High-level production of recombinant HBcAg virus-like particles in a mathematically modelled P. pastoris GS115 Mut+ bioreactor process under controlled residual methanol concentration

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Abstract
Recombinant hepatitis B core antigen (HBcAg) molecules, produced in heterologous expression systems, self-assemble into highly homogenous and non-infectious virus-like particles (VLPs) that are under extensive research for biomedical applications. HBcAg production in the methylotrophic yeast P. pastoris has been well documented; however, productivity screening under various residual methanol levels has not been reported for bioreactor processes. HBcAg production under various excess methanol levels of 0.1, 1.0 and 2.0 g L⁻¹ was investigated in this research. Results indicate that, under these particular conditions, the total process and specific protein yields of 876–1308 mg L⁻¹ and 7.9–11.2 mg gDCW⁻¹, respectively, were achieved after 67–75 h of cultivation. Produced HBcAg molecules were efficiently purified and the presence of highly immunogenic, correctly formed and homogenous HBcAg-VLPs with an estimated purity of 90% was confirmed by electron microscopy. The highest reported HBcAg yield of 1308 mg L⁻¹ and 11.2 mg gDCW⁻¹ was achieved under limiting residual methanol concentration, which is about 2.5 times higher than the next highest reported result. A PI-algorithm-based residual methanol concentration feed rate controller was employed to maintain a set residual methanol concentration. Finally, mathematical process models to characterise the vegetative, dead and total cell biomass (Xv, Xd and X), substrate (Glycerol and Methanol) concentration, reactor volume (V), and product (HBcAg) dynamics during cultivation, were identified. A rare attempt to model the residual methanol concentration during induction is also presented.

Keywords  Hepatitis B core antigen (HBcAg) • Pichia pastoris • Fed-batch bioreactor • Cultivation process modelling • Residual methanol PI control

Abbreviations
HBcAg  Hepatitis B core Antigen
DO  Dissolved Oxygen
OD  Optical Density at 590 nm
DCW  Dry Cell Weight
WCW  Wet Cell Weight
GMP  Good Manufacturing Practice
PID  Proportional, Integral and Derivative control parameters
PLC  Programmable Logic Controller
HPLC  High-Performance Liquid Chromatography
GC  Gas Chromatography
SCADA  Supervisory Control and Data Acquisition interface
Gly  Glycerol
MeOH  Methanol

List of symbols
Xv  Vegetative cell biomass concentration [g L⁻¹]
Xd  Dead (non-vegetative) cell biomass concentration [g L⁻¹]
X  Total cell biomass concentration [g L⁻¹]
S  Substrate concentration [g L⁻¹]
P  Product concentration [g L⁻¹]
V  Cultivation medium volume [L]
τI  Integral time constant [min.]
Kp  Proportional gain parameter [L² g⁻¹ h⁻¹]
Introduction

Hepatitis B virus core antigen (HBcAg) is a protein subunit of the hepatitis B virus icosahedral core shell. Intracellularly expressed in heterologous production systems such as bacteria, yeasts, plants or mammalian cell cultures, and the recombinant HBcAg molecules self-assemble in non-infectious virus-like particles (VLPs) [1]. These particles, being highly immunogenic and known to induce significant T-cell responses and high antibody titres in the exposed host organism [2], have been extensively studied and used in vaccine development [3]. Consequently, the efficient production of purified HBcAg-VLPs is a key aspect for both research and commercial applications.

HBcAg-VLPs have been recognised as a promising VLP carrier platform since the 1980s; naturally, they have also been extensively studied [4, 5]. Recent reviews illustrate that HBcAg-VLPs have successfully been employed as epitope carriers in different vaccine candidates against a myriad of diseases, contrary to its fraternal HBV surface antigen (HBsAg), still lack successful commercial examples [3, 6]. A promising HBcAg-VLP-based commercial vaccine candidate against malaria (MalariVax) was in development; however, it was discontinued after failing the sporozoite challenge [7]. On the contrary, a malaria vaccine candidate (Mosquirix™) based on HBsAg-VLPs was recently approved by the WHO for general use in Africa [8]. Several HBcAg-VLP-based vaccine patent applications can also be found against various diseases: melioidosis [9], hepatocarcinoma, and hepatitis B [10, 11], and HPV [12]. Despite the lack of successful commercial HBcAg-VLP-based vaccines, this platform still stands out as one of the frontrunners for future vaccine design.

A wide range of expression systems are available for recombinant protein production and each of them has their own advantages. The choice of a production platform is in large part dictated by the properties of the target protein—its structure and biological activity [13]. For the past 20–30 years, the P. pastoris expression system has been successfully employed for the production of various recombinant proteins for both research and industrial applications. Some of the main characteristics that make this methylotrophic yeast so suited for foreign protein expression include easy genetic manipulation, high-frequency DNA transformation, cloning by functional complementation, high levels of intra- and extracellularly expressed protein, and the ability to perform higher eukaryotic protein modifications (glycosylation, disulphide bond formation, and proteolytic processing) [14]. Also, relatively low levels of native secreted proteins allow for the use of simple purification strategies of the secreted recombinant proteins. When the economic aspects (high cell growth in minimal media and high product stability in prolonged processes) are taken into consideration, together with the powerful genetic techniques available, P. pastoris is clearly the system of choice for heterologous protein expression [15].

The two-stage cultivation on glycerol and methanol associated with Invitrogen’s “Pichia Fermentation Process Guidelines” is well documented and present in most AOX1-promoter-based P. pastoris cultivation strategies. However, recent trends advocate for a move away from standard
protocols towards a more conceptual approach, which allows for the development of process-specific strategies tailored both to the specific combination of product/genetic construct and the characteristics of the bioreactor equipment [16]. The standardized approach employs a constant substrate feed rate during methanol induction. Since cell biomass concentration keeps increasing, but the substrate feed rate remains constant, the amount of substrate per biomass (g DCW·h⁻¹) gradually decreases, resulting in a decreasing specific growth rate and, consequently, productivity [16].

