Response surface methodology-based optimization of production media and purification of α-galactosidase in solid-state fermentation by *Fusarium moniliforme* NCIM 1099

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Received: 25 June 2016 / Accepted: 23 November 2016 / Published online: 9 December 2016 © The Author(s) 2016. This article is published with open access at Springerlink.com

**Abstract** Response surface methodology was used to enhance the production of α-galactosidase from *Fusarium moniliforme* NCIM 1099 in solid-state fermentation. Plackett–Burman design was employed for selection of critical media constituents which were optimized by central composite rotatable design. Wheat bran, peptone and FeSO\(_4\)/7H\(_2\)O were identified as significant medium components using PB design. Further CCRD optimized medium components as wheat bran; 4.62 g, peptone; 315.42 µg, FeSO\(_4\)/7H\(_2\)O; 9.04 µg. RSM methodological optimization increased the enzyme production from 13.17 to 207.33 U/g showing 15.74-fold enhancement. The α-galactosidase was purified by 70% fractionation followed by DEAE anion exchange column chromatography which yields 23.33% with 28.68-fold purification. The molecular weight of α-galactosidase was 57 kDa which was determined by SDS-PAGE analysis. Purified enzyme has optimum pH of 4.0 and was found to be stable in wide pH range of 3.0–9.0. Its optimum temperature was 50°C, whereas its activity remains above 50% up to 2 h at 75°C. Hg\(^{2+}\) was found to be a potent inhibitor and Mg\(^{2+}\) acted as an activator of enzyme. No significant change was observed in enzyme activity for galactose concentration, ranging from 1 to 100 mM. The \(K_m\) values of enzyme for substrates \(p\)-nitrophenyl-α-D-galactopyranoside, melibiose and raffinose were 0.20, 1.36, and 3.66 mM, respectively. Low \(K_m\) and stability to various physiological conditions of enzyme represents its potential which can be exploited in various industrial applications.

**Keywords** α-Galactosidase · *Fusarium moniliforme* · Response surface methodology · Plackett–Burman design · Central composite rotatable design · SDS-PAGE

**Introduction**

α-Galactosidase (α-D-galactoside galactohydrolase EC. 3.2.1.22) is an exoglycosidase which catalyses hydrolysis of terminal α-1,6-linked galactosyl residue from a wide range of substrates, including oligosaccharides of raffinose family sugars, such as stachyose, melibiose, verbascose and polysaccharides of galactomannans, locust bean, and guar gum (Anisha and Prema 2006, Talbot and Sygusch 1990).

α-Galactosidase has been reported to various biotechnological and clinical applications like processing of soy-milk (Tzortzis et al. 2003), conversion of blood type (Goldstein et al. 1982), and in the treatment of Fabry’s disease (Fuller et al. 2004). Commercially, α-galactosidase is used in different industries like beet sugar industry for removing raffinose from molasses and sugar syrups (Sri-isha et al. 2010). Similarly, it is reported to increase the nutritional quality of legumes by hydrolyzing galactooligosaccharides, galactomannans, which is also responsible for improved gelling in guar gum (Bulpin et al. 1990). Due to the presence of α-galactosidases like raffinose and stachyose in soybeans, humans are incapable to digest soy-derived products (Silvestroni et al. 2002). The use of...
microbial α-galactosidase found to be a promising solution to degrade such non-digestible oligosaccharides in soy products, which opens an avenue for its use at industrial level (Cruz and Park 1982).

Over the period, fermentation techniques have achieved immense importance due to their economic, social, and environmental advantages. Solid-state (SSF) and submerged fermentations (SmF) are two types of fermentation techniques in which SSF has gained importance for the production of microbial enzymes, due to economical advantages, such as use of cheap and abundant agro-industrial waste as a substrate. It has special interest in the processes where the crude fermented product is used directly as the enzyme source (Pandey et al. 1999). Among various groups of microorganisms used in SSF, the filamentous fungi are the most exploited because of their ability to grow on solid substrates and production of wide range of extracellular enzymes (Archana and Satyanarayana 1997). SSF is superior to SmF due to various advantages like simple media composition (cost-effective constituents), aeration process, releases negligible quantity of effluent, thus reducing pollution concerns. In addition, they efficiently provide concentrated products.

Up till now, many researchers studied α-galactosidase from various perspectives using different microorganisms. A novel α-galactosidase was identified from the hyper-thermophile archaeon Sulfolobus solfataricus (Brouns et al. 2006). Cellular localization and purification were studied in variety of microorganisms, such as Saccharomyces carlsbergensis (Lazo et al. 1977), Pichia guilliermondii (Church and Meyers 1980), Candida javanica (Cavazzoni et al. 1987), Aureobasidium pullulans (Saha and Bothast 1998), and Pseudomonas sp. (Hema and Helen 2012).

