Pichia pastoris regulates its gene-specific response to different carbon sources at the transcriptional, rather than the translational, level

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Abstract

**Background:** The methylotrophic, Crabtree-negative yeast *Pichia pastoris* is widely used as a heterologous protein production host. Strong inducible promoters derived from methanol utilization genes or constitutive glycolytic promoters are typically used to drive gene expression. Notably, genes involved in methanol utilization are not only repressed by the presence of glucose, but also by glycerol. This unusual regulatory behavior prompted us to study the regulation of carbon substrate utilization in different bioprocess conditions on a genome wide scale.

**Results:** We performed microarray analysis on the total mRNA population as well as mRNA that had been fractionated according to ribosome occupancy. Translationally quiescent mRNAs were defined as being associated with single ribosomes (monosomes) and highly-translated mRNAs with multiple ribosomes (polysomes). We found that despite their lower growth rates, global translation was most active in methanol-grown *P. pastoris* cells, followed by excess glycerol- or glucose-grown cells. Transcript-specific translational responses were found to be minimal, while extensive transcriptional regulation was observed for cells grown on different carbon sources. Due to their respiratory metabolism, cells grown in excess glucose or glycerol had very similar expression profiles. Genes subject to glucose repression were mainly involved in the metabolism of alternative carbon sources including the control of glycerol uptake and metabolism. Peroxisomal and methanol utilization genes were confirmed to be subject to carbon substrate repression in excess glucose or glycerol, but were found to be strongly de-repressed in limiting glucose-conditions (as are often applied in fed batch cultivations) in addition to induction by methanol.

**Conclusions:** *P. pastoris* cells grown in excess glycerol or glucose have similar transcript profiles in contrast to *S. cerevisiae* cells, in which the transcriptional response to these carbon sources is very different. The main response to different growth conditions in *P. pastoris* is transcriptional; translational regulation was not transcript-specific. The high proportion of mRNAs associated with polysomes in methanol-grown cells is a major finding of this study; it reveals that high productivity during methanol induction is directly linked to the growth condition and not only to promoter strength.

**Keywords:** *Pichia pastoris*, Methylotrophic yeast, Crabtree-negative yeast, Polysome profiling, Microarray analysis, Transcriptome, Glucose repression, Carbon substrate repression, Methanol induction
Background

Pichia pastoris (syn. Komagataella sp.) is a methylo-trophic yeast that is widely used for the production of heterologous proteins and metabolites; it is also used as a model organism for the study of peroxisome biosynthesis and degradation, as well as for the analysis of protein secretion (see [1], and references therein). Its ability to use methanol as a carbon and energy source, its non-fermentative utilization of glucose and its efficient growth on glycerol are key metabolic features that make it attractive for bioprocess development.

Recently, Liang et al. [2] comprehensively annotated the P. pastoris transcriptome and identified novel untranslated regions (UTR), alternative splicing sites (AS), internal ribosome entry sites (IRES), upstream ATGs (uATGs) and upstream ORFs (uORFs). Transcriptional profiling of a recombinant strain harboring Rhizomucor miehei lipase (RML) under the control of the methanol-driven \( P_{\text{AOX1}} \) promoter revealed that cells grown on methanol induce genes involved in protein production and energy metabolism more than cells grown on glycerol. Methanol utilization takes place in peroxisomes; genes such as the alcohol dehydrogenases (\( \text{AOX1}, \text{AOX2} \)), formaldehyde dehydrogenase (\( \text{FDL} \)), dihydroxyacetone synthase (\( \text{DAS1}, \text{DAS2} \)) and peroxisomal genes (e.g. \( \text{PEXI} \)) were all found to be induced on methanol.

The specific growth rate of a culture, which was kept constant in the study by Liang et al. [2], is also known to play a fundamental role in gene regulation and consequently in protein production. High growth rates were previously suggested to be beneficial for protein production in \( P. \text{pastoris} \) due to the up-regulation of genes related to gene expression and translation, while catabolic processes (e.g. autophagy, transport to the peroxisome and mitochondrial degradation, many of them under the control of \( \text{TOR} \) signalling), were shown to correlate negatively with increasing growth rate [3].

Less is known about the specific regulation of carbon substrate utilization, with the notable exception of \( \text{Saccharomyces cerevisiae} \). Most studies in \( S. \text{cerevisiae} \) have been performed on glucose-grown cells under respiro-fermentative or fermentative growth conditions [4] or on non-fermentable carbon-sources such as glycerol or galactose. The shift from glucose to glycerol leads to extensive transcriptomic remodelling [5], a global translational down-regulation [6] and reduced growth rates. In contrast, the Crabtree-negative yeast, \( P. \text{pastoris} \), maintains its respiratory metabolism even under conditions of excess glucose (such as that used in batch cultivations) and exhibits similar growth rates and substrate uptake kinetics when grown on either glucose or glycerol [7]. Shifts from glycerol to methanol, which is metabolized even more slowly with lower maximal specific growth rates, are often used in bioprocesses that employ \( P. \text{pastoris} \).

Transcriptional regulators involved in glucose repression have been identified and studied in the methylo-trophic yeasts \( P. \text{pastoris} \) and \( \text{Hansenula polymorpha} \), and in the lactose-utilizing yeast \( \text{Kluyveromyces lactis} \) [8-13]. Glucose repression of methanol utilization genes is established as a feature of methylo-trophic yeasts such as \( \text{Candida boidinii}, \text{H. polymorpha}, \text{Pichia methanolica}, \) and \( P. \text{pastoris} \) [14], but the degree of repression/de-repression by different carbon sources is species-dependent. For example, different modes of regulation have been described for key enzymes of methanol metabolism pathways such as alcohol oxidase, dihydroxyacetone synthase and formaldehyde dehydrogenase (summarized in [14,15]). Understanding the molecular mechanisms underpinning the unique carbon substrate utilization properties of \( P. \text{pastoris} \) is now required in order to more fully understand this valuable host organism.

The regulation of gene expression is often analyzed at the level of transcription, although it is well established that altered transcript levels are not necessarily reflected by the corresponding protein levels [16]. For example, the protein level of more than 70% of \( S. \text{cerevisiae} \) protein-coding genes is transcriptionally regulated, but this drops to only about 50% in \( E. \text{coli} \) [17] and is even lower in humans [18]. In order to obtain a more complete view of the regulation of gene expression in \( P. \text{pastoris} \), we analyzed both transcriptional and translational responses of cells grown in glucose-, glycerol- or methanol-containing media. Microarray analysis was done on the total mRNA pool as well as on mRNAs that had been fractionated based upon ribosome occupancy. We adapted published methods for pol-some profiling [6,19]; translationally quiescent mRNAs were defined as being associated with single ribosomes (monosomes); actively-translated mRNAs with multiple ribosomes (polysomes) [20]. The hybridization of a microarray with these mRNA fractions as well as the total mRNA population provided insight into how efficiently individual mRNA translation and global transcriptional responses are affected by carbon source utilization.

