Tailoring nutritional and process variables for hyperproduction of catalase from a novel isolated bacterium *Geobacillus* sp. BSS-7

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**Abstract**

**Background:** Catalase (EC 1.11.1.6) is one of the important industrial enzyme employed in diagnostic and analytical methods in the form of biomarkers and biosensors in addition to their enormous applications in textile, paper, food and pharmaceutical sectors. The present study demonstrates the utility of a newly isolated and adapted strain of *Geobacillus* possessing unique combination of several industrially important extremophilic properties for the hyper production of catalase. The bacterium can grow over a wide range of pH (3–12) and temperature (10–90 °C) with extraordinary capability to produce catalase.

**Results:** A novel extremophilic strain belonging to genus *Geobacillus* was exploited for the production of catalase by tailoring its nutritional requirements and process variables. One variable at a time traditional approach followed by computational designing was applied to customize the fermentation process. A simple fermentation media containing only three components namely sucrose (0.55 %, w/v), yeast extract (1.0 %, w/v) and BaCl₂ (0.08 %, w/v) was designed for the hyperproduction of catalase. A controlled and optimum air supply caused a tremendous increase in the enzyme production on moving the bioprocess from the flask to bioreactor level. The present paper reports high quantum of catalase production (105,000 IU/mg of cells) in a short fermentation time of 12 h. To the best of our knowledge, there is no report in the literature that matches the performance of the developed protocol for the catalase production. This is the first serious study covering intracellular catalase production from thermophilic genus *Geobacillus*.

**Conclusions:** An increase in intracellular catalase production by 214.72 % was achieved in the optimized medium when transferred from the shake flask to the fermenter level. The extraordinary high production of catalase from *Geobacillus* sp. BSS-7 makes the isolated strain a prospective candidate for bulk catalase production on an industrial scale.

**Keywords:** *Geobacillus*, Catalase, Extremophilic, Hydrogen peroxide, Fermentation
waste disposal problems. The approach involving enzymatic degradation of \( \text{H}_2\text{O}_2 \) with catalase is devoid of the aforementioned drawbacks, and therefore, is a safer alternative to the chemical treatment approach.

Many strategies have been employed for the synthesis of catalase from microorganisms, which include optimization of critical factors, induction by \( \text{H}_2\text{O}_2 \), mutagenesis and other advanced genetic engineering approaches. However, all of these strategies failed to produce commercially viable catalase from microorganisms under adverse industrial process conditions [6, 7]. The harsh industrial conditions to which enzymes are subjected provide impetus for isolation of extremozyme producing extremophiles by employing specific strategies. Hence, the present study was carried out to explore catalase producing potential of an endospore forming newly isolated and adapted extremophilic bacterium \textit{Geobacillus} sp. BSS-7 under harsh industrial conditions. The strain is deposited at the Institute of Microbial Technology, Chandigarh under an accession no. MTCC 5873. The 16S rRNA sequence of this isolate has been submitted to the Genbank (NCBI) under an accession number KJ472212. In addition, this piece of research focuses on enhancing the microbial production of catalase by tailoring nutritional and multiple bioprocess parameters at the flask level (Fig. 1). The design of the study was further refined to improve the enzyme productivity using statistical tools. The scale up studies for catalase production was also carried out at bioreactor level. In the present investigations, a significant catalase yield of 105,000 IU/mg was achieved in a very short fermentation time of 12 h using above mentioned strategies. A comparative analysis of scientific literature reveals that the catalase yield in the present developed process is much better than other reported microbial sources like \textit{Rhizobium radiobacter} (30,420 IU/mg in 20 h) [8], \textit{Exiguobacterium oxidotolerans} T-2-2\textsuperscript{T} (16,000 IU/mg in 24 h) [9], \textit{Thermoascus aurantiacus} (5100 IU/mg in 20 days) [10] and \textit{Vibrio rumoiensis} (4092 IU/mg in 48 h) [11].

Results and discussion

Catalase production with respect to growth pattern of \textit{Geobacillus} sp. BSS-7

Catalase activity was recorded only in sonicated biomass of \textit{Geobacillus} sp. BSS-7 cells separated from the fermented broth and no extracellular catalase activity was found in the supernatant. This proves that \textit{Geobacillus} sp. BSS-7 produces intracellular catalase only. Production of biocatalysts is growth associated in nature and enzyme biosynthesis begins on attaining the log phase of growth [12]. It was analysed from the growth curve that \textit{Geobacillus} sp. BSS-7 reached its stationary phase after 36 h and thereafter, a decline in the growth and catalase activity was recorded. However, the maximum catalase activity of 22,100 IU/mg was obtained before the onset of stationary phase (Fig. 2a). Similar observations were also reported by other research groups where maximum catalase production has been achieved from \textit{Bacillus halodurans} LBK 261 [13] and \textit{Bacillus maroccanus} [14] after 16 and 24 h of incubation, respectively.