A myriad of different methanol feeding strategies have been reported with excellent recombinant product yields [16–18]. Two distinct groups can be identified—co-feeding and -stat strategies. Co-feeding strategies aim to supplement the methanol feed with additional substrates (glycerol, sorbitol, mannitol, etc.), whereas -stat strategies attempt to control the methanol feed in a way that a cultivation parameter (μ, DO, methanol concentration) remains at a pre-set value. One of the more popular approaches is the methanol-stat strategy, in which the residual methanol concentration is controlled at a set-point value during the methanol induction phase [17]. There are reports of this strategy leading to increased recombinant protein yields for several products; however, no reports can be found on the effect this strategy has on HBcAg production in P. pastoris.

The residual methanol concentration in AOX1-promoter-regulated P. pastoris systems is a critical factor to control, as it directly influences the production and proteolytic degradation of recombinant proteins, cell growth, lysis, and oxygen transfer; thus, an optimal methanol feeding strategy is crucial to maximize product yields [19]. Residual methanol concentrations between 2.0 and 3.5 g L⁻¹ are considered optimal for protein production with P. pastoris [19]; however, some authors report success with concentrations in 4.0–6.0 g L⁻¹ range [20, 21]. Monitoring and controlling the residual methanol concentration in the cultivation broth are crucial; therefore, a suitable methanol sensor and a methanol feed rate control algorithm should be used to ensure adequate methanol concentration control at a selected set-point.

Two types of control systems for residual methanol concentration control can be distinguished—open loop and closed loop. The open-loop systems do not require any online measurements (feedback); however, closed loop systems do. In this particular case, the methanol sensor on-line signal can be used as feedback for methanol feed rate control. Additionally, a control algorithm is necessary to successfully control the residual methanol concentration around set-point value with minimal deviation. For certain residual methanol level control, PI- [22] or PID- [23] control algorithms have been successfully employed.

Stringent requirements for producing biopharmaceutical products and their marketing in the corresponding regions, in accordance with good manufacturing practice (GMP) guidelines, are imposed by the EU [24], USA [25], and other developed countries. Since biotechnological microorganism cultivation processes are highly complex and their productivity is sensitive to optimal process parameter control, the above-mentioned directives require that, to ensure a high level of product quality and process repeatability (safety), advanced microorganism culture process control techniques must be employed. It is also mentioned that model-based methods facilitate an in-depth understanding of the process and expedite the performance of the procedures of cultivation process validation and revalidation [26]. Mathematical modelling of recombinant P. pastoris cultivation processes is well described; however, there are few examples modelling residual methanol concentrations during induction [27–31].

In the present work, we establish a P. pastoris cultivation procedure for high-level expression of HBcAg in a defined medium under various residual methanol levels controlled by the PI-feed rate control algorithm. A previously reported purification method [32] was employed with minor modifications, leading to superior results and producing HBcAg-VLPs with an estimated purity of 90%. Mathematical modelling was used to tune the Kp parameter for residual methanol PI-control algorithm. Suitable mathematical models and model parameters to model biomass growth, substrate (glycerol and methanol) consumption, and HBcAg formation were successfully identified.

**Materials and methods**

**Construction of an expression vector and selection of clones**

The construction of the expression vector pPIC-HBc and the selection of an HBc producer clone was described in [32].

**Experimental conditions**

Cultivation processes were carried out using a recombinant *Pichia pastoris* GS115 Mut⁺ strain. Batch cultivation and feeding media solutions used in this study were prepared according to the “*Pichia fermentation process guidelines*” by Invitrogen Corporation [33]: 1.9 l of Basal Salts Medium (26.7 ml L⁻¹ H₂PO₄ 85%, 0.93 g L⁻¹ CaSO₄, 18.2 g L⁻¹ K₂SO₄, 14.9 g L⁻¹ MgSO₄·7H₂O, 4.13 g L⁻¹ KOH, 40.0 g L⁻¹ glycerol and 4.35 ml L⁻¹ PTM₁ trace-element solution (0.02 g L⁻¹ H₂BO₃, 5 ml L⁻¹ H₂SO₄ 98%, 6.0 g L⁻¹ CuSO₄·5H₂O, 0.08 g L⁻¹ NaI, 3.0 g L⁻¹ MnSO₄·H₂O, 0.2 g L⁻¹ Na₂MoO₄·2H₂O, 0.5 g L⁻¹ Ca₃(PO₄)₂·2H₂O, 20.0 g L⁻¹ ZnCl₂, 65.0 g L⁻¹ FeSO₄·7H₂O, 0.2 g L⁻¹ biotin), was inoculated with 100 mL of inoculum grown in BMGY medium (10.0 g L⁻¹ yeast extract, 20.0 g L⁻¹ peptone, 100 mM...
potassium phosphate buffer, pH 6.0, 13.4 g L\(^{-1}\) yeast nitrogen base, 10.0 g L\(^{-1}\) glycerol, and 0.0004 g L\(^{-1}\) biotin) at 30 °C for 18–22 h in a shake flask at 250 RPM. Two feeding solutions were used—glycerol fed-batch solution (50% glycerol, 12 mL L\(^{-1}\) PTM, and methanol fed-batch solution (100% methanol, 12 mL L\(^{-1}\) PTM,).

The bioreactor vessel was filled with distilled water and sterilized at 121 °C for 30 min. BSM, BMGY, and glycerol fed-batch solutions were autoclaved separately at 121 °C for 30 min. PTM trace element and methanol fed-batch solutions were sterilized by filtration through a 0.2 µm filter.

Fermentations were carried out in a 5 l (2–4 l working volume) fully automated bench-top fermenter (Biotechniskais centrs, EDF-5.4/BIO-4, Latvia) schematically illustrated in Fig. 1. A calibrated pH sensor probe (Hamilton, EasyFerm Bio, Switzerland) was used to measure the medium pH during cultivation. Before the start of the cultivation process, the fermentation medium pH was adjusted at 5.0 ± 0.1 using a 28% NH\(_4\)OH solution, which was also used to maintain the set pH value during fermentation. The temperature was controlled at 30.0 ± 0.1 °C. The dissolved oxygen level was measured with a calibrated DO probe (Hamilton, OxyFerm, Switzerland) and kept above 30 ± 5% during fermentation by increasing the stirrer speed (200–1000 RPM) or inlet air enrichment with pure O\(_2\). Constant air or air/oxygen mixture at a flow rate of 3.0 slpm was maintained during all processes. A condenser was used to condense the moisture from outlet gasses and excessive foam formation was controlled by adding antifoam 204 (Sigma) when necessary.