Results and discussion

\( P. \text{pastoris} \) strain X-33 was cultivated in shake flasks under four different bioprocess conditions (Table 1): excess glycerol or glucose (batch culture conditions; these cells were harvested during exponential growth); limiting glucose (using slow glucose-releasing silica disks or feed beads in fed-batch mode, [21,22]); and periodic methanol addition (methanol induction conditions). Cells grown in excess glucose or glycerol or those grown in methanol had growth rates close to \( \mu_{\text{max}} \): 0.23 h\(^{-1}\) for the former and 0.1 h\(^{-1}\) for the latter conditions. Cells in limiting glucose conditions grew at \( \mu = 0.015 \text{ h}^{-1}\).

For polysome fractionation, cells were treated with cycloheximide, harvested and quickly chilled for sample preparation. Isolates were used for pol-some profiling to
obtain the profile data and to collect mono- and polysome fraction samples for mRNA extraction. mRNA was isolated from the fractionated and unfractionated isolates for microarray analysis; for each condition three biological replicates were analyzed.

The excess glucose condition, which is often used as a control for studies in *S. cerevisae*, was used as a control in our experiments.

Global transcript profiles are very similar for excess glucose or glycerol grown *P. pastoris* cells, while extensive transcriptional regulation is observed for cells grown on methanol or limiting glucose concentrations. Differentially expressed genes were identified from fold changes between total RNA samples (i.e. those from unfractionated isolates). Samples from the excess glucose condition were the control for all these experiments (cut-off criteria ±50% fold change and adjusted p-values < 0.05; [23]). Transcriptional fold changes for all genes are listed in addition file 1: Table S1. The data in Figure 1 show that cells cultured in excess glycerol (G) or glucose (D) have a very similar transcriptome with just 265 genes differentially regulated; in contrast 817 genes are differentially regulated in methanol-grown cells (M) and 2,822 are differentially regulated in glucose-limited cells (X) (Figure 1A). The corresponding Gene Ontology (GO) terms are listed in Additional file 2. A high correlation between the two excess carbon source condition transcriptomes (G and D) was also observed by principal component analysis (PCA), which showed a good correlation of the biological replicates of each condition (Figure 2). The methanol-grown and glucose-limited cells were also found to share many differentially-regulated genes and hence seem to be more similar to each other than to the two excess conditions (Figure 1B).

Further analysis (Figure 1B, C) revealed that only a small sub-set of genes are differently expressed in response to glycerol as carbon source (10% of the 148 up-regulated and 15% of the 114 down-regulated genes), while most of the regulated genes are shared either with both (56%) or at least one (approx. 30%) of the two other conditions (methanol induction or limiting glucose). We defined genes that are differentially regulated in excess glycerol conditions plus at least one other condition (either methanol induction or limiting glucose) to be subject to “glucose repression”. Genes that are differentially regulated in

### Table 1 *Pichia pastoris* cultivations in buffered synthetic media supplemented with different carbon substrates

| Condition          | ID | Start-OD<sub>600</sub> | Cultivation substrate | Cultivation time [h] | Harvest-OD<sub>600</sub> μ [h<sup>-1</sup>] | Bioprocess Step                  | Replicates |
|--------------------|----|------------------------|-----------------------|-----------------------|---------------------------------------------|----------------------------------|------------|
| Excess glucose     | D  | 0.1                    | 2% glucose            | 23.3                  | 10.0 (1.0)                                  | 0.23 (0.004)                     | Glucose batch | 3          |
| Excess glycerol    | G  | 0.1                    | 2% glycerol           | 23.3                  | 10.5 (1.3)                                  | 0.23 (0.001)                     | Glycerol batch | 3          |
| Methanol feed      | M  | 1.5                    | 0.5 and 0.6% methanol | 24.5                  | 8.6 (1.4)                                   | 0.10 (0.008)                     | Methanol shot/feed | 3          |
| Limiting glucose   | X  | 1.5                    | 0.25% glucose and feed beads | 16.8                  | 11.4 (0.6)                                  | 0.010 - 0.022                    | Glucose fed batch | 3          |

Cultures with different biomass densities were fed with appropriate amounts of carbon substrate in order that the cells could be harvested at a similar OD<sub>600</sub> (mean (sd)). Growth rates (μ) (mean (sd)) were recorded; the values were highly reproducible and reflect growth of typical bioprocess phases, as shown.

**Figure 1** Differentially expressed genes. The bar chart (A) shows the number of differentially expressed genes in excess glycerol (G), methanol (M) and limiting glucose (X) compared to the excess glucose condition. Venn diagrams illustrate the number of up-regulated (B) and down-regulated genes (C) in the conditions and intersections. Significantly-regulated genes were identified from total RNA fold changes compared to the excess glucose condition (cutoff ±50% fold change and adjusted p-values < 0.05; [23]).
response to methanol induction or limiting glucose conditions, but are not differentially regulated between the two excess conditions were defined as being subject to “carbon substrate repression”.

**Polysome-mRNA association is lowest in glucose-limited cells and highest in methanol-grown cells**

Isolates of cells subject to the different growth conditions in Table 1 were analyzed by polysome profiling, which characterizes the translational status of a cell according to the distribution of ribosomes across the mRNA pool. Profile curves showing the proportion of ribosomes that appear as individual sub-units (40S and 60S), monosomes or polysomes (where two or more ribosomes are associated with a given mRNA transcript) are shown in Figure 3. The ratios of the polysome to monosome peak areas (P:M ratios) in the profiles (Figure 3A) are presented in Figure 3B: mRNAs that are associated with polysomes are more highly translated than mRNAs associated with monosomes [20]. The P:M ratio is therefore established as a relative measure of translational activity at a cellular level [24,25]. In our experiments, triplicate cultures gave reproducible values for each of the different growth conditions.

Due to their similar transcript profiles, the two fastest growing conditions (excess glycerol and excess glucose, $\mu \sim 0.23 \text{ h}^{-1}$) were anticipated to have similar P:M ratios. However, the excess glycerol condition had a higher P:M ratio (Figure 3) suggesting higher translational activity compared to cells grown under conditions of excess glucose. The P:M ratio was highest in cells grown on methanol, although the specific growth rate was significantly lower ($\mu \sim 0.10 \text{ h}^{-1}$) compared to the excess glycerol and excess glucose conditions. The condition with the lowest specific growth rate (limiting glucose, $\mu \sim 0.015 \text{ h}^{-1}$) had the lowest P:M ratio.

The transcription of translation-related genes in *P. pastoris* was previously shown to be tightly connected to growth rate in glucose-limited chemostat cultivations [3]. We found that this was also true when we analyzed the total RNA of unfractionated, slow-growing cells cultivated under limiting glucose conditions ($\mu \sim 0.015 \text{ h}^{-1}$). Under these conditions, most ribosomal and translation-related genes were found to be expressed at a lower level (Additional file 1: Table S2). Strikingly, we found that those genes were equally expressed in slow-growing methanol fed cells ($\mu \sim 0.1 \text{ h}^{-1}$) compared to excess glucose and glycerol ($\mu \sim 0.23 \text{ h}^{-1}$), suggesting that the whole translation machinery is up-regulated despite the slow growth rate on methanol. The methanol induction-, excess glucose- and excess glycerol- conditions operated near $\mu_{\text{max}}$ for their respective condition, which means that they possess a similar $\mu/\mu_{\text{max}}$ ratio. Hence, the expression of growth-associated genes might respond to the ratio of $\mu/\mu_{\text{max}}$ rather than an absolute value of the specific growth rate ($\mu$).