The effect of \( \text{H}_2\text{O}_2 \) (0.05–1.0 mM) as an inducer of catalase enzyme was also investigated. It was found that there is no inducing effect of \( \text{H}_2\text{O}_2 \) on the enzyme synthesis. This clearly establishes the constitutive nature of the enzyme in \textit{Geobacillus} sp. BSS-7. Takebe and co-workers [15] have also reported constitutive nature of the catalase enzyme in \textit{Exiguobacterium oxidotolerans} cells.

Tailoring of nutrient requirements for catalase production at shake flask level

\textbf{Influence of carbon source}

Tailoring of microbial fermentations has been extensively used to increase the yield and productivities of large number of bioprocesses [16]. All the tested carbon sources were found to support bacterial growth and enzyme synthesis, however sucrose was adjudged as the best source for the catalase synthesis (29,000 IU/mg of cells) from \textit{Geobacillus} sp. BSS-7. The order of catalase production from tested carbon sources was sucrose > fructose > glucose > galactose > lactose > starch > maltose > mannose > raffinose > inositol (Fig. 2b). Raffinose, mannose and inositol were found to be ineffective for the catalase production in comparison to other studied carbon sources. It is pertinent to mention that enhanced catalase production from \textit{Alternaria alternata} [17], \textit{Septoria tritici} [18] and \textit{Rhodotorula glutinis} [19] has also been obtained by using sucrose in the production medium by other workers.

The concentration of sucrose in the production medium has a significant effect on the catalase activity. The increase in sucrose concentration in medium up to 1 % (w/v) causes an increase in the catalase production (33,600 IU/mg of cells) and thereafter, the enzyme activity declined (Fig. 2c).

\textbf{Influence of nitrogen source}

A comparable catalase production was observed when yeast extract, tryptone, peptone and beef extract were supplemented in the nutritional medium, however the maximum catalase production (33,600 IU/mg of cells) was observed with the yeast extract (Fig. 2d). It was also observed that complex nitrogen sources proved better than the inorganic ones for the production of enzyme. This may be due to the release of acidic ions from inorganic nitrogen sources after their utilization by microorganisms. The high acidic conditions may inhibit the
Bioreactor

Statistical approach

Qualitative estimation of catalase enzyme

Adaptation and enrichment of microbes (Temperature 50 °C, pH 10.0, 100 rpm, nutrient medium containing H₂O₂)

Microscopic examination for purity of the culture

Subculture from isolation streak

Isolated pure culture

Microbial bioprocess for catalase production

Soil to industrial applications: Schematic overview of microbial bioprocess for catalase production

Classical approach

Statistical approach

Bioreactor

Fig. 1 Soil to industrial applications: Schematic overview of microbial bioprocess for catalase production
**Fig. 2**  

a Growth curve of *Geobacillus* sp. BSS-7; influence of medium constituents on catalase production  
b carbon sources,  
c sucrose concentration,  
d nitrogen sources,  
e yeast extract concentration
growth and metabolic processes of the microbes and hence the synthesis of enzyme [20]. Yeast extract has also been adjudged as a good nitrogen supplement for the synthesis of catalase from a number of other microorganisms such as *Serratia marcescens* SYBC-01 [21], *Exiguobacterium oxidotolerans* [15] and *Septoria tritici* [18].

Further, the effect of yeast extract concentration on the catalase activity was also studied. A progressive increase in catalase activity (33,600 IU/mg) was recorded with increase of yeast extract concentration up to 0.5 % (w/v) and thereafter, it declined (Fig. 2e). The decrease in enzyme activity above 0.5 % (w/v) yeast extract concentration may be due to its complex nature as some of the ingredients of yeast extract may have caused toxic or inhibitory effect on enzyme synthesis at higher concentrations.

The higher concentration of carbon and nitrogen sources beyond optimum level may also have caused increase in the viscosity of fermentation medium due to the tremendous growth of bacterial biomass, resulting in depletion of oxygen and nutritional imbalance in the medium. This nutritional imbalance and oxygen depletion could have possibly disturbed the cellular machinery and translation processes resulting in lower catalase production.

### Influence of trace metal salts

The supplementation of the production medium with a variety of trace metal salts (Table 1) did not exert any effect on the production of catalase except BaCl₂. The results clearly revealed that only BaCl₂ at a concentration of 0.05 % (w/v) enhances the production of enzyme (36,400 IU/mg of cells). The increase in BaCl₂ concentration above 0.05 % (w/v) did not exert any significant influence on the enzyme synthesis (Table 1). In contrary, other trace metal salts such as CaCl₂, ZnSO₄, CuSO₄, MnSO₄, MgSO₄, FeSO₄, NiSO₄, CoCl₂, HgSO₄, AgNO₃, KI, Na₂SO₃, Na₂MoO₄ inhibit the enzyme production substantially. The addition of salts like NiSO₄, AgNO₃, CuSO₄, CoCl₂, HgSO₄ in trace amount led to the complete inhibition of enzyme synthesis and growth of bacterium. This may be due to the change in conformation of enzyme/protein to a less stable form caused by these salts by altering the enzyme surface charge.

