Potential Drug Targets in the Pentose Phosphate Pathway of Trypanosomatids

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ABSTRACT

The trypanosomatids, *Trypanosoma brucei*, *Trypanosoma cruzi* and *Leishmania spp*, are causative agents of important human diseases such African sleeping sickness, Chagas’ disease and Leishmaniasis, respectively. The high impact of these diseases on human health and economy worldwide, the unsatisfactory available chemotherapeutic options and the absence of human effective vaccines, strongly justifies the search for new drugs. The pentose phosphate pathway has been proposed to be a viable strategy to defeat several infectious diseases, including those from trypanosomatids, as it includes an oxidative branch, important in the maintenance of cell redox homeostasis, and a non-oxidative branch in which ribose 5-phosphate and erythrose 4-phosphate, precursors of nucleic acids and aromatic amino acids, are produced. This review provides an overview of the available chemotherapeutic options against these diseases and discusses the potential of genetically validated enzymes from the pentose phosphate pathway of trypanosomatids to be explored as drug target candidates.

INTRODUCTION

Trypanosomatids belong to the Euglenozoa phylum, Kinetoplastida class, Trypanosomatida order, and Trypanosomatidae family [1]. They are a group of flagellated protozoa parasites found primarily in insects but a few genera have a digenetic life-cycle involving a secondary host such vertebrates, invertebrates or plants [2]. Interestingly, as these parasites emerge from the most ancient eukaryotic lineages, with branches deeper than those from the younger metazoan kingdoms of fungi, plants and animals, they possess many peculiar characteristics [3]. The single branched mitochondrion containing a unique mitochondrial DNA structure called kinetoplast, the unidirectional gene clusters that are polycistronically transcribed, a RNA polymerase I-mediated transcription of protein coding genes, a RNA trans-splicing coupled to poly(A) addition, an extensive RNA editing of mitochondrial mRNAs, a compartmentalisation of energy metabolism with the glycolytic pathway and other enzymes sequestered in the specialized peroxisomes called glycosomes, and a redox metabolism based upon a unique thiol called trypanothione, are some of the unusual characteristics of trypanosomatids [4-9]. Trypanosomatids of the genera Leishmania and Trypanosoma are causative agents of humans diseases, with considerable morbidity and mortality, affecting more than 27 million people worldwide [10]. *Trypanosoma brucei*, *Trypanosoma cruzi* and *Leishmania spp*, are the causative agents of Human African trypanosomiasis, Chagas’ disease and Leishmaniasis, respectively. The geographic distribution of these diseases illustrates that they affect poor people living in the poorest regions of the world. In general the low access to ready diagnosis and affordable treatments, the absence of approved human vaccines and the use of the reference drugs for decades which induce severe side effects and leads to the emergence of drug-resistances makes these infections, high-impact neglected tropical diseases [10]. Since disease control relies mostly on chemotherapy, the search for new drugs becomes imperative.

In this review we will present the available treatment options for these diseases and discuss the potential of genetically validated enzymes of the pentose phosphate pathway to be explored as drug target candidates.
HUMAN AFRICAN TRYPANOSOMIASIS (HAT)

The causative agent and the clinical features. The causative agent of HAT, also known as sleeping sickness, is the extracellular flagellated trypanosomatid, *Trypanosoma brucei*. There are three *T. brucei* subspecies, of which two infect humans, *Trypanosoma brucei gambiense* (*T. b. gambiense*) and *Trypanosoma brucei rhodesiense* (*T. b. rhodesiense*), and one non-infective for humans, *Trypanosoma brucei brucei* (*T. b. brucei*), therefore routinely used in laboratories [11]. Different to humans, Nagana disease in animals result from infections with *T. congolense*, *T. vivax* or *T. b. brucei* [12]. Gambiense HAT, caused by *T. b. gambiense*, is considered an anthroponotic disease with a minor role for animal reservoirs, while rhodesiense HAT, induced by *T. b. rhodesiense*, is a zoonotic disease, affecting mainly animals; humans are only accidental hosts. Unlike *T. b. rhodesiense* infections, in which non-human vertebrates are the primary reservoir, representing <5% of HAT cases, *T. b. gambiense* infections represent >95% of the cases [13]. *T. b. gambiense* infections are found in West and Central sub-Saharan Africa, while *T. b. rhodesiense* causes infections in East sub-Saharan Africa [11]. The geographic barrier of the two forms of the disease coincides with the Rift Valley, with *T. b. gambiense and T. b. rhodesiense* present at the west and east side of the valley, respectively [14]. Curiously, Uganda is the only country where the two subspecies co-exist, however there still remains a spatial separation between the subspecies, mainly due to the different climatic and vegetative conditions required by the two species [15]. Although some patients are occasionally reported outside Africa [16], 36 sub-Saharan African countries are still considered the endemic areas for HAT[14].

HAT etiological agent is transmitted through the bite of an infected tsetse fly (genus *Glossina*). Thirty-one species have been described, however in nature, infections are carried almost exclusively by *Glossina palpalis* (for *T. b. gambiense*; distributed in the Atlantic coast from Senegal to Angola), *Glossina morsitans* (for *T. b. rhodesiense*; located mainly in East Africa) and *Glossina fuscipes* (for both; present in central Africa from Cameroon and Congo to the Rift Valley) [14]. The mammalian host becomes infected through the bite of an infected tsetse fly vector that injects infective metacyclic forms, covered with a variant surface glycoprotein (VSG). Actually, *T. brucei* parasites change the composition of the insect saliva, in a way that increases vector-host contact frequency and enhances the probability of parasite transmission [17]. Once inside the host, the metacyclic trypanosomes proliferate at the site of inoculation and then transform into slender forms, as they are carried by the draining lymph nodes to the bloodstream, where they replicate by binary fission [18]. Notwithstanding, the skin has recently been identified as an important but overlooked reservoir of these parasites [19, 20]. More recently, the adipose tissue forms (ATFs) were identified, as both replicative and infective parasites. These forms reside in the adipose tissue, the third major reservoir of *T. brucei* in the host, which is likely to underlie the weight loss experienced by the patients [21]. When parasitaemia in the host increases, the long slender trypomastigotes (proliferative forms) release a soluble factor called stumpy induction factor (SIF), which triggers via a quorum sensing mechanism, parasites differentiation into short stumpy forms (non-proliferative forms) [22]. Stumpy forms, which do not divide, are crucial for limiting parasitaemia within the host, and subsequent differentiation to procyclic forms when taken into a tsetse vector. During vector blood meal, stumpy forms expressing VSG are ingested and primed to differentiate into procyclic forms. After multiplication in vector midgut, procyclic trypomastigotes initiate a migration, that takes them through the peritrophic matrix, along the foregut to the proventriculus, and from there onwards through the mouthparts, salivary ducts.
and finally into the salivary gland [23]. In the proventriculus, procyclic trypomastigotes undergo extensive restructuring, coupled to an asymmetric division to produce one long epimastigote and one short epimastigote [24, 25]. The short epimastigote, coated with alanine rich protein (BARP), attaches to epithelial cells following arrival in the salivary gland [26]. In order to complete the life cycle, the attached epimastigotes undergo a final transformation into free metacyclic trypomastigotes, along with the acquisition of the VSG coat, for evasion in the mammalian host [23].

Clinically HAT evolves in two stages. The first, known as early or haemato-lymphatic stage, is defined by the presence of parasites in the lymph and blood systems, while the second, named as late or meningo-encephalitic stage, is characterized by the active invasion of trypanosomes in the central nervous system (CNS). The duration of the first stage depends on the T. brucei subspecies and varies from from weeks to months in acute T. b. rhodesiense infections, or from several months to years in chronic T. b. gambiense infections [14]. After inoculation, the formation of a trypanosomal chancre, characterized by local erythema, oedema, heat, tenderness and a lack of any suppuration [27], arises in about 50% of rhodesiense infections, but is rarely formed in gambiense infections. After 3–4 weeks, the chancre usually heals with overlying desquamation, sometimes with altered pigmentation [11]. In T. b. rhodesiense infections, which present poor demarcation between stages, pancarditis with congestive heart failure, pericardial effusion, and pulmonary oedema can cause fatalities. In contrast, T. b. gambiense infections show a more insidious development [11]. The first stage is characterised by general malaise, intermittent fever, headache, severe pruritus with scratching, skin lesions, mobile or rubbery lymphadenopathy, oedema of the face and extremities and, to a lesser extent, myocarditis, splenomegaly or hepatomegaly [27]. In the second stage general malaise worsens, headaches become more severe and patient sleep pattern is altered. Later this stage culminates in coma, severe organ failure and eventually death [11].

