Ablation of Succinate Production from Glucose Metabolism in the Procyclic Trypanosomes Induces Metabolic Switches to the Glycerol 3-Phosphate/Dihydroxyacetone Phosphate Shuttle and to Proline Metabolism

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Trypanosoma brucei is a parasitic protist that undergoes a complex life cycle during transmission from its mammalian host (bloodstream forms) to the midgut of its insect vector (procyclic form). In both parasitic forms, most glycolytic steps take place within specialized peroxisomes, called glycosomes. Here, we studied metabolic adaptations in procyclic trypanosome mutants affected in their maintenance of the glycosomal redox balance. T. brucei can theoretically use three strategies to maintain the glycosomal NAD+/NADH balance as follows: (i) the glycosomal succinic fermentation branch; (ii) the glycerol 3-phosphate (Gly-3-P)/dihydroxyacetone phosphate (DHAP) shuttle that transfers reducing equivalents to the mitochondrion; and (iii) the glycosomal glycerol production pathway. We showed a hierarchy in the use of these glycosomal NADH-consuming pathways by determining metabolic perturbations and adaptations in single and double mutant cell lines using a combination of NMR, ion chromatography-MS/MS, and HPLC approaches. Although functional, the Gly-3-P/DHAP shuttle is primarily used when the preferred succinate fermentation pathway is abolished in the Δpepck knock-out mutant cell line. In the absence of these two pathways (ΔpepckiRNAi/FAD-GPDHi mutant), glycerol production is used but with a 16-fold reduced glycolytic flux. In addition, the Δpepck mutant cell line shows a 3.3-fold reduced glycolytic flux compensated by an increase of proline metabolism. The inability of the Δpepck mutant to maintain a high glycolytic flux demonstrates that the Gly-3-P/DHAP shuttle is not adapted to the procyclic trypanosome context. In contrast, this shuttle was shown earlier to be the only way used by the bloodstream forms of T. brucei to sustain their high glycolytic flux.

Trypanosomes of the Trypanosoma brucei group are the etiological agents of human African trypanosomiasis, a parasitic disease that affects over 36 countries in sub-Saharan Africa (1). T. brucei is a unicellular eukaryote, belonging to the protist order Kinetoplastida, that undergoes a complex life cycle during transmission from the bloodstream of a mammalian host (bloodstream stages of the parasite) to the alimentary tract (procyclic stage) and salivary glands (epimastigote and metacyclic stages) of a blood-feeding insect vector, the tsetse fly.

In the glucose-rich environment of the mammalian bloodstream, the parasite relies solely on glucose to produce energy (for review see Ref. 2). The procyclic insect stage of T. brucei, our experimental model in this analysis, develops a more elaborate energy metabolism based on different carbon sources, including glucose, proline, and threonine (3, 4). Although proline is the major component of the fly hemolymph (5), the parasite prefers glucose when this carbon source is available (6, 7).

The procyclic trypanosomes convert glucose by aerobic fermentation into partially oxidized end products, such as succinate and acetate (Fig. 1) (for reviews see Refs. 8, 9). Most glycolysis takes place in specialized peroxisomes, called glycosomes (steps 1–5 and 8) (10). In the course of glycolysis, phosphoenolpyruvate is produced in the cytosol (steps 9–11), where it is located at a branching point. It can be converted into pyruvate (steps 12 and 13), which enters the mitochondrion to produce acetate (steps 23–25) (11, 12). Phosphoenolpyruvate can also reenter the glycosomes to be converted to succinate within that compartment (steps 14–17) or else, after conversion to malate, transferred to the mitochondrion to make succinate in that compartment (steps 18 and 19) (13, 14). Within the glycosomes, consumption and production of NADH are tightly balanced; NADH resulting from the reaction catalyzed by glyceraldehyde-3-phosphate dehydrogenase (step 8, colored in blue, Fig. 1) needs to be re-oxidized inside the organelle. It has been proposed that the glycosomal succinic fermentation pathway (steps in red, Fig. 1), which contains two NADH-dependent oxidoreductases (steps 15 and 17), is involved in this process (13). The glycosomal redox balance can also be theoretically...
maintained by the Gly-3-P/DHAP shuttle (steps in green, Fig. 1) (15). This pathway involves the following: (i) a glycosomal NADH-dependent glycerol-3-phosphate dehydrogenase (NADH-GPDH, step 6), which produces Gly-3-P from DHAP; (ii) a putative glycosomal exchanger, which exchanges Gly-3-P for DHAP; and (iii) the mitochondrial FAD-dependent glycerol-3-phosphate dehydrogenase (FAD-GPDH, step 29), which regenerates DHAP from Gly-3-P. Electrons produced by FAD-GPDH are ultimately transferred to O$_2$ via the mitochondrial respiratory chain (steps 31–34).

Gly-3-P could also be converted into glycerol in the glycosomes (steps 7, purple in Fig. 1), with a net production of one molecule of NAD$^+$ per molecule of glycerol excreted.

In a glucose-depleted environment, the procyclic trypanosomes modify their metabolism by increasing the rate of proline consumption compared with glucose-rich conditions. Succinate produced in the mitochondrion from proline metabolism is further converted into alanine (Fig. 2, A and B) (7, 16–18). The metabolic adaptation resulting from a repressive effect of glucose metabolism also affects the mode of ATP production, i.e. oxidative phosphorylation (step 35) versus substrate level phosphorylation (steps, 9, 12, and 26). In glucose-rich conditions, ATP is primarily produced by substrate level phosphorylation (4, 7, 19), whereas oxidative phosphorylation becomes essential in the absence of glucose (7, 20), although some differences may exist between strains (20).

Here, we further analyze the metabolic flexibility of the procyclic trypanosomes by abolishing a main branch of glucose metabolism. Abolition of the succinate production pathway in the phosphoenolpyruvate carboxykinase (PEPCK) gene knock-out mutant (Δpepck) induced successful metabolic adaptations that allowed the parasite to maintain its growth rate, although its rate of glucose consumption is 3.3-fold reduced. To understand these adaptations, we have generated a number of RNAi mutant cell lines in the PEPCK null background, which were analyzed by global metabolomics approaches.

**EXPERIMENTAL PROCEDURES**

**Growth and Maintenance of Trypanosomes**—The procyclic form of *T. brucei* EATRO1125.T7T was cultured at 27 °C in SDM79 medium containing 10% (v/v) heat-inactivated fetal calf serum (FCS) and 2.5 mg/ml hemin (21) or in a glucose-depleted medium derived from SDM79, called SDM80 (6). The SDM80 medium is supplemented with 9% (v/v) heat-inactivated fetal calf serum dialyzed by ultrafiltration against 0.15 M NaCl (molecular mass cutoff: 10,000 Da; Sigma F0392; glucose concentration, 1 mM) and 1% (v/v) heat-inactivated FCS (glucose concentration, 5 mM). The glucose concentration in the glucose-depleted medium (SDM80) is 0.15 mM as compared with 6 mM in the glucose-rich media. We recently used a new FCS batch, which caused an increase of the procyclic cell doubling time grown in glucose-rich conditions (10.5 versus 13.5 h).

3 The abbreviations used are: Gly-3-P, glycerol-3-phosphate; DHAP, dihydroxyacetone phosphate; FAD-GPDH, FAD-dependent glycerol-3-phosphate dehydrogenase; NADH-GPDH, NADH-dependent glycerol-3-phosphate dehydrogenase; PDH, pyruvate dehydrogenase; PEPCK, phosphoenolpyruvate carboxykinase; SDH, succinate dehydrogenase; SHAM, salicylhydroxamic acid.

