Abstract

The isolation of phytase using *Pichia Pastoris* under methanol/sorbitol co-feeding induction technique was investigated. The biological activity of extracellular phytase after optimization with co-substrates induction in 4 liters fermentor (NBS) increased to 13250 U/ml. This led to a 509 fold increases in comparison to the other type of phytase. This effect was studied via induction with sorbitol/methanol in fermentation by *Pichia pastoris* GS115 (Mut+) at 20 °C. The interference of by products; methylal, hexamine and (S)-(+)1,2-propanediol with release of phytase in *Pichia pastoris* under methanol induction were detected and cannot be repressed by methanol induction alone. The TLC was used for glycerin analysis under methanol/sorbitol induction and the results were lesser compare to that obtained during phytase production under methanol induction alone. This work showed the higher expression of heterologous proteins and by fed batch fermentation; the expression identified an advantage of producing a significant activity of phytase.

Practical applications

Plant derived products including sorbitol have been used as alternative medicines for the therapeutic treatment of various diseases, food supplements and could be used in many manufacturing processes. It serves as a culture media for bacteria, and helps to distinguish the pathogenic *E. coli* O157:H7 from its most other strains. Cells growing on methanol require high oxygen consumption. Sorbitol was used as an alternative cheap co-feeding for the production of proteins and is a non-repressing carbon source for AOX1 promoter with no effect on the level of r-protein at its induction phase. This report describes the isolation of phytase using *Pichia Pastoris* under methanol/sorbitol co-feeding induction techniques, and sorbitol showed to be a promising co-substrate, as it could enhance both cell growth and targeted protein productivity. This co-feeding and fed-batch induction technique was used for recombinant phytase production in a small and large scale production and the metabolites were analyzed.

Introduction

Sorbitol an isomer of mannitol can be isolated from the conversion of the carbonyl group of glucose to a hydroxyl group via the reduction reaction. This change allows the latter compound to be a carbohydrate classified as a polyol [1]. Sorbitol as a monosaccharide has four calories per gram and studies showed that it is a non-metabolite, since it is metabolically more inert than other saccharides [2,3]. It finds many applications in many manufacturing processes, as a pharmaceutical aid, toothpaste, sugar crystallization inhibitor and a dietary supplement [4]. In biotechnology, it is useful as a culture media for bacteria and allows to distinguish the pathogenic *Escherichia coli* O157:H7 from most other strains.
of E. coli [5]. In nature, sorbitol is dominantly found in various 
plants including algae and higher plants. For methanol 
induction alone is a high degree reductant releasing a larger 
amount of energy in form heat [6], and the process undergoes a 
significant effect upon a methanol induction phase at 
large scale. Cells growing on methanol, brings technological 
challenges as it requires high oxygen consumption, and the 
demand for safety precautions due to the use of a flammable 
and toxicity substrate [7]. Recent studies revealed that the use 
of methanol was cheap and common substrate at the time the 
P. Pastoris system was introduced [5,8]. Nowadays, sorbitol 
is an alternative available cheap co-feeding for the production of 
proteins and is a non-repressing carbon source for AOX1 
promoter as it does not affect the expression level of r-protein at 
its induction phase [1,2,4,11]. Many processes of fermentation 
are carried out following the method reported by [22,23]. On the other 
hand, the optimal temperature was measured at various 
temperatures using a 10mM buffer solution of Tris-HCl; pH 
7.5 mixed with sodium phytate and enzyme reactions. The 
enzyme reaction was initiated by the addition of the enzyme 
preparation and CaCl2 (1 mM). This process was 
followed by the measurement of the optimal pH of the activity 
of the enzyme utilizing a 25 mM solution of glycine-HCl; pH 
2.5, 0.1 mM solution of sodium acetate (pH 4.0-6.5) for 
purification [21], and 10 mM solution of Tris-HCl (pH 7.5-9.0). 
The determination of Km values of phytase was carried out following the method reported by [22,23]. On the other 
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preparation and CaCl2 (1 mM) for minute. The determination of Km, Vmax, Kcat and Kcat/Vmax was performed using both 
Michaelis-Menten formula and Lineweaver-Burk plot.