Substrate feed solutions were pumped using a pre-calibrated high-accuracy peristaltic pump (Longer-Pump, BT100–2 J, China).

The cultivations were initiated with a glycerol batch phase. After 18–24 h, all batch glycerol is consumed and glycerol fed-batch solution is fed into the reactor at a rate of 0.61 mL min\(^{-1}\) for 4 h and until an optical density of

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**Fig. 1** Schematic diagram of the bioreactor and control architecture
100–120 is reached. Then, the feed substrate is switched to methanol and the solution is fed into the reactor at a rate of 0.12 mL min⁻¹ for 5 h, then at a rate of 0.24 mL min⁻¹ for 2 h, and finally at a rate of 0.36 mL min⁻¹ up until 40 h of cultivation. At 40 h, a direct residual methanol concentration control using a PI-algorithm-based feed rate control is initiated at a selected set-point and carried out until the end of the process.

**Analytical measurements**

Cell growth was observed by off-line measurements of the culture optical density (OD) at a wavelength of 590 nm (GRANAT, KFK-2, St. Petersburg, Russia). Wet cell weight (WCW) and dry cell weight (DCW) measurements were determined gravimetrically. Biomass samples were placed in pre-weighted Eppendorf® tubes and centrifuged at 15,500 g for 5 min. Afterwards, the supernatant was discarded, and the cells resuspended in distilled water and centrifuged once more. The liquid phase was discarded and the remaining wet cell biomass weighted. Afterwards, samples were dried at 105 °C until constant weight was reached and the dry cell biomass weighted. The DCW measurement was selected as the most accurate cell biomass assessment with an average error of ±1.0 g L⁻¹ [34].

Glycerol and methanol concentrations were measured off-line using, respectively, HPLC (Agilent, 1100 HPLC) and GC (Agilent, 6890 N GC).

Protein samples taken during cultivation were analysed using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), with a 5% stacking and 15% separating polyacrylamide gel (PAAG), according to standard protocols. To visualize the separated protein bands, the gels were stained with 0.4% Coomassie Brilliant Blue (CBB) G-250 dye.

The signal around the set methanol concentration value makes it possible for the control parameters, \( K_p \) and \( \tau_I \), to remain constant throughout the cultivation. The \( K_p \) used was within the range of 0.02–0.05 [L² g⁻¹ h⁻¹], and, as it will be demonstrated further, was dependent on selected residual methanol concentration; the constant \( \tau_I = 10 \) min was used in the experiments.

**Mathematical modelling**

The macrokinetic model presented by Ren et al. [28] was employed to model cell biomass (\( X_c \), \( X_d \), and \( X \)) and glycerol concentrations (\( S_{\text{Gly}} \)) during HBcAg-producing \( P. \) *pastoris* cultivations. A non-monotonically increasing function equation was chosen to model the methanol concentration \( (\text{S}_{\text{MeOH}}) \); and the product accumulation dynamics (\( P_{\text{HBcAg}} \)) were estimated using a Luedeking–Piret model equation. MATLAB® 2018a was used for mathematical modelling.
Macrokinetic models describing the balances of energy and intracellular substances for both glycerol and methanol uptake phases were constructed in matrix form according to Ren et al. [28].

For the glycerol uptake phase

\[
\begin{pmatrix}
1 & K_{1,Gly} & 0 & 0 \\
-4K_{2,Gly} & 5 & -2 \\
-1 & K_{1,Gly} & -K_{2,Gly} & -\frac{1}{\tau_{\text{Gly}}} & 1 & 2P/O_{\text{Gly}} \\
1 & 0 & 0 & -1 & 0
\end{pmatrix}
\begin{pmatrix}
q_G \\
\mu \\
q_{Ac} \\
q_{O_2}
\end{pmatrix}
= \begin{pmatrix}
q_{Gly} \\
0 \\
m_{\text{ATP}_{\text{Gly}}} \\
0
\end{pmatrix}
\tag{6}
\]

The respective biomass growth rate (\(\mu\)) was determined using Eq. 6. To account for the characteristic lag phase in \(P.\) \(\text{pastoris}\) cultivations, the specific glycerol uptake rate (\(q_{\text{Gly}}\)) was first described using the metabolic regulator model and afterwards with the well-known Monod model. The switch between the two models was initiated by the minimum function

\[
\frac{dq_{\text{Gly},R}}{dt} = k_{1}(q_{\text{Gly}} + q_{\text{min,Gly}}) + (-k_{2} - \mu)q_{\text{Gly},R}
\tag{7}
\]

\[
q_{\text{Gly,M}} = \frac{q_{\text{max,Gly}} * S_{\text{Gly}}}{K_{S,\text{Gly}} + S_{\text{Gly}}}
\tag{8}
\]

\[
q_{\text{Gly}} = \min \{q_{\text{Gly,R}}, q_{\text{Gly,M}} \} * M_{\text{Gly}}.
\tag{9}
\]

For the methanol uptake phase

\[
\begin{pmatrix}
\frac{3}{1-\varphi} & \frac{3}{1-\varphi} - K_{1,\text{MeOH}} & 0 & 0 \\
\frac{6\varphi}{5\varphi+1} & \frac{6\varphi}{5\varphi+1} - K_{1,\text{MeOH}} - 4K_{2,\text{MeOH}} & 5 & -2 \\
-1 & -3K_{1,\text{MeOH}} - K_{2,\text{MeOH}} - \frac{1}{\tau_{\text{exp}}} & 1 & 2P/O_{\text{MeOH}} \\
1 & 0 & 0 & -1 & 0
\end{pmatrix}
\begin{pmatrix}
q_G \\
\mu \\
q_{Ac} \\
q_{O_2}
\end{pmatrix}
= \begin{pmatrix}
q_{\text{MeOH}} \\
0 \\
m_{\text{ATP}_{\text{MeOH}}} \\
0
\end{pmatrix}
\tag{10}
\]