For consistency, we always compared doubling times determined with the same FCS, such as described in Table 3. The FCS issue does not concern data obtained in glucose-depleted conditions, because no significant differences were observed with the different FCS batches.

**Gene Knock-out of PEPCK**—Replacement of the PEPCK gene (Tb927.2.4210) by the blasticidin and puromycin resistance markers via homologous recombination was performed with DNA fragments containing a resistance marker gene flanked by the PEPCK UTR sequences. Briefly, the pGEMt plasmid was used to clone an HpaI DNA fragment containing the blasticidin and puromycin resistance marker genes preceded by the PEPCK 5′-UTR fragment (514 bp) and followed by the PEPCK 3′-UTR fragment (597 bp). Primers used to produce PCR fragments are described in supplemental Fig. S1. The PEPCK knock-out was generated in the EATRO1125.T7T parental cell line, which constitutively expresses the T7 RNA polymerase gene and the tetracycline repressor under the control of a T7 RNA polymerase promoter for tetracycline inducible expression (22). Transfection and selection of drug-resistant clones were performed as reported previously (23). The first and second *PEPCK* alleles were replaced by blasticidin- and puromycin-resistant genes, respectively. Transfected cells were selected in glucose-rich SDM79 medium containing hygromycin B (25 μg/ml), neomycin (10 μg/ml), blasticidin (10 μg/ml), and puromycin (1 μg/ml).

**Inhibition of Gene Expression by RNAi**—Accession numbers of genes targeted by RNAi are as follows: F1β subunit of the F$_0$/F$_1$ ATP synthase (ATP$_e$-F1β, Tb927.3.1380), FAD-GPDH (Tb11.02.5280), E2 subunit of the pyruvate dehydrogenase complex (PDH-E2, Tb927.10.7570), and succinate dehydrogenase (SDH, Tb927.8.6580). The inhibition by RNAi of gene expression in the procyclic form (24) was performed by expression of stem-loop “sense/antisense” RNA molecules of the targeted sequences (22) introduced in the pLew100 or pLew79 expression vectors (kindly provided by E. Wirtz and G. Cross) (25) or the pT7F1-177 vector (kindly provided by B. Wickstead and K. Gull) (26), as described previously. These three vectors contain the plhemoycin resistance gene. Construction of the pLew-SDH-SAS, pLew-PDH-E2-SAS, and pLew-ATP$_e$-F1β used to produce the double mutants Δpepck/ΔRNAiSDH, Δpepck/ΔRNAiPDH-E2, and Δpepck/ΔRNAiATP$_e$-F1β, respectively, have been described before (7). The RNAiSDH-A1, RNAiPDH-E2, and RNAiATP$_e$-F1β cell lines have been generated before (7). The pLew-FAD-GPDH-SAS construct targets a fragment (from position 367 to 903 bp) of the FAD-GPDH gene. Briefly, a PCR-amplified 584-bp fragment, containing the antisense FAD-GPDH sequence with restriction sites added to the primers (supplemental Table S1), was inserted into the HindIII and BamHI restriction sites of the pLew100 plasmid. The separate PCR-amplified fragments containing the sense FAD-GPDH sequence were then inserted upstream of the antisense sequence, using HindIII and XhoI restriction sites (XhoI was introduced at the 3′-extremity of the antisense PCR fragment). The resulting plasmid (pLew-FAD-GPDH-SAS) contains a sense and antisense version of the targeted gene fragment, separated by a 50-bp fragment, under the control of the procyclic...
FIGURE 1. Schematic representation of glucose metabolism in the procyclic form of T. brucei. This figure describes the glycosomal NADH producing and consuming pathways, highlighted by a dashed circle and colored pathways. The glycosomal NADH-producing step is shown in blue; the glycosomal succinic fermentation pathway is shown in red; the Gly-3-P/DHAP shuttle and the associated complexes of the respiratory chain are shown in green; and the glycerol-producing step is shown in purple. Excreted end products from glucose metabolism are shown in black, red, green, or purple characters on a gray rectangle as background. ATP molecules produced by substrate level phosphorylation and oxidative phosphorylation are boxed and circled, respectively. Enzymatic steps targeted by RNAi are circled, and the PECK step, in which the gene has been deleted, is boxed; the name of the genetically manipulated enzymes is also indicated. Abbreviations used are as follows: 1,3BPAGA, 1,3-bisphosphoglycerate; c, cytochrome c; e, electrons; FBP, fructose 1,6-bisphosphate; FUM, fumarate; G3P, glyceraldehyde-3-phosphate; Gly3P, glycerol 3-phosphate; MAL, malate; OAA, oxaloacetate; PEP, phosphoenolpyruvate; PPR, pyruvate; ScCoA, succinyl-CoA; SUC, succinate. Enzymes used are as follows: steps 1, hexokinase; 2, glucose-6-phosphate isomerase; 3, phosphofructokinase; 4, aldolase; 5, triose-phosphate isomerase; 6, NADH-dependent glycerol-3-phosphate dehydrogenase; 7, glycerol kinase; 8, glyceraldehyde-3-phosphate dehydrogenase; 9, phosphoglycerate kinase; 10, phosphoglycerate mutase; 11, enolase; 12, pyruvate kinase; 13, pyruvate phosphate dikinase; 14, phosphoenolpyruvate carboxykinase (PECK, Tb927.2.4210); 15, malate dehydrogenase; 16, cytosolic fumarase; 17, glycosomal NADH-dependent fumarate reductase; 18, mitochondrial fumarase; 19, mitochondrial NADH-dependent fumarate reductase; 20, cytosolic malic enzyme; 21, mitochondrial malic enzyme; 22, unknown enzyme; 23, pyruvate dehydrogenase complex (PDH-including the E2 subunit, PDH-E2, Tb927.10.270); 24, unknown enzyme; 25, acetate-succinyl-CoA transferase; 26, succinyl-CoA synthetase; 27, rotenone-sensitive NADH dehydrogenase (complex I of the respiratory chain); 28, rotenone-insensitive NADH dehydrogenase; 29, FAD-dependent glycerol-3-phosphate dehydrogenase (FAD-GPDH, Tb11.02.5280); 30, succinate dehydrogenase (SDH, complex II of the respiratory chain, Tb927.8.6380); 31, ubiquinone; 32, SHAM-sensitive alternative oxidase; 33, complex III of the respiratory chain; 34, complex IV of the respiratory chain; 35, F$_{1}$F$_{0}$-ATP synthase (ATP$_{1}$- including the F$_{1}$ subunit, ATP$_{1}$-F$_{1}$, Tb927.3.1380).

Metabolic Flexibility in Procyclic Trypanosomes

The $\Delta$peck null mutant and the EATRO1125.T7T parental cell line have been transformed with expression plasmids described above. The RNAi-harboring single mutant cell lines were selected in glucose-rich SDM79 medium containing hygromycin B (25 $\mu$g/ml), neomycin (10 $\mu$g/ml), and phleomycin (5 $\mu$g/ml). For transfection of the $\Delta$peck cell line, blasticidin (10 $\mu$g/ml) and puromycin (1 $\mu$g/ml) were also included in the medium. Aliquots were frozen in liquid nitrogen to provide stocks of each line that had not been cultivated long term in medium. The selected cell lines were then adapted to the glucose-rich SDM80glu medium and the glucose-depleted SDM80 medium containing the same concentration of the five antibiotics.