**Despite the general transcriptional down-regulation of translation-related genes in *P. pastoris* cells grown in limiting glucose, the transcription of certain genes is induced**

Certain genes required for ribosome biogenesis and its regulation, RNA processing and translationally silent messenger ribonucleoprotein complexes (mRNPs) were highly expressed in *P. pastoris* cells grown in limiting glucose, as
determined by the analysis of total mRNA (Additional file 1: Table S2): RPS22A (protein component of the small (40S) ribosomal subunit, homologous to mammalian ribosomal protein S15A and bacterial S8, also up-regulated in methanol-fed cells); genes linked to ribosome association, interaction or biogenesis (TMA108, DOT6, GDE1, TMA64, PAS_FragB_0030, YMR295C, MTC1, YOR019W, MTG1); negative regulation of RNA polymerase III transcription and TOR signaling (KNS1); RRPE (ribosomal RNA processing element)-binding and glucose-induced transition from quiescence to growth (STB3); rRNA biogenesis (DOT6) and mitochondrial ribosome recycling (RRF1). Poly(A)-binding protein is also translation-associated, and the two genes are differently expressed (PAS_chr1-4_0283 is up- and PAB1 is down-regulated) in P. pastoris cells grown in limiting glucose. The gene encoding the translational activator GIS2 that was also up-regulated in limiting glucose, plays an important role as activator of mRNAs with internal ribosome entry sites [26]. It binds to a specific subset of mRNAs, associates with polysomes and localizes to RNA processing bodies (P bodies) and to stress granules. The role of cap-independent translation in physiological adaptation to stress in S. cerevisiae has been reported previously [27]. P bodies are used to store transcriptionally silent mRNPs [28], and glucose-limited P. pastoris cells were found to differentially express related genes. DHII1 (the gene product of which functions in de-capping and translational rePRESSION) was up-regulated, but PAT1 and EDC3, with a similar function, were down-regulated in glucose-limited cells. Hence, although limiting glucose decreases global translation, certain transcripts may be translated as a part of specific stress responses.

Growth conditions have a minimal influence on transcript-specific translational regulation

We next examined the fractionated mRNAs by microarray analysis. We normalized the abundance of each transcript in the polysome fraction to that of the total RNA, which we termed the “translational state”. In order to confirm the integrity of the RNA fractions, microarray signal intensities of the monosome, polysome and total RNA samples from the limiting glucose condition were compared as previously described [29]. The log10 intensity values of total RNA correlated with log10 of the sums of intensities in the monosome- and polysome-bound mRNA with a correlation coefficient of R2 = 0.963 (see Additional file 3). Translational states of individual transcripts for the excess glycerol, limiting glucose and methanol induction conditions were normalized to the excess glucose condition in order to identify transcripts with changed translational states (shown in Figure 4 and Additional file 4). This identified an increased or decreased abundance of transcripts that are actively translated in the polysome fraction. Translational states of individual genes ranged from 0.08-fold (in limiting glucose conditions) to 3.05-fold (in methanol). No transcripts were totally excluded from the polysome fractions, which is in agreement with a study published by Arava et al. [30].

Only 16 transcripts had different translational states (8 increased and 8 decreased) in response to excess glycerol compared to the excess glucose condition, while more differences were found for the glucose-limited and methanol-grown cells. In excess glycerol-grown cells, RPL2A, TEF2, RPS4B, ENO1, FBA1-1, RPL5, RPL11B and TDH3 had decreased translational states compared to cells grown in excess glucose. These genes are annotated with GO terms “biosynthetic/metabolic process” and “translation”. Both, the transcript level and the translational state was found to be decreased for transcripts of the glycolytic fructose 1,6-bisphosphate aldolase (FBA1-1), glyceraldehyde-3-phosphate dehydrogenase (TDH3) and phosphopyruvate hydratase (ENO1) in excess glycerol.
This suggests that specific translational down-regulation reinforces the transcriptional down-regulation of these genes in response to excess glycerol.

In methanol-grown cells, genes required for methanol utilization (MUT), were strongly up-regulated at the transcriptional level, but had a decreased translational state compared to excess glucose. Hence translational regulation appears to counteract the strong transcriptional up-regulation of most of these genes. Such "post-transcriptional buffering" has also been observed in two Saccharomyces species [31]. Neither significantly enriched GO terms nor other patterns could be found in the other gene groups with altered translational states.

Translational states are linked to ORF length and transcript abundance

We analyzed the translation states of individual transcripts compared to total mRNA for all growth conditions. Enriched gene groups were initially identified (Table 2); closer inspection revealed that the groups had closely correlated open reading frame (ORF) lengths, which has been reported previously for other organisms [32-34]. Liang et al. [2] identified P. pastoris gene ORFs, uORFs, UTRs and introns by sequencing, and found ORF lengths from 141 to 14853 bp, with an average of 1444 ± sd = 1032 bp (median of 1203 bp). We used this information to define three gene groups according to ORF length (Table 3): long and short genes, comprising the upper and lower quartile of all genes, and the remaining 50% of medium-length genes. Translation efficiency is also known to be affected by codon usage, so we included synonymous codon usage order (SCUO), which was obtained from the CodonO platform [35]; higher values indicate more codon bias, meaning less random codon use in a gene’s coding region. The three gene groups significantly differ in transcript level, translational states, codon usage bias (SCUO) and 5’UTR frequency: Short genes are highly transcribed (as measured by transcript abundance) and translated (high translational states), rarely possess a 5’UTR and have an enhanced codon usage bias (Table 3).

Statistical tests (Fishers exact test, chi square test and regression analysis) were used to verify these relationships.

Table 2 Translational regulation of functional gene groups for P. pastoris cells grown in excess glucose conditions

| Functional group | Genes in group | Significantly regulated genes | Average translational log₂ ratio of significantly regulated genes | Average ORF length of significantly regulated genes [bp] |
|-----------------|---------------|-------------------------------|-----------------------------------------------------------------|----------------------------------------------------------|
| Secretion: chaperones | 79 | 31 | 0.225 | 885 |
| Antioxidant | 21 | 7 | 0.160 | 476 |
| Transport(er) | 60 | 22 | 0.137 | 1669 |
| Pexophagy | 23 | 9 | −0.082 | 2302 |
| Autophagy | 69 | 25 | −0.117 | 1690 |
| Vacuole | 105 | 48 | −0.151 | 1781 |
| Mitochondria | 110 | 23 | −0.165 | 1541 |
| TCA | 20 | 10 | −0.339 | 1544 |
| Secretion: glycosylation | 46 | 28 | −0.344 | 1884 |

Average translational states and ORF length of functional gene groups for P. pastoris cells grown in excess glucose. Translational trends were similar in the other conditions.
ORF length was shown to have a negative correlation with transcript abundance (gene expression intensity by microarray) and codon usage bias, so short genes are more highly transcribed than longer ones (regression analysis, p-value < 1.5e−11) and more codon biased (non-linear regression, p-value < 2.2e−16). The correlation of ORF length with translational states and 5′UTR length was found to be significantly positive (p-value < 2.2e−16 for both). Hence, short genes are more highly translated and rarely have a 5′UTR, while longer genes are less highly translated and often possess a 5′UTR (Figure 5).

