MiniBEST). First, cDNA was synthesized using the PrimeScript™ RT reagent Kit with gDNA Eraser (TaKaRa) according to the supplier’s protocol. Primers for the amplification of MpFasA2, MpFasB2, MpPKS5, mppR1, mppA, mppB, mppC, mppD, mppE, mppR2 (GenBank Accession No. KC148521) and the actin gene (GenBank Accession No. AJ417880) are listed in Additional file 1: Table S1. The actin gene was used as a reference gene. Gene expression was monitored by RT-qPCR using SYBR Premix Ex TaqII (TaKaRa). RT-qPCR was performed using a LightCycler 96 (Roche, USA) with the following cycling program: pre-incubation at 95 °C for 30 s, followed by a two-step amplification (40 cycles of denaturation at 95 °C for 5 s, and annealing at 60 °C for 30 s) and dissociation curve analyses (at 95 °C for 10 s, annealing at 65 °C for 60 s, then collection of dissociation curves from 65 to 95 °C, with a final incubation at 97 °C for 1 s).

Statistical analysis

Each experiment was repeated in triplicate, at minimum. Numerical data are presented as the mean ± SD. The differences among the various treatments were analysed using one-way ANOVA. All statistical analyses were performed by using SPSS 22.0 software, and p < 0.05 and p < 0.01 were considered significant and highly significant, respectively.

Results

Water-soluble yellow pigment production from different H2O2 concentrations and adding times

Monascus ruber CGMCC 10910 has the potential to produce extracellular water-soluble yellow pigments with a maximum absorbance wavelength of 350 nm [20], and the extracellular water-soluble yellow pigments could reach a maximum yield of 147 AU350 under high glucose stress fermentation [21]. Further analysis was shown that the extracellular water-soluble yellow pigments mainly contained four kinds of yellow pigments (Additional file 2: Figure S1). The cell growth and pigment production achieved by adding various amounts of H2O2 to the medium on the 2nd day of M. ruber CGMCC 10910 fermentation under high glucose stress are shown in Fig. 1. The results showed that lower amounts of H2O2 could promote cell growth while higher amounts of H2O2 had an inhibitory effect on cell growth (Fig. 1a). Extracellular and intracellular pigments were enhanced under certain doses of H2O2 but were inhibited by high concentrations (Fig. 1a–c). The optimal amount of H2O2 was found to be 10 mM, which had a 19% increase in dry cell weight (DCW), and a 35 and 26% increase in extracellular and intracellular yellow pigment yield, respectively. Thus, 10 mM was chosen as the suitable concentration for the addition time experiments.

The optimized concentration, 10 mM H2O2, was added to the fermentation medium at various growth time points, including at day 0 (beginning of fermentation), the 1st day (early exponential phase), the 3rd day (middle exponential phase), and the 5th day (late exponential phase) during pigment fermentation. Cell growth and pigment production, including intracellular and extracellular yellow pigments, were inhibited when H2O2 was added at the beginning of fermentation (Fig. 2), and these parameters were marginally increased when H2O2 was added at the late exponential phase. However, when H2O2 was added at the middle exponential phase, cell growth and pigment production were much improved. The optimal H2O2 addition time was on the 3rd day (middle exponential phase), in which extracellular watersoluble yellow pigment yield reached approximately 209 AU350, 42% higher than that of the control (Fig. 2a, b), and intracellular yellow pigments also increased by 35% and reached a maximum of approximately 236 AU100 (Fig. 2c). The UV–visible spectra and mass spectra of intracellular yellow pigments were compared with literature data (Additional file 3: Figure S2) [40, 41], which were identified as the two well-known pigments, monascin and ankaflavin. Monascin was the mainly intracellular yellow pigment based on the HPLC profile (Additional file 4: Figure S3), and the maximum yield of monascin under HIO conditions was 467.75 µg/mL.
while dithiothreitol (DTT) acting as a reducing agent could reduce the extracellular ORP [24]. 10 mM of H$_2$O$_2$ was added to the medium on the 3rd day to create an extracellular oxidative environment, and 3 g/L DTT was added on the 3rd day to create an extracellular reducing environment. Extracellular ORP increased immediately after H$_2$O$_2$ addition and reached its highest level on day 4, then decreased to a low level, similar to the control, in the later phase of the fermentation (Fig. 3a). A low extracellular ORP was also promptly obtained when DTT was added to the medium. The cell growth rate increased under an oxidative environment, which resulted in a 19% increase in dry cell weight (DCW), while the DCW under a reducing environment increased only slightly by less than 5% (Fig. 3b). The glucose consumption rate increased immediately upon addition of H$_2$O$_2$, and the total glucose consumption was higher than that of the control. However, the glucose consumption rate under reducing conditions increased only on day 6 and resulted in the lowest residual glucose concentration (Fig. 3b).