Enzyme Assays—Sonicated (5 s at 4°C) crude extracts of trypanosomes resuspended in cold hypotonic buffer (10 mM potassium phosphate, pH 7.8) were tested for mitochondrial FAD-GPDH (EC 1.1.99.5) (15) and malate dehydrogenase (MDH, EC 1.1.1.40) (27).

Immune Sera Production and Western Blot Analyses—For the production of PDH-E2 antibodies, a recombinant fragment corresponding to the full-length PDH-E2 gene preceded by an N-terminal histidine tag (10 histidine codons) was expressed in the Escherichia coli BL21, using the pET16b expression vector (Novagen). Cells were harvested by centrifugation, and recombinant proteins were purified by nickel chelation chromatography (Novagen) according to the manufacturer’s instructions. Antisera were raised in rabbits or mice by three injections at 15-day intervals of 200 $\mu$g of PDH-E2–His recombinant nickel-purified proteins, electroeluted after separation on SDS-PAGE, and emulsified with complete (first injection) or incomplete Freund’s adjuvant.

Total protein extracts of procyclic form T. brucei (5 x 10$^{6}$ cells) were separated by SDs-PAGE (10%) and immunoblotted on Immobilon-P filters (Millipore) (28). Immunodetection was performed as described previously (28, 29) using as primary antibodies the rat antiserum against PECK diluted 1:1000 (gift from T. Seebeck, Bern, Switzerland), the rabbit antiserum against F1 moiety of the mitochondrial F$_{1}$/F$_{0}$-ATP synthase isolated from Crithidia fasciculata diluted 1:1000 (30) (gift from D. Speijer, Amsterdam, Netherlands), the heat shock protein 60 (hsp60) diluted 1:10,000, (31), or the E2 subunit of PDH diluted 1:500. Goat anti-rat, anti-rabbit, or anti-mouse IgG-per-
Metabolic Flexibility in Procyclic Trypanosomes

Determination of Glucose and Proline Consumption—To determine the rate of glucose and proline consumption, cells (incubated at 1–1.5 × 10⁷ cells/ml) were grown in 10 ml of SDM79, SDM80glu (6 mm glucose), or SDM80 (0.15 mm glucose) medium. Aliquots of each growth medium (500 µl) were collected 0, 1, 6, 9, 10, 23, and 24 h after incubation at 27 °C. The quantity of glucose present in the medium was determined using the “Glucose GOD-PAP” kit (Biolabo SA). Proline concentration was determined with a colorimetric assay as described previously (32) after deproteinization of the samples by perchloric acid treatment.

Analysis of Excreted End Products from Glucose and Proline Metabolism—5–8 × 10⁶ T. brucei procyclic cells were collected by centrifugation at 1400 × g for 10 min, washed once with phosphate-buffered saline (PBS), and incubated in 30 ml of PBS. For the analysis of glucose metabolism, the cells were maintained for 6 h at 27 °C in incubation buffer containing 110 µmol of d-[1-¹³C]glucose and 2 g/liter NaHCO₃, pH 7.4. For the analysis of L-proline metabolism, the cells were maintained in PBS, pH 7.4 (without NaHCO₃), containing 20 µmol of L-[4-¹³C]proline in the presence of 100 µmol of unenriched d-glucose. For NMR experiments, the supernatant was lyophilized, and ¹³C NMR spectra were collected at 125.77 MHz with a Bruker DPX500 spectrometer, as described before (7).

The supernatants analyzed by NMR were then lyophilized and dissolved in 1 ml of Milli-Q H₂O for further analyses by high performance liquid chromatography (HPLC). The amount of glycerol in samples was determined by using an HPLC system (HP 1100 Series, Agilent, Santa Clara, CA) coupled to a Shodex RI-101 refractive index detector. The analytical column (Aminex® HPX-87H, 300 × 7.8 mm, 9 µm) was maintained at 48 °C. The binary pump was operated isocratically with 5 mM H₂SO₄ at a flow rate of 0.5 ml/min for 40 min. The injection volume was 50 µl.

Analysis of Intracellular Metabolites—T. brucei procyclic cells grown in glucose-rich conditions were sampled by fast filtration (33). Briefly, 2 × 10⁶ cells were taken from the culture media, directly filtered on a vacuum (Whatman glass microfiber filters), and washed with 500 µl of culture media diluted 1:10 in PBS. The filters on which the cells were deposited were directly wrapped in aluminum paper and quenched in liquid nitrogen. Total sampling time was below 8 s. The filters were then transferred into 5 ml of boiling water. After 30 s of incubation, 200 µl of a uniformly ¹³C-labeled extract of E. coli was added as a quantification standard, and the solution was vortexed for 2 s. The extracts were immediately filtered (0.2 µm), chilled with liquid nitrogen, lyophilized, and dissolved in 200 µl of Milli-Q water prior to analysis. Three replicates were taken from each culture media, sampled, and analyzed separately.

The ion chromatography system (ICS 2000, Dionex, Sunnyvale, CA) was equipped with an EG50 potassium hydroxide eluent generator and a 2-mm ASRS-ULTRA II suppressor. Intracellular metabolites were separated at 29 °C on an IonPac-pack...
AS11 analytical column (250 × 2 mm, Dionex) as described previously (34).

Ion chromatography was coupled to a 4000 QTRAP triple quadrupole linear ion trap mass spectrometer (Applied Biosystems, Foster City, CA) operating in the negative ion mode. Multiple reaction monitoring was used to detect selective fragments of the precursor ions with a dwell time of 25 ms. The acquisition parameters are described in more detail in supplemental Tables S2 and S3. The method allowed the potential detection of 43 intracellular metabolites, including most intermediates from glycolysis and pentose-phosphate pathway, organic acids, and nucleotides (supplemental Table S2). In the trypanosome extracts, 31 metabolites showed signal-to-noise ratios consistent with accurate quantification of the metabolite pools (supplemental Table S3).

Cell Permeabilization and Measurement of Oxygen Consumption—A total of 3 × 10⁸ cells were collected from exponential phase growth, centrifuged at 10,000 rpm, and washed twice in buffer, pH 7.2, containing 240 mM mannitol, 100 mM KCl, 1 mM EGTA, 20 mM MgCl₂, and 10 mM KH₂PO₄. Oxidation rates were determined polarographically with a Clark-type oxygen electrode (Rank Brothers) at 25 °C in a glass vessel (final volume 2 ml). The Clark-type oxygen electrode response was calibrated in air-equilibrated buffer in the absence (100% O₂) and presence (0% O₂) of sodium dithionite. 100% O₂ concentration was taken as 240 μM (35). Permeabilization of whole cells was performed by the addition of 160 μg of digitonin to dilute cellular metabolites and consequently arrest the initial oxygen consumption, leaving the integrity of glycosomal and mitochondrial membranes intact (36). Respiratory chain response to substrate was tested by the consecutive addition of 12.5 mM Gly-3-P and 12.5 mM succinate, followed by KCN (6.25 mM) and SHAM (1.56 mM) to completely inhibit mitochondrial respiration.

