A dynamic method based on the specific substrate uptake rate to set up a feeding strategy for *Pichia pastoris*

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**Abstract**

**Background:** *Pichia pastoris* is one of the most important host organisms for the recombinant production of proteins in industrial biotechnology. To date, strain specific parameters, which are needed to set up feeding profiles for fed batch cultivations, are determined by time-consuming continuous cultures or consecutive fed batch cultivations, operated at different parameter sets.

**Results:** Here, we developed a novel approach based on fast and easy to do batch cultivations with methanol pulses enabling a more rapid determination of the strain specific parameters specific substrate uptake rate $q_s$, specific productivity $q_p$, and the adaption time ($\Delta t_{\text{adapt}}$) of the culture to methanol. Based on $q_s$, an innovative feeding strategy to increase the productivity of a recombinant *Pichia pastoris* strain was developed. Higher specific substrate uptake rates resulted in increased specific productivity, which also showed a time dependent trajectory. A dynamic feeding strategy, where the setpoints for $q_s$ were increased stepwise until a $q_s_{\text{max}}$ of 2.0 mmol·g$^{-1}$·h$^{-1}$ resulted in the highest specific productivity of 11 U·g$^{-1}$·h$^{-1}$.

**Conclusions:** Our strategy describes a novel and fast approach to determine strain specific parameters of a recombinant *Pichia pastoris* strain to set up feeding profiles solely based on the specific substrate uptake rate. This approach is generic and will allow application to other products and other hosts.

**Background**

Recombinant protein expression with biological hosts is one of the most examined key processes in the pharmaceutical industry. Numerous products like organic acids, antibiotics, enzymes and amino acids are produced heterologously by recombinant microorganisms. The methylotrophic yeast *Pichia pastoris* is one of the most important host organisms for this purpose.

Several of the published fermentation strategies for *P. pastoris* to date are based on the Invitrogen protocol http://tools.invitrogen.com. This protocol suggests constant feeding profiles for fed batch cultivations, but does not aim to improve production efficiency regarding time and yield or substrate consumption. Based on this protocol, different process strategies were developed to optimize recombinant protein production with *P. pastoris* in the past few years. A commonly used feeding strategy describes a feed forward regime based on a constant specific growth rate $\mu$ [1-5]. This strategy results in an exponential feeding profile and does not require complex instrumentation, but $\mu$ is also not controlled, and since the cells capacity may change over time, the feeding profiles consider a large safety margin. Another feeding strategy is based on a controlled $\mu$ and requires laborious continuous culture investigations and an effective computer controlled operation, based on established growth models and a feedback algorithm requiring expensive online measurement sensors for methanol [2,6-8]. Employing these strategies, the outcome regarding specific productivity and specific growth rate was diverse; some studies showed that the maximal specific productivity did not relate to the maximal specific growth rate [1,3,6,7], whereas another study showed a more or less growth associated productivity [9].

Due to these controversial findings, other parameters were analyzed for their possible correlation with the productivity in the past few years. Khatri and Hoffmann
analyzed the specific substrate uptake rate ($q_s$) and its association with the specific productivity ($q_p$) in fed-batch cultivations of *P. pastoris* and showed that lower $q_s$ resulted in higher $q_p$ [10,11]. In another study, Cunha et al. used a constant feeding rate of methanol during several fed-batch cultivations with different initial biomass contents to monitor $q_s$ over time [12]. Interestingly, Cunha et al. also showed increased $q_p$ at lower $q_s$ and clearly stated that $q_s$ was the most important induction parameter. However, despite the obvious effect of $q_s$ on $q_p$, feeding profiles based on $q_s$ have not been tested yet.

Regardless of which control parameter is chosen to set up feeding profiles for fed batch cultivations, strain specific parameters have to be determined. This can either be done by continuous cultures [13] or by several, consecutive fed batch cultivations, which are operated at different parameter sets [14,15]. These methods have the disadvantage of being very time-consuming and labor-intensive. A robust method to determine these strain specific data and to develop a suitable feeding strategy in a shorter time period is of high interest in biotechnology to speed up process development and to quantitatively screen industrial relevant strains.