Response surface methodology (RSM) is responsible for increased product formation which is the need of hour. Response surface methodological optimization can overcome the problems associated with classical optimization. RSM is an assortment of statistical techniques for designing experiments, building models, evaluating the interactive effects of variables, and pointing the optimum conditions (Shivam et al. 2009). RSM is widely used in bioprocess technology for optimization of remarkable types of fermentation media (Desai 2008, Sunitha et al. 1999). Optimization of α-galactosidase using RSM has been previously reported from various microorganisms like a mutant strain Aspergillus foetidus (Liu C et al. 2007), Streptomyces griseoloalbus (Anisha et al. 2007) and Aspergillus niger MRSS 234 (Srinivas et al. 1994), Rhizopus oryzae Strain SUK (Gajdhane et al. 2016), etc.

But till date, not even a single report is present on statistical optimization and production of α-galactosidase from Fusarium moniliforme NCIM 1099. Therefore, the objectives of the present work are 1. to determine the potency of Fusarium moniliforme NCIM 1099 for the production of α-galactosidase, 2. development of an optimal medium composition for the production of α-galactosidase in SSF using response surface methodology, and 3. enzyme extraction, purification, and characterization of purified α-galactosidase.

Materials and methods

Chemicals and media components

Para-nitrophenyl α-D-galactopyranoside (pNPG), raffinose, melibiose, and p-nitrophenol were purchased from Sigma-Aldrich (India), and all other sugars and chemical reagents were of analytical grade and procured from Himedia (India) and local suppliers.

Culture maintenance and evaluation for the α-galactosidase activity

Fusarium moniliforme NCIM 1099 was obtained from the National Collection of Industrial Microorganisms, NCL, Pune (M. S.), India, and maintained on potato dextrose agar (PDA). Different agro-industrial brans, such as wheat bran, rice bran, green gram bran, cowpea bran, soybean bran, black gram bran, sunflower bran, flax seeds bran, and red kidney beans bran, were studied for their effect on the production of α-galactosidase. The production media contained 5 g of solid substrate moistened with distilled water, in 250 ml Erlenmeyer flasks initially maintained at pH 7. The media were inoculated with 1% of inoculum collected from 48 h grown culture of Fusarium moniliforme. Inoculated production media were incubated under static conditions at 37 °C and enzyme production was checked after every 24 h for 10 days. Enzyme was extracted in 50 ml, 0.2 M McIlvaine buffer on a rotary shaker at 180 rpm for 15 min. The content was centrifuged and the supernatant was used as the enzyme source.

Optimization of media components for enhanced α-galactosidase production

Optimal media components for α-galactosidase production were screened by one factor-at-a-time method, keeping others constant (Liu C et al. 2007). In this experimental setup, effect of additional carbon sources, such as glucose, galactose, fructose, maltose, lactose, xylose, arabinose, sucrose, melibiose, cellulose, and starch, were studied at 0.01 g/gds (Sirisha et al. 2010). To study the effect of different nitrogen sources on α-galactosidase production, various organic (yeast extract, peptone, soybean meal, soy peptone, and urea) and inorganic compounds (ammonium...
sulphate, ammonium chloride, and potassium nitrate) at 0.03 g/gds were supplemented (Shivam et al. 2009). Effect of different essential minerals on α-galactosidase production was evaluated by supplementing FeSO₄·7H₂O, 0.001%; CuSO₄·5H₂O, 0.05%; MgSO₄·7H₂O, 0.05%; NaH₂PO₄, 0.05%; and KH₂PO₄, 0.05% in production medium (Siham and Hashem 2012). All experiments were conducted in three sets and data were presented as mean values with ±SD.

Media optimization using statistical tools

Plackett–Burman design

Wheat bran, peptone, FeSO₄·7H₂O, MgSO₄·7H₂O, inoculum age, inoculum size, incubation period, and pH were selected on the basis of classical method optimization studies which were further analyzed by Plackett–Burman design using Design Expert STAT Ease software 9.0 version (Minneapolis, USA). PB design was used to identify the critical media components influencing the production of α-galactosidase significantly. Each independent variable was tested for its low (−), high (+), and (0) middle values (Supplementary data Table 1). The experimental design with observed results is illustrated in Table 1. The PB design was based on the first-order model with the following equation.

\[ Y = \beta_0 + \sum \beta_i X_i, \]

where \( Y \) is the response for productivity of α-galactosidase, \( \beta_0 \) is the intercept, and \( \beta_i \) is the linear coefficient. Regression analysis was used to identify critical components with significant \( p \) value (<0.05) influencing the production of α-galactosidase. This model is inadequate to describe the interactions occurring between the variables and only used to screen and evaluate the critical variables affecting the production of α-galactosidase. Thus, the selected variables were further optimized by CCRD of response surface methodology.