Current therapeutic options. Since there is no vaccine against HAT and chemoprophylaxis is not recommended due to drugs toxicity, the only preventive measure rely on vector control. Vector control has been a way to limit mostly rhodesiense infections, although it was recently applied in gambiense HAT [13]. Vaccines are unlikely to be developed as a result of the different T. brucei defence strategies, like antigenic variation [28] and abrogation of B cell lymphopoiesis [29, 30]. In fact, several attempts have been made to come out with a vaccine, but this goal has never been achieved. The absence of an effective vaccine in combination with the knowledge about the obstacles in HAT vaccination field, strongly justifies an improvement of existing chemotherapy. HAT chemotherapy is still unsatisfactory, as it is lengthy, costly, toxic, ineffective against both stages of the disease and requires parenteral administration. If untreated, HAT is almost 100% fatal [31]. There are currently four licensed treatment regimes: pentamidine, suramin, melarsoprol and eflornithine. Nifurtimox-eflornithine combination therapy besides being used off-license, was placed on the WHO List of Essential Medicines [32].

Pentamidine. Pentamidine isethionate (Pentacarinat®) is an aromatic diamidine used against first-stage disease caused by T. b. gambiense infection. Pentamidine is unsuitable for the treatment of advanced disease, in part due to serum binding and tissue retention properties that reduce blood-brain barrier traversal [33]. Four mg/kg are given daily, or on alternate days by intramuscular injection for 7-10 days [34]. When given by intramuscular injection, site pain and transient swelling, abdominal pain and gastrointestinal problems, and hypoglycaemia (5–40%) are the most frequently reported adverse events. The drug enters
parasites principally via the adenosine transporter 2 (P2) aminopurine permease [35], however low and high affinity pentamidine transporters (LAPT1 and HAPT1, respectively) and a plasma membrane H+ ATPases, HA1-3, also contribute to uptake [36, 37]. Pentamidine action appears to be multifactorial possibly due to the binding to DNA [38]. It is also known to collapse the mitochondrial membrane potential, and inhibit the F1F0-ATPase [39]. Consequently, it is likely that the antitypansosomal activity of pentamidine is the result of selective accumulation, leading to multiple deleterious effects, rather than effects on a specific ‘diamidine target’ [40]. It is noteworthy that confirmed pentamidine failures are rare, likely due to low drug-resistance. Recently through genome-scale RNA interference target sequencing (RIT-seq) screens, aquaglyceroporins (AQPs) loss-of-function was linked to melarsoprol-pentamidine cross-resistance [37]. AQPs facilitate the transport of water and small neutral solutes across membranes in several organisms. *T. brucei* expresses three AQPs (TbAQP1-3) which are dispensable for viability and interestingly, for osmoregulation, yet making important contributions to drug-uptake, glycerol-transport and respiratory-inhibitor sensitivity [41, 42]. AQP2 is responsible for the uptake of both pentamidine and melarsoprol, in the case of the former, most likely by receptor-mediated endocytosis and has been linked to clinical cases of resistance for both drugs [41]. This was an important finding since the cross-resistance phenotype could not be explained based only in P2 and HAPT transporters.

**Suramin.** Suramin (Germanin®) is a sulphonated naphthylamine used for first-stage *T. b. rhodesiense* disease. Suramin is administered by intravenous injections every 3-7 days for 31 days (20mg/Kg, with a maximum of 1g per injection) [43]. Adverse drug reaction include acute and late hypersensitivity, nephrotoxicity, peripheral neuropathy, and bone marrow toxicity with agranulocytosis and thrombocytopenia [44]. This compound is selectively concentrated by trypanosomes through receptor-mediated endocytosis when conjugated with low-density lipoproteins (LDL) [45]. A high-throughput sequencing of a suramin-selected RNAi library identified a cohort of proteins contributing to drug efficacy [37]. In a proposed model, a 75 kDa ISG (ISG75) acts as a major receptor for suramin delivering the drug into the degradative arm of the endocytic pathway. Suramin is delivered to the lysosome by either the serum protein carrier being cleaved by cathepsin-L upon reaching the lysosome, or by ISG75 being degraded at the late endosome. Once free, suramin may inhibit lysosomal enzymes and may also escape into the cytoplasm via the major facilitator superfamly transporter (MFST), resulting in inhibition of other cellular processes [46]. This large negatively-charged polyanion appears to inhibit non-specifically many positively-charged enzymes, so it is difficult to conclude which may be the determinant of drug action [47]. More recently, it was shown that this compound inhibits a number of glycolytic enzymes [43], such as pyruvate kinase by binding the ATP site and that clearly the sensitivity to suramin is dependent on the uniquely active endocytic apparatus in *T. brucei* bloodstream forms [48, 49]. Resistance to suramin was reported in the 1950s [43]. The mechanisms for suramin resistance are not yet elucidated but may involve changes in ISG75 and in proteins of the endocytic apparatus, including AP-1 (adaptin complex-1), GLP-1 (Golg/ilysosomal protein-1), EMP70 (endosomal membrane protein 70), MFST, p67 (major lysosomal glycoprotein), and cathepsin-L [37].

**Melarsoprol.** Melarsoprol (Mel B, Arsobai®), a melaninophenyl arsenical synthesised in 1949, is still the most powerful trypanocide available to cure both stages of *T. b. gambiense* and *T. b. rhodesiense* HAT [50]. Importantly, is the only effective drug against late stage *T. b. rhodesiense* infections. It is usually administered intravenously and the recommended course is 10 days long with daily injections of 2.2 mg/kg [51]. Melarsoprol injections are extremely painful and toxic, and the most important reaction is a post-treatment reactive encephalopathy in 10% of patients, half of whom die, leading to an overall mortality of
about 5% [52]. Skin reactions, peripheral motoric, sensorial neuropathies and thrombophlebitis can also occur [53]. The parasites lyse rapidly when exposed to melarsoprol [54]. It is believed that melarsoprol is converted in the host to melarsen oxide, which is then transported by P2 transporter [55]. Melarsen oxide acts mainly through the formation of a stable adduct with trypanothione [T(SH)₂], known as MelT, which is an inhibitor of trypanothione reductase (TR), a central enzyme of the parasite thiol/disulfide redox balance [56, 57]. Additionally, melarsoprol also formed adducts with lipoic acid [58] and inhibits glycolysis [59]. In some T. b. gambiense HAT areas, melarsoprol treatment failures have already reached levels of 30% [60-62]. As mentioned above, the cross-resistance between pentamidine and melarsoprol was firstly associated with P2 transporter, once Tbat1 gene deletion and loss-of-function mutations were described in melarsoprol resistant strains generated in the laboratory [63-65], and the same mutations were also found in T. brucei spp. field isolates [66-68]. In addition, Tbat1-null trypanosomes were found to have lost the adenine-sensitive component of adenosine and melarsoprol-import [69, 70]. More recently, and as reported in pentamidine section, aquaglyceroporin 2 (AQP2) loss-of-function was linked to melarsoprol-pentamidine cross-resistance [37].