The extracellular water-soluble yellow pigment reaching maximum productivity on day 10 under oxidative conditions and reached 209 AU$_{350}$, which was 42% higher than that of the control. However, the maximum productivity of the control was reached on day 8. In contrast, the yield of extracellular water-soluble yellow pigments under reducing conditions was significantly decreased (Fig. 3c). Under oxidative conditions, the yield of extracellular water-soluble yellow pigments based on glucose consumption corresponded to approximately 1.45
AU/g, which was 37.6% higher than that of the control. The generation rate of intracellular yellow pigments was increased under oxidative conditions (Fig. 3d). The yield of intracellular yellow pigments under oxidative conditions based on glucose consumption corresponded to approximately 1.54 AU/g, which was 26.4% higher than that of the control. In contrast, the yield of intracellular yellow pigments under reducing conditions decreased in the later stages of fermentation and resulted in a lower overall yield (Fig. 3d). The yields of extracellular and intracellular yellow pigments per unit DCW under oxidative conditions were higher than that of the control, while those under reducing conditions were lower than the control (Table 1). Which demonstrated that the oxidative conditions improved yellow pigments productivity, while reducing condition played an opposite role. Therefore, extracellular oxidative conditions were better for yellow pigment production.

**Intracellular NADH/NAD+ levels under HIO conditions**

As shown in Fig. 4, the ratios of NADH/NAD+ under HIO and control conditions were increased and reached 1.3 on the 3rd day of fermentation. From the 3rd day to the 6th day, a large amount of energy was needed to transport intracellular nutrients to support the fast growth of mycelia and to biosynthesize metabolic intermediates. Consequently, the formation rate of NADH was less than its consumption rate during the growth stage, which resulted in a sharp decrease of the NADH/NAD+ ratio during this stage. The decrease rate of the
NADH/NAD+ ratio under HIO conditions was higher than that of the control and resulted in a lower NADH/NAD+ ratio in the later exponential phase and stationary phase of the fermentation process. These results indicate that the intracellular oxidoreduction status in M. ruber CGMCC 10910 was significantly influenced by H2O2 addition.

Expression of pigment biosynthesis genes under HIO conditions

The pigment biosynthetic gene cluster has been identified in Monascus spp. [31]. The expression of relative pigment biosynthetic genes under HIO conditions was analysed (Fig. 5). The MpkS gene encodes a pigment polyketide synthase, which is responsible for pigment biosynthesis. MpfasA2 and MpfasB2 encode a fungal
fatty acid synthase, which produces medium-chain (C8 and C10) fatty acyl moieties, and then a trichothecene 3-O-acetyltransferase (AT), encoded by the mppB gene, transfers the medium-chain (C8 and C10) fatty acyl groups onto the polyketide chromophore to complete pigment biosynthesis. mppA, mppC, and mppE encode oxidoreductases, of which a reductive enzyme encoded by mppE controls the biosynthesis of yellow pigments. The gene mppD encodes an amine oxidase/esterase. The genes mppR1 and mppR2 are regulatory genes for pigment biosynthesis [31, 32].

The expression levels of the genes MpFasA2, MpFasB2, MpPKS5, mppA, mppB, mppD, mppE, and mppR1 were significantly up-regulated (p < 0.01 or p < 0.05) under HIO conditions (Fig. 5). This corroborates findings regarding the intracellular and extracellular pigment production under oxidative conditions (Figs. 1, 2), indicating that H2O2 stimulates pigment production via up-regulation of transcript levels of these genes. However, the genes mppC and mppR2 were significantly down-regulated (p < 0.01 or p < 0.05) and were negatively correlated with pigment production. It has been reported that mppR2 is a negative regulatory factor [42].