### Optimization of process parameters at shake flask level

#### Influence of pH and temperature

The pH and temperature of the production medium are critical process variables in enzymatic fermentation processes and their influence on catalase synthesis by *Geobacillus* sp. BSS-7 was investigated. The results revealed an enhancement in catalase and biomass production with

| Table 1 Influence of metal salts on the production of catalase from *Geobacillus* sp. BSS-7 |
|-----------------------------------------------|
| **Metal salts** | **Catalase activity** | **Biomass** |
| (%) (w/v) | (IU/mg of cells) | (g/100 ml) |
| **Control** | 30,100 | 0.70 |
| CaCl₂ | 0.001 | 13,400 | 0.58 |
| | 0.01 | 21,400 | 0.67 |
| | 0.05 | 23,400 | 0.74 |
| | 0.10 | 20,600 | 0.63 |
| ZnSO₄ | 0.001 | 11,200 | 0.49 |
| | 0.01 | 24,500 | 0.70 |
| | 0.05 | 12,700 | 0.57 |
| | 0.10 | 8,700 | 0.37 |
| MnSO₄ | 0.001 | 6,500 | 0.29 |
| | 0.01 | 10,200 | 0.43 |
| | 0.05 | 15,900 | 0.59 |
| | 0.10 | 11,200 | 0.47 |
| MgSO₄ | 0.001 | 17,800 | 0.24 |
| | 0.01 | 19,900 | 0.31 |
| | 0.05 | 22,300 | 0.39 |
| | 0.10 | 12,300 | 0.19 |
| FeSO₄ | 0.001 | 9,100 | 0.41 |
| | 0.01 | 17,400 | 0.47 |
| | 0.05 | 12,800 | 0.37 |
| | 0.10 | 10,100 | 0.30 |
| BaCl₂ | 0.001 | 23,000 | 0.64 |
| | 0.01 | 31,200 | 0.72 |
| | 0.05 | 36,400 | 0.85 |
| | 0.10 | 16,700 | 0.36 |
| KI | 0.001 | 4,300 | 0.30 |
| | 0.01 | 10,400 | 0.41 |
| | 0.05 | 11,220 | 0.49 |
| | 0.10 | 13,400 | 0.13 |
| Na₂SO₃ | 0.001 | 900 | 0.09 |
| | 0.01 | 5,400 | 0.23 |
| | 0.05 | 8,300 | 0.34 |
| | 0.10 | 4,800 | 0.17 |
| Na₂MoO₄ | 0.001 | 13,400 | 0.50 |
| | 0.01 | 21,300 | 0.63 |
| | 0.05 | 20,200 | 0.49 |
| | 0.10 | 8,100 | 0.29 |
an increase in pH of the fermentation medium up to pH 7.0 (36,400 IU/mg of cells), and thereafter, it declined (Fig. 3a). The low enzyme production at lower and higher pH values may be due to alteration in the ionization of nutrient molecules, which thereby reduce their availability to the organism and hence disturb the ionic balance. In addition, drastic variations in pH can also harm the microbial cells by disrupting the plasma membrane and

Fig. 3 Influence of process parameters on catalase production from Geobacillus sp. BSS-7: a pH, b temperature, c inoculum age, d inoculum size, e agitation rate and f incubation time.
disturbing their metabolism. It is clear from the Fig. 3a that *Geobacillus* sp. BSS-7 was able to grow and produce catalase at a broad pH range. There are scarce reports on the production of catalase at higher pH (>8.0) values and these are from cultures of other genera i.e. *Bacillus* sp. SF [23] and *Oceanobacillus oncorhynchi* [26].

An increase in biomass along with catalase activity was observed with increase in temperature up to 60 °C (38,400 IU/mg of cells) and thereafter, a progressive decline in both biomass and catalase synthesis was recorded (Fig. 3b). However, the said strain was able to grow and produce catalase over a wide range of temperature from 10 to 90 °C. To best of our knowledge, this is the first report from genus *Geobacillus*, where bacterium is able to grow above 80 °C and below 30 °C.

The rate of enzymatic reaction in the cells increases with the increase in temperature till an optimum temperature is reached. Beyond the optimum temperature, the enzyme inactivation occurs due to protein denaturation, which ultimately affects the cell growth and productivity. The lower catalase production at high temperatures could also be due to the decreased oxygen solubility in the medium. Further, the gelling of plasma membrane at low temperature results in slowdown of transport processes and metabolism in the microbial cells. The majority of catalase producing microorganisms reported in the literature produce catalase below 50 °C, and that too at a neutral pH. There is only one study demonstrating the production of an enzyme at 65 °C by a bacterial strain *Bacillus* sp. SF [23].

