Expression of phosphofructokinase in *Neisseria meningitidis*

Gino J. E. Baart,1,2† Marc Langenhof,1,2 Bas van de Waterbeemd,1 Hendrik-Jan Hamstra,1 Bert Zomer,1 Leo A. van der Pol,1 E. C. Beuvery,3 Johannes Tramper2 and Dirk E. Martens2

1Netherlands Vaccine Institute (NVI), Unit Research and Development, PO Box 457, 3720 AL Bilthoven, The Netherlands
2Wageningen University, Food and Bioprocess Engineering Group, PO Box 8129, 6700 EV Wageningen, The Netherlands
3PAT consultancy, Kerkstraat 66, 4132 BG Vianen, The Netherlands

Neisseria meningitidis serogroup B is a pathogen that can infect diverse sites within the human host. According to the *N. meningitidis* genomic information and experimental observations, glucose can be completely catabolized through the Entner–Doudoroff pathway and the pentose phosphate pathway. The Embden–Meyerhof–Parnas pathway is not functional, because the gene for phosphofructokinase (PFK) is not present. The phylogenetic distribution of PFK indicates that in most obligate aerobic organisms, PFK is lacking. We conclude that this is because of the limited contribution of PFK to the energy supply in aerobically grown organisms in comparison with the energy generated through oxidative phosphorylation. Under anaerobic or microaerobic conditions, the available energy is limiting, and PFK provides an advantage, which explains the presence of PFK in many (facultatively) anaerobic organisms. In accordance with this, *in silico* flux balance analysis predicted an increase of biomass yield as a result of PFK expression. However, analysis of a genetically engineered *N. meningitidis* strain that expressed a heterologous PFK showed that the yield of biomass on substrate decreased in comparison with a *pfkA*-deficient control strain, which was associated mainly with an increase in CO2 production, whereas production of by-products was similar in the two strains. This might explain why the *pfkA* gene has not been obtained by horizontal gene transfer, since it is initially unfavourable for biomass yield. No large effects related to heterologous expression of *pfkA* were observed in the transcriptome. Although our results suggest that introduction of PFK does not contribute to a more efficient strain in terms of biomass yield, achievement of a robust, optimal metabolic network that enables a higher growth rate or a higher biomass yield might be possible after adaptive evolution of the strain, which remains to be investigated.

Abbreviations: dcw, dry cell weight; ED, Entner–Doudoroff; EMP, Embden–Meyerhof–Parnas; F6P, fructose 6-phosphate; FBA, flux balance analysis; FR, fold ratio; HGT, horizontal gene transfer; LP, linear programming; MCS, Monte Carlo simulation; NVI, Netherlands Vaccine Institute; PEP, phosphoenolpyruvate; PP, pentose phosphate, PTS system, phosphotransferase system.

The microarray data discussed in this paper are available under GEO accession number GSE18951.

Three supplementary tables, showing the relationships in bacterial species between PFK (encoded by *pfkA*), the glucose-specific PTS transporter EII*B*Glc (encoded by *ptsG*) and glucokinase, GK (encoded by *glk*), are available with the online version of this paper. Additional data are also available with the online version of this paper in the form of the Excel file Additional data.xls, which includes six worksheets. The first worksheet, named ‘model’, contains the simplified metabolic model. The second worksheet, named ‘abbreviations’, contains a list of abbreviations of the metabolites. The third worksheet, named ‘flux distributions’, contains the calculated flux distributions of all experimental datasets. The fourth worksheet, named ‘phylogenetic profiling’, contains the phylogenetic profile of 373 species, and is available as a separate PDF. The fifth worksheet, named ‘phylo reactions’, contains the relevant reactions list and the amino acid sequences used for the phylogenetic profiling. The sixth worksheet, named ‘transcriptome’, contains the gene expression data.
INTRODUCTION

The species *Neisseria meningitidis* (the meningococcus) is only found in humans, colonizing mucosal surfaces of the nasopharynx as a harmless commensal organism, and as such is carried by at least 5–10% of the adult population (Rosenstein *et al.*, 2001; Stephens *et al.*, 1983). Some strains are able to cross the mucosa into the bloodstream, from where they can cause the diseases meningitis and septicemia. There are different pathogenic *N. meningitidis* isolates, of which five serogroups (A, B, C, Y and W135) are responsible for most disease. Conjugate polysaccharide vaccines that offer protection against infection with meningococcal serogroups A, C, Y and W-135 are effective and have been widely used (Girard *et al.*, 2006; Trotter & Ramsay, 2007). However, a broadly protective vaccine against infection by serogroup B *N. meningitidis* is not yet available, although this serogroup contributes significantly to the burden of meningococcal disease in many industrialized countries, where both epidemic and endemic serogroup B infections occur (Zimmer & Stephens, 2006).

At the Netherlands Vaccine Institute (NVI), a cross-protective multivalent vaccine against serogroup B organisms, including different subcapsular porin A (PorA) proteins contained in outer membrane vesicles (OMVs), is currently being developed. The vaccine development process focuses on cultivation of the organism, extraction of OMVs and subsequent purification of the PorA-containing OMVs (Baart *et al.*, 2007b). Knowledge of the primary metabolism is essential for the development of a consistent cultivation step in the vaccine production process. For this reason, an *N. meningitidis* serogroup B genome-scale metabolic model has been built and used to study primary metabolism (Baart *et al.*, 2007a, 2008). Earlier studies on glucose utilization in *N. meningitidis* confirm the presence of enzymes related to the glycolytic Embden–Meyerhof–Parnas (EMP) pathway, the Entner–Doudoroff (ED) pathway and the pentose phosphate (PP) pathway (Holten, 1974, 1975; Jyssum *et al.*, 1961; Jyssum, 1962a, b, c) (Fig. 1). However, it has been found that the EMP pathway does not contribute to pyruvate synthesis, indicating that the EMP pathway is not functional. These observations are in accordance with the *N. meningitidis* serogroup B genome (Tettelin *et al.*, 2000), in which a gene encoding phosphofructokinase (PFK) is not present. In order to improve the efficiency of the cultivation step in the vaccine production process we extended *N. meningitidis* metabolism with PFK. Furthermore, we used a comparative genomics approach to establish which factors determine the presence or absence of PFK in bacteria.