**Phytase kinetic**

The kinetic determination of phytase was performed using various concentrations of the substrate of ranging 
from 0.2 to 5 mM in 10 mM of the solution of Tris-HCl buffer solution; pH 7.5 with CaCl2 (1 mM). This process was 
followed by the measurement of the optimal pH of the activity of the enzyme utilizing a 25 mM solution of glycine-HCl; 
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**Small scale and high cell density fermentation**

The process of fermentation at small scale was initiated 
in batch cultivation whereby the transfer of 20 ml pre-
inoculum was performed aseptically to a volume of 200 ml 
sterile medium in an Erlenmeyer baffled flasks; 500 ml which 
contained \( V_0 = 100 \) ml production medium, at 200 rpm and 
28 °C. The desired medium for laboratory scale experimental 
procedure contained 10 g/l of yeast extract, 20 g/l of glucose, 20 g/l of agar and 
0.1 g/l of Zeocin were utilized for the strain P. pastoris GS115 
growth that was transformed with recombinant pPICZαA-
phytase in presence of AOX1 promoter and incubated at 30 °C during 48 hours. In addition, a colony was inoculated into a 
volume of 5 ml YPD liquid medium containing Zeocin (100 μg/ 
ml), and grown via shaking at 250 rpm at 30 °C overnight. The 
culture of seed was then inoculated into the medium of BMGY 
containing 10 g/l of yeast extract, 20 g/l of peptone, 13.4 g/l of 
YNB, 4×10⁻⁵ g/l of biotin, 10 g/l of glycerol and a solution of 
0.1 M of potassium phosphate buffer whose is pH 6.0.

**Material and methods**

**Growth of a P. Pastoris GS115-pPICZαA-phytase**

The prepared medium of YPD containing 20 g/l of peptone, 
10 g/l of yeast extract, 20 g/l of glucose, 20 g/l of agar and 
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sterile medium in an Erlenmeyer baffled flasks; 500 ml which 
contained \( V_0 = 100 \) ml production medium, at 200 rpm and 
28 °C. The desired medium for laboratory scale experimental 
procedure contained 10 g/l. On the other hand, high cell 
density fermentation, P. pastoris was performed in batch mode
through sterilizing a basal salts medium (2.3 litres) in the NBS BioFlo 415 Benchtop SIP fermentor. The ingredients of BSM medium I: (g/l) CaSO₄·2H₂O; 0.46, K₂SO₄; 9.1, MgSO₄·7H₂O; 7.45, KOH; 2.06, Glycerol; 40, Histidine; 0.4 and 26.7 ml/l of H₃PO₄. The addition of the adjusted pH of the medium (pH = 4.8) with ammonium hydroxide (25%) and PTM1 (6 ml) trace salts/litre of basal salts medium was carried out aseptically (Clare, et al. 1991). The main ingredients of PTM1 solution: 6 salts/litre of basal salts medium was carried out aseptically medium l L: (g/l) CaSO₄·2H₂O; 0.46, K₂SO₄; 9.1, MgSO₄·7H₂O; 7.45, KOH; 2.06, Glycerol; 40, Histidine; 0.4 and 26.7 ml/l of H₃PO₄. The optimization of the repeated fed batch cultivation for the production of phytase at high cell density cultivation in Pichia Pastoris was performed using methanol/sorbitol co-feeding induction technique. The fed-batch cultivation time was converted to repeated fed-batch through harvesting 90% of the reactor volume, and recharging with fresh BSM medium.