Variation of PFK and G6PDH activities under HIO conditions

To understand the physiological consequences of M. ruber CGMCC 10910 caused by HIO conditions, the activities of key redox-dependent enzymes (PFK and G6PDH) in glycolysis and the pentose phosphate pathway (PPP) were analysed. As shown in Fig. 6a, PFK activity was inhibited between the 4th day and the 6th day under HIO conditions, which indicated that the glycolysis pathway was weakened when extracellular oxidative conditions were induced by H2O2. At the same time, G6PDH activity was enhanced between the 4th day and the 8th day under HIO conditions, which also indicated that the PPP was enhanced under HIO conditions (Fig. 6b). With the metabolic activities weakening, the PFK activity under HIO conditions became equal to that of the control on the 8th day, while G6PDH activity on the 8th day was also higher than that of the control and achieved levels similar to those of the control on the 10th day.

The responses of PFK and G6PDH activities to HIO conditions in this study suggest that there was a metabolic flux redirection from glycolysis to the PPP in M. ruber CGMCC 10910 fermentation, similar to that reported by other studies [28, 38, 39]. The higher G6PDH activity under HIO conditions suggested a higher PPP metabolic reaction rate, thus providing the pigment biosynthesis pathway with ample precursors.

Discussion

Monascus pigments are mixtures with multiple components [1, 10, 43]. It is difficult to determine the concentration of Monascus pigments with standard HPLC methods. Thus, the concentration of Monascus yellow pigments in this study was represented by the absorbance at their characteristic wavelengths (350 and 410 nm) [20, 21]. In this study, H2O2 was added at different fermentation time points to study the effects of H2O2 addition time on cell growth and yellow pigment production in M. ruber CGMCC 10910. Cell growth and yellow pigment production were both inhibited when H2O2 was added at the beginning of fermentation (Fig. 2), indicating the toxic effect of H2O2 on the conidia germination of M. ruber [44]. Cell growth and metabolic activities were slower at the late exponential phase; thus, there was not a significant effect on cell growth and yellow pigment production when H2O2 was added at this phase. However, extracellular water-soluble yellow pigment production was enhanced by more than 42%, and the DCW increased when H2O2 was added at the middle exponential phase (Fig. 2), a phase in which cell growth and metabolic activity were robust. The optimal amount of H2O2 for yellow pigment production was 10 mM; lower amounts of H2O2 were not sufficient to enhance yellow pigment production, while higher amounts of H2O2 had an inhibitory effect on cell growth and pigment production (Fig. 1). This kind of dose-dependent induction was also observed in other fermentation systems such as Taxus chinensis and Streptomyces hygroscopicus 5008 [39, 45]. Because the addition of H2O2 can improve the yield of yellow pigments, additional investigations using the M. ruber CGMCC10910 fermentation system were performed.

H2O2 is an electron acceptor, which can induce an extracellular oxidative environment (higher extracellular ORP) when added to the fermentation medium.
Meanwhile, DTT acts as a reducing agent and can decrease extracellular ORP, thereby creating reducing conditions in the extracellular environment [46]. In this study, H$_2$O$_2$ and DTT were used to modify the extracellular ORP to study the effect of extracellular ORP changes on the water-soluble yellow pigment production in *M. ruber* CGMCC 10910. The results showed that the yield of yellow pigments (both extracellular and intracellular) and cell growth under oxidative conditions were significantly increased (Fig. 3b–d), and it has been reported that *Monascus* pigment production was coupled with cell growth [47]; thus, the HIO condition increased yellow pigment production primarily by enhancing cell growth. In contrast, the reducing environment inhibited the biosynthesis of yellow pigments (Fig. 3), and the yield of intracellular yellow pigments under reducing conditions was decreased in the later phase of fermentation, which resulted in a lower yield. Based on the molecular
fermentation of *Monascus purpureus*, high stirring speed damages the mycelium and results in a lower pigments yield [50]. And the mycelium morphology of *Monascus anka* plays an important role in polyketides production [51]. Our study found that electron acceptors can be provided by *H₂O₂* and without increasing stirring speed, which would damage the mycelium morphology of *M. ruber* CGMCC 10910. Moreover, it was reported that yellow pigments are the reduction product of orange pigments [43], in which two of the C–C double bonds in the conjugated chain of two orange pigments are reduced to C–C single bonds and reducing power (NADH or NADPH) is required in this reduction reaction [12]. Thus, the ratio of NADH/NAD+ under oxidative conditions was lower than that of the control (Fig. 4).