PFK (EC 2.7.1.11) is a key enzyme in glycolysis and is known to be regulated by allosteric effectors. The regulation of enzymes by allosteric effectors is a major mechanism of metabolic control, since it couples the activity of an enzyme to changes in the concentrations of metabolites produced by reactions that are distant in the metabolic network. PFK catalyses the irreversible phosphorylation of fructose 6-phosphate (F6P) to fructose 1,6-bisphosphate (FBP) using ATP, and is allosterically inhibited by phosphoenolpyruvate (PEP) and activated by ADP (or GDP) when the energy demand of the cell increases (Blangy *et al.*, 1968). The ATP-PFK described above (PFK-1) is broadly distributed in Eukaryota and Bacteria, as indicated by homologous protein sequences and biochemical evidence (Bapteste *et al.*, 2003), and is encoded by the *pfkA* gene. A second type of ATP-PFK enzyme (PFK-2), found strictly in *Escherichia coli* and encoded by the *pfkB* gene, has a minor activity (~10% of the total) (Fraenkel, 1996) and is inhibited by ATP at low concentrations of F6P (Guzik & Babul, 1985). The inorganic pyrophosphate (PPi)-dependent PFK (PPi-PFK) catalyses the same reaction using PPi, in a reversible manner, and can thus function in both glycolysis and gluconeogenesis. PPi-PFK (EC 2.7.1.90) has a more limited distribution, but is also found in eukaryotes and bacteria (Bapteste *et al.*, 2003). ATP-PFK and PPi-PFK share a common ancestry, but phylogenies show a very complex evolutionary pattern (Muller *et al.*, 2001; Siebers *et al.*, 1998). Some Archaea contain a unique ADP-dependent PFK (ADP-PFK) that appears to be phylogenetically distinct and instead belongs to the glucokinase family of kinases (Ito *et al.*, 2001; Verhees *et al.*, 2001). Bapteste *et al.* (2003) addressed the taxonomic distribution of ATP- and PPi-dependent PFks and concluded that numerous horizontal gene transfer (HGT) events and substitution of amino acids in the catalytic sites appear to occur at a high rate. None of the above-mentioned types of PFK are present in *N. meningitidis*. If it is assumed that *N. meningitidis* once had a functional PFK and that during the evolution of the species this activity was lost (Bapteste *et al.*, 2003), the question of why *N. meningitidis* lost PFK activity remains. In particular, since glucose is present within nasopharyngeal tissue (Exley *et al.*, 2005), the presence of PFK can yield one additional ATP per unit of substrate consumed, which is the case for all substrates that enter metabolism at a level above F6P.

It has been shown that *Neisseria* species preferentially take up DNA from their own genus (Davidsen & Tonjum, 2006; Elkins *et al.*, 1991), and this process of DNA exchange by HGT has been implicated in the evolution of *argF* (Zhou & Spratt, 1992), alteration of *penA* resulting in penicillin resistance (Spratt *et al.*, 1992), pilin antigenic variation (Seifert *et al.*, 1988) and other genetic alterations. Since none of the *Neisseria* species of which the complete genome sequences are publicly available contain PFK homologues (Bentley *et al.*, 2007; Parkhill *et al.*, 2000; Peng *et al.*, 2008; Tettelin *et al.*, 2000; http://www.genome.ou.edu/gono_new.html), HGT events within and between *Neisseria* species can never result in PFK functionality. However, DNA exchange between phylogenetically divergent species is possible (Omelchenko *et al.*, 2003), as shown by the acquisition of *sodC* in *N. meningitidis* from *Haemophilus influenzae* by HGT (Kroll *et al.*, 1998). Since *H. influenzae* contains a gene encoding *pfkA*, the possibility for *N. meningitidis* to obtain this gene exists. Hence, the question of why *N. meningitidis* did not acquire the *pfkA* gene from, for instance, *H. influenzae* in order to enhance growth as in *E. coli* remains unanswered. In order to establish what
determines the presence or absence of PFK in a species we used phylogenetic profiling and the literature as a starting point. In addition, we engineered a recombinant *N. meningitidis* strain to determine the influence of heterologous expression of PFK on biomass yield and composition, and analysed the strain at the metabolic and transcriptome level.

**METHODS**

**Bacterial strains and growth conditions.** Competent *E. coli* cells (strain TOP10, Invitrogen) were used for cloning purposes. Transformants of *E. coli* were grown on solid or liquid Luria–Bertani medium (BD Biosciences) containing either 100 μg kanamycin ml⁻¹ or 100 μg ampicillin ml⁻¹, and, when necessary, 1 mM IPTG and X-Gal. *N. meningitidis* strain HB-1, a non-encapsulated, non-piliated variant of the group B isolate H44/76 (Bos & Tommassen, 2005), was used for genetic modification. Untransformed *N. meningitidis* HB-1 cells were grown overnight on GC medium base plates (BD Biosciences) supplemented with IsoVitaleX (BD Biosciences) or in liquid meningococcal medium (van der Ley et al., 1993), while HB-1 derivatives were grown on GC medium base plates containing kanamycin (100 μg ml⁻¹). All incubations were done in a CO₂ incubator at 37 °C in a humid atmosphere containing 5% CO₂. The final *N. meningitidis* HB-1 mutants (HB-1 + kanR and HB-1 + kanR + pKΔA) were used to generate bacterial stock cultures using chemically defined medium.
Expression of PFK in Neisseria meningitidis

(Baart et al., 2007a) to ensure starting material of consistent quality for the chemostat cultures. After the addition of glycerol (20%, v/v, final concentration), the stock cultures were stored at -135 °C, and when required, a 500 ml shake flask containing 150 ml medium was inoculated with 10 ml stock culture that contained 0.5 g biomass L⁻¹. After approximately 8 h of incubation at 35 °C with shaking at 200 r.p.m. in an aerobic humid atmosphere, the culture was used to inoculate the bioreactor. Bacteria were grown on minimal chemically defined medium (Baart et al., 2007a) in 3 l autoclavable ADI bioreactors (Applikon), operated in chemostat mode, with a working volume of 1.4 l. Temperature, pH, dissolved oxygen (DO) concentration and stirrer speed were 37 °C, 7.0, 30% and 600 r.p.m., respectively. The total gas flow rate was kept constant at 1.0 l min⁻¹. The oxygen concentration was controlled by changing the oxygen fraction in the gas flow using headspace aeration only. The growth rate was 0.13 h⁻¹. After at least four residence times, physiological steady state was assumed based on online measurements (constant DO signal, O₂ and CO₂ concentration in the off-gas) and offline measurements (constant optical density, glucose concentration). In steady state, four small samples were taken using a rapid sampling procedure as described below, and a large 1.0 l sample was taken, divided into portions and processed as described previously (Baart et al., 2007a). All chemostats were run in duplicate.