RESULTS

Deletion of the PEPCK Gene Stimulates Growth of the Parasite—We previously observed that knockdown of the PEPCK gene (Tb927.2.4210) by RNAi only slightly affected procyclic trypanosomes (7), although PEPCK activity has been considered crucial to maintain both the glycosomal ATP/ADP and redox balances. To study metabolic adaptations caused by the absence of PEPCK activity, we have generated three independent PEPCK knock-out mutant cell lines (Δpepck::BLA/Δpepck::PURO, named Δpepck) by replacing both PEPCK alleles by selectable markers in the procyclic EATRO1125.T7T cell line, which express the tetracycline repressor and T7 RNA polymerase. Deletion of both alleles was confirmed by PCR and by Western blot with an anti-PEPCK immune serum (Fig. 3 for the Δpepck-cl1 cell line). The three freshly selected Δpepck mutant cell lines grew faster compared with the EATRO1125.T7T wild type cell line (11% decrease of the doubling time). However, after a few weeks of growth in standard SDM79 medium, the doubling time of the Δpepck-cl1 cell line returned back to wild type levels (Fig. 3B and Table 1) (not done for the two other Δpepck clones). The same observation was made when the Δpepck-cl1 cell line was grown in glucose-depleted conditions (SDM80). Because the three analyzed clones showed the same genotype and initial growth phenotype, only the Δpepck-cl1 cell line will be further described here.

The next two sections describe the metabolic adaptations occurring in the Δpepck-cl1 cell line to compensate for the loss of the succinate fermentation pathway. The metabolic fate of glucose and proline in the Δpepck-cl1 cell line was investigated,

FIGURE 3. Analysis of the Δpepck mutant cell line. A shows a PCR analysis of genomic DNA isolated from the parental EATRO1125.T7T (WT) and Δpepck (Δ) cell lines. Amplifications were performed with primers based on sequences that flank the 5′ UTR and 3′ UTR fragments used to target the PEPCK gene depletion (black boxes) and internal sequences from the PEPCK gene (PCR products 1 and 2), the blasticidin resistance gene (BLASTR, PCR products 3 and 4), or the puromycin resistance gene (PUROR, PCR products 5 and 6). As expected, PCR amplification using primers derived from the PEPCK gene and drug-resistant genes were only observed for the parental EATRO1125.T7T (WT) and Δpepck (Δ) cell lines, respectively (DNA bands labeled with a star). B shows the growth curve of the EATRO1125.T7T and Δpepck cell lines incubated in SDM79 and the SDM80 medium either containing (SDM80-glu) or lacking (SDM80) glucose. Cells were maintained in the exponential growth phase (between 10⁶ and 10⁷ cells/ml), and cumulative cell numbers reflect normalization for dilution during cultivation. The inset shows a Western blot analysis of the parental (WT) and Δpepck (Δ) cell lines with the anti-PEPCK and anti-hsp60 immune sera.
**TABLE 1**

| Step(s) | Cell linea | Vector | ni/i| Activityc | Proteind | Doubling timee |
|---------|-------------|--------|-----|-----------|----------|---------------|
|         |             |        |     |           |          | Glu-rich | Glu-depleted |
| 14      | Δpepck (cl1) | plew100 | i   | 80         | ND       | ND          | 10.5 (13.5)  | 13.9         |
| 30      | Δpepck (cl1) | plew100 | i   | ND         | ND       | ND          | 13.4         | 30.1         |
| 14, 30  | Δpepck (cl1) | plew100 | i   | ND         | ND       | ND          | 13.4         | 30.1         |
| 35      | Δpepck (cl1) | plew100 | i   | ND         | ND       | ND          | 13.4         | 30.1         |
| 14, 35  | Δpepck (cl1) | plew100 | i   | ND         | ND       | ND          | 13.4         | 30.1         |
| 29      | FAD-GPDH    | plew79  | i   | ND         | ND       | ND          | 13.4         | 30.1         |
| 14, 29  | Δpepck (cl1) | plew79  | i   | ND         | ND       | ND          | 13.4         | 30.1         |
| 23      | FAD-GPDH    | plew79  | i   | ND         | ND       | ND          | 13.4         | 30.1         |

a The clone number is indicated in parentheses.

b ni means noninduced; i means tetracycline-induced.

c Relative enzymatic activities of the targeted protein(s) in the RNAi-harboring cell lines are expressed as a percentage of the activity detected in the parental EATRO1125.T7T cell line.

d Relative amount of the targeted protein(s) in the RNAi-harboring cell lines are expressed as a relative amount of the protein(s) detected by Western blotting in the parental EATRO1125.T7T cell line.

e WT means that the doubling time of the mutant cell line is similar to that of the parental cell line (EATRO1125.T7T).

f ND means not determined.

g ND means not detectable.

h Data are from Ref. 7.

i nd means not determined.

j † means not viable.

**TABLE 2**

| Glucose and proline consumption in procyclic T. brucei cell lines | Rate of glucose consumption | Rate of proline consumption |
|---------------------------------------------------------------|-----------------------------|----------------------------|
| Cell linea | mmol/h/mg protein | mmol/h/mg protein |
| EATRO1125.T7T  | 1.59 ± 0.20 | 0.21 ± 0.05 |
| EATRO1125.T7T (− Glu) | 0.48 ± 0.23 | 0.40 ± 0.12 |
| Δpepck        | NA* | 0.70 ± 0.20 |
| Δpepck (− Glu) | NA* | 0.67 ± 0.15 |
| Δpepck/FAD-GPDH.ni | 0.86 ± 0.05 | 0.27 ± 0.09 |
| Δpepck/FAD-GPDH.i | 0.79 ± 0.15 | 0.19 ± 0.08 |
| Δpepck/FAD-GPDH.ni | 0.29 ± 0.01 | 0.34 ± 0.00 |
| Δpepck/FAD-GPDH.i | 0.10 ± 0.01 | 0.53 ± 0.07 |
| Δpepck/PDH-E2.ni | 1.50 ± 0.20 | 0.18 ± 0.05 |
| Δpepck/PDH-E2.i | 1.60 ± 0.10 | 0.20 ± 0.04 |
| Δpepck/PDH-E2.ni | 0.68 ± 0.19 | 0.26 ± 0.11 |
| Δpepck/PDH-E2.i | 0.03 ± 0.03 | 0.50 ± 0.14 |

a i means tetracycline-induced; ni means noninduced.

b NA means not applicable.

c Cell line grown in SDM80glu.

determination of the nature of the metabolic end products, as well as the total amount of 13C label recovered in these compounds (7).

The PBS medium collected after 6 h of incubation with D-[1-13C]glucose was analyzed by 13C NMR spectroscopy to quantify the excretion of 13C-labeled products. In such conditions, around 743 nmol of 13C label was recovered in the supernatant of wild type cells (Table 3), with most of the label being detected in succinate (63.3%), acetate (25.0%), and lactate (7.4%). In comparison, much less label was recovered in the medium of the Δpepck mutant cells (208 nmol of 13C-enriched excreted molecules/h/mg of protein, see Table 3). This 3-fold reduction in 13C label recovery is consistent with the 3.3-fold reduction in glucose utilization in the Δpepck cell line. The spectrum of 13C-labeled products is also modified in the mutant cell line. In the parental EATRO1125.T7T cell, the major end product of glucose metabolism is succinate, which was barely detectable in the Δpepck mutant. This was also true for the downstream metabolites of succinate fermentation, namely malate and fumarate (Fig. 4, see Table 3). These data confirm that conversion of glucose into succinate proceeds uniquely via the PECK pathway in procyclic T. brucei. In addition, the NMR analyses showed increased signals of glycerol/Gly-3-P (the two compounds overlap in 13C NMR spectra) and β-hydroxybutyrate in the Δpepck cell line. This suggests a redistribution of flux toward the Gly-3-P/DHAP shuttle and acetate metabolism, respectively, when no glycosomal succinic fermentation is occurring (see below).