The cellular morphology of *Geobacillus* sp. BSS-7 at different temperatures (20, 50 and 80 °C) was also studied by Transmission Electron Microscopy (FEI, Tecnai, G2 F20, USA) and shown in Fig. 4a–c. Most of the cells were observed under active cell division when grown at 50 °C (Fig. 4b), whereas, bacterial cells with thick cell membrane were seen at 20 °C and only few cells were seen under division phase at this temperature (Fig. 4a). A noteworthy change in cellular morphology of *Geobacillus* sp. BSS-7 was observed when the cells were grown at 80 °C (Fig. 4c). The bacterial cells were seen as intracellular oval endospore forming rod shaped structures at this temperature. These endospores would have possibly the mechanism for *Geobacillus* sp. BSS-7 to survive under stress conditions. Limited literature is available on the cellular morphology of *Geobacillus* under extreme conditions but the presence of long endospore forming rod shaped cells of *G. thermoleovorans* at high temperatures was also observed by Marchant and co-workers [24, 25].

**Influence of inoculum age and size**

Inoculum age and size were observed to be significant parameters that exert a strong effect on the catalase production. The catalase production reached up to a maximum level of 38,400 IU/mg of cells with a 12 h old inoculum (Fig. 3c). In contrast, a lower catalase activity was obtained by using cells of inoculum age below 12 h, which could be due to the fact that bacterial cells might not have reached the exponential growth phase at that time. Keeping this in view, 12 h old culture was selected for further studies. An increase in catalase production (39,100 IU/mg of cells) was also recorded with the increase in inoculum size up to 2.5 % (v/v), and thereafter, it decreased (Fig. 3d). Hence, the inoculum size of 12 h old culture containing 1.2 × 10⁸ cells/mL was adjusted to achieve the maximum yield of enzyme.

A tremendous growth of bacteria with higher inoculum size results in nutritional imbalance due to stiff competition for nutrients between the cells. This ultimately leads to a decrease in the enzyme production capacity. On the other hand, low enzyme production below 2.5 % (v/v) inoculum is probably due to insufficient biomass for maximum utilization of nutrient molecules and production of catalase. An inoculum size of 2 % (v/v) for the production of catalase from *Pseudomonas putida* and *Bacillus subtilis* has been reported [22, 27].

**Influence of agitation rate**

The agitation of growth medium exerts a significant influence on the catalase synthesis as it affects the mixing of nutritional components and exchange of gases which further controls the availability of nutrients and oxygen to microbial cells. A progressive increase in enzyme activity (39,100 IU/mg of cells) was recorded with agitation rate up to 100 rpm, and thereafter, a decline in catalase synthesis was observed (Fig. 3e). Decreased enzyme activity at lower agitation rate may be due to inadequate mixing and supply of media components, and slow diffusion of gases. Whereas, decrease in catalase activity at higher agitation rate can be due to the influence of shear stress on bacterial cells. Similar to our observation, 100 rpm agitation rate has been reported suitable for catalase production from *Serratia marcescens* [28] and *Thermus brockianus* [29].

**Influence of incubation time**

The increase in catalase production was achieved with time and the maximum enzyme production (41,400 IU/mg of cells) was observed after 18 h of incubation (Fig. 3f). This trend may be due to an increase in the formation of oxygen radicals as a result of high metabolic rate during exponential growth phase. This enhanced production of reactive oxygen stimulates the catalase synthesis. However, they begin to form spores during late exponential phase (after 18 h) due to decrease in
nutrient availability which lead to a decreased metabolism, and hence, a decline in the catalase synthesis was recorded.

**Response surface optimization**
Tailoring of bioprocess components by traditional methods involves the change of one variable at a time which is extremely expensive and time consuming. In order to combat this drawback and determine the interaction between different experimental factors, a central composite response design was employed for the optimization of nutrient and process parameters. The observations of CCRD experiments to analyse the effect of six independent variables are shown in Table 2 along with the predicted and observed responses. The experimental results obtained were fitted in a second order polynomial equation. The fitted equation (in terms of coded value) for predicting catalase production after calculating the regression coefficient value was given successively regardless of the significance of the coefficients:

$$EA = +19,995.27 + 2312.83 \times A + 3989.95 \times B + 1430.80 \times C - 732.22 \times D + 3145.41 \times E$$
$$+ 621.15 \times F - 2849.29 \times A \times B + 3062.07 \times A \times CC + 4322.13 \times A \times D - 418.17 \times A \times E$$
$$- 83.70 \times A \times F + 5506.09 \times B \times C + 678.11 \times B \times D - 1862.18 \times B \times E + 2647.28 \times B \times F$$
$$- 533.24 \times C \times D + 3351.46 \times C \times E - 2339.07 \times C \times F + 439.34 \times D \times E - 1545.12 \times D \times F$$
$$+ 270.17 \times E \times F + 1944.21 \times A^2 - 2383.22 \times B^2 + 5516.38 \times C^2 + 8414.95 \times D^2 + 86.68 \times E^2$$
$$+ 1576.79 \times F^2$$