Analytical procedures. Dry biomass concentration was determined by fourfold measurements for each steady-state sample by centrifugation (8000 g) of 50–200 ml culture broth in preweighed tubes followed by drying (24 h, 80 °C). Before weighing, the tubes were cooled in a desiccator for at least 1 h and dry cell weight (dcw) was corrected for salts present in the medium. The O₂ and CO₂ concentrations in the exhaust gas from the chemostat cultures were measured with a mass spectrometer (Prima White Box 600, Thermo Electron). The volumetric oxygen transfer coefficient, kₐ, in the bioreactors was determined accurately at 37 °C using a steady-state set-up similar to that used by Dorresteijn et al. (1994). Glucose, lactate and ammonium in the culture supernatant were determined as described previously (Baart et al., 2007a). Acetate, ethanol and other possible metabolites present in the culture supernatant were determined by ¹H-NMR as described previously (Baart et al., 2007a). The molecular and macromolecular composition of biomass was determined experimentally as described previously (Baart et al., 2007a, 2008), and was very important in mathematical modelling of cellular metabolism (Lange & Heijnen, 2001). PFK activity was measured enzymically in cell-free extracts using a coupling enzyme system. The primers were diluted to a concentration of 1 μmol ml⁻¹ and were added to puReTag ready-to-go PCR beads (Amersham Biosciences) prior to the addition of E. coli cells. Using six combinations of mixtures containing various amounts of primers and E. coli cells, PCR was carried out (GeneAmp PCR system 9600, Perkin Elmer) using the following cycling conditions. Initial denaturation of the template DNA at 94 °C for 5 min followed by 30 cycles of 94 °C for 1 min, 55 °C for 30 s and 72 °C for 1 min, and completed with 7 min at 72 °C and 4 °C until further processing. From each reaction mixture, 10 μl was added to 4 μl loading buffer (1 mM EDTA, 0.25% bromophenol blue, 40% sucrose) and analysed on a 1% agarose gel following cycling conditions. Initial denaturation of the template DNA at 94 °C for 5 min followed by 30 cycles of 94 °C for 15 min, 55 °C for 30 s and 72 °C for 1 min, and completed with 7 min at 72 °C and 4 °C until further processing. From each reaction mixture, 10 μl was added to 4 μl loading buffer (1 mM EDTA, 0.25% bromophenol blue, 40% sucrose) and analysed on a 1% agarose gel using 1× TBE buffer (Bio-Rad) at 100 V for 30 min, to check for sufficient amplification of the DNA fragment. It was found that 10 μl primers in combination with 1–2 μl E. coli cells gave the best results (results not shown). The generated 1096 bp fragment was cloned into the pCR.2.1-TOPO plasmid, using the TOPO TA cloning kit (Invitrogen), according to the manufacturer’s protocol. The plasmids were purified from the culture using a miniprep kit (Promega). After digestion of the plasmid DNA (37 °C, 2 h) using HindIII and EcoRV, samples were analysed for the presence of the pfkA gene by agarose gel electrophoresis, and fragments were purified from the gel using the Promega Gel and PCR Clean-Up system according to the manufacturer’s protocol. Fragments were checked by sequencing using the ABI Prism DNA sequencing kit (Applied Biosystems) according to the manufacturer’s protocol. In order to obtain reliable sequencing results, a primer annealing at position 359 of the DNA fragment was designed (5'-GGCGGTGACGGTTCCTACAT-3') and sequencing was performed on an ABI Prism 310 genetic analyser (Applied Biosystems). Sequencing of eight pCR-TOPO-pfkA constructs resulted in one usable construct (pTOPO-pfkA).

Constructs. A pCF-linker plasmid, containing the sites necessary for DNA uptake and homologous recombination was available from previous studies (van der Ley et al., 1995). Since a usable construct has to contain a kanamycin resistance gene (kanR) next to the pfkA gene and both genes have to be cloned in the BamHI site of the pCF-linker plasmid, a construct was prepared containing both the kanamycin resistance gene and pfkA. Briefly, the procedure was as follows. First, the pTOPO-pfkA plasmid and a pBluescript vector,
which contains an ampicillin resistance gene (Stratagene), were digested by HindIII and EcoRV. Digestion of the pTOPO-pfkA by these enzymes resulted in an 1173 bp fragment containing pfkA which was purified as described above. Second, the purified fragment was ligated into the pBluescript vector (pBlue-pfkA) and transformation was done in medium containing ampicillin. Third, a kanamycin resistance gene was obtained from an available pUC4K vector (van der Ley et al., 1995) by digestion with PstI and subsequent purification. The kanamycin resistance gene was ligated into the pBlue-pfkA vector (pBlue-PK) and transformant selection was done in medium containing kanamycin. After purification of the resulting plasmid DNA, the orientation of the kanamycin resistance gene was checked by Xhol digestion. Plasmids containing the correct orientation were selected, yielding the required pfkA-kanR fragment. After ligation of the pfkA-kanR fragment into the BamHI site of pCF-linker, transformation of competent E. coli cells was carried out and transformant selection was done in medium containing kanamycin, yielding the final pCF-linker-pfkA-kanR plasmid (pCF-PK). For correct expression by the rmpM promoter, the orientation of the fragment was checked by Ndel and XbaI digestion. Plasmids containing the fragment in the correct orientation were selected and used for transformation of N. meningitidis HB-1. After transformation, a PCR was done with the primers for pfkA, which confirmed the presence of the inserted DNA fragment. The control strain was created by cutting both pCF-linker and pUC4K with BamHI. After purification of the fragments, kanR was ligated into pCF-linker, yielding the pCF-K plasmid. The orientation of kanR in this construct was checked by digestion with XbaI and CldI. This construct was used for transformation of N. meningitidis HB-1.

Transformation. The plasmid-constructs (pCF-PK and pCF-K) were cut using KpnI and the linear DNA was used for transformation. Overnight-grown N. meningitidis strain HB-1 was scraped from the gonococcal (GC) agar plate using a swab and resuspended in 10 ml meningococcal medium containing 10 mM MgCl₂. The suspension was diluted fivefold in the same medium. A 2.5 μl volume of linear construct DNA (1 μg ml⁻¹) was added to 2.5 ml of the diluted suspension. After transformation (37 °C, 3 h), 250 μl suspension was plated on GC agar plates containing kanamycin. Plates were stored overnight in a CO₂ incubator. Colonies obtained this way were picked and plated again on kanamycin-containing GC plates to get rid of untransformed cells, which remain alive but cannot grow in the presence of kanamycin. This procedure was repeated once to obtain a total of three plating steps on kanamycin-containing medium. The final N. meningitidis HB-1 mutants (HB-1 + kanR and HB-1 + kanR + pfkA) were used to generate stock-cultures.

Rapid sampling. When the chemostat cultures were run, a full-genome N. meningitidis microarray was not yet available at the NVI. For this reason we decided to quench the culture broth using a rapid sampling procedure followed by low-temperature storage to freeze bacterial metabolism and preserve RNA. The practical rapid sampling setup of Lange et al. (2001) was adjusted by addition of air filters (0.2 μm pore-size) at the relevant points to construct a closed system that was suitable for working with pathogens. In steady state, four samples of 5 ml were taken in 35 ml medium containing 10 mM HEPES, pH 7.5 (Lange et al., 2001) and stored at −80 °C until further analysis. The quenched samples were stored for 2 years prior to RNA isolation.