In this study, we developed a new and fast method to determine the strain specific data $q_s$ and $q_p$, as well as the adaptation time of the culture to the substrate methanol, based on easy to do batch cultivations with methanol pulses. The host *P. pastoris* and the product horseradish peroxidase (HRP) were used as a model system, for which we subsequently developed a novel feeding strategy solely based on the determined $q_s$ (Figure 1). Various fed batch cultivations, employing different $q_s$ feeding profiles, were carried out to characterize the strain, hence, to determine yields ($Y_{X/S}; Y_{CO2/S}$) and the specific productivity ($q_p$) to evaluate and improve the feeding strategy. To our knowledge this is the first time that different dynamic feeding profiles only based on $q_s$ were tested to develop a feeding strategy for *P. pastoris* aiming at increased productivity.

### Material and methods

#### Microorganism and recombinant protein

The *Pichia pastoris* strain KM71 H (arg4 aox1dar4) was transformed with a plasmid containing the gene for the horseradish peroxidase isoenzyme C1A (HRP) and was gratefully provided by Prof. Anton Glieder (Graz University of Technology, Austria). The phenotype of the strain corresponded to an AOX1-deficient clone which is characterized as Mut5 (methanol utilization slow) and HRP was secreted into the fermentation broth.

Horseradish peroxidase is a member of the plant peroxidase super family [16] and catalyzes the oxidation of many substrates using hydrogen peroxide, resulting in oxidized products and in the formation of two molecules of water per molecule of hydrogen peroxide [17-19]. It is a heme-containing glycoprotein with a molecular weight of approximately 44 kDa that has been studied for more than 100 years [19]. Horseradish peroxidase gains more and more industrial relevance in the past few years; it is used in waste water treatment [20,21], analytical diagnostics [22] and for the elimination of H$_2$O$_2$ from food and industrial wastewater [23].

#### Stability of the enzyme horseradish peroxidase in the presence of methanol

To check whether certain concentrations of methanol were affecting the stability of the enzyme horseradish peroxidase, HRP was diluted either in water or in BSM medium to a concentration of 1 U/mL and incubated in a waterbath at 28°C in the presence of up to 20 mg/mL methanol (which corresponds to 2.5% v/v) overnight. At several time points samples were taken and analyzed for catalytic activity and protein content.

#### Culture Media

**Preculture:** Yeast nitrogen base media (YNBM), per liter: potassium phosphate buffer (pH 6.0), 0.1 M; YNB w/o Amino acids and Ammonia Sulfate (Difco™), 3.4 g; (NH$_4$)$_2$SO$_4$, 10 g; biotin, 400 mg; glucose, 20 g.

**Batch/fed batch:** Basal salt media (BSM) [24], per liter: 85% phosphoric acid, 26.7 mL; CaSO$_4$·2H$_2$O, 1.17 g; K$_2$SO$_4$, 18.2 g; MgSO$_4$·7H$_2$O, 14.9 g; KOH, 4.13 g; C$_6$H$_{12}$O$_6$·H$_2$O, 44 g, Antifoam Struktol J650, 0.2 mL; PTM1, 4.35 mL; NH$_4$OH as N-source (see experimental procedure). Trace element solution (PTM1), per litre: CuSO$_4$·5H$_2$O, 6.0 g; NaI 0.08 g; MnSO$_4$·H$_2$O, 3.0 g; Na$_2$MoO$_4$·2H$_2$O, 0.2 g; H$_2$BO$_3$, 0.02 g; CoCl$_2$, 0.5 g; ZnCl$_2$, 20.0 g; FeSO$_4$·7H$_2$O, 65.0 g; biotin, 0.2 g, H$_2$SO$_4$, 5 mL.

Feed glucose, per liter: glucose, 250 g; PTM1, 12 mL, Struktol J650, 0.3 mL.
Feed methanol, per liter: methanol, 300 g; PTM1, 4 mL; Struktol J650, 0.3 mL, induction period was carried out in presence of δ-Aminolevulinic acid (δ-ALA), 1 mM.