**Eflornithine.** Eflornithine (DL-α-Difluoromethylornithine, DFMO; Ornidyl™) was developed in the 1970s [71]. Due to the unacceptable toxicity of melarsoprol, eflornithine is the first line treatment for the second stage T. b. gambiense disease [72-74]. It is administered intravenously. The high IC₅₀ and the low half-life dictate the use of large quantities of the drug (nearly 4 kg for a 50 kg patient) for the treatment regime of 56 infusions over 14 days [53]. It is logistically difficult to deliver and to administer in the rural areas leading to a continued reliance on melarsoprol in impoverished areas. Adverse drug reactions include bone marrow toxicity leading to anaemia, leucopenia, and thrombocytopenia (25–50%), gastrointestinal symptoms (10–39%), and convulsions (7%) [75]. Eflornithine mode of action is well understood as it irreversibly inhibits ornithine decarboxylase (ODC) an enzyme of the polyamines pathway [76, 77]. Eflornithine is only active against T. b. gambiense forms of HAT possibly due to a reduced rate of ODC turnover in this subspecie [78]. Identically the pronounced differences in ODC turnover rates between mammals and parasites explain drug selectivity [79]. American trypanosomes are totally refractory as they lack the molecular target. ODC inhibition by eflornithine leads to an increase in levels of ornithine, S-adenosylmethylionine (AdoMet) and decarboxylated S-adenosylmethionine (dcAdoMet), and a decrease in putrescine, spermidine (Spd) and T(SH)₂ [80-82]. As Spd conjugates with two molecules of GSH to form T(SH)₂ which protects the parasite from oxidative stress, the decrease in Spd and consequently in T(SH)₂ is therefore harmful for the parasites [9, 83]. Downstream of these metabolic changes, a generalized decrease in DNA, RNA and protein synthesis, including synthesis of VSG [84, 85] and morphological and biochemical changes like differentiation into stumpf forms can occur [86]. The fact that the trypanosomes are unable to undergo antigenic variation facilitates the death of the parasites by the host immune system [43]. Ultimately, the combination of distinctive parasite features, regarding the molecular target and polyamine metabolism, as well as the simultaneous interference in multiple cellular processes, render eflornithine a very successful drug. Resistance appears associated to a loss of the, amino acid transporter (TbaAT6) but whether this mechanism is found in field isolates awaits further investigation [33, 87-89].

**Eflornithine-nifurtimox.** The nifurtimox/eflornithine combination therapy (NECT) was advanced in 2009 [90, 91] for the treatment of late-stage T. b. gambiense, after showing equivalent to better efficacy than eflornithine alone in clinical trials [92]. NECT has not yet been tested against T. b. rhodesiense [93]. In combination, nifurtimox is given orally (daily intravenously. The high IC₅₀ and the low half-life dictate the use of large quantities of the drug (nearly 4 kg for a 50 kg patient) for the treatment regime of 56 infusions over 14 days [53]. It is logistically difficult to deliver and to administer in the rural areas leading to a continued reliance on melarsoprol in impoverished areas. Adverse drug reactions include bone marrow toxicity leading to anaemia, leucopenia, and thrombocytopenia (25–50%), gastrointestinal symptoms (10–39%), and convulsions (7%) [75]. Eflornithine mode of action is well understood as it irreversibly inhibits ornithine decarboxylase (ODC) an enzyme of the polyamines pathway [76, 77]. Eflornithine is only active against T. b. gambiense forms of HAT possibly due to a reduced rate of ODC turnover in this subspecie [78]. Identically the pronounced differences in ODC turnover rates between mammals and parasites explain drug selectivity [79]. American trypanosomes are totally refractory as they lack the molecular target. ODC inhibition by eflornithine leads to an increase in levels of ornithine, S-adenosylmethylionine (AdoMet) and decarboxylated S-adenosylmethionine (dcAdoMet), and a decrease in putrescine, spermidine (Spd) and T(SH)₂ [80-82]. As Spd conjugates with two molecules of GSH to form T(SH)₂ which protects the parasite from oxidative stress, the decrease in Spd and consequently in T(SH)₂ is therefore harmful for the parasites [9, 83]. Downstream of these metabolic changes, a generalized decrease in DNA, RNA and protein synthesis, including synthesis of VSG [84, 85] and morphological and biochemical changes like differentiation into stumpf forms can occur [86]. The fact that the trypanosomes are unable to undergo antigenic variation facilitates the death of the parasites by the host immune system [43]. Ultimately, the combination of distinctive parasite features, regarding the molecular target and polyamine metabolism, as well as the simultaneous interference in multiple cellular processes, render eflornithine a very successful drug. Resistance appears associated to a loss of the, amino acid transporter (TbaAT6) but whether this mechanism is found in field isolates awaits further investigation [33, 87-89].
A dose of 15 mg/kg, three times a day for 10 days) and the eflornithine infusions are reduced in frequency (daily dose of 400 mg/kg in 14 slow infusions, every 12 hours for 7 days). The administration of the combination therapy compared to eflornithine monotherapy is much easier to implement. NETC allows the use of lower doses of each drug, being logistically easier to transport due to the lower quantities required and the reduction in refrigeration costs [92]. NECT generates adverse events, mainly abdominal pain, vomiting and headache, however, the severity of these events is relatively low compared to previous treatments, and the majority of patients treated make a good recovery [90, 94]. With regard to resistance, a genome-scale RNAi library, to screen nilurtimox and eflornithine resistance, confirmed previous findings that nitroreductase loss-of-function is the major potential mechanisms of resistance to nilurtimox [88, 95].

**Fexinidazole.** Fexinidazole is in the development pipeline, being currently tested in a pivotal phase II/III comparative study in HAT endemic regions. If successful, it will provide patients with the first oral treatment for stage two sleeping sickness [96]. This compound, which came out from a screening of over 700 nitroheterocyclic molecules, is a 2-substituted 5-nitroimidazole that exhibits in vitro and in vivo activity against both T. b. rhodesiense and T. b. gambiense [97, 98]. It is rapidly metabolised through oxidation, forming at least two pharmacologically active metabolites, fexinidazole sulfoxide and fexinidazole sulfone, which presumably account for much of the trypanocidal activity [97]. Although the mode of action is still not fully clear, it is likely that fexinidazole acts similarly to other 5-nitroimidazole drugs [99, 100].

**CHAGAS DISEASE**

The causative agent and the clinical features. *T. cruzi* is the parasite responsible for Chagas’ disease. There are six discrete typing units (DTUs) of *T. cruzi* (TcI-TcVI), which have distinct geographical distributions, extensive genetic diversity and differential virulence and pathogenic characteristics. TcI, TcII, TcV and TcVI are the main agents of human Chagas disease in the Americas, and all are capable of causing cardiomyopathies, however, only TcII, TcV and TcVI have been so far associated to chronic digestive syndromes [101]. Despite the difficulty of an accurate determination of the number of cases, it is estimated 6 to 7 million chronically infected people worldwide and 10,000 to 14,000 deaths annually [102, 103]. Chagas’ disease is endemic in 21 Latin American countries [103]. Chagas was entirely confined mostly to Latin America, however nowadays it has spread to other continents, like United States, Canada, European countries, and some Western Pacific countries, due to a rise in population movements between Latin America and other continents [104-107], becoming a global health problem.

*T. cruzi* is transmitted to humans usually by the faeces of triatomine bugs. Parasites cannot penetrate intact skin and usually enter through skin microlesions originated from scratch the itching vector’s bite. Similar, to other trypanosomatids, the *T. cruzi* life cycle comprises different developmental forms including epimastigotes and metacyclic trypomastigotes in the insect vector while in the mammalian host, bloodstream trypomastigotes and replicative intracellular amastigotes [108]. Mammals’ infection occurs when infective metacyclic forms enter the mucosa and invade local reticuloendothelial and connective cells. Metacyclic trypomastigotes express an important adhesion molecule, GP82 (a surface glycoprotein of 82
kDa), which contributes to parasite internalization [109]. Inside the cytoplasm of host cells trypomastigotes differentiate into amastigotes that replicate by binary fission. When cells become full of amastigotes, they transform into trypomastigotes, bursting the infected cells. The released trypomastigotes then invade adjacent tissues, and spread via the lymphatics and bloodstream to distant body sites, infecting different cell types, like muscle and ganglion cells, where they undergo further cycles of multiplication [108]. The insect vector becomes infected when it ingests circulating trypomastigotes during a blood meal.