The metabolism of proline in the wild type EATRO1125.T7T procyclic cells is known to depend highly on the presence of glucose (7). Accordingly, in the absence of glucose, the parasite
TABLE 3

Excreted end products of \( ^{13}C \)-glucose metabolism by procyclic \( T. brucei \) cell lines

| Cell line \( a \) | \( ^{13}C \)-enriched excreted molecules \( ^{c} \) | Succinate | Acetate | Lactate | Malate | Fumarate | Alanine | \( \beta \)-Hyd | Pyruvate | Glycerol \( d \) | TOTAL |
|-----------------|-----------------------------------|---------|--------|--------|-------|---------|--------|------|--------|---------|---------|
| n \( b \) | mmol / h / mg of protein | | | | | | | | | | |
| EATRO1125.T77 \( e \) | | 5 | 470 ± 88 | 186 ± 21 | 55 ± 15 | 20 ± 1.7 | 4.0 ± 1.1 | 8 ± 2.1 | ND \( f \) | ND | ND | 743 ± 124 |
| \( \Delta pepck = r e v / F A D - G P D H \) c1l | | 4 | 1.2 ± 1.4 | 154 ± 54 | 18 ± 3.7 | ND | ND | 14 ± 8.7 | 8.5 ± 4.4 | ND | 6.3 ± 4.6 | 208 ± 58 |
| \( r e v / F A D - G P D H \) \( i \) | | 2 | 367 | 205 | 26 | 21 | i | ND | 3 | ND | 5 | 627 |
| \( \Delta pepck = r e v / F A D - G P D H \) D4.ni | | 3 | 1.6 ± 1.6 | 258 ± 5.4 | 36 ± 0.2 | ND | ND | 23 ± 9.8 | 14 ± 2.0 | ND | 23 ± 3.1 | 355 ± 13 |
| \( \Delta pepck = r e v / F A D - G P D H \) D4.i | | 3 | ND | 113 ± 13 | 5.3 ± 1.8 | ND | ND | 23 ± 2.8 | 1.0 ± 1.4 | ND | 70 ± 11 | 212 ± 19 |
| \( \Delta pepck = r e v / P D H - E 2 \) E11.i | | 3 | 0.6 ± 0.5 | 168 ± 88 | 33 ± 19 | ND | ND | 17 ± 7.9 | 7.4 ± 5.6 | 0.4 ± 0.5 | 14 ± 7.8 | 241 ± 129 |
| \( \Delta pepck = r e v / P D H - E 2 \) E11.ii | | 3 | 0.6 ± 0.8 | 27 ± 24 | 2.7 ± 1.9 | ND | ND | 22 ± 6.3 | 1.4 ± 2.0 | 10 ± 7.8 | 17 ± 9.3 | 81 ± 46 |

a \( i \) means RNAi cell lines were tetracycline-induced during 6–10 days depending on the cell line and the experiments; ni means non-induced RNAi cell lines.

b \( n \) means number of experiments. When only two experiments have been analyzed per cell lines, the deviation of the mean (difference between the values divided by 2) is lower than 20% of the mean.

c The result of a single experiment or the means of 2–5 experiments are presented; the results are shown ± S.D. when more than two experiments have been performed per cell line.

d Glycerol and Gly-3-P peaks cannot be distinguished by this NMR analysis.

Glycerol and Gly-3-P peaks cannot be distinguished by this NMR analysis.

e Data are from Ref. 7.

f ND means not detectable.

excretes mainly alanine (64.2% of label recovery from \( L-[^{1-13}C] \)proline) and glutamate (28.2%), with only small amounts of acetate (3.2%), \( \beta \)-hydroxybutyrate (3.1%), and succinate (0.5%) (Table 4 and Fig. 2 A). In the presence of glucose, proline utilization was significantly reduced, and its metabolic fate was also modified, \( i.e. \) succinate became a major end product, and alanine production was 30-fold reduced (Table 4 and Fig. 2 B). In the same glucose-rich conditions, the \( \Delta pepck \) cell line produced similar amounts of alanine and succinate (Table 4 and Fig. 2 C).

The alanine/succinate ratio observed in these conditions was 0.62. This was 5-fold higher than observed with wild type cells in the absence of glucose (0.11) but 7-fold lower than observed for the wild type cells in glucose-rich conditions (106). In conclusion, the \( \Delta pepck \) cell line shows, in the presence of glucose, an intermediary state between glucose-depleted and glucose-rich conditions in the wild type cells.

The intracellular metabolite content was analyzed by ion chromatography-MS/MS. In glucose-rich conditions, the levels of fumarate, malate, and succinate were significantly lower in the \( \Delta pepck \) mutant, compared with the wild type cells (supplemental Fig. S1). But the fact that significant extents of these metabolites could be detected in the \( \Delta pepck \) mutant, where the conversion of glucose into succinate no longer occurs, indicates that the utilization of proline could replenish the pool of dicarboxylic acids (fumarate, malate, and succinate) (Fig. 2 C).

To further ascertain the activation of proline metabolism when PEPCK is lacking, RNAi knockdowns of each of succinate dehydrogenase (SDH, \( step 30 \)) and the \( F_{0}/F_{1} \)-ATP synthase (\( step 35 \)) were performed in the \( \Delta pepck \) cell line. These two enzymatic steps are dispensable in glucose-rich conditions. However, in glucose-depleted conditions, SDH is a key step in the conversion of proline into alanine (see Fig. 2 B), and the \( F_{1} \) (ATP-eF\(_{1}\)) is essential for ATP production by oxidative phosphorylation (7). Considering that the increase in proline metabolism plays a critical role in the \( \Delta pepck \) cell line, the down-regulation of these two genes in the PEPCK null background was expected to strongly affect the cellular viability (see Fig. 2 C). Indeed, both double mutants showed a strong growth phenotype in glucose-rich conditions. After tetracycline induction, the \( \Delta pepck / \text{SDH} \) cell line (\( \Delta pepck/\text{SDH}.i \)) died within 8 days, whereas the uninduced mutant (\( \Delta pepck/\text{SDH}.ni \)) showed only a moderate growth phenotype (Fig. 5 A and Table 1). Three different \( \Delta pepck/\text{ATP}\_e F_{1}.i \) cell lines were analyzed, all of which showed a cessation of growth in glucose-rich medium, before a reversion to uninduced doubling time levels (Fig. 5 B). This reversion was concomitant with the re-emergence of the ATP-eF\(_{1}\) protein in Western blots of cell extracts (Fig. 5 B, inset). Altogether, these data confirmed that the \( \Delta pepck \) cell line switched to proline metabolism even in the presence of glucose.

Redox Balance within the Glycosomes—Current schemes of redox potential within the glycosomes show a tightly regulated balance of NADH production and usage, where the NADH produced by the glyceraldehyde-3-phosphate dehydrogenase is re-oxidized within the succinate fermentation pathway (2). In the \( \Delta pepck \) null mutant, where the latter pathway is no longer active, alternative process(es) must operate to maintain the glycosomal redox balance. Two processes could potentially play this role, namely the Gly-3-P/DHAP shuttle and glycerol fer-
mentation. To address the role of the Gly-3-P/DHAP shuttle, we compared knockdown effects of the FAD-GPDH gene in the wild type and Δpepck null backgrounds. Both the tetracycline-induced RNAi FAD-GPDH.i and Δpepck RNAi FAD-GPDH.i mutant cell lines showed a stable loss of the FAD-GPDH enzyme activity up to 7 days, even after 24 h of induction with tetracycline (Fig. 5, C and D, inset).