Base: NH₄OH, concentration was determined by titration with 0.25 M potassium hydrogen phthalate (KHP).

**Experimental Procedure**

**Preculture**

Frozen stocks (-80°C) were pre-cultivated in 100 mL of YNBM in 1000 mL shake flasks at 28°C and 200 rpm for max. 24 hours. Then, the preculture was transferred aseptically to the respective culture vessel. The inoculation volume was approximately 10% of the final starting volume.

**Batch cultivation and determination of qₛ**

Batch cultivations were carried out in a 1 L working volume glass bioreactor (Applikon, Netherlands). Basal salt media was sterilized in the bioreactor and pH was adjusted to pH 5.0 by using concentrated ammonia solution after autoclaving. Sterile filtered trace elements were transferred to the reactor aseptically. Dissolved oxygen (dO₂) was measured with a sterilizable polarographic dissolved oxygen electrode (Mettler Toledo, Switzerland). The pH was measured with a sterilizable electrode (Mettler Toledo, Switzerland) and maintained constant with a step controller using ammonia solution (1 to 2 M). Base consumption was determined gravimetrically. Cultivation temperature was set to 28°C and agitation was fixed to 1200 rpm. The culture was aerated with 1.25 vvm dried air and off-gas of the culture was measured by using an infrared cell for CO₂ and a paramagnetic cell for O₂ concentration (Servomax, Switzerland). Temperature, pH, dO₂, agitation as well as CO₂ and O₂ in the off-gas were measured online and logged in a process information management system (PIMS; Lucullus, Biospectra, Switzerland).

After the complete consumption of the substrate glucose, which was indicated by an increase of dissolved oxygen and a drop in off-gas activity, the first methanol pulse of a final concentration of 0.5% (v/v) was conducted with pure methanol (supplemented with PTM1, 12 mL/L of methanol). Following pulses were performed with 1% (v/v), before a last pulse with 2% (v/v) final concentration of methanol was carried out. To investigate the metabolic activity during methanol excess and also the dynamic behavior of the cell metabolism after methanol limitation for several hours, the pulse experiments were performed like this: after methanol was depleted after the “first” pulse (followed by off-gas analysis), an immediate “second” methanol pulse with the same concentration was conducted per day. After methanol depletion, methanol starvation was carried out for several hours before another so-called “first” pulse was applied (Figure 2).

For each pulse, at least two samples were taken to determine the concentrations of substrate and product as well as dry cell weight and OD₆₀₀ to calculate the specific substrate uptake rate qₛ.

**Fed batch cultivations**

Fed batch cultures were carried out in a 7.5 L (5 L working volume) glass bioreactor (Infors, Switzerland). Concentrated BSM medium (2-fold concentrated to supply necessary salts for high cell densities) was sterilized in the bioreactor and pH was adjusted to pH 5.0 by using concentrated ammonia solution after autoclaving. The initial volume was set to 1.5 L. Trace elements were filter sterilized and transferred to the reactor aseptically. Dissolved oxygen was measured with a sterilizable polarographic dissolved oxygen electrode (Hamilton, Switzerland). The pH was measured with a sterilizable electrode (Hamilton, Switzerland) and maintained constant using ammonia solution (3 to 5 M). Agitation was set to 1500 rpm. The culture was aerated with at least 1 vvm to avoid oxygen limitation. The dissolved oxygen signal was used to adjust air-in flow manually to keep levels >30% dO₂ at all time points. In case air flow was limited, pure oxygen was added. CO₂...
and O₂ were measured as described above. Base consumption and reactor weight were measured gravimetrically. The fed batch feed was measured and controlled using a gravimetrically based PID flow controller.

At several time points during fed batch cultivations, samples were taken and analyzed for accumulated methanol, biomass concentration (dry cell weight and optical density OD₆₀₀), protein content and enzymatic activity. Based on the total biomass content, feed rates were adjusted manually corresponding to the defined qs setpoint.