Microarrays. Samples were analysed using a full-genome N. meningitidis microarray. Based on the genome sequence of N. meningitidis serogroup B strain MC38 (Tettelin et al., 2000), a set of 2078 70-mer oligonucleotides was developed at Operon (Germany), covering 93% of all predicted ORFs. In addition, a single 70-mer pfkA oligonucleotide sequence from E. coli K-12 was added to the set. Oligonucleotide pellets were dissolved in 50% DMSO (v/v, in water) to a concentration of 20 μM and spotted in triplicate on UltraGAPS II coated slides (Corning), using the Omniprint 100 microarray spotter (GeneMachines). During steady state, duplicate RNA samples were taken from one chemostat for each bacterial strain. The third RNA sample was taken from the duplicate chemostats, giving a total of four chemostat cultivations. For each microarray sample, the quenched bacterial culture samples were used directly for RNA isolation. The amount of quenched bacterial culture used for RNA isolation was equal to 1.2 ml per OD₅₉₀ unit to ensure that an equal amount of bacteria was used as starting material. The samples were concentrated by centrifugation (3000 g, 15 min) and treated with Tris-EDTA buffer containing 0.5 mg lysozyme ml⁻¹ (Sigma-Aldrich) for 3 min. Total RNA was extracted with the SV Total RNA isolation system according to the manufacturer's protocol (Promega Benelux). Nucleic acid concentration was adjusted by precipitation and spectral analysis was used to determine final nucleic acid concentration and purity. RNA integrity was confirmed with the Bioanalyzer RNA6000 Nano assay (Agilent Technologies), according to the manufacturer’s protocol. RNA integrity was predicted by calculation of the RNA integrity number (RIN) (Schroeder et al., 2006).

Total RNA from three experimental samples per strain was reverse-transcribed to cDNA and labelled with Cy3 dye using the Chipshot Indirect Labeling and Clean-Up kit (Promega Benelux) according to the manufacturer's protocol, with one deviation: 2 μl random primer and no oligo-dT primer was used per reaction to reverse transcribe the RNA. Common reference samples, containing equal amounts of total RNA from all three experimental samples, were labelled with Cy5. The labelled and purified cDNA samples were pooled in Cy3/ Cy5 pairs and volumes were adjusted to 25 μl. An equal volume of hybridization buffer was added, to a final concentration of 25 % formamide, 5 × SSC and 0.1 % SDS. Samples were applied to the microarray slides and placed in a hybridization chamber (GeneMachines) for 16–20 h at 42 °C in the dark. Differential gene expression levels were calculated through comparison with a common reference sample, containing equal amounts of RNA from all experimental samples. The microarrays were scanned with a ScanArray Express microarray scanner (Perkin Elmer) and median fluorescence intensities were quantified for each spot using ArrayVision software (Imaging Research). The expression data were natural-log-transformed and quantile-normalized, and values of replicate spots were averaged. These data processing steps were done with the free statistical software R (WU), using an in-house-developed script. P values were calculated using one-way ANOVA statistical analysis. Fold ratio (FR) values were expressed as the natural log of the normalized signal difference between the two groups. A P value below 0.05 was used to select differentially expressed genes. To further select for biologically relevant effects, only genes that combined both a low P value with and an FR above 2.00 were included in the final results (see Supplementary Material, transcriptome worksheet).

Phylogenetic profiling. To determine the presence or absence of the pfkA gene and other genes related to glucose metabolism and glucose transport, in different species, the STRING database (http://string.embl.de/; von Mering et al., 2005) was used as a starting point. The protein sequences of the relevant genes of E. coli were used as an input. In cases where no or very low homology was found in N. meningitidis, although functionality has been indicated elsewhere (http://www.biocyc.org/; http://www.genome.ad.jp/kegg/), the protein sequences of the relevant genes of N. meningitidis were also used as input. In order to interpret the results obtained from the alignments more easily, the STRING scores were converted to a colour scale based on four colours using an in-house computer program running in Visual Basic (Microsoft). In this program the homology scores (bitscores) of the relevant protein sequence of a
species serve as an input. In the cases where no sequence homology was found, the colour red was assigned. In the cases where some homology was found but the bitscore was below the general threshold value of 60 (http://string.embl.de/), the colour orange was assigned. The genes that were present are indicated by the colour green and high homology (high scores) are indicated by a slightly darker green. The reliability of the colour rescaling was checked by cross-referencing using BLASTP and two randomly chosen protein sequences, which confirmed the results obtained.

Metabolic modelling. Methods for solving a metabolic network or set of linear equations have been discussed extensively elsewhere (Edwards et al., 1999, 2001; Forster et al., 2002; Vallino & Stephanopoulos, 1990; van Gulik & Heijnen, 1995). Briefly, first the genome-scale metabolic model (Baart et al., 1990) was simplified as described previously (Baart et al., 2008). Next, an acetaldehyde exchange reaction was added based on the measurement of acetaldehyde in the culture supernatant. In addition, for the HB-1 + kanR + pfkA strain, the irreversible PFK reaction was added to the model (see Supplementary material, model worksheet). For both cases, the primary measurements were translated to measured conversion rates using mass balances, and all measurement errors were translated to errors (i.e. variances) in measured conversion rates using Monte Carlo simulation (MCS), as described previously (Baart et al., 2007a).

The redundancy matrix expressing the redundancy relations between the measured exchange rates was calculated for each individual dataset and contained two independent equations. Inspection of these equations indicated a carbon and a nitrogen balance. Both metabolic models (HB-1 + kanR and HB-1 + kanR + pfkA) are partly under-determined. Linear optimization (i.e. linear programming, LP) was combined with MCS and used to find a single, unique, optimal value for the objective function maximization of ATP hydrolysis as described previously (Baart et al., 2008). Since ATP is an intermediate metabolite that cannot accumulate, maximization of the hydrolysis reaction automatically means that in the rest of the network ATP production is maximized. It is noteworthy that this single solution is not necessarily unique and that there can be more than one flux distribution that reaches the optimal value of the objective function.

In silico flux balance analysis (FBA). Prior to balancing and LP (FBA), in silico FBA simulations were done in order to determine the possible increase in yield as a result of the introduction of pfkA. The protein, fatty acid and biomass biosynthesis reactions in the in silico FBA simulations were based on the measured average composition which had been determined previously (Baart et al., 2008). The following external metabolites were allowed to freely enter and leave the system: ammonia, water, phosphate, thiosulfate, sulfate, carbon dioxide, oxygen and protons. In addition, amino acids, acetate, hydrogen sulfide, ethanol and extracellular protein were only allowed to leave the system. Glucose was the sole carbon source that was allowed to enter the system. The objective function used in in silico FBA was the maximization of biomass. This objective function was found to result in biologically meaningful predictions in nutrient-limited continuous E. coli cultures (Schuetz et al., 2007). In silico FBA was combined with MCS using previously determined substrate (Y_{x/s}, m_s) and energy parameters (Y_{ATP}, m_{ATP}) and their corresponding variances (Baart et al., 2008) as input for optimization at different chosen growth rates. The complete procedure was as follows. First, a growth rate was chosen and at this growth rate the more important it is to increase the yield of After completion of the simulations at the chosen growth rate, the mean value and the corresponding variance of each flux and exchange rate were calculated. Then the next growth rate was selected and the above described procedure was done again. After completion of the in silico FBA calculations for strain HB-1 + kanR, the stoichiometric matrix corresponding to strain HB-1 + kanR + pfkA was selected and the complete procedure was repeated using the same growth rates as an input. The Monte Carlo approach was used in order to estimate the errors in the calculated in silico yield values (Y_{x/s}^{insilico}) at the different growth rates in the two strains. The stoichiometric matrices used in this study were constructed from the set of reactions using a custom made computer program running in Visual Basic (Microsoft). Both MCS and FBA were executed in custom-made computer programs running in Matlab (version 6.5 r13; Mathworks).

