A \( \alpha \)-glycerophosphate Dehydrogenase is Present in *Trypanosoma cruzi* Glycosomes

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\( \alpha \)-glycerophosphate dehydrogenase (\( \alpha \)-GPDH-EC.1.1.1.8) has been considered absent in *Trypanosoma cruzi* in contradiction with all other studied trypanosomatids. After observing that the sole malate dehydrogenase can not maintain the intraglycosomal redox balance, GPDH activity was looked for and found, although in very variable levels, in epimastigotes extracts. GPDH was shown to be exclusively located in the glycosome of *T. cruzi* by digitonin treatment and isopycnic centrifugation. Antibody against *T. brucei* GPDH showed that this enzyme seemed to be present in an essentially inactive form at the beginning of the epimastigotes growth. GPDH is apparently linked to a salicylhydroxamic-sensitive glycerophosphate reoxidizing system and plays an essential role in the glycosome redox balance.

Key words: \( \alpha \)-glycerophosphate dehydrogenase - glycosome - *Trypanosoma cruzi*

*Trypanosoma cruzi*, like all known trypanosomatids, presents an unusual glucose metabolism. First, the 7-8 initial enzymes of the glycolytic pathway are compartmentalized in a microbody-type organelle, the glycosome (Opperdoes 1987, Taylor & Gutteridge 1987). Second, the oxidative degradation of glucose is not complete: only part of the glucose carbon appears as CO\(_2\) and the rest is excreted as reduced organic compounds, mainly succinate and alanine in *T. cruzi* epimastigotes (Cazzulo 1994). The incomplete breakdown of glucose into succinate is related to the presence in the glycosome of an auxiliary system, constituted by an ATP-producing phosphoenolpyruvate carboxykinase (PEPCK) and a malate dehydrogenase (MDH) (Urbina 1994), which transforms phosphoenolpyruvate (PEP) to malate by fixing CO\(_2\) and oxidizing reduced nicotinamide dinucleotide (NADH). Malate is eventually converted to succinate in the mitochondrion. This glycosomal system is present in all stages of the life cycle of *T. cruzi* (Urbina et al. 1993). Theoretically, MDH is able to reoxidize all the glycosomal NADH produced by glycolytic glyceraldehydephosphate dehydrogenase. Practically, for that to be so, all the PEP (2 moles per glucose) produced in the cytosol as a consequence of the glycosomal glycolytic activity, should re-enter the glycosome; and that is not possible due to the presence of a low cytosolic pyruvate kinase activity which converts part of the PEP to pyruvate. If the glycosomal membrane is considered as a true permeability barrier towards molecules such as oxidizing reduced nicotinamide dinucleotide (NADH), the existence of at least another glycosomal NAD\(^+\)-regenerating system, besides MDH, has to be postulated in order to account for the glycosome redox balance. The other dehydrogenase known to play a role in the redox balance of the glycosome of *T. brucei*, \( \alpha \)-glycerophosphate dehydrogenase (GPDH), has not been detected in *T. cruzi* (Taylor & Gutteridge 1987, Cazzulo et al. 1988) although it is present in all the other known trypanosomatidae (Tielens & Van Hellemond 1998). In these organisms, GPDH is part of a system which enables the reoxidation of glycosomal NADH by molecular oxygen through the classical dihydroxyacetone-phosphate/\( \alpha \)-glycerophosphate shuttle and a mitochondrial alternative \( \alpha \)-glycerophosphate oxidase or GPO (Tielens & Van Hellemond 1998). It is therefore noteworthy that different authors have observed an inhibition of oxygen consumption by salicylhydroxamic acid (SHAM) in *T. cruzi* epimastigotes (Felix et al. 1978, Stoppani et al. 1980, Engel et al. 1990), and even the direct oxidation of glycerocephosphate by mitochondrial extracts (Affranchino et al. 1985), an indication of the possible presence of a GPO system (generally referred as SHAM-sensitive alternative oxidase) and also of a \( \alpha \)-glycerophosphate generat-

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ing system. We therefore decided to have another look at *T. cruzi* GPDH (EC 1.1.1.8) activity.