Three different fed batch strategies were tested: fed batch A, where methanol was adjusted to a constant flow during the whole induction phase; fed batch B, where after the adaption time with a qs_adapt, a shift to a high substrate uptake rate of ~90% of qs_max was done and adjustments to the very same qs set point during cultivation were performed repeatedly; and fed batches C1 and C2, where after the adaptation period with qs_adapt, the methanol flow was stepwise increased up to qs_max. Values for qs_adapt and qs_max had been determined in batch experiments (vide supra). An overview of the fed batch cultivations and the corresponding settings is given in Table 1.

Analysis of growth- and expression-parameters

Dry cell weight (DCW) was determined by centrifugation of 5 mL culture broth (5000 rpm, 4°C, 10 min) in a laboratory centrifuge (Sigma 4K15, rotor 11156), washing the pellet with 5 mL deionized water and subsequent drying at 105°C to a constant weight in an oven. Optical density of the culture broth was measured using a spectrophotometer (U-1100 Hitachi, Japan) at a wavelength of 600 nm (OD₆₀₀). Correlation between dry cell weight measurement and OD₆₀₀ showed a coefficient of regression of R² = 0.997 over the full concentration range (data not shown) and could therefore be used for qs adaptation.

The activity of HRP was determined using a CuBiAn XC enzymatic robot (Innovatis, Germany). Cell free samples (10 μl) were added to 140 μl of 1 mM ABTS (2,2’ azino bis 3-ethylbenzthiazoline-6-sulphonic acid) prepared in 50 mM NaOAc buffer (pH 4.5). The reaction mixture was incubated at 37°C and was started by the addition of 20 μl of 0.075% H₂O₂. Changes of absorbance at 415 nm were measured for 80 seconds and rates were calculated. Calibration was done using commercially available horseradish peroxidase (Type VI-A, Sigma-Aldrich, P6782, Lot# 118K76703) as standard at six different concentrations (0.02; 0.05; 0.1; 0.25; 0.5 and 1.0 U/mL). Samples with high enzymatic activity were automatically diluted by the system. Protein concentrations were determined at 595 nm by the Bradford assay [25] using the BioRad Protein Assay Kit with BSA as standard.

Substrate concentrations

Concentrations of methanol were determined in cell free samples by HPLC (Agilent Technologies, USA) equipped with a Supelcoguard column, a Supelcogel C-610 H ion-exchange column (Sigma-Aldrich, USA) and a refractive index detector (Agilent Technologies, USA). The mobile phase was 0.1% H₃PO₄ with a constant flow rate of 0.5 mL/min and the system was run isocratic. Calibration was done by measuring standard points in the range of 0.1 to 10 g/l methanol.

Concentrations of glucose were determined in cell free samples by a commercial enzymatic assay kit using the CuBiAN XC enzymatic robot (Innovatis, Germany). Calibration was done with 4 standard points in the range from 0 to 3 g/l glucose. Samples with higher glucose concentration were diluted automatically by the system.

Data analysis

Measurements of biomass concentration, product concentration and substrate concentration were executed in duplicates: along the observed standard deviation for the single measurement, the error was propagated to the specific rates qs and qp as well as to the yield coefficients. The error of determination of the specific rates and the yields was therefore set to 10% and 5%, respectively.

Electrophoresis

To check the purity of the excreted HRP, electrophoresis was done with aliquots of supernatants obtained at different time points during the cultivation of P. pastoris.

### Table 1 Description of feeding strategies for fed batch cultivations based on the specific substrate uptake rate qs

| Fed batch name | Symbol | Description of the strategy |
|----------------|--------|-----------------------------|
| Fed batch A   | ▲      | conventional feeding strategy: long adaptation time and initially adjusted, constant flow rate |
| Fed batch B   | ◆      | short adaptation time (out of batch exp.) and adjustment to high uptake rate (90% of qs_max) with repeated readjustments |
| Fed batch C1  | ▼      | short adaptation time (out of batch exp.) and stepwise adjustment of qs until qs_max with repeated readjustments |
| Fed batch C2  | ●      | |

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expressing the hrp gene extracellularly as described by Laemmli et al. [26]. SDS-PAGE was performed using a 5% stacking gel and a 10% separating gel in 1× Tris-glycine buffer. Gels were run in the vertical electrophoresis Mini-PROTEAN Tetra Cell apparatus (Biorad; Vienna, Austria) at 150 V for about 2 h. Gels were stained with Coomassie blue. The protein mass standard used was the PageRuler Prestained Ladder (Fermentas; Vienna, Austria).