RESULTS AND DISCUSSION

In silico FBA

A functional glycolysis in N. meningitidis by inclusion of PFK theoretically leads to an increase in the biomass yield, as shown in Fig. 2. The possible increase in biomass yield is higher when the growth rate is lower, as expected. Since functional glycolysis only produces one additional ATP molecule to add to a total of about 20 ATPs in central metabolism, the effect of PFK when cells grow at a low growth rate becomes relatively more important, since the availability of ATP for biomass growth decreases with decreasing growth rates due to maintenance requirements. In other words, the more limiting energy becomes for growth the more important it is to increase the yield of

![Fig. 2](http://mic.sgmjournals.org)

**Fig. 2.** Percentage improvement of in silico biomass yield (Y_{x/s}^{insilico}) in the pfkA+ strain at various growth rates (μ) in comparison with the HB-1 + kanR strain. The grey lines represent the 95% confidence interval of the predictions (○). Since the substrate and energy parameters (see Methods) were estimated in the 0.04–0.16 range of growth rates, the mean line (black) is dashed outside this range.
ATP on a substrate and thus the more advantageous PFK becomes.

**Phylogenetic profiling**

Phylogenetic profiling (see Supplementary Material, phylogenetic profiling worksheet) reveals that many genomes of both γ- and δ-Proteobacteria, Acidobacteria, Firmicutes, Bacteroidetes, Actinobacteria, some Cyanobacteria and all Eukaryota contain a pfkA gene (or a pfkA homologue), while it is absent in most α-Proteobacteria, β-Proteobacteria and Chlamydiae genomes, and in Archaea (see Supplementary Material). Notably, all listed Firmicutes, δ-Proteobacteria and Acidobacteria contain a pfkA homologue. The presence of pfkA in all listed families, but not in Archaea, indicates that its origin is ancient. However, its phylogenetic distribution is patchy and complex. For instance, the genomes of two species within the listed β-Proteobacteria (i.e. *Rhodobacter ferrireducens* and *Chromobacterium violaceum*) contain a pfkA homologue, while all other species do not. Observations within all the described families of Bacteria show similar results. In other words, there always seem to be one or more species within a family that contains a pfkA gene, so phylogenetically based discrimination to reconstruct the evolutionary history of pfkA does not seem possible. However, the patchy phylogenetic distribution suggests, as indicated elsewhere (Bapteste et al., 2003), that pfkA can be lost during the evolution of a species. Phylogenetic profiling indicates that the genomes of most anaerobic organisms (e.g. *Clostridium* species) contain a pfkA gene. Since little energy is generated from substrate in anaerobically grown species, the relative gain in ATP yield increases significantly when the glycolytic pathway is present. This is in agreement with the obtained *in silico* FBA results, since at a low growth rate, an increased yield is predicted. The absence of a pfkA gene in anaerobic organisms goes together with the absence of genes related to the EMP pathway and/or the absence of glucose transport capabilities. In facultative anaerobic species, the presence or absence of a pfkA gene is more variable. For instance, the genomes of *Escherichia* species, *Shigella* species, *Salmonella* species, *Haemophilus* species, *Mammeinema succiniciproducens*, *Staphylococcus* species, *Corynebacterium* species, *Listeria* species, *Yersinia* species and *Vibrio* species contain a pfkA homologue, while the genomes of *Legionella* species, *Chlamydophila* species and *Chlamydia* species do not. In the genomes of the last three genera, additional genes related to the EMP pathway are also not present. Similar to facultative anaerobic species, the distribution of the pfkA gene in obligate aerobic species is variable; however, most obligate aerobic species seem to lack pfkA (e.g. *Burkholderia*, *Neisseria*, *Helicobacter* and *Bordetella* species). From an energy point of view, the lack of pfkA in an obligate aerobe seems obvious, since the contribution of PFK to the energy supply in an aerobically grown organism is limited in comparison with the energy generated through oxidative phosphorylation, which might explain the absence of pfkA in the *N. meningitidis* genome. In other words, under aerobic conditions, the substrate is more important for anabolism (i.e. supply of intermediates for growth), than for generating energy. Under anaerobic or microaerobic conditions the reverse is true; the substrate is more important for energy generation than for anabolism, and PFK provides an advantage. Therefore, in the case of a limiting energy supply, the objective of an organism seems to be to optimize the yield of energy from a substrate (maximize $Y_{ATP}$). In contrast, in the case of sufficient available energy, it seems that the strategy of organisms is to consume this energy as fast as possible (maximize $r_{ATP}$) in order to outcompete other micro-organisms living in the same habitat. However, there always seems to be a trade-off between high biomass yield and the high energy dissipation rates necessary for fast growth, as examined extensively by von Stockar and co-workers (Liu et al., 2007; von Stockar et al., 2006). Schuetz et al. (2007) found that under carbon (and energy) limitation in a chemostat, *E. coli* aims at maximization of biomass. In the case of an excess availability of carbon (and energy), the *E. coli* metabolism operates in such a way that a minimal number of biochemical reactions are used to generate the required ATP. This typical overflow metabolism (e.g. production of acetate) results in less efficient energy utilization of the carbon source. Nevertheless, the resulting suboptimal yields go hand in hand with dissipation of more energy and thereby enable higher catabolic rates (Schuetz et al., 2007), which might be a way to outcompete other organisms living in the same habitat. Furthermore, the excreted products such as acetate and ethanol may inhibit the growth of other organisms. This shows that *E. coli* has developed a mechanism that compromises between growth efficiency (i.e. biomass yield) and metabolic rate (i.e. growth rate), and the environmental conditions (i.e. habitat) determine which mechanism is preferred. There is more experimental evidence showing that at carbon substrate limitation, high-yield pathways are indeed adopted (Brauer et al., 2005; Postma et al., 1989; Vulic & Kolter, 2001).