(1)
where A, B, C, D, E and F represents sucrose, yeast extract, barium chloride, inoculum age, inoculum size and time, respectively. The statistical significance of Eq. (1) for response surface quadratic model was tested by ANOVA and results are presented in Table 3. The quadratic model in Eq. (1) with 27 terms have six linear, six
quadratic and fifteen two factorial interactions. The insignificant terms were neglected on \( p \) value basis (\( p \) values more than 0.0500 were excluded). Then, the model Eq. (1) was amended to reduced fitted model Eq. (2):

\[
EA = + 19,995.27 + 2312.83 \times A + 3989.95 \times B + 1430.80 \times C - 732.22 \times D
+ 3145.41 \times E + 621.15 \times F - 2849.29 \times A \times B + 4322.13 \times A \times D
- 83.70 \times A \times F + 5506.09 \times B \times C + 678.11 \times B \times D - 1862.18 \times B \times E
+ 2647.28 \times B \times F - 533.24 \times C \times D - 2339.07 \times C \times F + 439.34 \times D \times E
- 1545.12 \times D \times F + 270.17 \times E \times F + 1944.21 \times A^2 - 2383.22 \times B^2 + 5516.38 \times C^2
+ 8414.95 \times D^2 + 86.68 \times E^2 + 1576.79 \times F^2
\]

It is clearly depicted from Table 3 that the present model is significant. The significance was also recognized from a very low probability value for catalase production (\( P_{\text{value}} < 0.05 \)). The \( P \) values further indicate the pattern

| Factors | Catalase activity |
|---------|------------------|
| Sum of squares | df | \( p \) value | Coefficient estimate | Standard error | \( F \) value |
| Model | 4.857E+009 | 27 | 0.0041 | 19995.27 | 2563.98 | 11.76 |
| A | 9.897E+007 | 1 | 0.0021 | 2312.83 | 1968.55 | 49.7 |
| B | 2.955E+008 | 1 | 0.0053 | 3989.95 | 1968.55 | 20.08 |
| C | 3.788E+007 | 1 | 0.0124 | 1430.80 | 1968.55 | 10.04 |
| D | 9.992E+006 | 1 | 0.0134 | -732.22 | 1968.55 | 32.63 |
| E | 1.803E+008 | 1 | 0.0001 | 3145.41 | 1968.55 | 15.06 |
| F | 7.138E+006 | 1 | 0.0244 | 621.15 | 1968.55 | 1.34 |
| AB | 1.533E+008 | 1 | 0.0061 | -2849.29 | 2074.24 | 19.57 |
| AC | 1.562E+008 | 1 | 0.2321 | 3062.07 | 2074.24 | 0.81 |
| AD | 3.113E+008 | 1 | 0.0126 | 4322.13 | 2074.24 | 11.97 |
| AE | 2.914E+006 | 1 | 0.4421 | -418.17 | 2074.24 | 0.55 |
| AF | 1.167E+005 | 1 | 0.0403 | -83.70 | 2074.24 | 1.98 |
| BC | 5.052E+008 | 1 | 0.0093 | 5506.09 | 2074.24 | 12.54 |
| BD | 7.663E+006 | 1 | 0.0042 | 678.11 | 2074.24 | 2.62 |
| BE | 5.779E+007 | 1 | 0.0002 | -1862.18 | 2074.24 | 3.21 |
| BF | 1.168E+008 | 1 | 0.0065 | 2647.28 | 2074.24 | 17.57 |
| CD | 4.738E+006 | 1 | 0.0036 | -533.24 | 2074.24 | 20.12 |
| CE | 1.872E+008 | 1 | 0.2145 | 3351.46 | 2074.24 | 1.37 |
| CF | 9.117E+007 | 1 | 0.0003 | -2339.07 | 2074.24 | 17.32 |
| DE | 3.217E+006 | 1 | 0.0074 | 439.34 | 2074.24 | 5.21 |
| DF | 3.978E+007 | 1 | 0.0256 | -1545.12 | 2074.24 | 3.98 |
| EF | 1.216E+006 | 1 | 0.0001 | 270.17 | 2074.24 | 1.28 |
| A^2 | 5.006E+007 | 1 | 0.0215 | 1944.21 | 2326.81 | 6.02 |
| B^2 | 7.521E+007 | 1 | 0.0324 | -2383.22 | 2326.81 | 8.32 |
| C^2 | 4.030E+008 | 1 | 0.0339 | 5516.38 | 2326.81 | 8.11 |
| D^2 | 9.377E+008 | 1 | 0.0031 | 8414.95 | 2326.81 | 41.39 |
| E^2 | 99503.63 | 1 | 0.0470 | 86.68 | 2326.81 | 2.21 |
| F^2 | 3.292E+007 | 1 | 0.0669 | 1576.79 | 2326.81 | 1.24 |
| Residual | 9.321E+008 | 13 | 13 | 13 |
| Lack of fit | 9.321E+008 | 7 | 7 |
| Pure error | 0.00 | 6 |
| Core total | 6.045E+009 | 41 |