Results and Discussion
Stability of the enzyme horseradish peroxidase in the presence of methanol
To check whether the enzyme horseradish peroxidase denatures at certain concentrations of methanol, HRP was incubated in the presence of methanol concentrations of up to 2.5% (v/v) at 28°C overnight. No loss in catalytic activity and protein content was detected after this incubation, which is why methanol pulses in batch experiments with concentrations of up to 2.5% (v/v) were theoretically possible without any denaturing effects on HRP.

Determination of strain specific parameters by easy to do batch cultivations with methanol pulses
After depletion of glucose in batch cultivations, a first methanol adaption pulse with a final concentration of 0.5% (v/v) was applied. The adaptation time to the new substrate methanol (Δtimeadapt) was determined with 7 hours and was defined by the detection of a maximum in off-gas activity (Figure 2A).

The calculated carbon dioxide evolution rate (CER), signifying metabolic activity, allowed to distinguish different states within the methanol pulses with local minima and maxima (Figure 2). This metabolic behavior of the cells results from inhibition and regulation events in transient conditions, caused by intracellular components like e.g. produced H₂O₂, and transport actions, and has been described for various systems in literature before [27-29].

The high frequent determination of biomass, methanol and product concentrations allowed specific rate calculations for methanol uptake qₘ and productivity qₚ during the methanol pulses. Specific substrate uptake rates were calculated with 0.8 ± 0.08 mmol·g⁻¹·h⁻¹ in the adaptation period (qₘadapt) and with around 2 ± 0.20 mmol·g⁻¹·h⁻¹ as a maximum during pulses (qₘmax). No difference in the calculated qₘ between the “first” and the subsequent “second” pulse on a respective day was observed (Figure 2A). In contrast, the calculated values for qₚ (Figure 2B) were very different between the first and the subsequent second pulse. During the first pulse specific productivities of maximum 1.3 ± 0.13 U·g⁻¹·h⁻¹ were observed, whereas during the second pulse an increased specific productivity of up to 2.5 ± 0.25 U·g⁻¹·h⁻¹ was measured. This increased productivity during the second methanol pulse may be due to the fact that no adaption of the cells to methanol was necessary because all metabolic key functions for methanol assimilation remained in an active state. Thus, recombinant protein expression could start directly and energy was used more efficiently for product formation. This result shows clearly that Pichia cultures should be kept induced at all time to obtain maximal productivity. Interestingly, qₚ also increased over time for first and second pulses, respectively (Figure 2B). Methanol pulses, where the same final concentration of methanol was used, but which were conducted in the later phase of the batch cultivation, showed higher specific productivities compared to pulses before. Obviously, the culture exhibited a “memory effect” and thus a time-dependence of qₚ could be observed in the batch experiment.

As we show here, batch cultivations with methanol pulses allowed a fast identification of strain specific parameters, which are crucial for subsequent fed batch cultivations. The determined maximum specific substrate uptake rate qₘmax represents the upper end of the feed profile respective to qₘ. This novel method has the advantage of being less time-consuming and labor-intensive compared to the traditional methods, like continuous cultivations, and additionally allows a free choice of substrate, like e.g. the 2 to 3-fold cheaper glucose instead of glycerol.

Fed batch feeding strategy based on qₘ
Based on the batch results, we performed several fed batch cultivations with different feeding profiles based on the specific substrate uptake rate qₘ to find a feeding strategy for a recombinant P. pastoris strain.

Feeding profiles
After a batch phase on glucose as substrate (volume 1.5 L), an exponential fed batch cultivation with glucose yielded in biomass concentrations of up to 70 g/L in a volume of 2.5 L. At the end of this fed batch phase, a sample was taken to determine the current biomass concentration by measuring the OD₆₀₀ and the DCW.