Chagas’ disease comprises an acute and a chronic phase. The acute phase lasts between 1 to 2 months and usually passes unnoticed as the clinical symptoms are nonspecific and include fever, hepatosplenomegaly or lymphadenopathy. The inflammatory oedema at the entry site, known as chagoma in the skin, and Romaña’s sign in the eyelid, consist of specific symptoms [110]. Acute myocarditis and acute meningoencephalitis can be occasionally seen in young children aged 1-5 years, being in most cases fatal and abnormalities in the electrocardiogram are observed in about 50% of cases which usually disappear later in the course of the disease [111]. Following the acute phase, the immune response induces a reduction of the parasite load but do not prevent the development of the disease [108]. About 60% to 70% of patients never develop clinical manifestations [112], while in the remaining, up to 30% of chronically infected people develop cardiac alterations and up to 10% develop digestive, neurological or mixed alterations, usually 10 to 30 years after the initial infection [103]. The lesions of chronic Chagas heart disease are focal or extensive myocardial fibrosis, resulting from myocardial cell destruction due to direct parasite action, inflammatory response, and neuronal involvement. The gastrointestinal manifestations consist of progressive enlargement of the esophagus or colon caused by chronic inflammation and destruction of parasympathetic neurons [101]. Among all complications, progressive heart failure (70%) and sudden death (30%) are still the most frequent causes of death in patients with Chagas’ disease [113].

Current therapeutic options. Since there is no vaccine, Chagas’ disease control programs in endemic areas have been focused on preventing parasite transmission through household insecticide application. Additionally, once triatomine bugs live mostly in rural endemic countries especially in poor housing conditions, hence improved housing can decrease the spread of the disease. In non-endemic regions, where vectors are absent, control strategies include preventing $T. \ cruzi$ transmission from blood transfusion, organ transplantation, and mother to child, by screening donated blood and organs for the presence of parasites.

This disease is curable if treatment is initiated soon after infection, the current treatment options are limited to only two poorly tolerated nitro-heterocyclic drugs: benznidazole and nifurtimox. The toxicity of the available drugs [114], the emergence of benznidazole resistant $T. \ cruzi$ strains showing cross-resistance to nifurtimox [115], and the lack of alternative medications, underline the imperative need to develop new strategies for chemotherapy. The promising azoles compounds, specifically posaconazole and the ravuconazole prodrug E1224, were disappointing as they failed in recent clinical trials, with treatment failure in patients reaching 70% to 90%, as opposed to 6% to 30% failure for benznidazole-treated patients [112].

Benznidazole. Benznidazole, a 2-nitroimidazole (N-benzyl-2-nitroimidazole acetamide) is the reference drug against Chagas’ disease and is administered orally. For acute treatments, 5 to 7.5 mg/kg/day is used during 30 to 60 consecutively days, and divided in two or three daily doses. In the case of recent chronic infections, when the infection was acquired in the last 10 years, the treatment should be made 5 mg/kg/day during 30 to 60 days.
In accidental infections, the treatment if started immediately, last for only 10 to 15 days. Cases of late chronic infections without clinical manifestation or with mild cardiac or digestive manifestations should be treated during 60 to 90 days with the purpose to reduce the evolution of Chagas disease to more severe forms, fact that has not yet been proven [114]. Benznidazole adverse reactions include hypersensibility, dermatitis with cutaneous eruptions (between 7th-10th day of treatment), generalized oedema, fever, lymphadenopathy, articular and muscular pain, depression of bone marrow, thrombocytopenic purpura and agranulocytosis, polynuropathy, paraesthesia and polynuropitis of peripheric nerves [116]. The mode of action of this compound could involve covalent bond or other interactions of nitroreduction intermediates with parasite components [117] or binding to DNA, lipids and proteins [118].

**Nifurtimox.** Nifurtimox, another reference drug against *T. cruzi* infections, is a 5-nitrofuran (3-methyl-4-(5′-nitrofururylideneamine) tetrahydro-4H-1,4-tiazine-1,1-dioxide). As benznidazole, nifurtimox is given orally. In acute phase treatment and congenital cases its recommended 8 to 10 mg/kg/day during 30 to 60 consecutively days, divided in two or three daily doses, while in initial chronic infections the treatment should be made with 8 mg/kg/day during 30 to 60 days). The mode of action of nifurtimox involves generation of nitroanion radical by nitroreductases that, in the presence of oxygen, leads to the formation of reactive intermediates [119]. In trypanosomes the drug must undergo activation by nitroreduction, undertaken by NADH-dependent type I nitroreductases [95]. Recent evidences on the mode of action of nifurtimox showed that the formation of an open chain nitrile causes cellular death due to its interaction with a range of cellular targets [120].

**LEISHMANIASIS**

The causative agent and the clinical features. More than twenty *Leishmania* species and subspecies have the ability to cause Leishmaniasis [121]. The classification of *Leishmania* into subgenus *Leishmania*, which includes *L. donovani*, *L. infantum*, *L. major*, *L. tropica*, *L. aethiopica*, *L. mexicana*, *L. pifanoi*, *L. amazonensis* species, and subgenus *Vianna*, that comprises *L. braziliensis*, *L. guyanensis*, *L. panamensis* and *L. peruviana* species, was initially based on the vector gut parts colonized by the parasites, and later on was supported by DNA sequence-based phylogenetic analyses [122-124]. Some *Leishmania* spp., including *Leishmania major*, *Leishmania mexicana*, *Leishmania amazonensis* and *Leishmania braziliensis*, primarily cause tegumentary leishmaniasis, while others, like *Leishmania donovani* and *Leishmania infantum* disseminate to internal organs such as the liver, spleen and bone marrow causing visceral leishmaniasis. This later accounts for most of the deaths each year that are due to leishmaniasis [125]. *Leishmania* is prevalent in tropical and subtropical areas. This organism falls within two main groups, the Old World species (found in the eastern hemisphere; Europe, Africa and Asia), and the New World species (present in the western hemisphere; America). Species belonging to the subgenera of *Leishmania* are usually encountered in both Old and New Worlds, while others belonging to the subgenera *Vianna* are found only in the New World. Visceral leishmaniasis is caused by *L. donovani* in the Indian subcontinent and East Africa, and by *L. infantum* in other parts of Asia and Africa, Europe and the Americas. Tegumentary leishmaniasis can result by the infection of approximately 15 leishmania species: *L. major*, *L. tropica*, *L. aethiopica* and sometimes *L. infantum* in the Old World, and *L. braziliensis*, *L. amazonensis*, *L. guyanensis*, *L. panamensis*, *L. mexicana*, *L. pifanoi*, *L. venezuelensis*, *L. peruviana*, *L. shawi*, and *L. lainsoni* in the New World [126, 127]. Leishmaniasis is currently found in 88 countries, in which 72 are developing countries [128].
Interestingly, more than 90% of reported cases are located in 11 countries, India, Bangladesh, Nepal, Sudan, Brazil, Bolivia, Peru, Afghanistan, Iran, Saudi Arabia, and Syria [129]. This disease is associated with population displacement, poor housing, and lack of resources, therefore many cases are unreported. Nevertheless an estimated 1.3 million new cases and 20000 to 30000 deaths occur annually [130]. The geographical distribution of leishmaniasis has been increasing lately due to the incidence of HIV/Leishmania co-infections. It is estimated that HIV increases the risk of visceral leishmaniasis by 100-1000 times in endemic regions [131]. Moreover, the increased overseas travel, U.S. Gulf War veterans, the simultaneous encroachment of leishmaniasis and of HIV infection into rural areas have contributed to the increased appearance of leishmaniasis in developed countries [132, 133].

The natural transmission of leishmaniasis is mediated by the bite of an infected female sandfly upon blood feeding [134]. Within the order Diptera and the family Psychodidae, two sandflies genera, *Phlebotomus* in the Old World and *Lutzomyia* in the New World, are capable of transmitting the disease to man. About 70, from around 1000 known sandfly species, are competent leishmaniasis vectors [135] and only female sandflies can transmit the parasite as they need periodic blood meals in order to get the required nutrients for egg development. These vectors can be specific or permissive, whether they allow the development of one or several *Leishmania* species, respectively (Bates, 2008; Volf & Myskova, 2007). For instance, *Phlebotomus phlebotomus papatasi* and *P. Paraphlebotomus sergenti*, can only be infected by one species, whereas *Lutzomyia longipalpis* can transmit several parasite species [129]. Only on rare occasions, the vector requirement can be bypassed, such as in transplacental transmission, laboratory accidents or needle sharing [136-138]. In case of canine leishmaniasis, as parasites can be found in saliva, urine, semen and conjunctival secretions, as well as in blood, rare cases of horizontal transmission have been reported between dogs in the same household or kennel. Also rarely some ticks and canine fleas may also act as mechanical vectors [139].