Although the Δpepck RNAi FAD-GPDH.i double mutant showed a moderate growth phenotype upon tetracycline induction (Fig. 5D), the loss of FAD-GPDH in the Δpepck null background caused a number of metabolic alterations. First, glucose consumption was drastically reduced in the double mutant (16- and 5-fold, as compared with wild type and Δpepck cell lines, respectively). This was associated with an increase in proline consumption (Table 2). Second, the NMR analysis of incubation supernatants showed a reduced conversion of D-[1-13C]glucose into β-hydroxybutyrate, most likely as a result of the overall reduction in glycolytic flux (Fig. 4 and Table 3). Third, HPLC measurements of incubation supernatants showed that glycerol production was increased 14.4-fold in the Δpepck RNAi FAD-GPDH.i mutant upon tetracycline induction (Fig. 6). Glycerol was not detected in the wild type, Δpepck, RNAi FAD-GPDH.ni, and RNAi FAD-GPDH.i cell lines (data not shown). Finally, the ion chromatography-MS/MS analysis of intracellular metabolites showed that the intracellular amount of Gly-3-P was 2.5-fold increased in the Δpepck RNAi FAD-GPDH.i mutant compared with the uninduced cells (supplemental Fig. S1). These data indicate the concomitant accumulation of intracellular Gly-3-P and excretion of glycerol, which are, respectively, the substrate and product of glycerol kinase (step 7), in the Δpepck RNAi FAD-GPDH.i mutant. Altogether, this shows that glycerol fermentation was increased in the PEPC null background only when the FAD-GPDH was no longer active. This means that in the Δpepck mutant, reoxidation of glycosomal NADH mainly proceeds via the Gly-3-P/DHAP shuttle. These data also highlight the production of NAD⁺ by the glycosomal NADH-GPDH (step 6) in the absence of the glycosomal succinate fermentation branch.

FIGURE 4. Carbon-13 NMR spectra of metabolic end products excreted by procyclic cell lines incubated with D-[1-13C]glucose. For these NMR analyses, the parental EATRO1125.T7T, the Δpepck, and tetracycline-induced Δpepck RNAi FAD-GPDH.i cell lines were incubated with 4 mM D-[1-13C]glucose in PBS/NaHCO₃ buffer. The NMR spectra were obtained after addition of 15 μL of dioxane. Each spectrum corresponds to one representative experiment from a set of at least three. The boxed metabolites (succinate, malate, and glycero) are excreted metabolites produced in the glycosomes. The resonances were assigned as follows: Ala, alanine; Ace, acetate; β-Hyd, β-hydroxybutyrate; D, dioxane; Gly, glycerol; Lac, lactate; Mal, malate; Suc, succinate.

TABLE 4 Excreted end products of [4-13C]proline metabolism by procyclic T. brucei cell lines

| Cell line | G | n | Succinate | Alanine | Glutamate | Acetate | β-Hyd | Malate | Fumarate | Aspartate | Lactate | TOTAL |
|-----------|---|---|-----------|---------|-----------|---------|-------|--------|----------|-----------|---------|--------|
| EATRO1125.T7T | 3 | 0.5 | 64.2 | 28.2 | 3.2 | 3.1 | 0.3 | ND | 0.5 | ND | 24.5 | 227.4 |
| EATRO1125.T7T | 3 | 39.2 | 4.4 | 50.2 | 3.8 | 1.3 | 0.3 | ND | 0.1 | ND | 17.6 | 126.4 |
| Δpepck | 3 | 4.3 | 64.5 | 20.2 | 1.6 | 2.0 | 1.8 | ND | 5.6 | ND | 4.7 | 47.4 |
| Δpepck RNAi FAD-GPDH.E2 | 3 | 28.5 | 17.8 | 32.3 | 7.1 | ND | 6.8 | 0.7 | 4.9 | 1.9 | 23.6 | 260.1 |
| Δpepck RNAi FAD-GPDH.E2 | 3 | 32.2 | 13.0 | 27.3 | 10.3 | 3.3 | 6.2 | 1.1 | 3.2 | 3.5 | 22.9 | 259.4 |
| Δpepck RNAi FAD-GPDH.E2 | 3 | 14.3 | 35.3 | 17.0 | 8.2 | ND | 11.9 | 2.3 | ND | 10.9 | 32.5 | 362.1 |

| Cell line | G | n | % of 13C-enriched excreted molecules |
|-----------|---|---|----------------------------------|
| EATRO1125.T7T | 3 | 0.5 | 64.2 | 28.2 | 3.2 | 3.1 | 0.3 | ND | 0.5 | ND | 24.5 | 227.4 |
| EATRO1125.T7T | 3 | 39.2 | 4.4 | 50.2 | 3.8 | 1.3 | 0.3 | ND | 0.1 | ND | 17.6 | 126.4 |
| Δpepck | 3 | 4.3 | 64.5 | 20.2 | 1.6 | 2.0 | 1.8 | ND | 5.6 | ND | 4.7 | 47.4 |
| Δpepck RNAi FAD-GPDH.E2 | 3 | 28.5 | 17.8 | 32.3 | 7.1 | ND | 6.8 | 0.7 | 4.9 | 1.9 | 23.6 | 260.1 |
| Δpepck RNAi FAD-GPDH.E2 | 3 | 32.2 | 13.0 | 27.3 | 10.3 | 3.3 | 6.2 | 1.1 | 3.2 | 3.5 | 22.9 | 259.4 |
| Δpepck RNAi FAD-GPDH.E2 | 3 | 14.3 | 35.3 | 17.0 | 8.2 | ND | 11.9 | 2.3 | ND | 10.9 | 32.5 | 362.1 |

a i means tetracycline-induced; ni means noninduced.
b Cells were grown in SDM79 and incubated in PBS/[4-13C]proline containing (G) or not ( ) 3.3 mM D-glucose.
c Number of experiments for each cell line are shown.
d The results are shown ± S.D.
e Data from Ref. 7.
f ND means not detectable.
Metabolic Flexibility in Procyclic Trypanosomes

A. Log of cumulative cell number * ml\(^{-1}\)

- EATRO1125.T7T
- Δpepck\(^{RNA}\) SDH-H3.ni
- Δpepck\(^{RNA}\) SDH-H3.i

B. Log of cumulative cell number * ml\(^{-1}\)

- EATRO1125.T7T
- Δpepck\(^{RNA}\) ATP\(_{e}\) F1β-D4.ni
- Δpepck\(^{RNA}\) ATP\(_{e}\) F1β-D4.i

C. Activity

- WT
- Δ

D. Activity

- WT
- Δ

E. PDH-E2

- WT
- Δ

Days post induction

Days

Activity

Log of cumulative cell number * ml\(^{-1}\)

Days

Log of cumulative cell number * ml\(^{-1}\)
In contrast, none of the metabolic perturbations found in the ΔpepckRNAi/FAD-GPDH.i double mutant were observed in the ΔpepckRNAi/FAD-GPDH.i single mutant cell line, indicating that the Gly-3-P/DHAP shuttle is not critical for glycolysis when the PEPC enzyme is active (Tables 2 and 3, Fig. 5D, and supplemental Fig. S1). In conclusion, these data demonstrate that the Gly-3-P/DHAP shuttle is primarily used in the absence of glycosomal succinic fermentation to maintain the glycosomal redox balance.