Based on the calculated specific methanol uptake rate during adaptation (qₘadapt) from the batch experiment, which was around 0.8 ± 0.08 mmol·g⁻¹·h⁻¹, we used a little bit lower qₘadapt of 0.5 mmol·g⁻¹·h⁻¹ methanol for the fed batch cultivations during the adaption time. After Δtimeadapt, three different feeding strategies (fed batches A, B and C1/C2) were tested and compared (Figure 3 Table 1).

For fed batch A, which described a conventional feeding strategy as in the Invitrogen protocol, the methanol feed was adjusted to a substrate uptake rate of 1.0 mmol·g⁻¹·h⁻¹. The adjustment just happened at the
beginning and no further readjustments of the feeding rate were performed during the cultivation. Consequently, as biomass still increased, a drop down of $q_s$ over time was observed (Figure 3).

In the other fed batch experiments B, C1 and C2 adjustments of $q_s$ were done after each sampling, based on OD$_{600}$ measurements. As shown in Figure 3 the effective $q_s$, determined by off-line sampling, and the designated $q_s$ in these fed batch cultivations were very similar. Thus, a frequent determination of the actual biomass concentration in the reactor vessel is necessary, to be able to adjust the feeding rate to the chosen $q_s$. The development of robust measurement devices for the online determination of the biomass concentration, as e. g. FTIR, capacity probes or soft sensors, could allow an online adjustment of the feeding rate to $q_s$, by an automatic control system in the future, and thus the labor-intensive, frequent sampling procedure would be unnecessary.

**Maintenance metabolism vs. cell growth**

The relationship between $q_s$ and the specific growth rate was tested and found to be linearly correlated (data not shown). Based on this, the maintenance coefficient for this *P. pastoris* strain could be determined with around 0.5 mmol·g$^{-1}$·h$^{-1}$ (which equals 0.016 g·g$^{-1}$·h$^{-1}$). Very similar results were reported by Jahic *et al.* for fed batch cultures using *P. pastoris* expressing a fusion protein [5] with a maintenance coefficient of 0.013 g·g$^{-1}$·h$^{-1}$. When we adjusted $q_s$ to values higher than 1.0 mmol·g$^{-1}$·h$^{-1}$, we observed a significant increase in $Y_{X/S}$ and a corresponding decrease in $Y_{CO2/S}$ (Figure 4). That means that at $q_s$ of <1.0 mmol·g$^{-1}$·h$^{-1}$ most of the energy is obviously used for maintenance metabolism of the cells and thus for the production of CO$_2$ and not for cell growth or protein production. Interestingly, the same effect on biomass yield was observed by D’Anjou *et al.* using a continuous culture system with a mixed feed strategy [13]. However, we were able to extract this information from time-efficient, dynamic experiments equivalently. Consequently, we employed specific substrate uptake rates of >1.0 mmol·g$^{-1}$·h$^{-1}$ to guarantee a sufficient amount of energy for both, maintenance and cell growth.

**Required methanol concentration for induction - a principle question**

When producing recombinant proteins with *P. pastoris*, the principle question is how much methanol has to be fed to guarantee a fully induced AOX promoter. A lot of different studies have examined this topic, with different results. D’Anjou *et al.*, for example, reported that a methanol concentration between 1 and 2 g/L should be maintained in the culture to guarantee fully induced heterologous protein production [13]. Another study of the methanol concentration and its effect on *P. pastoris* Mut$^3$ strains was done by Kupcsulik and Sevella, who showed that the specific productivity of a recombinant human serum albumin expression system showed a maximum at 0.45 g/L of methanol [30]. In contrast to those studies, Cregg reported an even stronger induction of the AOX promoter in limited conditions [31], which
was the basis for several following studies applying a limited methanol supply for induction [1-3].

For all described fed batch strategies in our study, no significant methanol accumulation was detected (detection limit 0.1 g/L). We clearly show that high residual concentrations of methanol during the production phase are not required for the induction of the \textit{P. pastoris} Mut\textsuperscript{a} strain, as even higher specific productivities were obtained in limited fed batches with constant substrate uptake rates, compared to the batch cultivations with high methanol concentrations applied in the pulse-experiments (Table 2).