During its life cycle *Leishmania* parasite goes through two main stages: one extracellular, the promastigote stage within invertebrate hosts (phlebotomine sandfly), and one intracellular, the amastigote stage in vertebrate hosts. During blood feeding by infected sandflies, infective non-dividing metacyclic promastigotes are regurgitated and delivered to the host, together with immunomodulatory parasite-derived proteophosphoglycans and various salivary components [125, 140, 141]. In the local environment, the parasites are then phagocytosed by one of several possible cell types, including neutrophils, macrophages and dendritic cells. Inside the cells, metacyclic promastigotes evolve into spherical intracellular forms, amastigotes, which undergo replication by binary fission. When too many amastigotes are present, the cells disrupt allowing reinfection of local phagocytes. Taken up by another sandfly, upon a blood meal, amastigotes are then converted into promastigotes in the midgut. The transmission cycle is then completed when procyclic promastigotes differentiate again into metacyclic promastigotes, which are located ready for transmission at the sandfly stomodeal valve (an invagination of the foregut into the midgut) [125].

Leishmaniasis is associated with a wide spectrum of clinical manifestations. The clinical outcome of a *Leishmania* infection depends upon a multifaceted association of factors among the three main players involved: hosts, parasites, and vectors. The disease has three different manifestations, cutaneous (CL), mucocutaneous (MCL) and visceral (VL), that differ in symptoms and consequences [142].

**Cutaneous leishmaniasis.** CL is the most common and most widespread. It is characterized by painful skin sores located at the site or within close proximity of a sand fly
There is an emergence of one or several ulcer(s) or nodule(s) in the exposed areas of the skin. The ulcers are generally self-healing with no need for specific treatment, however they leave disfiguring scars [143]. The clinical manifestations are quite variable with regard to the formation of the ulcers, the level of inflammation and the healing time, reflecting 1) the diversity of Leishmania species that can cause CL; 2) the host status [144, 145]. CL has a worldwide distribution, but most cases (~95%) occur in the Americas, the Mediterranean basin, the Middle East and Central Asia. CL is endemic in 70 countries, but strikingly, more than two thirds of CL cases take place in 6 countries: Afghanistan, Algeria, Brazil, Colombia, Iran and the Syrian Arab Republic. Estimates point to 0.7 to 1.3 million new cases worldwide annually [130].

Mucocutaneous leishmaniasis. MCL is a severe form of CL, characterized by destructive ulcerations of the mucosa, extending from the nose and mouth to the throat cavities and surrounding tissues. This extensive tissue destruction is accompanied by an uncontrolled immune cell infiltration and inflammation [146]. Moreover, the consequent facial disfigurement profoundly burdens the patients with a lifelong social stigma. The current view is that the occurrence of MCL is determined by the host genetic background, but the severity of the pathology may be exacerbated by parasite-driven factors [147]. Almost 90% of the cases occur in South America, particularly in Bolivia, Brazil and Peru and is caused by species of the Viannia subgenera [130]. In contrast to the cutaneous form, self-healing is rare, the lesions are difficult to treat and potentially fatal due to opportunistic bacterial secondary infections [143].

Visceral leishmaniasis. VL is the most severe, with a mortality rate of essentially 100% if untreated. It is characterized by irregular bouts of fever, substantial weight loss, swelling of spleen and liver, and anaemia [129]. L. donovani is the causative agent of VL in Asia and East Africa, whereas L. infantum equally leads to visceral compromise but in the Mediterranean Basin and South America [135]. Differently, while dogs are the most important reservoirs of VL caused by L. infantum, in the case of L. donovani, the disease is mostly anthroponotic [143]. Another striking difference is that L. donovani is capable of causing disease in people of all ages, although children and young adults are the most affected. In the case of L. infantum, small children used to be the main age group, however, as HIV (Human Immunodeficiency Virus)/L. infantum co-infections emerged, nowadays adults actually represent almost half of VL cases in Europe [130, 143]. Post Kala-azar Dermal Leishmaniasis (PKDL) is a resurgence of the visceral form, months or even years after treatment and is characterized by macular, maculopapular, and nodular rashes [129]. The cause, risk factors and pathogenesis of PKDL are largely unknown, although it has been associated with several host factors [148] and as drug related phenomenon [149].

Current therapeutic options. As antileishmanial vaccines are still under development [150], current leishmaniasis control strategies rely upon case management (detection and treatment), reservoir and vector control, and chemotherapy. Active case detection and surveillance, accompanied by preventing reinfection measures, through targeting the socio-economic and cultural factors, should be implicated in the attempts to reduce disease transmission. With respect to reservoir control, infected dogs are most frequently treated with the drugs meglumine antimoniate, alopurinol, amphotericin B, or a combination of meglumine antimoniate and alopurinol. However, these therapies are the also used against human leishmaniasis and usually do not prevent relapse of the disease or eliminate parasite carriage. Similar to the above mentioned trypanosomatid diseases, new drugs and treatment strategies are necessary to achieve a consistent parasitological cure [151]. Sandflies eradication is widely put in practice by the use of chemicals, environmental management and personal
protection [134]. Nevertheless, the major control strategy is still based on chemotherapeutic treatments which are expensive, toxic and associated with high relapse and resistance rates. The leishmaniasis chemotherapy depends on some factors including type of disease, parasite species and geographic location [130]. For the past seven decades, chemotherapy has mostly relied on pentavalent antimonials, the standard first line treatment for both visceral and cutaneous leishmaniasis, but the emergence of resistance has limited their usefulness. Alternative therapies, like amphotericin B and its lipid formulation, miltefosine and paromomycin, are available but their use is limited due to toxicity or high cost [152]. Once all these drugs are parenteral, the discovery of effective oral agents is crucial [153].

**Pentavalent antimonials.** Pentavalent antimonials were developed in 1945 [154] and this family of compounds, represented by sodium stibogluconate (Pentostam®) and meglumine antimonate (Glucantime®), remains the first line therapy against leishmaniasis in most of the world's regions, with few exceptions, like Bihar state [155]. The antimonial-based therapies have disadvantages that include the long-term treatment and parenteral administration under close medical supervision due to high toxicity. Their high toxicity represents an important limitation in the use of these drugs by pregnant women, the elderly, and individuals with cardiac disease, renal disease, or liver alterations [156]. These drugs are produgs that require biological reduction to trivalent antimonials for antileishmanial activity. The bioactivation can occur in the host cell or the parasite: glutathione, present in the cytosol of macrophages, cysteine or cysteinylglycine thiols, found mostly inside the host lysosomes, as well as trypanothione, the most abundant thiol in *Leishmania* parasites, can actively reduce SbV [157, 158]. Additionally, SbIII enters the parasites via aquaporin 1 (AQP1) [159] and *Leishmania* enzymes such as thiol dependent reductase 1 (TDR1) and antimoniate reductase have also been implicated in SbV bioactivation [160]. Despite the mechanism of action not being fully understood, SbIII is well established as the active species that ultimately leads to parasite death [161]. There is some evidence supporting zinc finger proteins and trypanothione reductase as possible molecular targets [162, 163]. Inclusively, by exposing the parasites to trivalent antimonials, an efflux of glutathione and trypanothione from promastigotes and isolated amastigotes was observed, suggesting an interference with the parasites' redox state [164]. Additionally, these drugs appear to inhibit parasite glycolysis and fatty acid β-oxidation, however no specific targets on these pathways have been identified [165, 166]. Another possible mode of action is based on SbV's capacity to form complexes with ribonucleosides, which can inhibit purine transporters or directly interfere with the purine salvage pathway in *Leishmania* [167].