* Symbols A, B, C, D, E, F are the same as mentioned in Table 5
of interaction between the coefficients on the basis of the significance of each coefficient. Smaller the $P$ value, the more significant is the corresponding coefficient. The coefficient of determination ($R^2$) presents a measure of the variability in the actual response values. This variability can be described by the experimental variables and their interactions.

A value of 1 suggests an ideal case at which 100 % of the variation in the experimental value can be explicated by the model. The matching quality of the values acquired by the model was analysed with respect to coefficient of determination ($R^2$). The mathematical tuning of those values which generated $R^2 = 0.9876$ (98.76 %) for catalase production have revealed that the model is unable to describe only 1.24 % of all the effects, thereby representing it as the best statistical model. The coefficient estimates of Eq. (2) are shown in Table 3. The model $F$ values of 11.76 for catalase production imply that the model is significant, where $F$ value represents the ratio of mean squares due to regression and mean squares due to error. Adequate precision is a measure of signal to noise ratio (value greater than 4 is desirable) and this ratio of 7.301 indicates an adequate signal for catalase activity. The lack of fit value for catalase activity (9.321E + 008) relative to pure error (Table 3) specifies that the experimental data attained fitted well with the model.

The three dimensional response surface graphs (Fig. 5a–d) based on the final model were plotted by keeping four variables at their optimum value while varying the other two variables within their experimental range. The contour curves were used for analysing the effects of different variables on the catalase production. Statistically optimal values of variables were obtained from these curves while moving along the major and minor axis of the curves. The response at the central point depicts the highest degree of achievable catalase synthesis for that set of variables. Response surface curves were obtained by keeping the catalase production on the Z-axis against any two independent factors while holding the other independent factors at their ‘0’ level.

The 3D curve and a contour plot of calculated response surface obtained from the interaction between sucrose and yeast extract is presented in Fig. 5a. The highest catalase production of 39,400 IU/mg of cells was achieved when the concentration of yeast extract and sucrose was kept at 1.0 % (w/v) and 0.55 % (w/v), respectively. Thereafter, a decrease in enzyme activity was observed with the increase in either the yeast extract or sucrose. This may be due to the inhibitory effect of both nitrogen and carbon source at higher concentrations. The interaction between inoculum size and age of inoculum indicated that inoculum level above ‘0’ level (5 %, v/v) showed the best response with 6 h old inoculum culture for the catalase production (Fig. 5b).

Further, response curves were also varied at different levels of BaCl$_2$ along the axis which describes that there is a significant interaction between inoculum size and BaCl$_2$. The response shown in Fig. 5c depicts the combined effect of above said variables in the fermentation media for catalase production (43,000 IU/mg of cells). A sharp increase in the catalase production was also achieved after 12 h of incubation by using computational tools as compared to that of 18 h at the flask level (Fig. 5d).

Validation of the model
Catalase production as envisaged by the final quadratic model along with the corresponding observed values suggests that there is a good agreement between the predicted and experimental values. The optimum values for each factor [sucrose (0.55 %, w/v), yeast extract (1.0 %, w/v), BaCl$_2$ (0.08 %, w/v), inoculum age (6 h), inoculum size (5 %, v/v) and fermentation time (12 h)] obtained by differentiation of quadratic model for achieving maximum catalase production were used for additional experiments to validate the accuracy of the model. The predicted optimal catalase production corresponding to these values was found to be 49,780 IU/mg of cells. The maximum catalase production of 48,900 IU/mg of cells obtained proves an excellent agreement between the predicted and experimental values, which further validates this statistical model. A considerable increase (118.11 %) in catalase production was obtained at the shake flask level using statistical tools.