Epimastigotes (EP strain) were cultivated, as already described by Concepcion et al. (1999), in LIT medium supplemented with 10 mM glucose and harvested at mid-exponential phase (around 0.6 OD at 600 nm) or as pointed out in the following experiments: (1) GPDH activity was determined on whole epimastigote extracts in both directions in order to avoid misinterpretations due to other NAD+/NADH-utilizing systems (Niesel et al. 1982). Only one extract out of nine (corresponding to independent cultures) did not present any detectable activity. In the other extracts GPDH specific activity varied from about 0.3 U/mg protein (two cases) down to 0.005 U/mg protein depending on the culture, with a mean value of about 0.07 to be compared with 0.75 and 0.15 U/mg protein for phosphoglucone isomerase (PGI) and hexokinase (HK) respectively. Fig. 1A shows that GPDH specific activity was not detectable at the beginning of the exponential phase, increased during growth. However an antiserum raised against *T. brucei* GPDH evidenced a maximum of cross-reacting protein at the beginning of the growth. Put together, these two observations indicated that most of the enzyme expressed during early exponential phase was inactive (Fig. 1B); (2) *T. cruzi* epimastigotes were permeabilized with increasing amounts of digitonin as already described by Concepcion et al. (1999). GPDH was liberated at the same digitonin concentrations as for HK, the classical glycosomal marker (Fig. 2), in agreement with a glycosomal localization. Phosphoglucone isomerase was liberated in a biphasic manner as expected for an enzyme presenting a dual location (Concepcion et al. 1999); (3) glycosomes were purified by differential and isopycnic centrifugations according to Concepcion et al. (1998, 1999). GPDH activity distribution was absolutely identical to those of PEPCK and HK by differential (not shown, see Concepcion et al. 1998) and isopycnic centrifugations (Fig. 3). The western blot performed in the presence of the anti-GPDH antibody exhibited the same distribution as the enzymatic activity (not shown), indicating an essentially exclusive glycosomal location of GPDH. In this experiment, the purified glycosomes presented latencies of 92% for GPDH, 87% for HK and 80% for PGI; (4) an estimation of GPO activity was obtained by determining SHAM inhibition of the oxygen consumption rate of whole cells in a Warburg respirometer (each 3 ml vial contained 1 mg of cell protein and 3 mM glucose; the center well contained 0.3 ml of 20% potassium hydroxide). The experiments were performed in the presence and absence of SHAM (1.2 mM) and/or cyanide (0.1 mM) on nine independently grown epimastigote cultures (harvested between 0.6 and 0.9 O.D. at 600 nm). SHAM inhibited oxygen consumption by about 23% as an average but with variations between 0% (one case) and 57%. Not only was the inhibitory effect of SHAM generally not additive with that of cyanide, but in two cases an antagonistic effect was observed. Antimycin inhibited between 90 and 100%. Glucose consumption [measured as residual glucose according to Bergmeyer (1983)] was also inhibited by SHAM between 30 and 60%, but not by cyanide. These results differed somewhat from those of Engel et al. (1990) who only observed SHAM inhibition on antimycin A inhibited cells and from those of Felix et al. (1978), but agree with them in indicating the likely presence of a GPO system in *T. cruzi* epimastigotes.

This work evidences that *T. cruzi* can synthesize, as do all other studied trypanosomatids (Tielens & Van Helmond 1998), a glycosomal
of the oxidative systems (Engel et al. 1990), that seems unlikely since we have shown here that GPDH activity may or may not be detectable in the same strain grown under the same conditions. Harvesting time is evidently an essential factor: our results, but also those of Felix et al. (1978), show that GPDH is expressed differentially during epimastigotes growth, with a maximum in some moment of the late exponential phase. However, it is likely that essentially uncontrolled medium factors are also involved in this apparent incongruity. However the differential expression of the GPDH gene might not be the unique factor at stake. Anti-GPDH antibody evidenced higher levels of the GPDH protein at the beginning of the growth curve than at the end, contrarily to what was observed with GPDH activity, thus indicating a possible post-translational activation and a possible explanation for the variable levels of GPDH activity. The variability of GPDH expression/activity may be correlated to pyruvate kinase (PK) activity since more glycosomal GPDH would be necessary to maintain the redox balance in situations with relatively high PK activity and therefore less PEP re-entry in the glycosome. A 5-fold increase in PK specific activity was observed (unp. obs.) during late exponential phase of epimastigotes growth. In any case even low GPDH specific activities are compatible with an auxiliary role in maintaining the redox balance since PK activity is generally low. The glycerol production observed in T. cruzi amastigotes (Sanchez-Moreno et al. 1995) confirms the presence of a GPDH gene in T. cruzi. This gene is homologous to that of T. brucei as shown by the cross-reaction observed with an antiserum raised against T. brucei GPDH. The existence of at least two NADH-reoxidizing systems in the glycosome of T. cruzi gives possible explanations to several observations that could only be rationalized on the ground of a permeable glycosome membrane; for example, that the glycolytic activity was only little affected by a drastic inhibition of PEPCK (Urbina et al. 1990).
Fig. 3: glycerolphosphate dehydrogenase distribution profile after isopycnic centrifugation. A post-small granule fraction was run on a linear sucrose gradient (0.3 to 2.5 M) and 20 fractions were collected from the density gradient (density varied from 1.02 to 1.28 g.cm⁻³). The experiment (repeated four times) was performed as described by Concepcion et al. (1998). The determination of enzymatic activities was performed according to Bergmeyer (1983) or as described in the previous experiments.

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