Central composite rotatable design for media optimization

Further optimization of media was done using three critical media components, wheat bran, peptone, and FeSO₄·7H₂O, which were selected from PB Design and were coded at five levels (−2, −1, 0, +1, +2) (Supplementary data Table 2). Design of matrix includes 20 experimental runs which consist of 12 random points, 6 center points, and 2 axial points. Three-dimensional (3D) surface plots were developed by respective data of experiments and quadratic equation was derived as shown below.

\[
Y = \beta_0 + \beta_1A + \beta_2B + \beta_3C + \beta_1\beta_1A^2 + \beta_2\beta_2B^2 + \beta_3\beta_3C^2 + \beta_1\beta_2AB + \beta_1\beta_3AC + \beta_2\beta_3BC \]

where \( Y \) is the response enzyme activity in units; \( A, B, \) and \( C \) are the coded independent variables; \( \beta_1, \beta_2, \) and \( \beta_3 \) are linear coefficients; \( \beta_0 \) was the intercept term; \( \beta_1\beta_1, \beta_2\beta_2, \) and \( \beta_3\beta_3 \) are the quadratic coefficients; \( \beta_1\beta_2, \beta_1\beta_3, \) and \( \beta_2\beta_3 \) are the interactive coefficients. The CCRD design was developed using Design Expert STAT Ease experimental design and was shown in Table 2.

Analytical techniques

Extraction of enzyme

For enzyme extraction, a known quantity of the fermented matter was mixed with McIlvaine buffer 30 ml (pH 7) by

Table 1 Plackett–Burman experimental design representing observed and predicted responses for production of α-galactosidase enzyme

| Trial | Coded value | Actual response (U/g) | Predicted response (U/g) |
|-------|-------------|-----------------------|-------------------------|
|       | A B C D E F G H I J K |                      |                         |
| 1     | -1 1 1 1 1 -1 -1 1 -1 1 | 41.26                 | 40.98                   |
| 2     | -1 -1 1 1 -1 1 1 -1 1 -1 | 27.13                 | 26.65                   |
| 3     | 1 1 -1 -1 1 1 1 -1 1 -1 | 53.24                 | 52.14                   |
| 4     | 1 -1 1 1 -1 1 1 -1 1 -1 | 45.04                 | 44.87                   |
| 5     | 1 1 1 -1 1 -1 1 1 -1 1 | 45.8                   | 46.76                   |
| 6     | -1 -1 1 1 1 -1 1 -1 1 -1 | 31.6                  | 32.58                   |
| 7     | 0 0 0 0 0 0 0 0 0 0 0 | 61                     | 61                      |
| 8     | 1 -1 -1 -1 -1 1 1 -1 1 -1 | 32.56                 | 31.87                   |
| 9     | 1 1 -1 -1 -1 1 1 -1 1 -1 | 37.64                 | 35.54                   |
| 10    | -1 1 1 -1 -1 1 1 -1 1 -1 | 38.53                 | 38.69                   |
| 11    | -1 -1 1 1 -1 1 -1 1 1 -1 | 49.86                 | 50.03                   |
| 12    | 1 -1 1 1 1 -1 1 -1 1 -1 | 41.96                 | 40.08                   |
| 13    | -1 -1 -1 -1 -1 1 1 -1 1 -1 | 19.73                 | 18.65                   |
shaking on a rotary shaker (180 rpm, 15 min) and cen-
trifuged at 10,000 \( g \) for 10 min (4 \( ^{\circ} \mathrm{C} \)) and the supernatant 
was used as crude enzyme.

**Enzyme assay**

Assay of \( \alpha \)-galactosidase was carried out by the method of 
Dey with some modifications (Dey and Wallenfels 1974). One 
milliliter of reaction mixture contains 0.2 ml of enzyme, 0.2 ml of 0.02 M citrate phosphate buffer pH 5.0, and 0.6 ml of 1 mM \( p \)-nitrophenyl \( \alpha \)-d-galactopyranoside (\( p \)NPG) solution in the same buffer. The mixture was 
incubated for 10 min at 50 \( ^{\circ} \mathrm{C} \) and the reaction was arrested 
by the addition of 3 ml of 0.5 M sodium carbonate solu-
tion. The amount of \( p \)-nitrophenol liberated was measured 
by the absorbance at 405 nm. One unit of enzyme activity 
was defined as the quantity of enzyme that liberated 
1 \( \mu \)mol of \( p \)-nitrophenol per min per ml of enzyme under 
assay conditions. The specific activity was articulated as 
units of enzyme activity per mg of protein content.

**Purification of enzyme**

The extracellular enzyme was subjected to 70% ammonium 
sulphate precipitation. The precipitate was collected by 
centrifugation at 8000 rpm for 15 min at 4 \( ^{\circ} \mathrm{C} \) then re-dis-
solved in 10 ml 0.02 M citrate phosphate buffer of pH 5.0, 
and dialyzed against the same buffer. The dialyzed enzyme 
was concentrated and subjected to DEAE cellulose, previ-
ously equilibrated with McIlvaine buffer (20 mM, pH 4.0). 
The enzyme was eluted with linear gradient of 0.1–0.5 M 
NaCl at a flow rate of 20 ml/h and fractions of 5 ml were 
collected. All the fractions were checked for \( \alpha \)-galactosidase 
activity and protein content was measured at 280 nm. The 
active fractions having high \( \alpha \)-galactosidase activity were 
pooled, stored at 4 \( ^{\circ} \mathrm{C} \), and used for further enzyme study.