In the past few years, following the growing and alarming emergence of antimonial resistance in endemic areas, many studies have been conducted to dissect the molecular mechanisms. Inclusively, microarray technology and proteomic screening have been employed in order to elucidate these mechanisms [168-170]. Those include loss of AQP1 activity [159], extrusion of trypanothione/SbIII complexes by ATP-binding cassette (ABC) transporters, sequestration of SbIII in vacuoles [171] or downregulation of MAPK1 that is a negative regulator of P-glycoprotein type efflux pumps [172]. Therefore, the mechanism of resistance is multi-factorial, including a decrease in uptake, an increase of efflux/sequestration and the modulation of the parasite thiol metabolism [173, 174]. Despite all these drawbacks, antimonial drugs remain highly effective in areas where resistance has not yet emerged [175].

**Amphotericin B.** Amphotericin B was originally developed as a systemic antifungal drug that targets ergosterol-like sterols. With ergosterol being one of the most abundant sterol in *Leishmania* promastigotes and amastigotes surface membranes [176, 177], amphotericin B treatment leads to an increased permeability and an influx of ions [176-178]. As mentioned
above, due to the increased emergence of resistances to pentavalent antimonials in India, amphotericin B deoxycholate in a dose of 0.75-1mg/kg for 15 to 20 infusions daily or on alternate days, produced cure rates of 97%, is the drug of choice in this region, despite intravenous administration and its high toxicity requiring close medical supervision [179, 180]. It causes adverse effects that include nausea, vomiting, fever, hypokalemia, renal failure, anaemia, leukoencephalopathy and heart problems [181]. The toxic events were substantially reduced by the emergence of amphotericin B lipid formulations. Even though, in many undeveloped countries or impoverished communities the use of lipid formulations was until recently impractical due to their high cost and the need of close monitoring due to the risk of serious, even fatal, side-effects [129]. Three amphotericin B lipid formulations are commercially available: liposomal amphotericin B (AmBisome®), amphotericin B colloidal dispersion (Amphocil®) and amphotericin B lipid complex (Albacete®). AmBisome® is the best tolerated [179] and it is the first-line treatment of visceral leishmaniasis in the United States and southern Europe, representing the treatment of choice for immunocompetent patients in the Mediterranean basin and the preferred drug for HIV/VL co-infection. In immunosuppressed patients a total dose of 40mg/kg AmBisome® spread over 38 days is recommended [182], however in immunocompetent individuals different dose regimens have been used [183-185] [183, 184, 186, 187]. Apart from the direct effect of amphotericin B on the parasites through formation of aqueous pores in the plasma membrane, AmBisome® also appears to interact with the sterols in the membrane of macrophages, preventing Leishmania entry as well [188, 189].

A few cases of resistance have been reported but the underlying mechanisms are poorly understood [190]. Laboratory acquired resistance has been associated to ergosterol deficiency in the parasite membrane or the enzyme S-adenosyl methionine transferase, which is involved in the ergosterol synthesis [191, 192]. Changes in the plasma membrane sterol profile have also been reported in L. donovani clinical isolates, alongside with an upregulation in ATP-binding cassette transporters and the thiol metabolic pathway [193], and cross-resistance to miltefosine uptake [194]. Additionally, a large scale proteomic study has identified several proteins that were differentially expressed in amphotericin resistant parasites, including enzymes involved in metabolic pathways (glycolysis, TCA), transcription and translation, ROS scavenging and also heat shock proteins [193, 195].

**Miltefosine.** One of the biggest efforts in drug discovery against leishmaniasis was to find an efficacious oral drug. Miltefosine (Impavid®, hexadecylphosphocholine) is an alkylphosphocholine that belongs to the structural class of alkylphospholipids, in which the phosphocholine is esterified with a long-chain alcohol [196]. It was initially used as an antineoplastic, but later became the first oral antileishmanial drug to treat visceral leishmaniasis [153]. This drug has also been reported for the treatment of cutaneous leishmaniasis, however two different studies, one from northern Ethiopia and the other from Spain, showed that miltefosine is less effective in HIV co-infected patients [197-201]. The recommended therapy regimen is a single oral dose of 50mg for patients with less than 25kg body weight or a twice daily dose of 50mg over a period of 28 days for patients weighing more than 25kg [185]. This drug is highly effective with mild and temporary toxicity that includes gastrointestinal, hepatic, and renal side effects. Additionally, miltefosine is teratogenic therefore strictly forbidden in pregnant women [156, 200, 202, 203].

Miltefosine induces the parasite’s killing through an apoptosis-like process that involves mitochondria membrane depolarization due the inhibition of cytochrome C oxidase [204-207]. This drug also alters the biosynthesis of lipids, through perturbing alkyl-lipid metabolism, glycosylphosphatidilyinositol anchor biosynthesis [208], and ether-lipid remodeling through the
inhibition of the alkyl-lyso-phosphatidylcholine specific acyl coenzyme A acyltransferase [209]. Overall, it seems that the intracellular accumulation of the drug, which is mediated by two transporters, is detrimental for its microbicidal action [205]. Miltefosine is prone for resistance emergence: 1) the treatment may often be unsupervised and lack of compliance can lead to subtherapeutic doses in the patients, 2) long half-life, as it can still be detected in the blood 5 months after the treatment [197].

Miltefosine resistant parasites are easily generated in the laboratory, and due to the widespread use of miltefosine, the fear of resistance is eminent [210]. However, no human cases had been reported up until recently [211]. Initial studies demonstrated that the resistance phenotype is associated to a decreased accumulation of the drug in the cell due to an increased drug efflux and a decreased of drug uptake [210, 212-214]. Later, Clos et al. have isolated a gene from *L. infantum* that, upon over-expression, confers protection not only against miltefosine, but also against trivalent antimony [215]. In a recent study, a transcriptomic analysis of resistant versus susceptible strains to miltefosine revealed differential expression of genes related to DNA repair/replication machinery, protein translation and folding, lipid metabolism, antioxidant defense and transporters activity [216]. Miltefosine is certainly the most cost-effective option to treat leishmaniasis in the areas of antimonial resistance, however, its use as first line therapy is inevitably limited by its teratogenicity and high potential for resistance development [175].

**Paromomycin.** Paromomycin (Aminosidine) is a broad-spectrum aminoglycoside antibiotic belonging to the neomycin family that in 1960s was found to have leishmanicidal activity. It consists of three amino sugars bonded by a glycosidic linkage and a ribofuranosyl subunit in the central position [217]. It was licensed in 2007, in India, as an effective, well tolerated and affordable treatment for visceral leishmaniasis [218]. A 21-day course treatment using an intramuscular injection of 20 mg/kg of body weight per day can be effectively used [219, 220]. However, clinical trials in East Africa using 15-mg/kg regimen for 21 days, showed much lower efficacy, particularly in Sudan where the cure rate was <50%. Even the increased dose of 20 mg/kg for 21 days gave only an 85% cure rate, insufficient for consideration as a monotherapy [221]. In the case of cutaneous leishmaniasis, paromomycin has been formulated for topical treatment of CL, offering the advantage of easier administration, fewer side effects and cost-effectiveness in comparison to systemic treatment [222]. This drug is used in a methylbenzethonium chloride ointment (Leshcutan) as topical treatment [223]. Moreover, in a phase 2, placebo controlled study, carried out in Tunisia and France, the third generation aminoglycoside ointment WR279,396, a hydrophilic formulation of 15% paromomycin plus 0.5% gentamicin] was reported to be safe with a cure rate of 94% [224]. Ototoxicity and local pain upon injection are the main side effects [218] Paramomycin binds to the 30S ribosomal subunit and interferes with the initiation of protein synthesis. Paromomycin also interferes with mitochondrial membrane potential, inhibiting respiration and appears to affect plasma membrane fluidity and permeability [225]. No clinical resistance has been reported in the field, however aminoglycosides do have strong potential for resistance development, and it has been induced in vitro [226-228]. In the latter, a decrease in the uptake, coupled with drug efflux have been implicated [228, 229]. Once this drug is inexpensive it has been accepted as a first-line alternative drug in the setting of resistance to classical antileishmanial drugs. Moreover, the combination of antimonial drugs with paromomycin results in a highly efficacious regimen accompanied by a reduction of the side effects of both drugs [230, 231].