Biomass and total protein analysis

The control of cell biomass was performed by the method of turbidometric determination or light scattering phenomenon. The Optical density (OD₆₀₀nm) was utilized to represent the level of cells corresponding to the quantity of the scattered light by the culture broth. The turbidity or light scattering was measured at the wavelength of 600nm using the spectrophotometer (Hitachi U-2000). The colorimetric Bio-Rad protein assay based on the Bradford method was used to determine the total amount of protein in culture liquid. This assay focuses on the color change of Coomassie Brilliant Blue G-250 dye while binding to primarily basic such as arginine and amine group of the protein. The quantity of protein was measured in the format of micro-plate, as determined by the assay kit and the spectrum absorbance peaks recorded at 595 nm using an ELISA reader (Labsystems Multiskan MS).

GC-MS analysis of extract

The analytical measurement using GC-MS for extract analysis from repeated fed batch fermentation of recombinant Phytase (pPICZA-phytase) was performed at high cell density cultivation in Pichia Pastoris under methanol/sorbitol co-feeding induction with in methanol. The latter procedure was carried out using a Clarus 500 Perkin Elmer gas chromatography equipped with an Elite-5 capillary column of phenyl: 5%, dimethyl polysiloxane; 95% (30 nm X 0.25 mm ID X 0.25 μmfd) and mass detector turbo-mass gold which was performed in El mode. The inert helium gas was used as the carriers at a flow rate of 1ml/min, and the injector was operated at 290 °C. The oven temperature condition was set as follows; 50 °C at 8 °C/min to 200 °C (5 min) at 7 °C/min to 290 °C (10 min). The data analysis of mass spectrum (GC-MS) was treated by means of the database of National Institute Standard and Technology (NIST), having more than 62,000 patterns. The mass spectrum of the unknown analyte was compared with the spectrum of the known sample stored in the (NIST) library and the name, molecular weight and structure of the components of the test materials were ascertained.

### Results and discussion

During glycerol batch phase, a culture grows whereas the level of dissolved oxygen (DO) lowers and it is in line with the work done by Carly, et al. [24]. When the amount of glycerol in the medium was totally used the DO rose significantly. These results showed the final step of the starting batch phase, and the second step of glycerol 98 % fed-batch was initiated (Figure 1). The method of continuous cultivation was also carried out first to maintain the volume by replacing the old medium by fresh nutrient medium. Its role concentrates on the maintenance of a constant product quality as well to minimize the release of by-products formed during methanol induction alone. The results showed that the repeated fed-batch culture bearing an external separator of phytase formation at high cell density cultivation in Pichia pastoris under methanol/sorbitol co-feeding induction techniques enhanced the activity at about 49 times in comparison to methanol induction alone in a reactor (7 liters) (Figure 2). The volume was maintained and by the time the fermentation process reached a certain stage after which is not effective at all, 90% of medium was taken off from the vessel and replaced by fresh nutrient medium. In this case no culture was removed until the final stage of the batch, while the decrease in volume led to an increase in the productivity. An effective repeated fed-batch process under methanol/sorbitol feed as carbon source indicated a significant increase in OD₆₀₀nm about 5.6 times in comparison with that methanol induction alone. The optimization of phytase formation under methanol/sorbitol co-feeding techniques at lower temperature enhanced the activity at high level [25]. Heterologous expression of phytase in P. pastoris at low temperatures (2 °C) minimized extracellular proteolysis and

**Table 1: Parameters for High Cell Density Fermentation of Pichia pastoris (Pichia Fermentation Process Guidelines, Invitrogen).**