In *N. meningitidis*, glucose uptake proceeds via glucokinase (NMB1390). In the genomes of *Staphylococcus* and *Streptococcus*, other known inhabitants of the nose and throat, pfkA is present while glucokinase is absent, and glucose uptake proceeds via the PEP: carbohydrate phosphotransferase system (PTS system). PTS-mediated substrate uptake involves the transfer of the phosphoryl group of the high-energy metabolite PEP to the carbohydrate through a cascade of phosphotransfer proteins (Deutscher et al., 2006; Postma et al., 1993). The significantly decreasing transcription level of pfkA in a PTS-deficient *E. coli* mutant (ΔptsHerr) in comparison with the wild-type (Flores et al., 2005) suggests that there is a correlation between PFK and PTS. Since a functional EMP pathway can generate twice as much PEP from glucose than the ED pathway, the presence of PFK and the PTS system indeed seems to go hand in hand. To illustrate the possible correlation between PFK and the PTS system, Fig. 3 shows Venn diagrams for pfkA, glk (glucokinase, GK) and ptsG (glucose-specific PTS transporter EIIBGlc) in γ- and β-
Proteobacteria, Firmicutes and Actinobacteria. No \textit{ptsG} homologs have been found in \textit{\alpha}- and \textit{\delta}-Proteobacteria, Acidobacteria, Chlamydiae, Bacteroidetes and Cyanobacteria species, whereas \textit{pfkA} homologues are present in some of these species (see Supplementary Tables S1–S3). From the Venn diagrams (Fig. 3) it can be calculated that in 157 of the 210 cases, PFK and EIIB\textsubscript{Glc} are correlated (75 %), meaning that either they are found together or neither is present, or stated differently, in the 156 cases where either PFK or EIIB\textsubscript{Glc} is present, they are present together in 103 cases, which indicates that they are correlated. PFK and GK are correlated in 22 % of the cases, and GK and EIIB\textsubscript{Glc} are correlated in 40 % of the cases. In 38 of the 210 cases shown in the Venn diagrams (18 %), PFK, EIIB\textsubscript{Glc} and GK are correlated. When the 312 bacterial species listed in the phylogenetic profile are taken into consideration, PFK and EIIB\textsubscript{Glc} are correlated in 69 % of the cases (see Supplementary Tables S1–S3). Notably, phylogenetic profiling indicates that the E1 PTS protein (encoded by \textit{ptsI}), is present in 92, 20 and 96 % of the Bacteria, Eukaryota and Archaea, respectively, which clearly indicates more (ancient) functional roles for the PTS system.

Since the PTS system is typical for anaerobic and facultative aerobic organisms (Postma \textit{et al.}, 1993), the lack of a PTS system in \textit{Neisseria meningitidis} seems obvious. Although, during infection, pathogenic \textit{N. meningitidis} encounters host anatomical sites with low oxygen levels, which indicates that anaerobic metabolism is required (Bartolini \textit{et al.}, 2006; Nassif \textit{et al.}, 1999), one can argue that this requirement is not a main determinant of \textit{N. meningitidis} evolution; in particular, natural evolution of \textit{N. meningitidis} takes place in the aerobic nasopharyngeal mucosa. The possibility cannot be ignored that other main nutrients for growth (for example lactate), which are not processed via F6P and are present the natural habitat of \textit{N. meningitidis}, might also be determinants of \textit{pfkA} deficiency in \textit{N. meningitidis}. In conclusion, for growth on hexose sugars, whether a micro-organism is strictly anaerobic or strictly aerobic seems to be the most important factor determining the presence or absence of PFK, and is correlated with the presence or absence of a PTS system, respectively, and thus can also explain the absence of PFK in \textit{N. meningitidis}, which is an aerobic bacterium. A second factor, which can also be important, is growth on substrates that enter metabolism below the PFK level, such as lactate.

**RNA integrity and comparison of expression profiles**

Gene expression profiles were analysed by taking three RNA samples for each strain (HB-1 + \textit{kanR} and HB-
1 + kanR + pfkA). Steady-state duplicate RNA samples were taken from one chemostat and one RNA sample from a second chemostat (chemostat duplicate), giving a total of six RNA samples for two strains. The RNA integrity number scores (Schroeder et al., 2006) of the RNA samples ranged from 9.4 to 9.8, indicating that the RNA was of good quality and suitable for use in DNA microarray analysis. This shows that the rapid sampling protocol used in this study followed by storage of the quenched biomass at −80 °C prevents RNA degradation for a long period of time (at least 2 years). Gene expression correlation within the chemostat duplicates was high and similar to the correlation between the chemostat duplicates (all R² values were between 0.940 and 0.972). Therefore, all RNA samples from each strain were used as independent replicates during statistical analysis. The averaged gene expression differences indicate that the heterologous pfkA gene is expressed in strain HB-1 + kanR + pfkA. Introduction of the pfkA gene in N. meningitidis induced a change in the gene expression level of the pfkA gene (FR = 2.6, P = 0.013) and high gene expression changes (FR > 3.7, P < 0.003) in the expression of only seven genes. These genes were upregulated in strain HB-1 + kanR + pfkA and include the pilE gene (NMB0018) and six pilS genes (NMB0020, NMB0021, NMB0022, NMB0023, NMB0025 and NMB0026). These virulence genes are phase-variable, i.e. are subject to reversible on/off switching of the expression state (Davidsen & Tonjum, 2006). Therefore, it is possible that the ‘on’ expression state of these genes was selected during strain construction and that the upregulation of these genes is not the result of heterologous pfkA expression. Hence, no large effects related to heterologous expression of pfkA were observed in the transcriptome.

Physiological and metabolic observations

Successful expression of PFK in N. meningitidis HB-1 + kanR + pfkA was indicated by measuring the PFK activity in cell-free extracts using a coupling enzyme assay. In the HB-1 + kanR + pfkA cell lysate, the measured PFK activity was 440 U l⁻¹. If it is assumed that all enzyme was liberated during the procedure to obtain cell-free extract, this translates into a specific PFK activity of 0.015 mol (g dw)⁻¹ h⁻¹ [0.353 μmol (mg protein)⁻¹ min⁻¹]. Notably, this observed PFK activity was significantly lower (133-fold) than the measured PFK activity level in E. coli (47 μmol (mg protein)⁻¹ min⁻¹] grown in batch culture on LB medium (Ogawa et al., 2007). This may be due to the strength of the rmpM promoter, the presence of allosteric inhibitors as explained in the Introduction, or to active fructose-1,6-bisphosphatase in the N. meningitidis cell extract, which might interfere with the activity measurement (Uyeda & Luby, 1974). Interference is unlikely in the E. coli cell extract, since fructose-1,6-bisphosphatase is not required for growth on hexoses and pentoses (Fraenkel & Horecker, 1965).

The macromolecular biomass composition was determined on the two duplicate aerobic glucose-limited chemostat cultures and is shown in Table 1. In strain HB-1 + kanR + pfkA, the RNA content was significantly higher and the LPS content lower, whereas no significant changes were observed for the other components. The cellular amino acid composition of the two strains did not differ significantly (Table 2). We observed small differences in fatty acid composition (Table 3). In particular, the unsaturated hexadecenoic acid (C16:1) content in strain HB-1 + kanR + pfkA was somewhat higher, while the dodecanoic acid (C12:0), hydroxydodecanoic acid (C12:0-3OH) and octadecanoic acid (C18:0) content was somewhat lower. Overall the changes in the biomass composition were minimal.

The redundancy matrix, expressing the redundancy relations between the measured exchange rates, was calculated for each individual dataset and contained a carbon and a nitrogen balance. The residuals obtained after multiplication of the redundancy matrix by the measured exchange rates could be explained on the basis of random measurement variances with test values that were lower than the 95 % chi-squared critical value of 5.992 (van der Heijden et al., 1994). In addition, the individual carbon and nitrogen balances could be closed for all datasets. Hence, the carbon- and nitrogen-containing exchange rates were balanced.