\textbf{Extracellular protein production and specific productivity} $q_p$

To follow the formation of excreted horseradish peroxidase (HRP) during the induction phase, SDS-PAGE analyses with cell-free supernatants were performed. SDS-PAGE analysis (Coomassie staining, Figure 5) of the cultivation broth taken at different time points did not show a distinct band for HRP, but rather a smear between 60 - 65 kDa. The difference between the reported molecular weight of around 44 kDa for HRP and the observed molecular weight in this study results from the high degree of glycosylation of the recombinant protein expressed in \textit{P. pastoris} with mannose-type oligosaccharides, as described previously for HRP by Morawski \textit{et al.} [32].

The protein content in the cultivation broth increased over time from 0.052 mg/mL after the fed-batch phase with glucose to 0.243 mg/mL at the end of the cultivation, which were around 91 hours of induction. As shown in Figure 5 the increase of the protein content in the cultivation broth is ascribed to an increased amount of HRP, which constituted the majority of secreted proteins in the cultivation broth.

Based on the determined biomass content and the enzymatic activity, the specific productivity $q_p$ during the different fed batch cultivations was calculated. The lowest $q_p$ was obtained in fed batch A, which described a conventional feeding strategy, where also the lowest $q_s$ was used and was not adapted over time (Figure 6). In fed batch B, which considered the actual biomass concentration and where the $q_s$ of 1.75 mmol·g\textsuperscript{-1}·h\textsuperscript{-1} was adapted regularly over time and thus the cells were kept in a certain physiological state, a 2-fold increase in $q_p$ was observed. Clearly, a higher $q_s$ resulted in a higher $q_p$, which disagrees with the results obtained by Khatri and Hoffmann and Cunha \textit{et al.} [10,11], who stated that lower $q_s$ resulted in higher protein production.

However, a dynamic, stepwise feeding strategy resulted in an even higher productivity compared to the other strategies tested (Figure 6). This dynamic feeding strategy considered $q_{s_{max}}$, which had been determined in the batch experiment, as the highest possible substrate uptake rate of the cells, as well as the yield coefficients and the maintenance coefficient as the lower end of efficient energy usage. The feed profile was set up in a way to head off as quickly as possible from the maintenance state of the cells (hence $q_s$ was set to values ≥1.0 mmol·g\textsuperscript{-1}·h\textsuperscript{-1}), before $q_s$ was dynamically adapted in steps to $q_{s_{max}}$. In fed batch cultivations C1 and C2, where this dynamic feeding strategy was applied, a $q_s$ of around 11 ± 0.11 U·g\textsuperscript{-1}·h\textsuperscript{-1} was determined, which represents a 5.5-fold increase compared to fed batch A.

\textbf{Time adaptation}

The fact that there was still a more than 2-fold difference in $q_p$ between fed batches C1/C2, where $q_s$ was increased stepwise, and fed batch B, where immediately after the adaption period a high $q_s$ of 1.75 mmol·g\textsuperscript{-1}·h\textsuperscript{-1} was applied, indicated another factor being crucial for

| Experiment* | Methanol concentration | Specific substrate uptake rate $q_s$ [mmol·g\textsuperscript{-1}·h\textsuperscript{-1}] | Specific productivity $q_p$ [U·g\textsuperscript{-1}·h\textsuperscript{-1}] |
|-------------|------------------------|-------------------------------------------------|-------------------------------|
| Batch with methanol pulses | from 0 to 16 g/L | ~ 2 | 25 ± 0.25 |
| Fed batch A | limited | < 1 | 20 ± 0.20 |
| Fed batch B | limited | 1.75 | 5.0 ± 0.50 |
| Fed batch C1 | limited | stepwise up to 2 | 10 ± 0.10 |
| Fed batch C2 | limited | stepwise up to 2 | 11 ± 0.11 |