**Imidazole derivatives, Allopurinol and Sitimaquine.** The first oral compound used against leishmaniasis was ketoconazole, an imidazole derivative that inhibit ergosterol...
biosynthesis. This compound, as well as itraconazole and fluconazole, which are also imidazole derivatives, present variable success rates against different causative species of CL and in general are not very promising as monotherapies against VL [232, 233]. They may have some interest in the context of a combination therapy against VL [234].

Allopurinol, a hypoxanthine analogue and xanthine oxidase (XO) inhibitor, inhibits purine anabolism in Leishmania [235]. Similarly to the imidazole derivatives, it is not valuable as a monotherapy, but in combination it displays some success in CL but not as promising in VL [236, 237].

Sitamaquine (8-aminoquinoline) was specifically developed for VL treatment, actually in CL the results were disappointing. In high concentrations it affects parasite motility, morphology and growth, however, its mechanism of action is unclear. It seems to be inserted in biological membranes through electrostatic interactions and accumulates in acidocalcisomes [238]. More studies are still required to further understand its mode of action, adverse side-effects and resistance-associated mechanisms, so it can be properly assessed as reasonable alternative to the already established VL treatments [239].

THE PENTOSE PHOSPHATE PATHWAY OF TRYPANOSOMATIDS: DRUG TARGET CANDIDATES

As mentioned in the previous section, there is still an urgent need for novel drugs against trypanosomatids. A common aim in drug discovery is to find proteins that are essential for parasite survival and/or infectivity and absent or sufficiently different at a structural level from the host [240]. This sections addresses the efforts that have been made to explore the particularities of the pentose phosphate pathway of trypanosomatids and to identify and validate potential drug targets allowing subsequent identification of lead compounds (Table 1). Trypanosomatids have unique organelles in order to respond to the needs inherent to their specific lifestyle within their host. Among those are glycosomes, peroxisome related organelles, which contain enzymes of important metabolic pathways such as glycolysis, pentose phosphate pathway, fatty acid β-oxidation, gluconeogenesis, purine salvage and biosynthesis of pyrimidines, ether lipids biosynthesis and early parts of the mevalonate pathway [241]. The compartmentalization of different pathways in the glycosomes is an enigma, but in general is thought to prevent the accumulation of toxic intermediates as well as enable a fast metabolic adaptation to environmental changes [241, 242]. Apart from glycolysis, glucose is metabolized by the PPP (Fig. 1), whose role in protozoa and their interaction with their hosts has recently become a very attractive and viable drug target. It is a key metabolic pathway that relies upon glucose and is classically divided in two branches: an oxidative branch and a non-oxidative branch. Some of the resulting products and intermediates (ribose-5-phosphate - R5P, glyceraldehyde-3-phosphate - G3P, fructose-6-phosphate - F6P) and cofactors (NADPH) are used to synthesize nucleic acids and lipids, as well as to maintain redox homeostasis [243]. PPP does not necessarily act as a cycle, as the enzymatic reactions can be adjusted according to the cells demands [243]. In most organisms, this pathway localises to the cytosol, but in trypanosomatids it localises between the glycosomes and the cytosol [5]. The oxidative branch is considered unidirectional, and comprises of three enzymatic steps glucose-6-phosphate dehydrogenase (G6PDH), 6-phosphogluconolactonase (6PGL), and 6-phosphogluconate dehydrogenase (6PGDH) (Fig 1), for the synthesis of ribulose 5-phosphate (Ru5P), with the concomitant production of two moles of NADPH per mole of glucose 6-phosphate (G6P) consumed [244]. Until now the oxidative branch was subjected to a higher number of studies compared to the non-oxidative
PPP, however the latter deserves further investigation, since it supplies the parasites with ribose-5-phosphate (R5P), required to produce RNA and DNA, and erythrose 4-phosphate (E4P), a precursor of aromatic amino acids and vitamin B6 [245]. In the bidirectional non-oxidative branch, the Ru5P from the oxidative branch, can be converted into either R5P or xylulose 5-phosphate (X5P). Thereafter two enzymes, transketolase (TKT) and transaldolase (TAL), are responsible for the relative complex interconversion reactions between R5P/X5P and the glycolytic intermediates glyceraldehyde 3-phosphate/fructose 6-phosphate via reshuffling of monophosphate sugars (Fig 1). Through the sharing of these intermediates, TKT and TAL act as an interface between glycolysis and the PPP, allowing rapid interconversion of these important metabolites depending upon the cells requirements [245].

Oxidative branch.

**Glucose-6-phosphate dehydrogenase.** Glucose-6-phosphate dehydrogenase (G6PDH; Fig. 1) is the first enzyme of the oxidative branch of PPP, catalysing the oxidation of G6P to 6-phosphogluconolactone (6PGL) as NADP+ is reduced to NADPH (reviewed in [246]). This enzyme shares approximately 50% identity to its human homologue [247]. It localises mostly to the cytosol, although there is a small fraction compartimentalized in the glycosomes, although a typical peroxisomal targeting sequence (PTS) is absent [248, 249]. T. brucei and L. mexicana have a single copy sequence, whereas T. cruzi has five copies, two of them being pseudogenes. The crystal structure of the TcG6PDH revealing the molecular basis of substrate and cofactor recognition was recently obtained showing that differences in the cofactor binding site compared to the human homologue allowing future exploration towards the design of new competitive NADP+ inhibitors [250].

This enzyme has been genetically validated as a drug target in T. brucei, using RNAi, probably due to the depletion of nucleotide and NADPH pools [251]. Moreover, a critical role in anti-oxidant defence has been demonstrated for G6PDH in both T. cruzi and Leishmania [252, 253].

**Phosphogluconolactonase.** The second step of this pathway is catalysed by 6-phosphogluconolactonase (6PGL; Fig. 1), which hydrolysates 6-phosphogluconolactone (6PGL) into 6-phosphogluconate (6PG) [248]. In T. brucei, it localises mostly in the cytosol, but around 15% localises to the glycosomes, and a PTS-1 signal peptide can be found [249, 254]. It has only 20% identity to the human homologue and its crystal structure has been solved. Although its substrate, 6-phosphogluconolactone, can spontaneously hydrolyse, in conditions of high NADPH demand, it may accumulate especially inside the glycosomes. Due to its electrophilic character, it may irreversibly inactivate key metabolic enzymes. Therefore, 6PGL may have a detrimental role to prevent the accumulation of this PPP intermediate. However, its essentiality has not been addressed in any trypanosomatid [247].

**Phosphogluconate dehydrogenase.** The last step is catalysed by 6-phosphogluconate dehydrogenase (6PGDH; Fig. 1), which is responsible for the oxidation and decarboxylation of 6-phosphogluconate (6PG) to Ru5P, while again reducing NADP+ to NADPH [255]. It has only around 35% homology to the human counterpart, however there is a high conservation of the residues involved in the substrate and coenzyme binding, challenging the design of selective inhibitors [247, 256]. Similarly to the first two enzymes of the pathway, it localises predominantly to the cytosol, and to a less extent to the glycosomes [257]. In T. brucei, deletion of 6PGDH leads to the accumulation of 6-phosphogluconate, which inhibits phosphoglucone isomerase and consequently glycolysis. This is probably exacerbated in vivo with the increase of G6P flow through the PPP. Moreover, it remains to be investigated
whether a decrease in R5P production also plays a role in the observed defect [247]. 6PGDH essentiality has not yet been addressed in Leishmania and T. cruzi.

In summary, the oxidative branch is therefore of critical importance to these parasites, as it supplies not only Ru5P, a precursor of the non-oxidative branch, but also NADPH, an essential molecule for lipid synthesis, defence against oxidative stress by regenerating GSH, among other important cellular processes [247].

Non-oxidative branch.