The measured (balanced) C-mol yield of biomass on substrate, Y_x/s in strain HB-1 + kanR + pfkA was found to be 0.45 ± 0.03 C-mol C-mol⁻¹, while Y_x/s in strain HB-1 + kanR was found to be 0.52 ± 0.03 C-mol C-mol⁻¹. Notably, predictions obtained from in silico FBA, at the same growth rate of 0.13 h⁻¹, indicated a small Y_x/s

## Table 1. Measured macromolecular composition (% w/w) of N. meningitidis strains HB-1 + kanR (pfkA⁻) and HB-1 + kanR + pfkA (pfkA⁺) (mean ± sd)

<table>
<thead>
<tr>
<th>Component</th>
<th>pfkA⁻</th>
<th>pfkA⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein</td>
<td>67.14 ± 7.85</td>
<td>69.41 ± 9.18</td>
</tr>
<tr>
<td>RNA</td>
<td>5.87 ± 1.47</td>
<td>9.97 ± 2.49</td>
</tr>
<tr>
<td>DNA</td>
<td>1.79 ± 0.27</td>
<td>1.78 ± 0.27</td>
</tr>
<tr>
<td>Phospholipids*</td>
<td>11.22 ± 0.16</td>
<td>10.69 ± 0.14</td>
</tr>
<tr>
<td>LPS†</td>
<td>5.83 ± 0.16</td>
<td>4.88 ± 0.12</td>
</tr>
<tr>
<td>Peptidoglycan‡</td>
<td>2.50 ± 0.25</td>
<td>2.50 ± 0.25</td>
</tr>
<tr>
<td>Total</td>
<td>94.3 ± 8.0</td>
<td>99.2 ± 9.5</td>
</tr>
</tbody>
</table>

*The overall phospholipid composition used in the present study was based on the values provided by Rahman et al. (2000), including 11 % phosphatidate (PA), 71 % phosphatidylethanolamine (PE) and 18 % phosphatidylglycerol (PG).
†LPS was quantified based on the measured amount of C12:0-3OH and possesses uncertainty (Baart et al., 2008).
‡Peptidoglycan in N. meningitidis was estimated based on E. coli (Neidhardt & Umbarger, 1996), and the composition was based on the average of the peptidoglycan structures present in N. meningitidis (Antignac et al., 2003). For modelling purposes a relative mean standard deviation of 10 % was assumed.
increase for strain HB-1 + kanR + pfkA. This might explain why the pfkA gene is not obtained by HGT, since it is initially unfavourable for biomass yield. The lower Yx/s in strain HB-1 + kanR + pfkA means that less carbon is assimilated in biomass, which was compensated mainly by an increase in CO₂ production (i.e. 0.52 ± 0.06 C-mol C-mol⁻¹ for strain HB-1 + kanR + pfkA and 0.44 ± 0.04 C-mol C-mol⁻¹ for strain HB-1 + kanR), while the production of by-products (i.e. ethanol, acetate, acetaldehyde, lactate, amino acids and proteins) was low and similar in the two strains (i.e. 0.03 ± 0.01 C-mol C-mol⁻¹ for strain HB-1 + kanR + pfkA and 0.05 ± 0.01 C-mol C-mol⁻¹ for strain HB-1 + kanR).

As explained in Methods, a single solution for the underdetermined parts (see Supplementary Material, model worksheet) of both metabolic networks was calculated using maximization of ATP yield (maxATP) as an objective function. Both the ED pathway and the EMP pathway are underdetermined, which means that the calculated fluxes for these parts are determined by the objective function. Due to the thiosulfate conversion reaction (reaction 51 in the model), the oxidative phosphorylation reaction (reaction 71 in the model) is also underdetermined. However, based on the thiosulfate concentration in the medium (0.38 mM), we calculate that the maximum oxygen consumption rate for this reaction is less than 1% of the measured oxygen consumption rate, which means that the measured oxygen consumption rate determines the rate of the oxidative phosphorylation reaction and therefore ATP production from this source. It was found that in the HB-1 + kanR + pfkA strain, 24% of the total ATP was formed in the EMP pathway, the citric acid cycle and the acetate production reaction, whereas in the HB-1 + kanR strain, 16% of the total ATP was formed in the ED pathway, the citric acid cycle and the acetate production reaction. In both strains, the remainder of the ATP was formed by oxidative phosphorylation (see Supplementary data, flux distributions worksheet). As expected, using this objective function, the HB-1 + kanR strain processes glucose 6-phosphate solely through the ED pathway, since the ED pathway can yield slightly more energy per molecule of glucose consumed than the PP pathway (Baart et al., 2008). It was found that the ATP hydrolysis flux (i.e. the flux that was maximized) was twice as high in the HB-1 + kanR + pfkA strain (see Supplementary data, flux distributions worksheet). Even if it is assumed that the EMP pathway is not active in the PFK-positive strain due to a PFK activity that is too low, and that glucose is metabolized through the ED pathway, the ATP hydrolysis flux is still 1.6 times higher in the PFK-positive strain. This means that in the PFK-positive strain more ATP is produced, which is directly related to the higher oxygen consumption measured for this strain. The

### Table 2. Measured cellular amino acid composition (% w/w) of N. meningitidis strains HB-1 + kanR (pfkA⁻) and HB-1 + kanR + pfkA (pfkA⁺) (mean ± sd)

<table>
<thead>
<tr>
<th>Component</th>
<th>pfkA⁻</th>
<th>pfkA⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspartate</td>
<td>5.68 ± 0.26</td>
<td>5.62 ± 0.27</td>
</tr>
<tr>
<td>Glutamate</td>
<td>7.78 ± 0.43</td>
<td>7.84 ± 0.45</td>
</tr>
<tr>
<td>Serine</td>
<td>3.79 ± 0.19</td>
<td>4.01 ± 0.21</td>
</tr>
<tr>
<td>Histidine</td>
<td>2.13 ± 0.14</td>
<td>2.05 ± 0.16</td>
</tr>
<tr>
<td>Glycine</td>
<td>5.45 ± 0.27</td>
<td>5.51 ± 0.28</td>
</tr>
<tr>
<td>Threonine</td>
<td>4.64 ± 0.21</td>
<td>4.46 ± 0.20</td>
</tr>
<tr>
<td>Alanine</td>
<td>8.75 ± 0.40</td>
<td>9.14 ± 0.42</td>
</tr>
<tr>
<td>Arginine</td>
<td>7.85 ± 0.33</td>
<td>8.05 ± 0.35</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>3.72 ± 0.14</td>
<td>3.91 ± 0.15</td>
</tr>
<tr>
<td>Valine</td>
<td>6.6 ± 0.32</td>
<td>6.59 ± 0.30</td>
</tr>
<tr>
<td>Methionine</td>
<td>2.67 ± 0.15</td>
<td>2.57 ± 0.21</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>4.54 ± 0.16</td>
<td>4.45 ± 0.17</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>5.11 ± 0.22</td>
<td>5.08 ± 0.22</td>
</tr>
<tr>
<td>Leucine</td>
<td>8.32 ± 0.32</td>
<td>8.16 ± 0.33</td>
</tr>
<tr>
<td>Lysine</td>
<td>6.96 ± 1.11</td>
<td>6.88 ± 1.16</td>
</tr>
<tr>
<td>Proline</td>
<td>3.72 ± 0.31</td>
<td>3.64 ± 0.30</td>
</tr>
<tr>
<td>Asparagine</td>
<td>4.35 ± 0.23</td>
<td>4.31 ± 0.24</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>1.98 ± 0.54</td>
<td>1.91 ± 0.54</td>
</tr>
<tr>
<td>Cysteine</td>
<td>0.96 ± 0.33</td>
<td>0.83 ± 0.32</td>
</tr>
<tr>
<td>Glutamine</td>
<td>4.97 ± 0.39</td>
<td>5.00 ± 0.41</td>
</tr>
</tbody>
</table>