Lowry method was used for the protein quantification and 
bovine serum albumin was used as standard (Lowry et al. 
1951). Sodium dodecyl sulphate-polyacrylamide gel elec-
trophoresis (SDS-PAGE) was performed according to earlier 
protocol (Phugare et al. 2011) with 4% stacking gel and 12% 
separating gel using a vertical gel electrophoresis system 
(GeNei, Bangalore, India). Protein bands were visualized by 
silver staining and protein marker (6.4–205 kDa, Genei, 
India) were used for molecular weight determination of 
purified \( \alpha \)-galactosidase.

**Characterization of \( \alpha \)-galactosidase**

**Effect of pH and temperature on enzyme activity 
and stability**

To estimate the optimum pH for \( \alpha \)-galactosidase activity, 
the various buffers were used in varying pH ranges like

| Trial | Coded value | Actual response (U/g) | Predicted response (U/g) |
|-------|-------------|-----------------------|--------------------------|
| 1     | 0 0 0      | 189.21                | 189.41                   |
| 2     | 1 0 0      | 175.75                | 174.24                   |
| 3     | 1 0 1      | 179.14                | 177.63                   |
| 4     | 1 1 1      | 178.63                | 178.02                   |
| 5     | 1 1 0      | 174.33                | 177.17                   |
| 6     | 0 0 0      | 186.76                | 189.41                   |
| 7     | 0 0 0      | 189.17                | 189.41                   |
| 8     | 0 1 0      | 150.52                | 149.92                   |
| 9     | 0 0 0      | 188.44                | 189.41                   |
| 10    | 0 0 0      | 192.33                | 189.41                   |
| 11    | 0 0 0      | 163.08                | 162.55                   |
| 12    | 0 0 0      | 160.34                | 163.24                   |
| 13    | 0 0 0      | 154.45                | 153.02                   |
| 14    | 0 0 0      | 190.22                | 189.41                   |
| 15    | 0 0 0      | 169.08                | 167.51                   |
| 16    | 0 0 0      | 171.21                | 169.79                   |
| 17    | 0 0 0      | 166.14                | 165.54                   |
| 18    | 0 0 0      | 159.84                | 158.47                   |
| 19    | 0 0 0      | 162.57                | 165.47                   |
citrate phosphate buffer (for pH 3–6), phosphate buffer (for pH 7–8), and glycine-NaOH buffer (for pH 9–11), at 50 °C. Its pH stability was studied by pre-incubation in buffer solutions (pH 3.0–10.0) for 24, 48, and 72 h at 50°C. Aliquots were withdrawn and assayed for residual α-galactosidase activity. Residual activity was expressed as percentage of activity as compared to untreated enzyme. The effect of temperature on the enzyme activity was evaluated using different incubation temperatures ranging from 10 to 80 °C and keeping other physicochemical parameters unaltered. Thermostability of purified enzyme was determined by pre-incubating the enzyme at 0, 55, 60, 65, 70, and 75 °C for a time interval of 30–210 min. In each of the panels, the activity under optimum condition was set as 100%.

Effect of different metal ions, reagents, galactose, and organic solvent on enzyme

The effect of different metal ions, such as Cu²⁺, Mg²⁺, Ca²⁺, Co²⁺, Ba²⁺, and Hg²⁺, each at 1 mM concentration on enzyme activity was studied. Similarly reagents like EDTA, β-mercaptoethanol, and urea (each at 1 mM) were also studied for their effect on enzyme activity. To check the feedback inhibition, different concentrations of galactose (1–100 mM) incubated with purified enzyme for 30 min at 37 °C and further assayed under standard conditions. Organic solvents tolerance (for methanol, butanol, n-propanol, glycerol, and ethyl alcohol) of the enzyme was investigated by determining its residual activity under standard assay conditions after incubating enzyme with organic solvent in 1:1 ratio for the period of 1–6 days, respectively.

Enzymatic hydrolysis of melibiose

To 500 µl of 1% solution (w/v) of melibiose in 20 mM citrate phosphate buffer (pH 4.0), 100 µl of α-galactosidase (32 U) was added. Mixtures were incubated at 50 °C and analyzed by TLC on silica plates using n-butanol/ethanol/water (10:8:7) as a solvent system. Sugars were detected by spraying the dried chromatograms with 1% orcinol (w/v) in ethanol containing 10% sulfuric acid (v/v).

Substrate specificities and kinetic parameters

The relative substrate specificities of the α-galactosidase towards synthetic (pNPG) and natural substrates (Melibiose and Raffinose) were determined under standard assay conditions. For kinetic studies, the initial rate of hydrolysis of various glycosides at different concentrations (pNPG: 1–10 mM, Melibiose and Raffinose: 1–20 mM) was measured under standard assay conditions, and the kinetic constants $K_m$ and $V_{max}$ were determined from Lineweaver–Burk plot. All data obtained from the enzyme kinetic assays and Microsoft Office Excel were used for plotting.