Scale up studies
To further improve the catalase production on a pilot scale, the effect of some critical parameters such as agitation, air supply and fermentation time were investigated at bioreactor level. The key parameter in bioreactor studies i.e. the air supply was varied from 0.25 to 1.25 vvm to achieve the maximum enzyme yield. The optimum volume of air supply was found to be about 0.75 vvm with a dissolved oxygen concentration of 62.5 % saturation, at which maximum catalase production of 85,400 IU/mg of cells was obtained (Fig. 6a). Low biomass production and catalase synthesis at lower aeration rate (less than 0.75 vvm) was recorded due to oxygen limitation conditions. This major increase in catalase production might be due to continuous air supply to the bacterium because oxygen transfer rate is a critical factor that influences the overall cell metabolism. However, it was observed that high aeration rate above optimum level have caused metabolic shift in favour of cell growth which proved detrimental for enzyme synthesis. The low enzyme production at higher aeration beyond 0.75 vvm may also be due higher enzyme inactivation caused by irreversible oxidation of amino acid residues of the enzyme structure [30].
It was also observed that there was an increase in catalase production on increasing the rate of agitation up to 150 rpm (105,000 IU/mg of cells), and thereafter, enzyme production decreased significantly. An increase in dissolved oxygen concentration from 32.4 to 74.4 % saturation was also observed with enhanced agitation rate from 50 to 300 rpm (Fig. 6b). The higher growth and highest catalase synthesis was achieved at 150 rpm with a dissolved oxygen concentration of 70.4 % saturation. This might be due to continuous supply of air and even nutrient distribution facilitated by agitation. Two factors, namely oxygen supply and shear stress generally act in opposite direction, where oxygen supply enhances biomass production but shear stress lowers the enzyme productivity. The shearing effect preponderates the effect of oxygen supply at a higher agitation speed (>150 rpm) and induces cell disruption and enzyme inactivation. Hence, low biomass and low enzyme activity was observed at higher agitation rates beyond optimum level. Lower enzyme activity at a higher shear stress may also be due to its detrimental effect on enzyme structure [31].

Further, the effect of fermentation time was studied with an interval of 3 h up to 24 h of fermentation. It was found that the maximum catalase production
(105,000 IU/mg of cells) was achieved after 12 h of incubation in the bioreactor (Fig. 6c). The decrease in enzyme production by bacterial cells on reaching the stationary phase after 12 h can be ascribed to nutrient depletion. A significant enhancement in catalase production (105,000 IU/mg of cells) was achieved in the bioreactor
when compared to shake flask level (48,900 IU/mg of cells). This extraordinary enzyme production in a very short fermentation time of 12 h is a significant achievement, which has not been achieved by any method reported in the literature till date. A comparative analysis of methodologies involving catalase production from microorganisms (Table 4) clearly establishes that the developed methodology provides a simple and economical medium that employs only three constituents. Contrary to it, all the reported methodologies use complex multi component media for different microorganisms for synthesis of catalase [9, 13, 32].

Conclusions

One factor at a time traditional approach and response surface methodology was applied for tailoring nutritional and process requirements of Geobacillus sp. BSS-7. A hyperproduction of catalase was achieved by customizing media components (sucrose 0.55 %, w/v; yeast extract 1.0 %, w/v; BaCl2 0.08 %, w/v) and process variables (inoculum age 6 h; inoculum size 5 %, v/v and fermentation time of 12 h) at an optimum level. High similarity recorded between the predicted and experimental responses reflected the accuracy and applicability of statistical model used. A record 214.72 % increase in catalase production on implementing the bioprocess from the flask to bioreactor level proves the significance of multiple process variables in the studied fermentation process. The hyperproduction of catalase in a bioreactor is attributed to improved aeration and uniform distribution of nutrients. This is the first report on the hyper catalase production from any microbial source in a short incubation time using simple and economical media. The findings of this work would be useful for catalase production from Geobacillus sp. BSS-7 on a commercial scale to meet industrial demands.

Methods

Microorganism and culture conditions

Geobacillus sp. MTCC 5873, an extremophilic isolate of our laboratory (Enzyme Biotechnology Laboratory, Punjabi University, Patiala, INDIA), was grown in a medium (prepared in 50 mL distilled water) containing peptides digest of animal tissue (5.0 g/L), NaCl (5.0 g/L), beef extract (1.5 g/L) and yeast extract (1.5 g/L) for preparation of seed culture at 50 °C for 12 h on a rotary shaker inoculated with 5 % (v/v) inoculum culture. Further, a short term preservation of the isolate was carried on sterilized nutrient agar slants containing the aforementioned media supplemented with agar powder (15.0 g/L) by subculturing at fortnight intervals. The long term storage of the isolate was done at −80 °C in a 15 % glycerol solution.