The respective biomass growth rate (\(\mu\)) values were determined by solving the equation. The specific methanol uptake rate (\(q_{\text{MeOH}}\)) was calculated according to the non-monotonically increasing function first described by Jackson and Edwards [36]

\[
q_{\text{MeOH}} = \frac{q_{\text{max,MeOH}} * S_{\text{MeOH}}}{K_{S,\text{MeOH}} + S_{\text{MeOH}} + (S_{\text{MeOH}}/K_{i})} * M_{\text{MeOH}}.
\tag{11}
\]

As recombinant product biosynthesis under the AOX1 promoter takes place only during the methanol induction phase, product accumulation was modelled only during this phase and is described by the following equation:

\[
q_P = \mu_{\text{MeOH}} * Y_{\text{PX}}.
\tag{12}
\]

Finally, a bioreactor model was constructed to estimate the vegetative, dead and total cell biomass (\(X_v, X_d, \) and \(X,\) respectively), and substrate (\(S\) and product (\(P\)) concentrations. A model describing the liquid media volume (\(V\)) is also included, as the reactor volume directly influences the concentration of all the aforementioned parameters

\[
\frac{dX_v}{dt} = \mu X_v - K_d X_v - \frac{F_S + F_{\text{NH3}} + F_{\text{AF}} - F_{\text{exp}}}{V} X_v
\tag{13}
\]

\[
\frac{dX_d}{dt} = K_d X_v - \frac{F_S + F_{\text{NH3}} + F_{\text{AF}} - F_{\text{exp}}}{V} X_d
\tag{14}
\]

\[
X = X_v + X_d
\tag{15}
\]

\[
\frac{dS_{\text{Gly}}}{dt} = -q_{\text{Gly}} X_v - \frac{F_S + F_{\text{NH3}} + F_{\text{AF}} - F_{\text{exp}}}{V} S_{\text{Gly}} + \frac{F_S}{V} S_{F,\text{Gly}}
\tag{16}
\]

\[
\frac{dS_{\text{MeOH}}}{dt} = -q_{\text{MeOH}} X_v - \frac{F_S + F_{\text{NH3}} + F_{\text{AF}} - F_{\text{exp}}}{V} S_{\text{MeOH}} + \frac{F_S}{V} S_{F,\text{MeOH}}
\tag{17}
\]

\[
\frac{dP}{dt} = q_P X_v - \frac{F_S + F_{\text{NH3}} + F_{\text{AF}} - F_{\text{exp}}}{V} P
\tag{18}
\]

\[
\frac{dV}{dt} = F_S + F_{\text{NH3}} + F_{\text{AF}} - F_{\text{smpl}} - F_{\text{exp}}.
\tag{19}
\]

**Model parameter sensitivity analysis**

To evaluate the model parameters that have the most impact on modelling results, a model parameter sensitivity analysis procedure was carried out. The value of one selected parameter was increased by 5%, while others remained constant, and the effect on modelled variables was estimated by evaluating the difference of the new and reference root-mean-square errors (RMSE_{5\%} and RMSE_{0}, respectively) using the following formula:

\[
\text{Err}_{5\%} = \frac{\text{RMSE}_{5\%} - \text{RMSE}_{0}}{\text{RMSE}_{0}} * 100\%.
\tag{20}
\]

**Downstream processing of HBcAg**

To compare the outcome of recombinant HBcAg-VLPs produced under different cultivation conditions, a
well-established protocol from Freivalds et al. with some modifications was used [32].

Briefly, 4.0 g of frozen yeast cells were first suspended in 16 mL of lysis buffer (20 mM Tris–HCl, 100 mM NaCl, 0.1% Triton X-100, pH 8.0) and disrupted with a French press (4 cycles, 10’000 psi). The supernatant was separated by centrifugation (30 min, 15,500g, at 15 °C) and pH was adjusted to 8.0 using 0.5 M NaOH. The supernatant was then incubated at 65 °C for 1 h and subsequently centrifuged (15 min, 15,500g). Solid ammonium sulfate was added to the solution up to a saturation of 40%, incubated at 4 °C for 30 min, and centrifuged once more (15 min, 15,500g). The precipitate was then dissolved in 2 mL of 20 mM Tris–HCl buffer (pH 8.0) and loaded onto a gel filtration Sepharose 4 Fast Flow (50 mL bed volume) column in 20 mM Tris–HCl buffer (pH 8.0) and loaded onto an anion-exchange Fractogel DEAE column (20 mL bed volume) connected to an ÄKTA chromatography system (Amersham Biosciences, UK). The fractions containing HBcAg-VLPs were isolated, pooled, and loaded onto an anion-exchange Fractogel DEAE column (20 mL bed volume) connected to an ÄKTA chromatography system (Amersham Biosciences, UK). The column was preequilibrated with 20 mM Tris–HCl pH 8.0, 100 mM NaCl, and run at 5.0 mL min⁻¹. Column-bound proteins were eluted by a linear gradient with the same buffer containing 1 M NaCl.

Purity and quality of HBC VLPs were assessed by SDS-PAGE and electron microscopy and the concentration established by the Bradford assay method.