Transcriptional regulation responding to different carbon sources correlates with expression of corresponding transcription factors

As mentioned above, excess glucose was used as a calibrator to calculate the transcriptional regulation in the other conditions (see Additional file 1: Table S1 for respective values for all genes). Concerning global transcriptional control systems, we could identify P. pastoris gene expression responding to glucose repression, carbon catabolite repression elicited by excess glucose and glycerol, as well as control by methanol availability. Limiting glucose triggers extensive transcriptional responses due to carbon limitation and low growth rate, which correlate well with the regulation patterns described by Rebnegger et al. [3] recently. Corresponding to the important role of glycogen metabolism in slow growing conditions [36], we found genes encoding glycogen synthase (GSY2), phosphoglucomutase (PGM2) and other glycogen metabolism genes (UGP1, NTH1, ATH1, GLG1, GLC3, GLC7) up-regulated in limiting glucose.

Glucose repression signalling is mainly mediated through the central kinase Snf1, which controls the expression of important transcription factors such as Mig1, Sip4, Rds2, Cat8 and Adr1 [37], thereby playing an important role in the utilization of non-fermentable carbon sources in S. cerevisiae [38]. We found the transcripts of many genes involved in catabolite (de)repression to be induced in limiting glucose, especially CAT8-2, which is about 39-fold up-regulated compared to excess glucose (and about 7-fold up-regulated on methanol). In addition, almost all genes that are reported to be controlled by CAT8 in S. cerevisiae [39] are also up-regulated.

Interestingly, 2 homologs of Mig1 are found in the P. pastoris genome, one of which is about 9-fold up-regulated in response to methanol and limiting glucose (MIG1-1), while the second one is down-regulated on all other tested carbon sources compared to glucose (MIG1-2);
it is possible that it acts as a carbon catabolite or glucose repressor similar to CRE1 in *Trichoderma reesei* [40] or CREA in *Aspergillus nidulans* [41].

The homologue of *S. cerevisiae* Activator of Ferrous Transport, AFT1, was found to have induced expression levels in excess glycerol, methanol and limiting glucose conditions and has been reported to play a role in the regulation of carbon repressed genes in *P. pastoris* recently [42]. The transcription factors PAS_chr4_0324, CTH1, PAS_chr1-1_0422, PAS_chr3_1209, PAS_chr1-1_0122 were related to excess conditions.

Among the most strongly-induced genes in methanol and limiting glucose conditions, several transcription factors are present (Table 4). Of these, the Zn(II)2Cys6 zinc cluster protein PAS_chr3_0836, which has an 80-fold higher transcript level on methanol and 120-fold higher transcript level under limiting glucose compared to excess glucose, has significant sequence homology to *H. polymorpha* MPP1 [43]. Mpp1 was suggested to be the master regulator of methanol-responsive genes in *H. polymorpha* [43,44]. Since PAS_chr3_0836 is also located in a similar chromosomal arrangement (next to *DAS1/2; PAS_chr3_0832 and PAS_chr3_0834*) to *H. polymorpha*, we propose that it is the *P. pastoris* homologue of *H. polymorpha* Mpp1. PpMXR1 encoding a transcription factor that is necessary for the activation of many genes in response to methanol [8] is induced in all three conditions compared to excess glucose. We suggest that PpMXR1, similar to its *S. cerevisiae* homolog ADRI, is needed for the activation (de-repression) of genes for alternative carbon sources including the MUT genes that are repressed in the presence of excess glucose and glycerol, but that Mpp1 is the transcriptional activator of peroxisomal import and matrix proteins required for methanol utilization in *P. pastoris*. This awaits experimental verification in future.

Other previously-characterized transcription factors acting on methanol metabolism, *ROP* (repressor of phosphoenolpyruvate carboxykinase; PAS_chr3_0554, [10]) and *TRM1* (positive regulation of methanol, PAS_chr4_0203) are induced only on methanol, but not on limiting glucose, confirming their specific involvement in methanol metabolism (reviewed by [15]).

**Glucose and carbon catabolite repression regulate the expression of genes involved in glycolysis, gluconeogenesis and the metabolism of alternative carbon sources**

The expression of genes related to carbon source uptake and initial metabolism is strongly regulated at the level of transcription. The respective transcriptional control of genes such as glucose sensors and transporters (low- and high-affinity), hexokinase, and glycerol- and methanol utilization are shown in Table 5 and Figure 6.

We found glycolytic *P. pastoris* genes involved in upper and lower glycolysis to have lower expression levels in all three conditions compared to excess glucose. Glycolytic genes are known to be weakly regulated at the level of transcription in *S. cerevisiae* [45], but transcriptional regulation has been previously described for Crabtree-negative yeasts such as *P. pastoris* and *K. lactis*, and was assumed to coincide with their limited glucose uptake [46,47]. As expected, the genes encoding the key gluconeogenic enzymes fructose-1,6-bisphosphatase (*FBP1*) and PEP carboxykinase (*PCK1*) are less expressed in excess glucose (compared to the other conditions). The transition between those two pathways is associated with *Gid2/Rmd5*-dependent ubiquitin-proteasome linked elimination of the key enzyme fructose-1,6-bisphosphatase [48]. *Vid24/Gid4*, encodes a previously-identified key regulator of *GID2/Rmd5* that is strongly up-regulated in methanol fed cells. A hypothetical gene (PAS_chr1-1_0399), also strongly induced on methanol and limiting glucose, could encode the homolog of Rmd5; it contains a C3HC4 RING finger domain.

In *S. cerevisiae*, fermentative glucose- or catabolite-repressed growth is described for cells grown on excess glucose [49]. Upon glucose depletion or in the presence of non-fermentable carbon sources, such as glycerol or ethanol, extensive reprogramming of gene expression allows *S. cerevisiae* to take up alternative carbon sources and enhances activity of the glyoxylate cycle, the tricarboxylic acid (TCA) cycle and gluconeogenesis [5]. We found several *P. pastoris* genes encoding enzymes involved in the metabolism of alternative carbon sources to be less expressed during growth on glycerol, methanol and/or limiting glucose (Additional file 1: Table S3 and Figure 6). Among them, the non-annotated ORFs PAS_chr4_0338, PAS_chr4_0339 and PAS_chr4_0341 could be identified to be homologs of *LRA1*, 2 and 4. The encoded enzymes are part of the alternative pathway of L-rhamnose catabolism present in *Pichia* (*Scheffersomyces*) *stipitis* [50] and most probably allow *P. pastoris* to utilize rhamnose as sole carbon source [51]. Interestingly, *PpLRA2* and 4 flank an uncharacterized fungal-specific Zn2/Cys6 transcription factor (PAS_chr4_0340), which is up-regulated in response to methanol and limiting glucose (Table 4). Increased transcript levels in comparison to excess glucose can also be seen for many TCA cycle genes, isocitrate lyase (*ICL1*) involved in the glyoxylate cycle (Figure 6) and genes involved in channeling alternative carbon sources into the TCA cycle (e.g. the cytosolic aldehyde dehydrogenase isoforms *ALD4-2* and PAS_chr4_0470). Interestingly, some genes encoding proteins present as isoenzymes such as *ACO1/2*, *IPD1/2* and *ACS1/2* are oppositely regulated in all the de-repressed conditions.