Statistical analysis

Obtained results were mean of three determinants. Microsoft Office data’ analysis tool pack was used to do ANOVA. Difference of $p \leq 0.05$ was considered as significant.

Results

Classical optimization of various parameters for enzyme production

Among different substrates used, wheat bran was the suitable substrate in SSF with 13.17 U/g activity (Supplementary data Fig. 1). To study the effect of pH of the medium on enzyme production, experiments were performed with media of different pH and incubated for 96 h. Enzyme production progressively increased with increase in pH of the medium from 5.0 to 7.0 and maximum production (13.44 U/g) was recorded at pH 7.0. The enzyme yield decreases if the medium pH was higher than this. In fermentation, size of inoculum is an important biological factor which determines biomass production. Maximum enzyme production was obtained when SSF medium was inoculated with 1% of spore suspension. Enzyme production yield was optimum with 55% of moisture level which is a critical factor of SSF. Maximum enzyme production was 16.55 U/g following the study of all physiological parameters.

Effect of carbon and nitrogen sources on α-galactosidase activity

The carbon source employed in microbial enzyme production is one of the most important factors of SSF. In this study, cultivations were performed with wheat bran along with additional carbon sources. Combinational effect of wheat bran with that of pure carbon sources does not showed any amendment in enzyme production. Hence, wheat bran itself fulfills the requirement of carbon source and also acts as most significant substrate; hence, further experiments were carried out with it (Supplementary data Fig. 2).

The type of nitrogen source is equally important as a nutritional requirement of organism. The effects of organic and inorganic nitrogen sources were further studied at constant carbon source (wheat bran). In this study, peptone was found to be the most effective for α-galactosidase
production (25.49 U/g) which was followed by yeast extract (23.47 U/g), meat extract (23.60 U/g), and soybean meal (23.60 U/g), while inorganic nitrogen sources failed to increase the α-galactosidase production (Supplementary data Fig. 3).

**Effect of minerals on enzyme production**

Apart from carbon and nitrogen sources, many other essential minerals, such as magnesium, calcium, and trace elements, are required to support active cellular function. Accordingly, an effect of minerals on the production of α-galactosidase was studied with the inclusion of best carbon and nitrogen sources in the production medium. Results demonstrated that FeSO₄·7H₂O (28.15 U/g) was the most effective mineral inducing α-galactosidase activity which was followed by MgSO₄·7H₂O (26.6 U/g), whereas KH₂PO₄ (25.3 U/g), NaH₂PO₄ (25 U/g), MnSO₄·H₂O (24.7 U/g), CuSO₄·5H₂O (23.14 U/g), and CaCl₂·2H₂O (16.52 U/g) showed negative effect on α-galactosidase production.

**Statistical medium optimization studies**

**Plackett–Burman design**

Exploration of the most effective nutrient components influencing the yield of α-galactosidase production was carried out using PB design. Among eight investigated variables, three variables, viz., wheat bran, peptone, and FeSO₄·7H₂O, were enmarked as the most effective ones influencing the yield of enzyme with their significant p values where the values of “Prob > F” less than 0.05 specify model terms are significant, while remaining components show p values (0.1) which are above the significant level; hence, those components are considered to be insignificant.

The main effect of medium components, standard analysis of variance (ANOVA), regression coefficient, F values, and p values of variables examined in this study, and are illustrated in Table 3. The model F value 18.67 implies that the model is significant and there is only 0.06% chance that this large value could occur due to noise; in addition, the pre-determined R² 0.72 was in reasonable agreement with the adjusted R² 0.83. The adequate exactitude measures signal-to-noise ratio. A ratio greater than 4 is desirable and 12:68 ratio indicated an adequate signal; thus, this model can be used to navigate the design space. The final equation was derived in terms of actual factors which revealed the yield of enzyme as function of independent variables represented as follows:

\[
\text{Enzyme activity (units)} = 38.28 + 3.59A + 6.11B + 5.55D. \tag{3}
\]

Using PB experimental design, only effective critical components affecting enzyme production were identified, but it does not examine the interactive effects between selected critical components which were further studied by employing central composite design (CCRD) of RSM.

**Central composite rotatable design**

The CCRD design experiment was developed using three factors at two-level factorial. Based on the quadratic regression analysis, the CCRD results substantiated the subsequent second-order polynomial equation. The equation is as follows,

\[
\text{Enzyme activity (U/g)} = 189.41 + 7.23A + 1.47B + 3.60C - 2.90AB + 1.95AC + 1.85BC - 9.66A^2 - 8.62B^2 - 6.30C^2. \tag{4}
\]