**Functional Analysis of FAD-GPDH**—FAD-GPDH is the first step of the respiratory chain involved in electron transfer from Gly-3-P to molecular oxygen (see Fig. 1). To further confirm the involvement of FAD-GPDH in the Δpepck null background, it was important to demonstrate that knockdown of FAD-GPDH by RNAi significantly affected oxygen consumption from Gly-3-P. Thus, we compared oxygen consumption of wild type and mutant cell lines in the presence of Gly-3-P. intact trypanosome cells are unable to use Gly-3-P as a respiratory substrate, because this compound cannot cross the plasma membrane. Therefore, we selectively permeabilized the plasma membrane by a mild digitonin treatment, without affecting mitochondrial integrity, as described previously (36). The wild type cells incubated in the absence of respiratory substrates showed a high rate of endogenous respiration, which dramatically decreased in the presence of digitonin (Fig. 7A). This suggests that the consumption of an unknown endogenous carbon source was abolished when the subcellular compartments were disconnected. In these conditions, the plasma membrane has been successfully permeabilized, because addition of 12.5 mM Gly-3-P restored oxygen consumption. A subsequent addition of
Metabolic Flexibility in Procyclic Trypanosomes

12.5 mM succinate did not further stimulate oxygen consumption. Succinate is one of the preferred respiratory substrates of procyclic trypanosomes (37), which is oxidized by complex II of the respiratory chain (SDH, step 30), indicating that Gly-3-P is also a good respiratory substrate in these experimental conditions. As expected, addition of both 6.25 mM KCN (inhibitor of the terminal oxidase, step 34) and 1.56 mM SHAM (inhibitor of the alternative oxidase, step 32) completely inhibited oxygen consumption (Fig. 7A). The Δpepck and uninduced Δpepck/RNAi pepck cell lines showed the same behavior, although the rate of oxygen consumption was ~2-fold lower compared with the parental cell line (Fig. 7A). In contrast, the tetracycline-induced Δpepck/RNAi FAD-GPDH/H9004 cell lines showed only a 4-fold reduction of the rate of oxygen consumption in the presence of Gly-3-P, compared with uninduced cells. The addition of succinate increased the respiration rate to levels comparable with that of the Δpepck and uninduced Δpepck/RNAi FAD-GPDH/H9004 cell lines. This result shows that in the induced double mutant, the reduced oxygen consumption observed in the presence of Gly-3-P was not due to a reduced capacity of oxygen consumption but the direct result of the loss of FAD-GPDH. In conclusion, these experiments demonstrate that, as expected, the FAD-GPDH function was considerably reduced upon tetracycline induction of the Δpepck/RNAi FAD-GPDH cell line.

Inhibition of Both the Succinate and Acetate Branches Is Detrimental for Glucose Metabolism—Glucose metabolism in procyclic trypanosomes mainly results in succinate and acetate production. To further alter glucose metabolism, the production of both end products was abolished by generating a RNAi mutant targeting the pyruvate dehydrogenase (PDH, step 23) in the PEPCK null background. In glucose-rich conditions, the RNAiPDH-E2 single mutant showed no growth phenotype upon tetracycline induction (7). The Δpepck/RNAi PDH-E2 i cell line survived tetracycline induction, with only a moderate increase (1.5-fold) of its doubling time (Table 1 and Fig. 5E). However, the Δpepck/RNAi PDH-E2.i double mutant showed important metabolic differences compared with wild type cells and the Δpepck mutant. First, glucose was no longer the main carbon source used in the Δpepck/RNAi PDH-E2.i cell line, because only a residual consumption was detected in glucose-rich conditions (2 and 6% of the wild type and Δpepck rates of consumption, respectively) (Table 2). Second, the incapability of the Δpepck/RNAi PDH-E2.i cell line to metabolize glucose was confirmed by NMR. The conversion of d-[1-13C]glucose into 13C-labeled end products was decreased 3- and 9-fold compared with the Δpepck and wild type cell lines, respectively (Table 3). Third, pyruvate, the PDH substrate, became a major excreted end product of glucose metabolism (accounting for 12.6% of total end products, with a 25-fold increase after tetracycline induction). Acetate was still detectable because of the impossibility to completely switch off gene expression by RNAi. Finally, as observed for the Δpepck mutant, the Δpepck/RNAi PDH-E2.i cell line switched to proline metabolism as shown by the increased rate of proline consumption (Table 2) and the increased amount of alanine produced from L-[4-13C]proline (Table 4). Altogether, these data show that the inhibition of acetate production in the absence of the succinate branches induced the arrest of glycolysis, compensated by a switch to proline (as observed for wild type procyclic cells grown in glucose-depleted conditions).

DISCUSSION

Kinetoplastids, including trypanosomatids and bodonids, are the only known organisms performing glycolysis in the specialized peroxisome organelles, called glycosomes. Within glycosomes, NADH consumed by the glyceraldehyde-3-phosphate dehydrogenase (step 8) needs to be regenerated by NADH-dependent oxidoreductases, to maintain the organellar redox (NADH/NAD+) balance. The prothetic form of T. brucei contains three different glycosomal pathways for re-oxidation of NADH into NAD+; i.e. succinic fermentation (steps 14–17), glycerol production (steps 6 and 7), and the Gly-3-P/DHAP shuttle (steps 6, 29, and 31–34).

Succinate Fermentation Is the Preferred Route for Glycosomal Reoxidation of NADH—Our data demonstrated that the contribution of the Gly-3-P/DHAP shuttle in the wild type environment is low (and possibly not significant). The RNAi knockdown of FAD-GPDH (step 29) resulted in no distinct phenotype discernable from the wild type cells, in terms of growth rate, rates of proline/glucose consumption, or metabolome analyses using NMR, HPLC, or ion chromatography-MS/MS. Similarly, the wild type parasites and the RNAiFAD-GPDH cell line did not use the glycerol production pathway, because glycerol is not detectable in the extracellular medium of cells incubated in the presence of d-[1-13C]glucose. Consequently, glycosomal succinic fermentation alone is sufficient for maintaining the glycosomal redox balance in wild type procyclic cells, as proposed previously (13). This contrasts with the bloodstream forms of T. brucei grown in aerobic conditions, which use only the Gly-3-P/DHAP shuttle to maintain the glycosomal redox balance.

Abolition of the Glycosomal Succinic Fermentation Stimulated Growth of the Parasite—This surprising behavior was observed in three independent Δpepck cell lines. This clearly indicates that, in the absence of this key glycosomal metabolic branch, the parasites develop successful metabolic alternatives. We have identified the following three main adaptations in response to the PEPCK gene deletion, (i) use of the Gly-3-P/DHAP shuttle to maintain the glycosomal redox balance; (ii) down-regulation of glucose metabolism, and (iii) switch to proline metabolism. Each of these adaptations will be individually addressed below.

The growth stimulation observed in the Δpepck cell lines is probably a consequence of these metabolic adaptations. Indeed, an increase of proline metabolism, although glucose is still consumed (albeit with a reduced rate), may stimulate the biosynthetic pathways by increasing the production of both ATP and biosynthetic precursors. This is consistent with the growth stimulation observed for different wild type procyclic strains grown in glucose-rich medium supplemented with N-acetyl-d-glucosamine (a glucose transport inhibitor), which also induces a switch to proline metabolism (38, 39). However, after a couple of weeks in the glucose-rich medium, the Δpepck cell line analyzed here reverted to normal growth, suggesting that the mutant reached a more stable metabolic steady state by reducing its metabolic activity.
Maintenance of the Glycosomal Redox Balance—Our data demonstrate a hierarchy between the three different glycosomal pathways for the regeneration of NAD⁺. As already mentioned, succinate fermentation is the first NAD⁺-production pathway used by the wild type procyclics to maintain the redox balance. When succinate fermentation is inactivated (Δpepck), the Gly-3-P/DHAP shuttle is primarily used instead. Glycerol production, which is not detectable in wild type cells, is also increased in the Δpepck mutant, but its contribution remains marginal (2.8% of the excreted metabolites). When both the Gly-3-P/DHAP shuttle and succinate fermentation were blocked (Δpepck/ΔpepckRNAiFAD-GPDHi), glycerol became a major end product of glucose metabolism (21.4% of excreted metabolites), indicating that glycerol production is significantly used only in the absence of the two other alternatives.