Respiration is repressed in excess glucose conditions during fermentative growth in *S. cerevisiae* [5,52,53], thus respiration-associated functions such as oxidative phosphorylation, mitochondrial electron transport and ATP generation are induced upon glucose depletion. Unlike
**Table 4 Transcriptional regulation of transcriptional regulators**

| Short name Pp    | Description                                                                 | G-D logFC | G-D adjPV | M-D logFC | M-D adjPV | X-D logFC | X-D adjPV |
|------------------|------------------------------------------------------------------------------|-----------|-----------|-----------|-----------|-----------|-----------|
| PAS_chr4_0340    | Fungal specific transcription factor domain; Zn2/Cys6 DNA-binding domain     | 0.35      | *         | 0.72      | ***       | 1.50      | ***       |
| CAT8-2           | Zinc cluster transcriptional activator; necessary for derepression of a variety of genes under non-fermentative growth conditions in *S. cerevisiae* | −0.07     | 2.72      | ***       | 5.27      | ***       |           |
| YAP1             | Basic leucine zipper (bZIP) transcription factor; required for oxidative stress tolerance | 0.27      | 1.13      | ***       | 1.64      | ***       |           |
| PAS_chr1-4_0516  | Putative transcription factor                                               | 0.94      | 7.81      | ***       | 7.86      | ***       |           |
| MPP1             | Fungal Zn2/Cys6 DNA-binding domain; homolog to *Hansenula polymorpha*        | 0.90      | ***       | 6.34      | ***       | 6.99      | ***       |
| AFT1             | Transcription factor, possibly involved in carbohydrate metabolism           | 2.17      | ***       | 3.68      | ***       | 5.16      | ***       |
| YPR022C-3        | Putative transcription factor                                               | 1.57      | ***       | 2.33      | ***       | 4.55      | ***       |
| PAS_chr3_048     | Helix-loop-helix DNA-binding domain                                         | 0.06      | 0.29      |           | 3.69      | ***       |           |
| ADR1/MXR1        | Carbon source-responsive zinc-finger transcription factor, required for transcription of the glucose-repressed gene ADH2, of peroxisomal protein genes, and of genes required for ethanol, glycerol, and fatty acid utilization | 1.34      | ***       | 1.61      | ***       | 2.16      | ***       |
| RSF2/ROP         | Zinc-finger protein; involved in transcriptional control of both nuclear and mitochondrial genes in *S. cerevisiae* | −0.10     | 1.85      | ***       | −0.24     |           |           |
| PpTRM1           | Zn(II),Cys6-type transcription factor involved in the positive regulation of methanol utilization genes in P. pastoris and C. boidinii | −0.14     | 0.74      | ***       | 0.34      | *         |           |
| SNF1             | AMP-activated serine/threonine protein kinase; found in a complex containing Snf4p and members of the Sip1p/Sip2p/Gal83p family; required for transcription of glucose-repressed genes, thermotolerance, sporulation, and peroxisome biogenesis in *S. cerevisiae* | 0.39      | **        | 0.61      | **        | 1.42      | ***       |
| SNF2             | Catalytic subunit of the SWI/SNF chromatin remodeling complex; involved in transcriptional regulation; contains DNA-stimulated ATPase activity | 0.13      | 0.40      | **        | −0.37     | **        |           |
| SNF4             | Activating gamma subunit of the AMP-activated Snf1p kinase complex           | 0.19      | 0.35      |           | 0.76      | ***       |           |
| MIG1-1           | Transcription factor involved in glucose repression in *S. cerevisiae*; regulated by the Snf1p kinase and the Gic7p phosphatase; | 0.57      | *         | 1.09      | **        | 3.09      | ***       |
| MIG1-2           | Transcription factor involved in glucose repression in *S. cerevisiae*; regulated by the Snf1p kinase and the Gic7p phosphatase; | −0.76     | **        | −1.23     | ***       | −0.56     | ***       |
| SIP2             | One of three beta subunits of the Snf1 kinase complex in *S. cerevisiae*    | 0.00      | −0.14     |           | 0.65      | ***       |           |
| RDS2             | Transcription factor involved in regulating gluconeogenesis and glyoxylate cycle genes; member of the zinc cluster family of proteins; confers resistance to ketoconazole in *S. cerevisiae* | −0.07     | 0.20      |           | 0.83      | ***       |           |
| PAS_chr1-3_0274  | Fungal specific transcription factor; Zn2/Cys6 DNA-binding domain            | 0.11      | 0.29      |           | 0.90      | ***       |           |
| PAS_chr4_0324    | Fungal specific transcription factor; Zn2/Cys6 DNA-binding domain            | −3.07     | ***       | −2.99     | ***       | −3.47     | ***       |
| CTH1             | Member of the CCH zinc finger family                                         | −2.54     | ***       | −2.81     | ***       | −2.92     | ***       |
| PAS_chr1-1_0422  | Myb/SANT-like DNA-binding domain                                            | −0.13     | −0.57     |           | −2.56     | ***       |           |
| PAS_chr3_1209    | Helix-loop-helix DNA-binding domain                                         | 0.16      | −0.21     |           | −2.56     | ***       |           |
| PAS_chr1-1_0122  | Helix-loop-helix DNA-binding domain                                         | −0.93     | −0.57     |           | −2.33     | ***       |           |

Log2 fold changes and adjusted P-values (* adjPV < 0.1; ** adjPV < 0.05; *** adjPV < 0.01) are shown (see Additional file 1: Table S1 for detailed data). Up-regulated genes are in bold letters, down-regulated genes in bold and italics.

*S. cerevisiae*, Crabtree-negative yeasts are dependent on respiratory processes even in excess glucose. Consequently, the expression of mitochondrial genes is not induced in the presence of non-fermentable carbon-sources in *P. pastoris* (Additional file 1: Table S4). However, several subunits of respiratory complex I [54], which is not present in *S. cerevisiae*, appear to be de-repressed.

Methanol utilization and peroxisomal genes are subject to carbon substrate repression

Unexpectedly, the transcript levels of most genes involved in methanol utilization (MUT) are not only highly induced in methanol-grown cells but also in glucose-limited cells (Table 6). The transcript level of *AOX1* is almost equally high in both conditions. This observation correlates well
with pre-induction expression from the AOX1 promoter in the glycerol-fed batch prior to methanol addition [55-57], and high Aox1 protein levels in glucose-limited chemostats [58,59]. Repression of AOX1 expression was previously determined in P. pastoris grown on glucose, glycerol, ethanol and acetate [60], with glycerol repression being specific for P. pastoris AOX1/2, but not for alcohol oxidase genes in related yeasts such as H. polymorpha or C. boidinii [14].