*all experiments were performed in the presence of different concentrations of the inducer methanol.
the specific productivity besides $q_s$. Therefore, the specific productivity $q_p$ was plotted against the induction time, to analyze possible time-dependent effects (Figure 7). Fed batch A and B were characterized by a linear increase of $q_p$ over time (fed batch B was characterized by a significant steeper slope), whereas fed batches C1 and C2 even showed an exponential increase of the specific productivity. It became obvious that the productivity was directly correlated to the induction time for all experiments conducted in this study, even in the batch experiments (Figure 2). Plantz et al. have recently reported an influence of the induction time on $q_p$ as well. They showed increasing product yields for the recombinant production of an interferon with $P$. pastoris in the first period of induction, but a subsequent decrease in product formation and a shift of the energy transfer towards biomass growth during the later phase of the cultivation [8]. In contrast to the study of Plantz et al., we clearly showed an increase of $q_p$ over the whole induction time for all experiments in this study (Figure 7).

Furthermore, fed batches C1 and C2 were characterized by an exponential increase of $q_p$ over time, in contrast to a linear increase for fed batches A and B (Figure 7). Obviously, a feeding strategy starting with a lower $q_s$ at the beginning of the induction phase, followed by a subsequent increase with a stepwise, dynamic feeding profile until $q_{s \text{ max}}$ (fed batch C1 and C2), was superior to a feeding strategy, where immediately after the $\Delta t_{\text{timeadapt}}$ a high $q_s$ was applied and kept constant (fed batch B). When the cells had the time to undergo a dynamic, physiological adaptation to the current culture condition before $q_s$ was increased, a boost in $q_p$ was observed, which probably cannot be triggered by constant or exponential feeding regimes (Table 2).

Summarizing, we showed a clear time dependent trajectory for specific product formation, as $q_s$ increased over induction time for all experiments in this study. A tool for the early identification of this effect is of utmost importance in order to consistently compare different feeding profiles.

Conclusions
In this study, we developed a fast and easy to do method based on batch cultivations with methanol pulses to determine strain specific parameters of a $P$. pastoris Mut3 strain expressing the enzyme horseradish peroxidase. A subsequently developed dynamic feeding strategy solely based on $q_s$, where the cells on the one hand had time to adapt to culture conditions, but were then challenged again repeatedly by a stepwise increase of $q_s$ up to $q_{s \text{ max}}$ resulted in the highest $q_p$ compared to the other strategies tested. Hence, dynamic feeding profiles turned out to be a valuable method to boost the specific productivity. This calls for increased use of dynamic process conditions even for industrial feed profiles. We strongly believe that the strategy presented here can be successfully applied on other microbial expression systems, which is why we are currently testing the applicability of our novel, dynamic approach on other expression systems, like $E$. coli, and other products.

Abbreviations
$\Delta t_{\text{timeadapt}}$: time for adaptation of the culture to the new substrate (methanol) [h]; $\mu$: specific growth rate [h$^{-1}$]; CER: carbon dioxide evolution
rate $[\text{mmol}\cdot \text{L}^{-1}\cdot \text{h}^{-1}]$; HRP: horseradish peroxidase; Mut$^+$: methanol utilization slow phenotype; PID: proportional-integrative-derivative controller; $q_m$: specific productivity of horseradish peroxidase $[\text{U}\cdot \text{g}^{-1}\cdot \text{h}^{-1}]$; $q_{\text{sub}}$: specific substrate uptake rate $[\text{mmol}\cdot \text{g}^{-1}\cdot \text{h}^{-1}]$; $q_{\text{w}}$: yield coefficient of methane; $C_{\text{m}}$: substrate uptake rate $[\text{mmol}\cdot \text{L}^{-1}\cdot \text{h}^{-1}]$; pm: rounds per minute; vvm: volume gas flow per volume liquid per minute; $Y_X/S$: yield coefficient of biomass respective to methanol $[\text{C}\cdot\text{mol}:\text{C}\cdot\text{mol}^{-1}]$.

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Authors’ contributions
CD designed and performed the experiments and analyzed data. OS performed some experiments. CD and OS wrote the paper, CH conceived the study and supervised research. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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