The interactive effects of medium components deduced by standard analysis of variance (ANOVA), regression coefficient, F values, and p values of variables were examined, and are illustrated in Table 4. The model F value 65.10 implies that the model is significant and there is only 0.1% chance that model F value, this large could occur due to noise. The values of “prob > F” less than 0.05 showed model terms (A, C, AB, AC, A², B², and C²) are significant and values greater than 0.1 indicated model terms are nonsignificant. The pre-determined R² 0.89 was in reasonable agreement with the adjusted R² 0.97 which depicted adequacy of model to predict response. Adequate precision measures signal-to-noise ratio, precision ratio greater than four is desirable, and the ratio is 22.64. Therefore, model can be used to navigate the design space. The “Lack of Fit F value” of 2.52 implies that the lack of fit is not significant relative to the pure error. Non-significant lack of fit is good which confirmed that the model equation was adequate to predict the enzyme yield. The value of coefficient of variation (CV% = 1.42) revealed the precision and reliability of the model.
Interaction of variables

Graphical representations of a regression equation [three-dimensional (3D) response surface plots and their respective two-dimensional (2D) contour plots] are simple. They provide a crucial contribution for understanding the interactions between the two variables and finding their optimum levels (Jhample et al. 2015). The circular order of counter plots indicates that the mutual interactions among corresponding variables are non-significant, while the elliptical order specifies significant interactions between corresponding variables (Bhagwat et al. 2015). Two interacting variables were kept at optimum levels and the remaining variables were kept at zero level. The resulted contour and 3D response surface plots can be used to predict the yield of enzyme at different concentration of variables. 3D response surface plots and contour plots are verified in subfigures of Fig. 1. Interactions between wheat bran and peptone were showed in Fig. 1a1 and a2 which were moderate significant with nearly elliptical order of contour plots indicating their influence on yield of enzyme. When concentration of both sources increases, the enzyme yield was also considerably increased but further increase in concentration resulted in reduction of enzyme yield. In Fig. 1b1 and b2, contour plots displayed nearly circular in nature; hence, the interactions between FeSO$_4$·7H$_2$O and wheat bran have less significant effect on enzyme production, while the contour plot in Fig. 1c1 and c2 showed highly elliptical nature which suggests most significant interactions among FeSO$_4$·7H$_2$O and peptone.

The optimum conditions for the maximum production of $\alpha$-galactosidase were determined by the response surface analysis and regression equation. The predicted values of the regression equation closely agreed with the experimental values. Experimental model was validated by testing batch experiments under optimal operating conditions. Results of three repeated experiments were compared. The $\alpha$-galactosidase activity obtained from the experiments was very close to the response predicted by the regression model proving the validity of the model. At these optimized conditions, the maximum $\alpha$-galactosidase activity was found to be 192.33 U/g.

Experimental model validation

The processed parameters were set to achieve a maximum response involving optimum concentration of variables: wheat bran; 4.62 g, peptone; 315.42 g, FeSO$_4$·7H$_2$O; 9.04 g; at these optimized conditions, the maximum $\alpha$-galactosidase activity was found to be 207.33 U/g.

Purification of $\alpha$-galactosidase

The $\alpha$-galactosidase from *Fusarium moniliforme* was purified to homogeneity with 28.68-fold and 23.33% yield (Table 5). The elution pattern is shown in Fig. 2a. The enzyme applied for molecular weight determination using SDS-PAGE showed a single band confirming its purity with molecular weight of 57 kDa (Fig. 2b).

Effect of pH and temperature on enzyme activity and stability

Purified $\alpha$-galactosidase is assayed for relative and residual activity at varying pH and temperature values, keeping other parameters constant. The optimum pH of enzyme was...
found to be 4.0, and in case of stability, enzyme was stable at wide pH range of 3.0–9.0 (Supplementary data Fig. 4a, 4b). The relative activities at pH 3 and 11 were 94.48 and 68.40%, respectively. The enzyme retained 100% activity when preincubated for 72 h in pH 4.0 at 50°C, whereas it retained 48.29% activity when kept in buffer of pH 10. Enzyme was having temperature optima of 50°C (Fig. 3a), whereas it was stable from 55 to 70°C, while it showed more than 50% of residual activity when preincubated at 70°C for nearly 3 h. Figure 3b shows the temperature-activity profile for α-galactosidase.

Effect of different metal ions, reagents, galactose, and organic solvents on enzyme

Among all the tested minerals, Mg²⁺ enhances the enzyme activity and it was unchanged with, Ca²⁺, Ba²⁺, K⁺, and Na⁺, whereas 100% activity was lost by Hg²⁺, indicating it as a strong inhibitor of α-galactosidase. Cu²⁺ and urea had a little inhibitory effect on enzyme, while EDTA and β-mercaptoethanol did not affect enzyme activity at all. The

| Enzyme fraction                  | Volume (ml) | Protein conc. (mg) | Activity (units) | Specific activity (U/mg) | Purification (fold) | Yield (%) |
|----------------------------------|-------------|--------------------|------------------|--------------------------|---------------------|-----------|
| Culture filtrate                 | 100         | 51.6               | 300              | 5.81                     | 1                   | 100       |
| Ammonium sulphate precipitation  | 10          | 2.77               | 230              | 83.03                    | 14.29               | 76.66     |
| DEAE-cellulose chromatography    | 7           | 0.42               | 70               | 166.66                   | 28.68               | 23.33     |
galactose tolerance of the enzyme was determined by estimating relative activity under standard assay conditions which was 91.94% for 100 mM galactose concentration (Supplementary data Table 3). In organic solvent stability study, relative activity of enzyme after 144 h of incubation was 99.78 and 93.6% for acetone and methanol, respectively, while glycerol had 79.75% (Supplementary data Table 4).