Although it was assumed that some MUT genes might also be regulated by catabolite de-repression [15], the extent of this regulatory pathway has not been shown experimentally in P. pastoris. Early observations reported that the mRNA levels of AOX1 upon de-repression was only 1-2% of the methanol-induced mRNA levels [61], while FLD expression was assumed not to be under glucose repression control [62]. On the contrary we see a high level of de-repression in cells grown on limiting glucose (Table 6). This contradiction might be explained by the fact that in our set up, the cells are actively growing, while previous experiments employed glucose-exhausted stationary-phase cells for studies of de-repression. Upon (constant)

| Short name | Description | G-D logFC | G-D adjPV | M-D logFC | M-D adjPV | X-D logFC | X-D adjPV |
|------------|-------------|-----------|-----------|-----------|-----------|-----------|-----------|
| PpHKT1     | P. pastoris major low affinity glucose transporter (major facilitator superfamily) | −1.31 |  −3.34 | *** | −0.82 | * |
| ITR2       | Myo-inositol transporter | −0.40 | * | −0.88 | * | −0.62 | *** |
| PAS_c034_0021 | Major facilitator superfamily, related to STL1 | −0.59 | ** | 0.10 | 0.55 | *** |
| PAS_chr2-1_0006 | Major facilitator superfamily, Quinate permease (Quinate transporter) - similar to S. stipitis | −0.06 |  −0.80 | 0.01 |
| YBR241C    | Putative transporter, member of the sugar porter family | 0.12 |  −0.16 | 0.26 |
| PpHKT2     | P. pastoris putative low affinity glucose transporter of the major facilitator superfamily | −0.10 |  −0.10 | −0.09 |
| STL1-1     | Glycerol proton symporter of the plasma membrane, subject to glucose-induced inactivation in S. cerevisiae | 0.08 |  −0.11 | 1.23 | *** |
| STL1-2     | Glycerol proton symporter of the plasma membrane, subject to glucose-induced inactivation in S. cerevisiae | −0.27 |  0.40 | 2.08 | *** |
| SNF3       | P. pastoris plasma membrane glucose sensor Gss1, regulates glucose transport | 0.16 |  0.44 | 1.60 | *** |
| PAS_chr3_1076 | Glycerol proton symporter of the plasma membrane, related to RGT2 | 0.37 |  0.65 | ** | 0.62 | ** |
| PAS_chr3_1099 | Glycerol proton symporter of the plasma membrane, related to STL1 or RGS2 | 0.34 |  0.80 | ** | 1.33 | *** |
| MAL31      | Maltose permease, high-affinity maltose transporter (alpha-glucoside transporter) | 0.09 |  0.81 | *** | 0.68 | *** |
| GTH1       | P. pastoris major high affinity glucose transporter; similar to K. lactis HGT1 | 0.17 |  1.09 | *** | 6.14 | *** |
| PpHGT1     | P. pastoris high affinity glucose transporter - similar to K. lactis HGT1 | 0.59 |  0.86 | ** | 4.91 | *** |
| PAS_chr4_0828 | Myo-inositol transporter with strong similarity to the major myo-inositol transporter Itr1p, member of the sugar transporter superfamily | 2.35 |  *** | 3.65 | *** | 7.30 | *** |
| HXX1       | Hexokinase isoenzyme 1; a cytosolic protein that catalyzes phosphorylation of glucose during glucose metabolism; expression in S. cerevisiae is highest during growth on non-glucose carbon sources | 0.30 |  −0.21 | 1.69 | *** |
| HXX2       | Hexokinase isoenzyme 2; catalyzes phosphorylation of glucose in the cytosol; predominant hexokinase during growth on glucose in S. cerevisiae | −0.12 |  0.18 | 0.03 |
| GLK1       | Glucokinase; catalyzes the phosphorylation of glucose at C6; expression regulated by non-fermentable carbon sources in S. cerevisiae | −0.99 |  ** | −2.58 | *** | −0.34 |

Log2 fold changes and adjusted P-values (* adjPV < 0.1; ** adjPV < 0.05; *** adjPV < 0.01) are shown (see Additional file 1: Table S1 for detailed data). Up-regulated genes are in bold letters, down-regulated genes in bold and italics.
Figure 6 (See legend on next page.)
methanol addition e.g. in fed batch or chemostat, MUT gene transcript levels are on average 55-fold higher compared to glucose-limited growth conditions (unpublished data). However, our data highlight that different degrees of carbon catabolite repression are acting on individual MUT genes; for example DAS1/2 are less de-repressed than AOX1/2. This strongly points towards – yet unidentified – transcriptional regulators being involved in induction/repression of the individual MUT genes in addition to the global methanol regulator PpMXR1 (summarized by [15]). Induction of Peroxisomal protein synthesis was observed in S. cerevisiae grown on glycerol as sole carbon source [5], which appears to be different from the situation in P. pastoris. In the present study, up-regulation of peroxisomal gene transcript levels occurs in glucose-limited and methanol-grown cells but not in excess glycerol (Table 6), which may also be associated with the specific repression exerted by glycerol on MUT gene expression; it might be speculated that the zinc cluster protein Cat8-2 (Table 4) is the responsible transcription factor for this.

Peroxisomal processes such as methanol utilization and beta-oxidation are associated with the formation of H$_2$O$_2$, requiring the action of antioxidants. YAPI, the oxidative stress response transcription factor, and many of its target genes [63] were found to be significantly up-regulated in methanol-grown cells and/or more pronounced in limiting glucose. While it was previously shown that Yap1 is required for ROS detoxification and sufficient growth on methanol [64], the strong up-regulation of YAPI in glucose-limited conditions was unexpected. Interestingly, starvation is linked to the expression of genes encoding oxidative stress functions in bacteria and yeast [65,66]. The protective effect of antioxidants is proposed to have a beneficial effect in cells with nutrient limitation.

The expression of fatty acid beta-oxidation genes is up-regulated in P. pastoris cells responding to limiting glucose

Peroxisomal protein expression and fatty acid oxidation were previously reported to be regulated by Snf1 kinase through Adr1 action [67,68]. At least three other transcription factors act in concert with Adr1 in S. cerevisiae [68], but two of them – Oaf1 and Pip2 – cannot be found in P. pastoris. Instead, the putative fungal specific transcription factor PAS_chr1-3_0274 (Zn2/Cys6 domain) represents a homolog to FarA/B, the transcriptional activators of fatty acid utilization in Aspergillus spp., and C. albicans and Y. lipolytica Ctf1 [69]. The elevated transcript levels of PAS_chr1-3_0274 in limiting glucose are reflected by the strong induction of fatty acid utilization genes (e.g. all genes involved in beta-oxidation FAA2, FOX2, POT1, POX1, ECI1, SPS19, PA1 and PA2 have on average 100-fold higher transcript levels in limiting glucose, while only having approximately 2-fold higher transcript levels on methanol or glycerol in comparison to excess glucose). A similar regulation pattern was also observed for the non-annotated genes PAS_chr2-1-0249, PAS_FragB_0022, PAS_chr2-2_0403 and PAS_chr1-1_0108, indicating a possible involvement in beta-oxidation. Indeed, PAS_FragB_0022, PAS_chr2-1-0249 and PAS_chr1-1_0108 contain predicted PTS1 targeting signals [70], the latter having strong sequence homology to the peroxisome-targeted non-specific lipid transfer protein Pox18 present in Candida tropicalis and Candida maltosa [71,72]. Additionally, many genes connected to synthesis and degradation of triacylglycerol (TAG; metabolic pathway based on [73]) are regulated mainly in response to limiting glucose, which probably leads to the accumulation of free fatty acids which can then be degraded by beta-oxidation. Genes encoding fatty acid synthases (FAS1, FAS2) needed for de novo fatty acid
biosynthesis are only up-regulated in methanol-grown cells, while all sterol biosynthesis genes with the exception of ERG10, which encodes the first step of the pathway (acetyl-CoA C-acetyltransferase), are down-regulated in limiting glucose. Potential interaction partners which are also strongly induced in glucose-limited and methanol-grown cells could be the putative transcription factor SLT2 (PAS_chr1-4_0516) and MPPI, which was previously described to regulate peroxisomal matrix proteins and peroxins in Hansenula polymorpha [43].