Results

HBcAg production

Scientific literature lacks successful examples of bioreactor cultivations at increased residual methanol concentrations for HBcAg production with P. pastoris; however, it is standard practice when producing HBcAg in flasks. Whether induction at increased residual methanol concentrations can improve recombinant HBcAg biosynthesis yields in P. pastoris was the main hypothesis investigated during this research. A series of five experiments was performed and three residual methanol levels during the induction phase were investigated — 0.1 (limiting), 1.0 and 2.0 g L⁻¹. The experiments can be summarized as follows:

- Exp. 1—according to the Invitrogen protocol; after adaptation to growth on methanol (process 30th hour), constant methanol feed solution addition with the rate of 0.36 mL min⁻¹.
- Exp. 2—after adaptation to growth on methanol (process 32th hour), constant methanol feed solution addition with the rate of 0.4–0.5 mL min⁻¹.
- Exp. 3—after adaptation to growth on methanol (process 40th hour), methanol feed rate control according to the PI algorithm, methanol set-point — 1.0 g L⁻¹; \( K_p = 0.05 \) L² g⁻¹ h⁻¹, \( \tau_I = 10 \) min.
- Exp. 4—after adaptation to growth on methanol (process 40th hour), methanol feed rate control according to the PI algorithm, methanol set-point — 2.0 g L⁻¹; \( K_p = 0.05 \) L² g⁻¹ h⁻¹, \( \tau_I = 10 \) min.
- Exp. 5—after adaptation to growth on methanol (process 40th hour), methanol feed rate control according to the PI algorithm, methanol set-point — 2.0 g L⁻¹; \( K_p = 0.02 \) L² g⁻¹ h⁻¹, \( \tau_I = 10 \) min.

The biomass growth, methanol concentration, methanol feed rate, and HBcAg accumulation dynamics during these experiments are presented in Fig. 2.

Under various residual methanol levels, differing amounts of HBcAg (858–1308 mg L⁻¹) were recovered after 67–75 h of cultivation (40–45 h of induction). Interestingly, opposite to the reports of increased residual methanol levels promoting other recombinant product biosynthesis, the highest HBcAg concentration of 1308 mg L⁻¹ was produced in Exp. 1, which was performed in accordance to the Invitrogen guidelines and without any residual methanol concentration control. Although the higher cell concentration reached in this experiment attributed to an increased overall process productivity, the specific HBcAg yields per DCW were also higher than in other experiments — 11.2 mg g⁻¹. The higher methanol feed rate (0.4–0.5 mL min⁻¹) and a slightly higher reached cell biomass in Exp. 2 did not amount to an increased final HBcAg concentration or specific yield — 996 mg L⁻¹ and 8.3 mg g⁻¹, respectively.

Cultivation in Exp. 3 was carried out under a controlled residual methanol concentration of 1.0 g L⁻¹. The PI-algorithm-based methanol concentration control performed admirably and was able to maintain the residual methanol concentration at 1.0 g L⁻¹ with an average deviation of ±0.3 g L⁻¹. After 72 h of cultivation, 1 L of culture was removed to assess how the control would adapt to the new conditions. After a sharp methanol concentration increase, the feed rate values were decreased and increased again when the excess residual methanol was consumed. At approx. 78 h, a sudden methanol concentration spike was noted. It is unclear what exactly caused the cell methanol uptake to suddenly drop as such. Although the methanol feed was automatically stopped, it took about 12 h for the cells to consume the excess methanol. The results from this experiment also indicate that the optimal cultivation length for maximum HBcAg production is around 70–75 h, as an overall yield decrease can be noted the next day (92 h). Yet, a lower biomass concentration (92.0 g L⁻¹) and a lower both total and specific productivity were achieved in this
experiment in comparison to Exp. 1 – 876 mg L\(^{-1}\) and 9.6 mg g\(^{-1}\), respectively.

Exp. 4 was performed at a residual methanol concentration of 2.0 g L\(^{-1}\); however, the same control parameter values were used as in Exp. 3 and this led to a very poor methanol concentration control with an average deviation of approx. ± 1.0 g L\(^{-1}\). This illustrates the importance of process-specific control parameter tuning. Consequently, a very poor production was shown by SDS-PAGE analysis and a decision was made not to purify the end culture. This experiment was repeated with more suitable PI-control parameter values (\(K_P = 0.02 \text{ L}^2 \text{ g}^{-1} \text{ h}^{-1}\)) in Exp. 5 and a much better residual methanol concentration control was achieved – 2.0 ± 0.39 g L\(^{-1}\). Consequently, a much better productivity was achieved – 944 mg L\(^{-1}\) and 7.9 mg g\(^{-1}\). Although this productivity is comparable to the one achieved in Exp. 1 and was achieved in 67 h (instead of 75 h), around 20% more methanol was used, which would impact process overall economic feasibility. Taking this into consideration, Exp. 1 still seemingly has the upper hand production-wise.

The gel electrophoresis analysis from Exp. 1 in Fig. 3 illustrates the accumulation dynamics of HBcAg during methanol induction. Significant HBcAg accumulation can already be noted in the sample taken just 3.5 h after the start of methanol induction. Despite the thick HBcAg band of the 48th h sample, the specific yield of recombinant protein appears to be approx. 2–3 times higher for the end culture (75th process hour).

Previously processed samples were loaded onto a size-exclusion chromatography column packed with Sepharose 4 Fast flow resin. In laboratory scale, we used a 10 mm Tricorn column, packed at 550 mm with a flow rate of 0.30 mL min\(^{-1}\). HBcAg elution profile is presented in Fig. 4, where the second peak represents the HBcAg VLP fractions (3–7).

SDS-PAGE gel showing HBcAg bands and immunodiffusion titres after each purification step is shown in Fig. 5. Finally, the collected fractions from the size-exclusion chromatography are merged and loaded onto an anion-exchange column packed with a Fractogel DEAE resin. In
Fig. 3 SDS-PAGE analysis of HBcAg accumulation dynamics from Exp. 1

Fig. 4 Sepharose 4FF size-exclusion chromatography profile of HBcAg elution (Exp. 1, 75 h). Black line represents protein elution; red line—conductivity

Fig. 5 SDS-PAGE analysis of HBcAg purification steps (Exp. 1, 75 h)
At laboratory scale, we used a column with a bed volume of 20.0 mL and a flow rate of 4.0 mL min⁻¹. The HBcAg is bound to the column and eluted by a linear salt gradient. The HBcAg elution profile is shown in Fig. 6. Fractions 2 and 3 are collected and pooled for final HBcAg concentration determination and electron microscopy imaging.

The HBcAg concentration in the final fraction was analysed using the universal Bradford assay. It was estimated that the specific HBcAg yield reached 11.2 mg g⁻¹ and the total yield was 1308 mg L⁻¹. The estimated product purity is roughly 90%. The presence of purified, correctly formed, homogenous HBcAg-VLPs is confirmed by the electron microscopy images in Fig. 7. As is apparent from the electron microscopy images, the purified HBcAg-VLPs from samples after 48 h and 75 h of cultivation indicate similar quality.