The biosynthesis of *Monascus* pigment requires many primary metabolites, such as acetyl-CoA, malonyl-CoA, NADH, and NADPH [52]. NADPH acted as reducing power used for polyketide and lipid biosynthesis [31], which was mainly produced by PPP, which is a branch pathway of glycometabolism [29]. The activities of key enzymes involved in the glycolysis pathway and PPP were analysed in this study. The activity of PFK under HIO conditions was lower than that of the control group (Fig. 6a), which indicates that PFK was allosterically inhibited by higher metabolite concentrations in the glycolysis pathway under HIO conditions [53]. G6PDH activity under oxidative conditions was higher than that of the control group (Fig. 6b), which indicated that the metabolites involved in PPP under oxidative condition were significantly up-regulated. These results showed that *M. ruber* CGMCC 10910 redirects its metabolic flux from the glycolysis pathway to the PPP, and more energy (NADPH) could be produced for polyketide and lipid biosynthesis [28, 35, 36], which is consistent with reported studies that show that metabolic flux was altered from the glycolysis pathway to the PPP, and an abundance of precursors were available for validamycin A biosynthesis in *Streptomyces hygroscopicus*. 5008 fermentation in response to various oxidant treatments [38, 39]. Moreover, analysis of the relative expression levels of the pigment biosynthetic genes showed that the genes *MpFasA2, MpFasB2, MpPKS5, mppA, mppB, mppD, mppE*, and *mppR1* were significantly up-regulated under HIO conditions (*p* < 0.01 or *p* < 0.05) (Fig. 5). Thus, HIO conditions increase yellow pigment accumulation via up-regulation of the transcript levels of relative genes and improving the precursor concentrations for pigment biosynthesis.

**Conclusions**

Extracellular oxidative conditions caused by *H₂O₂* addition at suitable fermentation time points can improve the yield of extracellular water-soluble yellow pigments in...
submerged fermentation with *M. ruber* CGMCC 10910. Cell growth and extracellular water-soluble yellow pigment production of *M. ruber* CGMCC 10910 under HIO conditions were enhanced, while these parameters were inhibited in reducing conditions induced by dithi-othreitol. Intracellular ORP (NADH/NAD+ ratios) was decreased by H2O2 addition; fewer metabolites were used to balance NADH/NAD+, and more energy could thus be directed to cell growth and yellow pigment production instead of unusable metabolite production. The transcription levels of relevant pigment biosynthesis genes were up-regulated under HIO conditions; moreover, the activity of G6PDH was increased while PFK activity was inhibited. HIO conditions increased yellow pigment accumulation via up-regulation of the transcription levels of relevant genes and improving the precursor concentrations of yellow pigment biosynthesis through redirection of metabolic flux, which provides a potential improved strategy for producing water-soluble *Monascus* yellow pigments.

**Additional files**

**Additional file 1:** Table S1. Primers for RT-qPCR analyzing pigments biosynthetic genes.

**Additional file 2:** Figure S1. Mass spectra and UV–visible spectra of extracellular yellow pigments detected by LC–MS and HPLC-PDA.

**Additional file 3:** Figure S2. Mass spectra and UV–visible spectra of intracellular yellow pigment monascin and ankaflavin detected by LC–MS and HPLC-PDA.

**Additional file 4:** Figure S3. HPLC-PDA chromatogram of intracellular yellow pigments and monascin standard curve detected by HPLC-PDA.

**Abbreviations**

ORP: oxidoreduction potential; DTT: dithiothreitol; HIO: H2O2 induced oxidative; PPP: pentose phosphate pathway; AU: absorbance units; DCW: dry cell weight; PFK: 6-phosphofructokinase; G6PDH: glucose-6-phosphat dehydrogenase.

**Authors’ contributions**

TH planned and carried out the experiments, analysed the data and wrote the manuscript; HLT and FJL assisted in carrying out experiments; GC reviewed the manuscript; ZQW participated in the data analysis and finalized the manuscript. All authors read and approved the final manuscript.

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**Competing interests**

The authors declare that they have no competing interests.

**Availability of data and materials**

We conducted the experiments and generated the data. All data are shown in figures, tables and additional data.

**Consent for publication**

Not applicable.

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