| Short name | Description                                      | G-D logFC | G-D adjPV | M-D logFC | M-D adjPV | X-D logFC | X-D adjPV |
|------------|--------------------------------------------------|-----------|-----------|-----------|-----------|-----------|-----------|
| AOX1       | Alcohol oxidase (Pichia pastoris)                | 0.28      | 7.00      | ***       | 6.64      | ***       |           |
| AOX2       | Alcohol oxidase (Pichia pastoris)                | 0.39      | 7.44      | ***       | 7.48      | ***       |           |
| CTA1       | Catalase A, breaks down hydrogen peroxide in the peroxisomal matrix | 1.48  | *  | 5.45      | ***       | 6.11      | ***       |
| DAQ2       | Dihydroxyacetone kinase, required for detoxification of dihydroxyacetone (DHA) | –0.18 |   | 4.13      | ***       | 2.97      | ***       |
| DA51       | Dihydroxyacetone synthase variant 1             | 0.21      | 8.91      | ***       | 4.72      | ***       |           |
| DA52       | Dihydroxyacetone synthase variant 2             | 0.10      | 8.78      | ***       | 4.89      | ***       |           |
| FDX1       | NAD(+) dependent formate dehydrogenase, protect cells from formate | 0.44      | 8.74      | ***       | 8.75      | ***       |           |
| FGH1       | S-formylglutathione hydrolase; involved in the detoxification of formaldehyde | 0.65      | 5.25      | ***       | 4.86      | ***       |           |
| FLD        | glutathione-dependent formaldehyde dehydrogenase | 0.34      | 4.56      | ***       | 3.89      | ***       |           |
| PEX1       | AAA-peroxin                                      | 0.50      | *  | 2.56      | ***       | 2.75      | ***       |
| PEX10      | Peroxisomal membrane E3 ubiquitin ligase        | 0.33      | 3.64      | ***       | 4.19      | ***       |           |
| PEX11      | Peroxisomal membrane protein                     | 1.01      | ***       | 5.40      | ***       | 5.57      | ***       |
| PEX12      | C3HC4-type RING-finger peroxin and E3 ubiquitin ligase | 0.36  | **  | 2.50      | ***       | 3.75      | ***       |
| PEX13      | Integral peroxisomal membrane protein           | 0.55      | *  | 4.39      | ***       | 3.90      | ***       |
| PEX14      | Peroxisomal membrane peroxin                    | 0.23      | 3.14      | ***       | 3.90      | ***       |           |
| PEX17      | Peroxisomal membrane peroxin                    | –0.26     | 2.26      | ***       | 2.96      | ***       |           |
| PEX19      | Chaperone and import receptor for newly-synthesized class I PMPs | –0.07   | 0.75      | ***       | 2.10      | ***       |           |
| PEX2       | RING-finger peroxin and E3 ubiquitin ligase     | 0.75      | ***       | 3.48      | ***       | 3.63      | ***       |
| PEX20      | Peroxin 20                                       | 0.74      | ***       | 1.03      | ***       | 3.97      | ***       |
| PEX22      | Putative peroxisomal membrane protein           | 0.11      | 0.55      | *  | 0.85      | ***       |           |
| PEX25      | Peripheral peroxisomal membrane peroxin         | –0.19     | 1.09      | ***       | 3.29      | ***       |           |
| PEX28      | Peroxisomal integral membrane peroxin           | 0.04      | 0.23      | 1.55      | ***       |           |           |
| PEX29      | Peroxisomal integral membrane peroxin           | –0.24     | –0.16     |     | 0.48      | ***       |           |
| PEX3       | Peroxisomal membrane protein (PMP)              | 0.37      | **       | 2.27      | ***       | 1.30      | ***       |
| PEX30      | Peroxisomal integral membrane protein           | 0.10      | 0.09      | 0.47      | ***       |           |           |
| PEX31      | Peroxisomal integral membrane protein           | 0.36      | 0.93      | *  | 2.29      | ***       |           |
| PEX4       | Peroxisomal ubiquitin conjugating enzyme        | 0.76      | ***       | 2.03      | ***       | 4.45      | ***       |
| PEX5       | Peroxisomal membrane signal receptor            | 0.29      | 4.63      | ***       | 4.87      | ***       |           |
| PEX6       | AAA-peroxin                                      | 0.82      | ***       | 3.53      | ***       | 2.62      | ***       |
| PEX7       | Peroxisomal signal receptor                     | –0.22     | 0.30      | 1.98      | ***       |           |           |
| PEX8       | Intraperoxisomal organizer of the peroxisomal import machinery | 0.42  | **       | 2.93      | ***       | 3.52      | ***       |
| PEX26      | Peroxisomal membrane protein                    | 0.94      | ***       | 3.16      | ***       | 4.63      | ***       |
| PEX11C     | Ortholog of PEX11                               | 0.36      | 3.45      | ***       | 1.53      | ***       |           |

Log₂ fold changes and adjusted P-values (* adjPV < 0.1; ** adjPV < 0.05; *** adjPV < 0.01) are shown (see Additional file 1: Table S1 for detailed data). Up-regulated genes are in bold letters, down-regulated genes in bold and italics.

Conclusions

Our current knowledge of translational regulation comes from studies on S. cerevisiae cells [74-77], where stress conditions have been found to induce a global translational down-regulation that is mediated by translation initiation factors (eIFs). The specific regulation of defined mRNAs is dependent on regulatory UTR-binding protein complexes and miRNAs [78]. A significant finding emerging from this work is that the response of P. pastoris to different carbon sources (glycerol, glucose and
methanol) is regulated mainly at the transcriptional level. Furthermore, we found translational regulation to be global rather than transcript-specific in the analyzed conditions.

Strikingly, cells grown on excess glycerol or glucose have a very similar transcriptome in contrast to the situation in *S. cerevisiae*, which undergoes extensive changes when shifting between those two catabolites [5,6]. We have also identified genes that are subject to glucose repression in *P. pastoris*. Global gene regulation patterns in glucose-limited cells differ strongly from cells grown in excess glycerol, which is a de-repressing carbon source. While this may be partly associated with the reduced growth rate of glucose-limited cells, transcriptional de-repression of genes of the methanol utilization pathway, peroxisome biogenesis and fatty acid β-oxidation is specific to glucose-limited growth (apart from methanol induction). The transcription factor(s) responsible for this regulatory function remain(s) to be identified.