**Mathematical modelling**

Along with the Process Analytical Technology (PAT) and Good Manufacturing Practice (GMP) directive-driven requirements for knowledge-based in-depth fermentation process understanding, some typical large-scale culture problems can also be mediated by successfully employing...
fermentation process mathematical modelling. Namely, to select an appropriate substrate feeding rate profile that provides optimal cell biomass growth and avoids substrate-induced catabolite repression, while also mitigating the very high oxygen demand and metabolic heat production during the methanol induction phase.

When modelling *P. pastoris* cultivations, the goal should be to refrain from using batch-dependant parameters, and, if possible, to identify comprehensive parameter values that provide suitable accuracy, while remaining constant for all batches. The former approach also sometimes requires time-intensive parameter-identification procedures for every batch, whereas the latter does not.

Here, we present mathematical models to portray cell biomass (*X_v*, *X_d*, and *X*), glycerol (Gly) and methanol (MeOH) concentrations, HBcAg accumulation (*P*), and reactor volume (*V*) change dynamics during five HBcAg-producing *P. pastoris* cultivations. Experimental data were used to evaluate model precision.

To our knowledge, this is the first reported attempt to model HBcAg accumulation during a *P. pastoris* bioreactor cultivation and the residual methanol concentrations in this particular range. The modelled cultivation experiments are presented in Fig. 8. Identified model parameters and their respective reference values are compiled in Table 1.

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**Fig. 8** Modelling results from 5 HBcAg-producing *P. pastoris* cultivations. Vertical blue dashed line indicates the onset of the glycerol fed-batch phase, and red line indicates the start of methanol induction.
Model quality was estimated by means of root-mean-square error (RMSE) and normalized root-mean-square error (NRMSE) values. Calculated values are presented in Table 2.

The model parameter sensitivity analysis results are presented in Table 3. Table values indicate the deviation in each respective modelled variable RMSE, caused by a 5% increase of said model parameter.

The results from the model parameter sensitivity analysis procedure indicate that both glycerol and methanol modelled concentrations are very sensitive to most model parameter values. This can be explained by the relatively low residual concentrations of the substrates in the medium. As for the parameters that have the most impact on modelling results, constants $k1$ and $k2$ jump out. Such high sensitivity can be explained, first of all, because they directly influence the biomass (X) concentration, which in turn has a major effect on most other modelled variables. Second, because minor deviations from the biomass growth profile in the early cultivation hours can have a snowball effect in the late process hours.

**Discussion**

Historically, HBcAg-VLPs have played a key role in vaccine development and, consequently, the efficient production and purification of these particles can play a major role in commercial vaccine production. There are many reports of successfully producing these particles in various recombinant expression platforms; however, *P. pastoris* stands out due to its ability to reach high cell densities on minimal media, perform post-translational modifications, and drive high-level expression of recombinant proteins. The substrate feeding strategy during the methanol induction phase for recombinant product biosynthesis is recognised to have a significant

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**Table 1** Identified model parameters

| Symbol | Unit | Glycerol phase | Methanol phase |
|--------|------|----------------|---------------|
| $K1$  | mol g$^{-1}$ | 0.001 | 0.001$^b$ | 0.0015 |
| $K2^*$ | mol g$^{-1}$ | 0.0127 | 0.009–0.015$^b$ | 0.017 |
| $Y_{ATP}$ | g mol$^{-1}$ | 10.50 | 10.50$^b$ | 10.50 |
| $\phi$ | – | – | 0.25 |
| $P/O$ | mol mol$^{-1}$ | 1.50 | 1.50$^b$ | 2.13 |
| $K_S$  | g L$^{-1}$ | 0.30 | 0.30$^b$ | 0.30 |
| $mATP$ | mol g$^{-1}$ h$^{-1}$ | 0.0015 | 0.0015$^b$ | 0.0001 |
| $q_{min}$ | mol g$^{-1}$ h$^{-1}$ | 0.0006 | 0.0006$^b$ | – |
| $q_{max}^*$ | mol g$^{-1}$ h$^{-1}$ | 0.067 | 0.053–0.062$^b$ | 0.008 |
| $K_i^*$ | g L$^{-1}$ | – | – | 11.0 |
| $k1$  | h$^{-1}$ | 0.40 | 0.40$^b$ | – |
| $k2^*$ | h$^{-1}$ | 0.255 | 0.20–0.29$^b$ | – |
| $F_{exp}^*$ | L h$^{-1}$ | 0.0066 | – | 0.0066 |
| $S_F$  | g L$^{-1}$ | 500 | – | 790 |
| $K_d^*$ | h$^{-1}$ | 0.0012 | – | 0.0012 |
| $Y_{RX}^*$ | g g$^{-1}$ | – | – | 0.091 |
| $M$  | g mol$^{-1}$ | 92.09 | 92.09$^f$ | 32.04 |

$^a$ Determined experimentally; $^b$ Jahic et al. [27]; $^c$ Ren et al. [28]; $^d$ Niu et al. [29]; $^e$ Barrigon et al. [30]; $^f$ Zhang et al. [31]; $^g$ Pubchem database [37]
influence on the product yield and process productivity. Despite this, the effect of residual methanol concentration on HBcAg production with *P. pastoris* in bioreactor cultivations has not been extensively studied and reported. In this particular research, three residual methanol levels (0.1, 1.0, and 2.0 g L\(^{-1}\)) during the HBcAg-producing *P. pastoris* cultivation methanol induction phase were investigated.