| Parameter | Reason |
|-----------|--------|
| Temperature (28 - 20 °C) | To get biomass and protein concentration |
| Dissolved oxygen (> 20%) | P. pastoris needs oxygen to metabolize glycerol and methanol. |
| pH (4.8) | Important when secreting protein into the medium and for optimal growth. Adjusted using 25% NH₄OH / 88% H₃PO₄ |
| Agitation (200 to 800 rpm) | Maximizes oxygen concentration in the medium. |
| Aeration (0.1 to 1.0vvm for glass fermenters) | Maximizes oxygen concentration in the medium which depends on the vessel. |
| Antifoam (the minimum needed to eliminate foam) | PPG was used to a minimum level because excess foam may cause denaturation of phytase and it also reduces headspace. |
| Carbon source (variable rate) | Must be able to add different carbon sources at different rates during fermentation |
Phytase enzyme kinetic (Kcat/Km) that shows how efficiently an enzyme converted a substrate into a product was 4.62*10^7 M^-1s^-1, whereby the substrate declined from 5 mM to 0.2 mM. In this process, 50% reduction time was reached after 3 minutes of reaction rate which is little different from a theoretical upper limit of 10^9 - MS [28]. An enzyme that works close to this value is termed superefficient and also its reduction time is reached after a period of 2.5 minutes. Other report showed that the expression of an enzyme at lower temperature is applied to promote the yield in P. pastoris, although the fermentation period takes a long time at higher temperature. The results from this report indicates that growing a culture first at 28 °C during Glycerol Batch Phase (GBP) increased the level of biomass prior to induction time. The higher activity was achieved within lesser cultivation time compare to the other studies. Many reports on phytase production showed that phytases with broader substrate generally had low specific activities. Despite considerable economic interest, low yield and high cost of enzyme formation are the main limiting factors in the use of phytase in animal diet. An important idea stresses on the point that sorbitol lowers the accumulation of toxic metabolite-formaldehyde in fermentation process using Pichia pastoris (Mut+) by repressing the formaldehyde dissimilatory pathway [29]. The obtained glyc erin was the only byproducts produced during phytase formation; as shown via GC-MS data. On the other hand, the following compounds such as dimethyl-Aetalacetal-of-formaldehyde (methylal), methenamine (Hexamine) were confirmed as by products coming from the reaction of formaldehyde and ammonia. Another compound that was present consists of (S)-(+)-1,2-propanediol which is a product of glyc erin and cannot be repressed by the use of methanol only (data not shown). The latter compounds are interfering with phytase production in Pichia pastoris under the induction of methanol. The results obtained from TLC analysis showed that the concentration

![Figure 1: Growth curve analysis during Glycerol Batch and Fed-Batch Phases.](https://doi.org/10.29328/journal.abb.1001024)

![Figure 2: Phytase activity under sorbitol/methanol induction.](https://www.heighpubs.org/hjb)

![Figure 3: Evaluation of optimum temperature for recombinant Pichia pastoris expressed phytase (A), Evaluation of optimum pH for recombinant Pichia pastoris expressed phytase (B). Lineweaver Burke plot of recombinant Pichia pastoris expressed phytase (C) and the measurement of half time required for substrate consumption (D).](https://www.heighpubs.org/hjb)
of glycerin obtained under methanol/sorbitol induction technique was smaller than that produced during phytase formation under methanol induction. This indicates that there was too much production of formaldehyde under methanol induction process. Sorbitol as a low degree reductant, non-repressing source of carbon for pAOX1 and energy supplier reduces oxygen consumption rate by increasing heat or methanol usage so that this alcohol may be strongly utilized for phytase production. Co-feeding of sorbitol minimizes the oxidation flux in the peroxisome, oxygen consumption and heat production as its heat of reaction is highly less than that of methanol. These parameters are of high importance for high cell-density cultures using recombinant P. Pastoris strains in case of large scale formation.

Conclusion

During this study, the mating factor (-factor) preproleader had been considered for the secretory expression of phytase whereby a modified plasmid for phytase expression had been constructed. This process was performed through putting stop a codon before polyhistidine and C-my epitope tag so that none nature factor from recombinant phytase could be removed. The enzymatic reactions of sorbitol and the impact of glycerin on recombinant protein production are very important tools for portion determination of metabolism that could benefit from future studies. The process of phytase recombinant in presence of sorbitol/methanol is limited by knowledge gaps but this hypothetical process may be discussed by SBO (Systems Biology Ontology) expression omitted process by: Sink metabolism to know how to predict what will occur in organic reaction mechanism and FBA (Flux Balance Analysis) to predict cell growth with different carbon sources. Research to expand our understanding of sorbitol metabolism and bio-chemistry needs to be understood whereby the technique of integrating and establishing biochemical knowledge into computational reconstructions is the future perspectives.

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