Enzymatic hydrolysis of melibiose

Hydrolysis of melibiose was examined by thin-layer chromatography. The melibiose was digested for 20 min with purified enzyme suggesting efficiency of enzyme to hydrolyze non-digestible oligosaccharide that is melibiose. It was observed that melibiose hydrolyzed to glucose and galactose (Fig. 4).

Determination of kinetic parameters

For determination of $K_m$, the effect of substrate concentration on the rate of hydrolysis of $p$-nitrophenyl-$\alpha$-$\beta$-galactopyranoside, melibiose, and raffinose was investigated with standard assay procedure. The apparent Michaelis–Menten constants ($K_m$) of the enzyme for pNPG, melibiose, and raffinose were 0.20, 1.36, and 3.66 mM, and maximum velocities ($V_{max}$) were 1.0, 0.76, and 1.42 $\mu$mol/min/mg, respectively (Supplementary data Fig. 5).

Discussion

Solid-state fermentation is cheap, simple, and cost-effective technique, and due to all these advantages, it may be used for production of many novel industrial products (Pandey et al. 1999, Archana and Satyanarayana 1997, Maurice 1998, Viniegra-Gonzalez et al. 2003, Beniwal 2010). The pH of the medium drastically affects the
production of enzymes, but in this study, we observed that no significant difference was found in the enzyme yield with the fluctuation of pH of media (data not shown). This could be explained by the fact that wheat bran possesses excellent buffering capacity for the media (Pandey et al. 1999). Similar observations were made by Nampoothiri et al. for chitinase production in wheat bran-based SSF medium (Nampoothiri et al. 2004). In general, fungal cultures are habituated to grow at low moisture content equally in this study. Fusarium moniliforme showed maximum enzyme production at 55% of moisture level. The metabolic activities of the culture and consequently product synthesis were drastically affected at lower and higher moisture levels. Lower moisture leads to reduced solubility of the nutrients, whereas a lower degree of substrate swelling and higher water tension occurred (Bhatti et al. 2007). Similarly, higher moisture contents were reported to cause decreased in porosity, gas exchange capacity, loss of particle structure, development of stickiness, and reduction in gas volume (Lonsane et al. 1985).

In the current study, it was observed that wheat bran itself acted as a best carbon source, while peptone acted as a nitrogen source. In case of α-galactosidase from Aspergillus foetidus, wheat bran and soybean meal were reported as the best sources of carbon and nitrogen, respectively (Liu et al. 2007). Furthermore, Fe$^{2+}$ was potent enhancer for enzyme production, followed by Mg$^{2+}$ in our study, whereas, in case of α-galactosidase from Aspergillus parasiticus MTCC-2796, its activity was found to be enhanced by Mn$^{2+}$ and Mg$^{2+}$ (Shivam et al. 2009). In our study, decrease in the production of α-galactosidase was observed in the presence of Ca$^{2+}$; similar results were observed by Liu CQ in case of α-galactosidase from Aspergillus foetidus, where Ca$^{2+}$ decreased the production which was due to its inhibitory effect on mycelial growth and proliferation (Liu et al. 2007).

It is noteworthy that using the simplest carbon and nitrogen sources, we have succeeded in optimizing the media components using RSM giving improved enzyme yield. The experimental results of PB stated that wheat bran, peptone, and FeSO$_4$·7H$_2$O were critical medium components, and significantly influenced the production of enzyme. Media optimization studies were carried out using CCRD where critical medium components were further optimized and were investigated for significant influence on the yield of enzyme through interactions among them. Results implicated that interactions between wheat bran and peptone were most significant ones followed by wheat bran and FeSO$_4$·7H$_2$O, whereas interactions between peptone and FeSO$_4$·7H$_2$O were less significant. Model evaluation revealed that the experimental and predicted values were very close, indicating that experimental design is effective for process optimization. Using PB design and CCRD of response surface methodology, the yield of α-galactosidase was considerably enhanced from 13.17 to 207.33 U/g which was significantly higher as compared to Streptomyces griseoalbatus which showed maximum activity of 117 U/g in a RSM optimized media (Anisha et al. 2007).