### Table 3. Measured cellular fatty acid composition (% w/w) of N. meningitidis strains HB-1 + kanR (pfkA⁻) and HB-1 + kanR + pfkA (pfkA⁺) (mean ± sd)

<table>
<thead>
<tr>
<th>Component</th>
<th>pfkA⁻</th>
<th>pfkA⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dodecanoic acid (C12:0)</td>
<td>7.99 ± 0.33</td>
<td>6.93 ± 0.33</td>
</tr>
<tr>
<td>3-Hydroxydodecanoic acid (C12:0-3OH)</td>
<td>7.54 ± 0.26</td>
<td>6.83 ± 0.24</td>
</tr>
<tr>
<td>3-Hydroxytetradecanoic acid (C14:0-3OH)*</td>
<td>3.31 ± 0.27</td>
<td>3.50 ± 0.12</td>
</tr>
<tr>
<td>Tetradecanoic acid (C14:0)</td>
<td>8.53 ± 0.26</td>
<td>7.72 ± 0.25</td>
</tr>
<tr>
<td>Hexadecenoic acid (C16:1 trans-9)</td>
<td>20.92 ± 0.83</td>
<td>23.29 ± 0.75</td>
</tr>
<tr>
<td>Hexadecenoic acid (C16:1 cis)</td>
<td>0.40 ± 0.02</td>
<td>0.46 ± 0.02</td>
</tr>
<tr>
<td>Hexadecanoic acid (C16:0)</td>
<td>39.53 ± 1.4</td>
<td>38.98 ± 1.43</td>
</tr>
<tr>
<td>Octadecenoic acid (C18:1 cis)</td>
<td>10.52 ± 0.44</td>
<td>11.40 ± 0.72</td>
</tr>
<tr>
<td>Octadecanoic acid (C18:0)</td>
<td>1.26 ± 0.06</td>
<td>0.89 ± 0.06</td>
</tr>
</tbody>
</table>

*The C14:0-3OH quantity was calculated from the corresponding measured C12:0-3OH quantity, as explained previously (Baart et al., 2008).
expected effect on biomass yield due to the extra ATP produced is abolished by an ATP-consuming process, which can be stress, protein turnover or futile cycling in combination with fructose bisphosphatase, resulting in the loss of ATP. To compensate for this, more ATP must be generated from glucose, resulting in more O2 consumption and CO2 generation, and a decrease in biomass yield. The biomass composition of both strains is similar, which indicates that it is unlikely that more ATP is required for HB-1 + kanR + pfkA biomass formation. Notably, the measured PFK activity in the HB-1 + kanR + pfkA strain is sufficient to explain the additional ATP turnover and the calculated PFK flux. In the HB-1 + kanR + pfkA strain, the calculated PFK flux is 0.002 mol g⁻¹ h⁻¹ and the increase in the ATP hydrolysis reaction upon PFK introduction is 0.008 mol g⁻¹ h⁻¹ (see Supplementary data, flux distributions worksheet). If we assume that this additional ATP hydrolysis is caused by futile cycling through PFK and phosphatase, then the total PFK flux is 0.01 mol g⁻¹ h⁻¹, which is 32% lower than the measured specific PFK activity (0.015 mol g⁻¹ h⁻¹) in the HB-1 + kanR + pfkA strain and shows that the calculated flux is feasible. Hence, futile cycling upon introduction of pfkA in N. meningitidis seems the most plausible ATP-consuming process.

No significant differences between the maximum growth rates of the two strains were observed during the batch phase preceding the chemostat phase (results not shown). Theoretically, a higher yield would be expected, and due to the above-mentioned trade of between high biomass yield and high energy dissipation rates a lower maximum growth rate would be expected. As a result of futile cycling, ATP is lost (and energy is dissipated), more CO2 is produced and a lower yield of biomass on substrate is found. The extra dissipation of energy may explain why a lower growth rate was not found, which is thus in line with the theory. Adaptive evolution to achieve a robust, optimal metabolic network that enables a higher growth rate (Hua et al., 2007) or a higher yield might be possible.

Conclusions

The phylogenetic distribution of PFK indicates that it is lacking in most obligate aerobic organisms. In many (facultative) anaerobic organisms, PFK is present and goes hand in hand with the presence of a PTS system. A competitive advantage under anaerobic conditions seems to be the most important factor for the presence of PFK. In the case of limiting energy availability due to the absence of oxidative phosphorylation, the objective of an organism seems to be to optimize the yield of energy on substrate (maximize YATP/A), favouring PFK. In the case when sufficient energy can be liberated from the substrate by oxidative phosphorylation, the objective of an organism seems to be to optimize the rate of energy generation (maximize rATP) in order to outcompete other microorganisms living in the same habitat. Since the natural evolution of N. meningitidis takes place in the aerobic nasopharyngeal mucosa, the absence of pfkA in the N. meningitidis genome can be explained. In silico FBA predicts an increase in biomass yield upon inclusion of PFK. However, experimentally, a lower biomass yield is found in the presence of PFK than in the control strain without PFK. We hypothesize that this is caused by active phosphatases, creating futile cycles and the dissipation of energy. This makes it clear that no initial advantage is gained when PFK is expressed in N. meningitidis, which indicates that possible HGT events to obtain the pfkA gene are unlikely.

ACKNOWLEDGEMENTS

We thank Jeroen Pennings for his help with the microarray analysis and data processing, Mariene Willemsen, Elly Verhagen and Jan van den Ijssel for practical assistance, Alex de Haan for HPLC and GC measurements, Martin Hamzink for NMR measurements, Marcel Hoefnagel for sharing his knowledge of the determination of phosphofructokinase activity in cell-free extracts, Jo Maertens for his work on sequence analysis and Christoph Schoen for sharing his view on this subject. We also acknowledge the Gonococcal Genome Sequencing Project, and B. A. Roe, S. P. Lin, L. Song, X. Yuan, S. Clifton and D. W. Dyer, supported by USPHS/NIH grant #AI38399.

REFERENCES


Edited by: J. Moir