Finally, we have shown that translational regulation is global rather than transcript-specific for *P. pastoris* cells in different growth conditions. Cells growing on methanol exhibited the highest P:M ratio – which might also account for the superior protein production capacities observed in this condition. Despite the lower growth rate, transcription of genes encoding ribosomal constituents and parts of the translational machinery is not affected on methanol, indicating an increased global translation which is also reflected in the degree of polysome-associated mRNAs in the polysome profiles. The high abundance of methanol utilization enzymes [14] in combination with peroxisome proliferation [79] increases the burden on the translation machinery in methanol-grown cells. Indeed, *P. pastoris* has increased cellular protein content during methylotrophic growth (Buchetics, Russmayer et al. manuscript in preparation).

**Methods**

**Yeast strain and growth conditions**

*Pichia pastoris* wildtype (X-33, HIS4*, Mut*, Invitrogen) was used for this study. In liquid culture, cells were cultivated in shake flasks at 25°C on a rotary shaker at 180 rpm. YP media without carbon source (20 g L⁻¹ peptone and 10 g L⁻¹ yeast extract) and synthetic media (buffered M2 minimal media, pH set to 6.0, see Delic et al. [80]) with carbon source were used for pre- and main cultures, respectively. Four different cultivation strategies (Table 1) were applied for the analysis of distinct growth phases: carbon excess (starting with 2% glycerol or glucose), methanol induction (repeated batch) or glucose-limitation (12 mm glucose feed beads, Kuhner, CH).

Cultivations with excess glycerol and glucose were inoculated to an OD of 0.1 and started with 2% carbon source, while methanol fed and glucose-limited cultivations were started with an OD of 1.5 and 0.5% or 0.25% carbon source, respectively. For the cultivation on methanol, another pulse of 0.6% methanol was given after 16 hours, about 8 hours before harvesting the culture. Limiting glucose was applied by using glucose feed beads, which are polymer particles releasing glucose at a non-linear rate of 1.63 · t²/3 g per disc. In order to generate a growth rate of about 0.015 h⁻¹, 9 feed beads were added to 40 mL culture. The cells were harvested after 16 hours, at which time point the beads liberate 5.32 mg glucose per hour. Growth rate is calculated considering the average biomass concentration (3.3 g/L DCW), the average glucose feed rate (5.32 mg/h) and the low substrate yield coefficient Yₜₜ (0.37 g/l) at low growth rates (see [3]). Assuming that any of the three variables would deviate up to 35%, the growth rate would still be within the range of 0.010 – 0.022 h⁻¹. All cultivations were performed in triplicates and harvested at an OD of about 10 (Table 1).

**Polysome isolation and analysis**

The method for polysome isolation and analysis for *P. pastoris* was adapted from previously published methods [6,19]. RNA is prone to degradation, so working with pre-cooled and RNase-free materials is required. Polysomes were fixed by the addition of 0.1 mg cycloheximide (fresh solution of 10 mg/mL DEPC water) per mL main culture (at an OD₆₀₀ ~ 10, synthetic M2 media). The cultures were incubated for another 15 minutes on the shaker and then rapidly chilled by pouring into a 50 mL falcon tube containing 10 mL frozen DEPC-treated water and by using an ice water bath. Then the cells were recovered by 2 centrifugation steps (5300 × g, 4°C, 5 minutes) and a washing step with 10 mL cold lysis buffer (10 mM Tris–HCl pH 7.5, 0.1 M NaCl, 30 mM MgCl₂, 50 µg/mL cycloheximide, 200 µg/mL heparin, 1% DEPC) in between. Resuspended cells (500 µL cold lysis buffer, or more if too dense) were mixed with about 1 mL baked acid washed glass beads in ribolyzer/breaking tubes and applied in a Fast Prep (pre-cooled to ~80°C, Thermo Fisher Scientific, UK) for 3 minutes at 50 RPM. The lysate was transferred into fresh RNase-free tubes, cleared by centrifugation (13 K RPM, 4°C, 15 min) and analyzed using a Nanodrop spectrophotometer (Thermo Fisher Scientific, UK).

Sucrose gradients were prepared by stacking and freezing (~80°C) of each 2 mL 50%, 40%, 30%, 20% and 10% sucrose (in sucrose gradient buffer: 50 mM NH₄Cl, 50 mM Tris-OAc pH7, 12 mM MgCl₂) in ultracentrifuge tubes. Gradients (stored at ~80°C, thawed o/n at 4°C) were carefully loaded with polysome isolate corresponding to 150 µg RNA and centrifuged at 38 K RPM and 4°C for 2 hours in a SW40 Beckman rotor. The gradient station (Biocomp, CAN) was cleaned with ethanol (70%) and DEPC-treated water prior to gradient analysis, then blanked with water and used at a speed of 0.34 mm/s. The profile was
recorded and fractions were collected. ImageJ was used to calculate P:M ratios from the profiles, which is a measure of cellular translational activity.

**RNA isolation**

Monosome and polysome fractions (each about 5 mL) were separated according to the live polysome profile and collected in ice-cold tubes containing 15 mL 6 M guanidine hydrochloride (resulting in ~4 M final concentration), mixed with 2.5 volumes ice-cold 100% ethanol and precipitated o/n at ~20°C. Tubes were centrifuged at 3400 x g and 4°C for one hour, supernatant was removed entirely (apply short spin for residual liquid) and pellets were carefully air-dried for 5 minutes (this step can be repeated to pool material from 2 or more gradients). In order to isolate total RNA, polysome isolate corresponding to 150 µg RNA was directly mixed with guanidine hydrochloride and processed as described above. RNA was purified from the pellets using RNeasy mini kit (Qiagen, DE). Therefore, 100 µL DEPC-treated water was used for resuspension, mixed with 350 µL buffer RLT and further processed according to the manufacturer’s protocol. In the last step, 70 µL RNAse-free water was used to elute the RNA and the sample quality was checked by Nanodrop spectrophotometer and bioanalyzer analysis or gel electrophoresis.

**Microarray & data analysis**

In-house *P. pastoris* DNA microarrays (Agilent platform, AMAD-ID: 034821, design and general processing as described by [23]) were used. cRNA synthesis, hybridization and scanning were done according to the Agilent protocol for 2-color expression arrays. Each sample was hybridized against an RNA reference pool sample in dye swap. The microarray data were not background normalized. Within the arrays, loess-normalization was done for the color-effect. Quantile normalization was done between the arrays, the limma package (R-project) was used to calculate fold-changes, and p-value correction was done for multiple testing using the false discovery rate controlling method of [81]. Raw microarray data are provided in Additional file 5. Venn diagrams were created using the web-based tool Venny [82] and gene ontology (GO) term enrichment analysis was conducted with GO term finder and Saccharomyces Genome Database (SGD) annotations.

Principal component analysis was performed with the Excel plug-in XLSTAT.

Synonymous codon usage order (SCUO) analysis was performed online using the CondonO platform [35].

The statistical analysis was done in R using the standard functions fisher.test, chisq.test, and lm for the regression [83]. The implementation of the Fisher test obtains the p-values directly if a 2 by 2 table is present [84], otherwise a network implementation based on FEXACT was used [85]. For the group comparisons a test on normality was performed (Shapiro-Wilk-test) and Wilcoxon-Rank tests were performed since normality was not given.
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