Down-regulation of Glucose Metabolism in PEPCK Null Environment—The consumption of glucose was significantly reduced (3.3-fold) in the Δpepck cell line, compared with wild type cells. In parallel, the intracellular amounts of the glycolytic intermediates glucose 6-phosphate and fructose 6-phosphate were also reduced in the Δpepck cell line, which is consistent with a reduced glycolytic rate (supplemental Fig. S1). The reduced glucose consumption in the Δpepck cell line is not due to the inability of the Gly-3-P/DHAP shuttle to maintain the redox balance, because both the glycosomal and mitochondrial parts of the shuttle are functional in procyclic cells (15, 40). In parallel, we confirmed the activity of the shuttle in the PEPCK null environment by showing that the rate of oxygen consumption in the presence of Gly-3-P is ~10-fold reduced in the Δpepck/ΔpepckRNAiFAD-GPDHi mutant, compared with wild type cells (Fig. 7). Altogether, these data suggest that, although the Gly-3-P/DHAP shuttle is functional, it is not well adapted to sustain a relatively high glycolytic flux in the procyclic trypanosome metabolic context.

Glycerol production is definitively not a good alternative for maintaining the glycosomal balance in procyclic trypanosomes, because glucose metabolism is almost abolished when both succinate fermentation and the Gly-3-P/DHAP shuttle are inactivated (16-fold reduction in the Δpepck cell line, compared with wild type cells). This may be related to the kinetic properties of the glycerol kinase. The conversion of Gly-3-P to glycerol is thermodynamically unfavorable and is only feasible at high Gly-3-P concentrations and a high ADP/ATP ratio (41). However, in the experimental conditions used for NMR investigations, the Δpepck/ΔpepckRNAiFAD-GPDHi mutant incubated with glucose as the sole carbon source was forced to consume glucose at a rate similar to that observed for the Δpepck cell line (212 versus 208 nmol of [13C]-enriched molecules excreted/h/mg of protein) (Table 3). In these conditions, the double mutant mainly converts glucose into acetate and glycerol (53.2 and 32.9% of the excreted end products, respectively) (Table 3). Interestingly, the nature of glucose metabolism in the Δpepck/ΔpepckRNAiFAD-GPDHi cell line is highly similar to that observed in the bloodstream forms grown anaerobically, except that the procyclic mutant further converts pyruvate into acetate. Although this double mutant cell line still has the capability to perform glycolysis, it switches to proline metabolism when placed in glucose-rich conditions, with almost no glucose consumed.

Activation of Proline Metabolism in the Δpepck Cell Line Grown in Glucose-rich Conditions—The absence of a growth phenotype in the Δpepck cell line is primarily due to the capacity of the parasite to switch to proline metabolism, as observed previously for wild type cells grown in glucose-depleted conditions (6, 7). This means that the metabolism becomes more oxidative in the Δpepck mutant. Accordingly, the Δpepck/ΔpepckRNAiATPε-F1βi and Δpepck/ΔpepckRNAiSDHi double mutants were not able to grow in glucose-rich conditions, although such a phenotype was observed for the Δpepck/ΔpepckRNAiATPε-F1βi and Δpepck/ΔpepckRNAiATPε-F1βi single mutants in glucose-depleted conditions (7). However, the metabolism of the Δpepck mutant in glucose-rich conditions does not closely mimic that of the wild type cells in glucose-depleted conditions. First, proline is consumed at a lower rate by the Δpepck mutant (0.40 versus 0.70 µmol consumed/h/mg of protein, respectively). Second, the ratio of alanine/succinate excreted from proline metabolism is considerably larger in glucose-depleted conditions (wild type and Δpepck mutant cell lines) compared with glucose-rich conditions (wild type cells) (106–15.1 versus 0.11). Again, the Δpepck cell line grown in glucose-rich conditions shows an intermediate value (0.65). This incomplete switch to proline metabolism is probably related to the remaining consumption of glucose observed in the Δpepck cell line. Interestingly, the Δpepck/ΔpepckRNAiATPε-F1βi double mutant expresses similar metabolic parameters (0.50 µmol of proline consumed/h/mg of protein, with alanine/succinate = 2.4), although its glucose consumption is almost abolished. This raises a number of questions such as the following. What is the maximum glycolytic flux compatible with the maintenance of the highest rate of proline consumption or how much glycolytic flux is required to fully repress proline metabolism? It has been shown previously that the metabolic adaptation is a graded response rather than a binary switch and that the repressive effect of glucose on proline metabolism seems to be related to the overall metabolic status of the cells rather than any allosteric action of glucose on its own (39). Alternatively, the presence of high concentrations of extracellular glucose (6 mM) might influence proline metabolism, even in the absence of a significant glycolytic flux. The two first steps of the proline metabolic pathway (proline uptake and dehydrogenation) are up-regulated upon glucose depletion (6), suggesting that their activity is controlled by glucose. However, the precise nature of this control is unknown, and further investigations would be needed to fully take apart the components of proline regulation.

Acetate Production in the PEPCK Null Environment—The production of acetate is similar in the Δpepck and wild type cell lines (154 versus 186 nmol of [13C]-enriched molecules excreted/h/mg of protein, respectively). Acetate is produced from glucose, the consumption of which is ~3.5-fold higher in the wild type cells compared with the Δpepck mutant. These data suggest that acetate production could be at its maximal capacity in the wild type parasite, because the higher glycolytic flux in this strain does not result in significant increase in acetate production. This hypothesis is consistent with the observation that glucose is converted into β-hydroxybutyrate in all cell lines lacking the PEPCK gene (Table 3). β-Hydroxybutyrate is usually produced in conditions where acetyl-CoA accumulates. Its
production by Δpepck mutants may result from a redistribution of carbon fluxes toward the acetate branch, which would become saturated, as a consequence of a limiting capacity. The absence of β-hydroxybutyrate production in the wild type cells is consistent with this model, because the succinification pathways redirect part of carbon flow toward succinate production and thus may prevent acetyl-CoA accumulation. Considering this flux redistribution, one would anticipate the observed abolition of glucose consumption when the acetate branch is inactivated in the PEPCK null background (Δpepck/RNAi/PDH-E2.i) (Table 2).

In conclusion, we show that the succinate fermentation, used by the procyclic trypanosomes to maintain the glycosomal redox and ATP/ADP balances, is not essential for the growth of the parasite. However, it is required to maintain a relatively high glycolytic flux. The other glycosomal NADH-consuming pathways (Gly-3-P/DHAP shuttle and glycerol production) are functional but not adapted to sustain a glycolysis-based metabolism in the procyclic cells. The flexibility provided by the redundant pathways might prove important under certain biological conditions that the parasite encounters in its complicated life cycle, such as during the migration from one tissue or an organ to another. In contrast, acetate production, the other major branch of glucose metabolism, is essential for the procyclic trypanosomes to generate ATP in the mitochondrion (11, 12) as well as to feed the cytosolic fatty acid biosynthesis through the so-called “acetate shuttle” (42). At least two different enzymatic activities account for the last step of the acetate branch (12). Identification of the acetate production enzymes will certainly prove helpful to fully understand all the roles played by this key pathway.

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