Excellent recombinant HBcAg yields were achieved in all cultivations; however, the best yield was produced in Exp. 1—according to the Invitrogen guidelines and without residual methanol concentration control during induction (1308 mg L\(^{-1}\) and 11.2 mg g\(_{DCW}\)\(^{-1}\)). When the residual methanol concentration was controlled at 1.0 or 2.0 g L\(^{-1}\), lower HBcAg yields were registered—876 mg L\(^{-1}\) and 9.6 mg g\(_{DCW}\)\(^{-1}\) in Exp. 3, and 944 mg L\(^{-1}\) and 7.8 mg g\(_{DCW}\)\(^{-1}\) in Exp. 5, respectively. Additionally, approx. 10–20% more methanol was used in Exps. 3 and 5, which should also be taken into account when evaluating the economic feasibility of a production process. Also, residual methanol concentration control requires both the technical possibility and the implementation and tuning of a suitable methanol feed rate control algorithm. Overall, it is evident that in this particular case, residual methanol concentration control is not feasible. A review of reported *P. pastoris*-produced HBcAg yields is presented in Table 4.

Unfortunately, experimental screening at higher residual methanol levels (4.0–6.0 g L\(^{-1}\)) was not possible, due to the insufficiently low temperature of the tap water used for cooling. Consequently, the bioreactor’s cooling system was unable to successfully maintain the temperature below 32 °C. To avoid this setback, an additional cooling system should be employed, for example, a circulation chiller. Another possible option to avoid cooling issues would be to grow the cells to lower densities; however, this would also lead to lower HBcAg yields. An example can be seen in Exp. 3—where lower cell densities lead to lower overall HBcAg yields, despite the second highest specific HBcAg yield.

The overall HBcAg yield in Exp. 1 is the highest reported in both scientific and patent literature to our knowledge. The second highest reported yield by Freivalds et al. [32] utilizes the same HBC producer clone and a similar purification procedure. There are two main differences. First, BMGY was selected as the cultivation medium. In general, production in minimal (salt) medium, such as BSM, is preferred over rich (complex) BMGY medium, as it eases subsequent product purification and improves overall process economic feasibility. Also, the ability to reach high cell densities and high-level production in minimal medium is one of the main advantages the yeast *P. pastoris* presents. Second, the authors did not employ a glycerol fed-batch phase, but added a glycerol–methanol mixture to facilitate faster adaptation to methanol consumption. While this may indeed facilitate better cell adaptation to methanol consumption, the absence of a glycerol fed-batch phase leads to lower cell biomass concentrations and, consequently, overall process yields. Although the specific HBcAg yields are comparable (11.2 vs 10.0 mg g\(_{DCW}\)\(^{-1}\)), we reached an almost 2.5 times higher biomass concentration and, consequently, a 2.5 times higher overall HBcAg yield.

The residual methanol concentration in Exps. 3–5 was controlled by altering the methanol feed rate according to the values calculated by the PI algorithm. This control solution proved to be able to control the residual methanol concentration at set-point value with sufficient accuracy. The main advantage being relatively easy implementation. This controller, however, is quite sensitive to the selected parameter.
values, as can be seen in Exps. 4 and 5; therefore, a procedure for optimum control parameter value identification is necessary. The control was able to adapt to unexpected situations—for example, when approx. 1 L of culture was removed during Exp. 3 (71 h)—although a brief spike in the residual methanol concentration was noted, it was corrected shortly after. At times, however, the control proved capable to control the methanol concentration at the set-point value with sufficient accuracy and can be employed in *P. pastoris* cultivation processes.

The purification procedure used in this research allowed for a rapid and efficient HBcAg VLP purification. When purifying recombinant proteins, an important aspect to consider is the time required for the purification procedure. In this case, the whole procedure can be carried out in 1 day. By following this purification protocol, highly immunogenic, correctly formed and homogenous HBcAg-VLPs with an estimated purity of 90% can be achieved.

As indicated earlier, process mathematical modelling can play an important role in recombinant product development and manufacture and, although there are several reports of successful *P. pastoris* cultivation process models, there is little information on modelling the residual methanol concentration during the induction phase. In this particular research, we present a process model that characterises cell biomass (\(X_v\), \(X_d\) and \(X\)), substrate (Gly. and MeOH) concentration, reactor volume (\(V\)), and product (HBcAg) accumulation dynamics during cultivation. The experimental data accumulated during these cultivations were used to estimate the model accuracy.

Good results were achieved when modelling the cell biomass (\(\pm 6.05 \text{ g L}^{-1}\)) and glycerol (\(\pm 1.65 \text{ g L}^{-1}\)) concentrations. In prior research, using a dielectric spectroscopy probe and an off-gas analyser, we have observed that, during the start of the methanol induction phase—when significant methanol accumulation in the reactor takes place—a reduction in overall cell viability can be noted [34]. The experimental dry cell biomass measurements account also for dead (non-vegetative) cells; therefore, separate models were introduced to account for both vegetative cell (used in substrate and product calculations) and total cell (used for correlation with DCW measurements) biomass concentration. To estimate the dead cell fraction and offset the effect of these non-viable cells present on modelled cell biomass values, an empirical parameter \(K_d\) was introduced. This allowed to significantly improve the accuracy of cell biomass estimation, especially in the late induction phase, when the dead cell fraction composes a significant part of total cell biomass. However, some underestimation can still be noted. This is likely because the term \(K_d\) does not remain constant during the whole cultivation and is influenced by both the residual methanol concentration and cell exposure (induction) time.

The residual methanol concentration, however, proved difficult to model. Although the model was able to successfully