Table 4 Comparison of intracellular catalase production from Geobacillus sp. BSS-7 with other microorganisms

| Microbial source | Production media (g/l)* | Process parameters | Catalase production Reference |
|------------------|-------------------------|--------------------|-----------------------------|
|                  | pH | Temp. (°C) | Time (h) | Inoculum size | Agitation (rpm)/stationary |                  |
| Geobacillus sp. BSS-7 | 7.0 | 60 | 12 | 5 % (v/v) | 150 | 105000 ± 2563 IU/mg Present study |
| Rhizobium radiobacter Strain 2-1 | (–) | 30 | 20 | (–) | 130 | 30420 ± 1083 IU/mg [8] |
| Serratia marcescens SYBC08 | CSTL (33.8), citric acid (30) | 7.0 | 30 | 40 | 4 % (v/v) | 400 | 20289 IU/ml [28] |
| Exiguobacterium oxidotolerans T-2-2 | Pep (8.0), YE (3.0), Sodium succinate (5.0), Aminolevulinic acid (2.0 mM), Tween 60 (1.0) | 7.5 | 27 | 24 | (–) | 60 | 16000 IU/ml [9] |
| Psychrobacter piscatorii | Pep (8), YE (3), NaCl (5), H2O2 (10 mM) | (–) | 27 | (–) | (–) | 12000 IU/mg [33] |
| Bacillus halodurans LBK 261 | Glu (3), YE (5), Pep (5), KH2PO4 (1), NaCl (1.5), NaNO3 (10), CaCl2 2H2O (1.7), FeSO4 7H2O (1.3), MnCl2 4H2O (1.51), ZnSO4 7H2O (0.25), H3BO3 (2.5), CuSO4 5H2O (0.12), Na2MoO4 2H2O (0.12), Co(NO3)2 6H2O (0.23), H2SO4 (2.5 ml) | 10 | 55 | (–) | (–) | 200 | 275 IU/g [13] |

Glu glucose; Pep peptone; YE yeast extract; CSTL corn steep liquor powder; Suc sucrose; (–) Data not available
Shake flask optimization of medium constituents and process parameters

The production of intracellular catalase was carried out in a fermentation medium (50 mL) containing the yeast extract (0.5 %, w/v) and glucose (1.0 %, w/v) having pH of 7.0 on a rotary shaker (Certomat M, Germany) at 100 rpm for 24 h at 50 °C, unless otherwise specified. The individual effect of various carbon sources (glucose, fructose, sucrose, maltose, lactose, starch, mannose, raffinose, galactose, inositol) at 0.5 % (w/v) concentration was investigated to study their effect on catalase synthesis and biomass production.

Various nitrogen sources (Urea, NH₄SO₄, NH₄NO₃, NaNO₃, KNO₃, meat extract, yeast extract, malt extract, beef extract, tryptone and peptone) at 0.5 % (w/v) concentration were also added in the medium to study their influence on enzyme and biomass production. The effect of concentration of selected carbon and nitrogen source was also investigated. The medium was also supplemented with different metal salts (CaCl₂, ZnSO₄, CuSO₄, MnSO₄, BaCl₂, MgSO₄, FeSO₄, NiSO₄, CoCl₂, HgSO₄, AgNO₃, KI, Na₂SO₃, and Na₂MoO₄) independently to study their influence on enzyme production. The effect of various bioprocess parameters such as pH of the medium, temperature, inoculum age and size, agitation rate, and fermentation time on catalase production was also investigated.

Statistical design for selection of critical variables

A central composite rotatable design (CCRD) with six critical variables having 42 runs was used to determine the optimum combination of variables by studying their response patterns [34]. The selection of critical components (Table 5) of the medium was made on the basis of observations recorded using one variable at a time classical approach as discussed in previous section. The statistical analysis of the response patterns was made using the Design Expert 7.0.3 statistical software (Stat-Ease Inc., MN, USA). The experimental design with values in actual and coded form using Design Expert Program is listed in Table 5. The results of the final design were validated using analysis of variance (ANOVA) along with with Fischer test by testing its significant effect (P < 0.05).

Bioreactor studies

Bioreactor studies were carried out on a laboratory scale stirred tank reactor of 1.5 L capacity with working volume of 1.0 L (Biolab, B. Braun, Germany) having Ruston type impeller with six blades. Different parameters such as aeration (0.25–1.25 vvm), agitation (50–300 rpm) and fermentation time (3–24 h) were varied to achieve maximum production of catalase. To control foam formation, sterilized olive oil (0.002 %, v/v) was added at the start of process system.

Recovery of catalase and enzyme assay

The biomass from the fermented broth was harvested by centrifugation (REMI, CPR-30 Plus) at 5000 g for 10 min at 4 °C for the extraction of intracellular catalase. The bacterial cells were disrupted (25 kHz, 10 min, pulse ON 25 s, OFF 05 s) by sonication (Vibra-Cell VCX 130, Newtown, USA) at 4 °C. The slurry was then centrifuged to separate the cell debris and supernatant thus obtained was analyzed for catalase activity. The estimation of catalase was carried out by the established method [35]. Catalase activity was calculated in terms of percent degradation of H₂O₂ and one unit (IU) of enzyme activity was expressed as an amount of enzyme that decomposes one micromole of H₂O₂ per minute under standard conditions.

Authors’ contributions

This study was designed and conceived by Dr. Balwinder Singh Sooch (BSS). Further, experimental work was performed by Mr. Baljinder Singh Kauldhar (BSK). Both the authors have written the research paper. Both authors read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

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