In the current study, enzyme was purified with 28.68 purification fold and 23.33% yield. In earlier reports, white rot fungus Pleurotus florida gave only 0.17% yield with 36-fold purification (Ramlingum et al. 2007). Similarly, Penicillium purpurogenum gave 2% yield with purification fold of 9.44 (Ramlingum et al. 2010). Molecular weight of purified α-galactosidase in our study was found to be 57 kDa. Puchart et al. reported α-galactosidase with similar molecular weight from Thermomyces lanuginosus (Puchart et al. 2000). Experimental results substantiated that enzyme could work at broad range of pH and temperature. The pH stability profile indicated that the purified enzyme was stable in the pH range of 3.0–9.0 but underwent a decrease in activity above 10. Maximum α-galactosidase activity was found in the acidic pH range and decreases in the alkaline range (Rodriguez et al. 2008). The effect of pH on the activity of α-galactosidase may be attributed to its amino acids and active site which undergoes protonation or deprotonation and the conformational changes induced by amino-acid ionization (Sabu et al. 2005). Similarly, α-galactosidase from white rot fungus Pleurotus florida and Monascus pilosus was stable from pH 4.0 to 9.0, and pH 3.0 to 8.0, respectively (Wong et al. 1986). Till date, values reported on temperature stability of enzyme were up to 65 °C, but in our study, it was up to 75 °C which increases the applicability of this enzyme. However, thermostability of our enzyme was low as compared to that of α-galactosidases from extremophilic marine bacteria Thermotoga neapolitana (King et al. 1998) and Thermotoga maritime (Comfort et al. 2007). D-galactose is reported as a strong inhibitor of α-galactosidase, but our enzyme showed tolerance to d-galactose up to 100 mM concentration, similar results were reported for α-galactosidase from Streptomyces griseoalbus (Anisha GS 2007), and is an important from industrial point of view, while galactose tolerance is an significant character which improves the effectiveness of α-galactosidases in liberating galactose residues from oligomeric and polymeric α-galactosides.

Activity of enzyme was completely inhibited by Hg$^{2+}$, similar reports were observed in case of α-galactosidase from Cucurbita pepo plant and various fungal sources like Torulaspora delbrueckii IFO, G. lucidum, and white rot fungus Pleurotus florida (Gaudreault and Webb 1983, Oda and Tonomura 1996, Ramlingum et al. 2007). It was observed that EDTA has no negative effect on enzyme activity, while on the other hand, it showed inhibitory
effect on Cucurbita pepo (Gaudreault and Webb 1983). It was interesting to note that α-galactosidase was stable in all organic solvents used, while stability was found to be maximum in acetone. Similar results were reported for α-galactosidase from Talaromyces flavus (Simerska et al. 2003).

Earlier, it is reported that the activity of α-galactosidase is inhibited by the higher concentrations of substrate p-nitrophenyl-α-D-galactopyranoside (Wong et al. 1986); in contrast to these results, in our study, there is no any inhibitory effect which was found with high concentration of pNPG. While studying the kinetic parameters, it was found that $K_m$ for pNPG, melibiose, and raffinose were 0.20, 1.36, and 3.66 mM, respectively. α-galactosidase from both the Monascus pilosus as well as Cicer showed $K_m$ of 0.8 mM for pNPG (Wong et al. 1986, Singh and Kayastha 2012), whereas it was around 1.07 for tomato α-galactosidase (Okutucu et al. 2010) and 0.27 for Humicola sp. α-galactosidase (Kotwal et al. 1999). In case of Aspergillus oryzae and Aspergillus niger, both of them showed $K_m$ of 5.5 mM for raffinose (Kulkarni et al. 2006), while in case of Humicola sp., it was 1.45 mM for raffinose (Kotwal et al. 1999). This study revealed that our enzyme has high affinity for synthetic substrate than natural substrate raffinose which is a larger and more bulky trisaccharide. Melibiose was hydrolyzed to glucose and galactose by enzyme under study, and was proved by TLC analysis, similar results were reported by α-galactosidase from Azotobacter vinelandii (Wong 1990).

**Conclusion**

Microbial enzymes are more efficient, significant, cheap, and easily available as compared to plants and animal enzymes. Therefore, nowadays, industrially applicable microbial enzyme production is thirst area of research. Our present investigation emphasized on production of α-galactosidase from potent microbe Fusarium moniliforme NCIM 1099 by response surface methodology in solid-state fermentation. It is significantly noted that statistically optimized medium with simplest carbon and nitrogen sources showed noteworthy increase in the production of α-galactosidase enzyme. The extracellular α-galactosidase was isolated from Fusarium moniliforme NCIM 1099 and was purified to its highest purity. Considering the characteristics of α-galactosidase produced from Fusarium moniliforme NCIM 1099, such as ability to hydrolyze nondigestible sugars, broad pH, temperature activity, and stability; opens an avenue for its use in various food industries; thus this strain could be an excellent candidate for the industrial production of α-galactosidase.

**Acknowledgements** Sanjivani B. Gajdhan is thankful to the Department of Microbiology, Shivaji University Kolhapur, for awarding Departmental Research Fellowship to support the present research work. Prashant K. Bhagwat, one of the authors, is thankful to UGC, Govt. of India for awarding BSR meritorious fellowship for doctoral research. Corresponding author would like to thank UGC-MRP with sanction Grant No. F. No. 41-1282/2012 (SR)

**Compliance with ethical standards**

**Conflict of interest** All the authors declare that they have no conflict of interest.

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