| Table 4 Reported purified HBcAg yields |
|----------------------------------------|
| HAg yield | Host | Strain | Media* | MeOH (g L\(^{-1}\)) | Cultivation length (h) | Scale | Refs. |
|----------|------|--------|--------|----------------|------------------------|-------|------|
| 1308.0 | *P. pastoris* | GS115 Mut\(^*\) | BSM | Limiting | 75 | 5L bioreactor | This work |
| 944.0 | *P. pastoris* | GS115 Mut\(^*\) | BSM | 0.2% | 67 | 5L bioreactor | This work |
| 876.0 | *P. pastoris* | GS115 Mut\(^*\) | BSM | 0.1% | 71 | 5L bioreactor | This work |
| 700.0 | *P. pastoris* | GS115 Mut\(^*\) | BMGY | Limiting | 92 | 10L bioreactor | [32] |
| 398.0 | *E. coli* | BL21 | M9Cas | – | 16 | Shake flasks | [38] |
| 120.0 | *P. pastoris* | GS115 Mut\(^*\) | BMGY/BMMY | 0.5% | 92 | Shake flasks | [39] |
| 80.0 | *P. pastoris* | GS115 | BMGY/BMMY | 0.6% | 114 | Shake flasks | [40] |
| 79.0 | *E. coli* | BL21 | 2YT | – | 24\(^*\) | Shake flasks | [41] |
| 69.0 | *P. pastoris* | GS115 Mut\(^*\) | BMGY | Limiting | 72 | 2L bioreactor | [42] |
| 64.0 | *P. pastoris* | GS115 Mut\(^*\) | BMGY | 0.5% | 144 | Shake flasks | [41] |
| 58.1 | *E. coli* | BL21 | 2YT | – | 24\(^*\) | Shake flasks | [43] |
| 55.2 | *E. coli* | BL21 | LB | – | 24\(^*\) | Shake flasks | [44] |
| 50.0 | *P. pastoris* | GS115 Mut\(^*\) | BMGY | Limiting | 96 | 2L reactor | [45] |
| 24.7 | *E. coli* | JM109 | LB | – | – | Shake flasks | [43] |
| 8.7 | *P. pastoris* | GS115 | BMGY/BMMY | 0.5% | 114\(^*\) | Shake flasks | [43] |
| 2.5 | *E. coli* | BL21 | 2YT | – | 24\(^*\) | Shake flasks | [45] |

\(^*\)Estimated; \(^*\)BSM Basal Salts Medium, BMGY Buffered Glycerol Complex Medium, M9Cas Trp-deficient M9 minimal medium supplemented with casamino acids, BMMY Buffered Methanol Complex Medium, 2YT Yeast extract and Tryptone medium, LB Luria Bertani medium.
account for methanol concentration peaks at the start of residual methanol concentration control, the overall accuracy could still use more improvement (± 1.26 g L⁻¹). The model also somewhat underestimated the residual methanol concentration during cultivations. Looking at other residual methanol concentration modelling results, the model presented by Ren et al. was unable to account for a fast accumulation of methanol up to 10.0 g L⁻¹ [28]. The model, however, was able to accurately estimate the methanol concentration in a methanol-limited cultivation. Although the model presented by Jahic et al. included methanol concentration estimation, they opted to represent the methanol concentration using experimental measurements, perhaps, thus, indicating that the model was not able to accurately estimate the accumulated methanol concentration up to nearly 2.0 g L⁻¹ during methanol adaptation [27]. A more sophisticated approach is presented by Niu et al., as their model was able to account for methanol accumulation during initial adaptation (approx. 6.4 g L⁻¹) and at the end of the cultivation (approx. 16 g L⁻¹) [29]. Although the model is successful at estimating the residual methanol concentration at the start and the end of the methanol induction, it seemingly fails to account for a small methanol spike in the middle of cultivation. While this presented approach seems promising, it is also complex, as it incorporates real-time exhaust gas analysis and a dynamic parameter-identification procedure, which lessens the potential for practical application. Even though the accuracy of our presented model could still be improved, it demonstrates a rare attempt to model residual methanol concentration with sufficient accuracy and low complexity. The ability of the model to accurately predict increases (spikes) in methanol concentration can be very useful during both laboratory and industrial cultivations, as rapid detection of such spikes may help avoid deviating from the optimal growth trajectory or whole batch discard in general.

Recombinant HBcAg accumulation dynamics were also modelled (± 112.1 mg L⁻¹) and follow the experimentally determined yield values. However, the model could not account for the high productivity in Exp. 1, which could indicate that some other unknown factor could possibly have influenced the high yield in this particular experiment. A significant factor to note is also the fact that this measurement is very dependent on the estimated cell biomass concentration. Ensuring a higher cell biomass estimation would also positively impact this measurement in this case.

Overall, we have presented a relatively simple and robust bioreactor model to successfully portray cell biomass (X_v, X_d, and X) concentration, substrate (Gly. and MeOH), reactor volume (V), and product (HBcAg) dynamics during a P. pastoris GS115 cultivation at both limiting (0.1 g L⁻¹) and residual (1.0 and 2.0 g L⁻¹) methanol concentrations. Relatively easy implementation and parameter identification enables this approach to also be suitable for practical applications. The presented model could also be further used for model-based optimization and software-sensor development of the investigated process.

Conclusions

In this particular research, three residual methanol levels (0.1, 1.0, and 2.0 g L⁻¹) during the HBcAg-producing P. pastoris cultivation methanol induction phase were investigated. Increased residual methanol concentration did not have a profound effect on HBcAg accumulation dynamics. Considering the economic aspects of the production process, bioreactor cultivation under limiting methanol concentration, according to the Invitrogen cultivation guidelines, is preferable. The highest reported HBcAg VLP yield of 1308 mg L⁻¹ and 11.2 mg gDCW⁻¹ is presented, which is about 2.5 times higher than the next highest result (489 mg L⁻¹) reported by Freivalds et al. [32].

We also demonstrate that the PI-algorithm-based residual methanol concentration control was able to maintain the selected set-point concentrations in the explored range; however, control quality was sensitive to the values of selected control parameters K_p and r_i.

The produced recombinant HBcAg molecules were successfully purified using a modified purification protocol, and the presence of highly immunogenic, correctly formed and homogenous HBcAg-VLPs with an estimated purity of 90% was confirmed using electron microscopy.

A relatively simple and robust bioreactor model to portray cell biomass (X_v, X_d and X), substrate (Gly. and MeOH), reactor volume (V), and product (HBcAg) dynamics during a P. pastoris GS115 cultivation at both limiting (0.1 g L⁻¹) and residual (1.0 and 2.0 g L⁻¹) methanol concentrations is presented. A rare attempt to model residual methanol concentration during the induction phase is also presented. Relatively easy implementation and parameter identification enables this approach to also be suitable for practical applications. The presented model could be further used also for model-based optimization and software-sensor development of the investigated process.

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Declarations

Conflict of interest The authors declare no conflicts of interest.

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