Ribose 5-phosphate isomerase. Ribose-5-phosphate isomerase (RPI; Fig. 1) is responsible for the interconversion of R5P in Ru5P. There are two types of RPI enzymes, type A RPI (RPIA) is represented in all life kingdoms, contrasting with the type B (RPIB), restricted to some bacteria and protozoans [258]. An adverse phenotype was observed in E. coli [258] and also humans [259] upon RPI deficiency, suggesting a critical conserved role through evolution. Trypanosomatids possess a type B RPI, which is absent from humans that instead have a structurally unrelated type A and therefore possibly an ideal drug target [260-265]. Recombinant enzymes from L. donovani [262], L. infantum [261], L. major [261], T. brucei [263] and T. cruzi [263, 264] have been formally demonstrated to have in vitro isomerase activity by catalysing the interconversion of R5P and Ru5P. The Km values for both R5P and Ru5P were similar between Leishmania and trypanosomes enzymes, however, Kcat values are considerably higher for Leishmania, in both direct and inverse reactions [261-264]. A decrease in Km and an increase in Kcat were consistently observed for Ru5P in comparison to R5P, suggesting that the conversion of Ru5P into R5P is favoured, which might be explained by the important role of R5P as a building block for nucleic acid synthesis.

Regarding RPI localization, LmRPIB possesses a PTS-2 signal sequence (–RVALGCDHA–), which is conserved in L. infantum and L. donovani [249]. However, recent proteomic analysis of L. donovani glycosomes failed to detect RPIB in these organelles [266]. The same analysis detected HGPRT (PTS-1), aldolase (PTS-2), as well as some enzymes of PPP that are upstream RPIB, namely putative G6PD (PTS-1), putative 6PGDH (non-identified), or enzymes downstream RPIB, such as TKL (PTS-1), putative RPE (PTS-1) and putative TAL (non-identified signal peptide), or other related proteins like putative ribokinase (PTS-2 signal peptide). In both T. brucei and L. infantum RPIB shows a dual localization despite preferentially located in the cytosol compared to the glycosomes [261, 263]. The amount of RPIB that is in the cytosol or the glycosomes, as well as whether the distribution changes throughout the parasite life cycle needs further investigation.

In T. brucei, RNAi induction against RPIB led to a significant decrease of bloodstream forms proliferation in vitro but more importantly it dramatically affects the infectivity of the parasites leading to a decrease in parasitaemia and a prolonged mice survival [261]. Phenotypic reversion was achieved by complementing induced RNAi clones with an ectopic copy of the T. cruzi gene suggesting ultimately a conserved role of RPIB among different Trypanosoma species. Moreover, RPIB is as an essential gene as T. brucei is resistant to complete RPIB gene removal and mice infected with single knockout mutants showed prolonged survival upon infection [261]. RPIB gene essentiality for parasites survival was also demonstrated in L. infantum as null mutants generation was only possible when an episomal copy of RPIB gene was introduced and the latter was retained both in vitro and in vivo in the absence of drug pressure [261]. Importantly, the inability to remove the second allele of RPIB gene in sKO mutants complemented with an episomal copy of RPIB carrying a mutation that abolishes isomerase activity suggests the essentiality is due to its metabolic function [261]. This is
Ribulose 5-phosphate epimerase. Another player in the PPP non-oxidative branch is ribulose-5-phosphate epimerase (RPE; Fig. 1), which interconverts Ru5P and X5P. In *T. brucei* RPE activity was detected in procyclics, but not in parasites isolated from mice [260]. *Leishmania* and *T. cruzi* encode for two isoenzymes that differ in the presence and absence of a PTS [271, 272], and the first has recently been detected in *L. donovani* glycosomes [266]. Additionally, its activity has been detected in *L. mexicana* promastigotes and was 3 fold higher than its competitor enzyme, RPIB [273]. In *T. cruzi* despite the high sequence similarity between the two isoenzymes the cytosolic TcRPE1 shows a specific activity 80-100-fold greater than the glycosomal isoenzyme TcRPE2 [271]. Furthermore, the subcellular localization of TcRPE2 is absolutely dependent on the presence of its targeting signal, since over-expression of a truncated mutant lacking the putative glycosomal targeting signal, not only remains in the cytosol, but also negatively affects parasite growth, extending its doubling time *in vitro* [271].

RPE has not been subjected to a targeted study in any trypanosomatid to address whether it is essential for survival. However, considering that the genome-wide RNAi screening in *T. brucei* using RITseq indicated that mRNA depletion did not cause any fitness cost, and that its activity had not been detected in parasites recovered from the mice (Alsford et al, 2011; Cronin et al, 1989; Kovárová et al, 2016), this indicates that RPE may not qualify as a drug target in *T. brucei*. In *Leishmania* and *T. cruzi*, this may be different and requires further research.

Transketolase. Transketolase (TKT; Fig. 1) is an enzyme capable of catalysing two distinct reversible reactions that involve the transfer of a two carbon unit: (i) from xylulose 5-phosphate (X5P) to R5P, producing glyceraldehyde 3-phosphate (G3P) and sedoheptulose 7-phosphate (S7P), and (ii) from xylulose 5-phosphate (X5P) to erythrose 4-phosphate (E4P) to generate glyceraldehyde 3-phosphate (G3P) and fructose 6-phosphate (F6P), using thiamine diphosphate as a cofactor [274]. This enzyme may prevent the accumulation of toxic metabolites or modulate NADPH production by redirecting sugar phosphates towards the oxidative branch of the pathway [247]. In trypanosomatids, at least two residues involved in the substrate and cofactor binding are not conserved when comparing to the human counterpart [247]. TKT is expressed in all developmental stages of *T. cruzi* and its activity has been reported in several *Leishmania* species [247]. In particular, in *L. mexicana* it displays dual localization, in cytosol and glycosomes [275]. TKT activity has not been detected in *T. brucei* parasites recovered from the mice [260], very similarly to RPE. Apparently a great extent of the non-oxidative branch of PPP appears dispensable for *T. brucei* bloodstream forms contrarily to insect stage [247, 276]. This metabolic adaptation may be employed in order to channel sugar metabolism exclusively into R5P and NADPH generation, to support the high proliferation rate of the bloodstream forms [247, 276]. Since *T. cruzi* and *Leishmania* present a fully operative PPP, the role of TKT in these parasites still awaits to be unravelled.
TKT is expressed in all developmental stages of *T. cruzi* and its activity has been reported in several *Leishmania* species [247]. In particular, in *L. mexicana* it displays dual localization (cytosol and glycosomes) [275]. Since *T. cruzi* and *Leishmania* present a fully operative PPP, the role of TKT in these parasites still awaits further investigation.

**Transaldolase.** Transaldolase (TAL, Fig. 1) catalyses the reversible transfer a three-carbon fragment, dihydroxyacetone, from sedoheptulose 7-phosphate (S7P) to glyceraldehyde 3-phosphate (G3P), producing fructose 6-phosphate (F6P) and erythrose 4-phosphate (E4P) [247].

A gene annotated as TAL was cloned from *T. brucei* and the expressed protein was shown to be capable to form O8P using R5P and F6P as acceptor and donor substrates, respectively [276]. This indicates TAL as a likely source of the higher carbon content carbohydrates referred to earlier [243]. TAL activity has been detected in *Leishmania* [247] and further studies need to be undertaken to better understand its role in both *Leishmania* and *T. cruzi*.

It is also noteworthy that organisms, like yeast, have an alternative NADP-independent pathway for R5P synthesis, designated riboneogenesis. This pathway transforms glycolytic intermediates into sedoheptulose-1,7-biphosphate, by a combined action of the TKL and an aldolase (ALD). Subsequently, sedoheptulose-1,7-biphosphate, through the action of a sedoheptulose-1,7-bisphosphatase (SBP), is converted into S7P, which can in turn be a substrate of TKL to ultimately generate R5P [277]. Genes encoding a putative sedoheptulose-1, 7-bisphosphatase (SBP) are present in the genomes of both *T. brucei* and *T. cruzi*, but absent in *Leishmania* [272]. Importantly, it has not been formally demonstrated that this pathway operates in trypanosomes.

Overall the components of the non-oxidative branch of PPP are more heterogeneous, comprising members that do not possess a mammalian homologue (RPIB) and others that are developmentally regulated and specifies-specific dispensable (RPE and TKT) [247].

**INTERFERING WITH PPP: DO TRYPANOSOMATIDS COPE OR NOT?**

The oxidative branch of PPP is of great importance to *Leishmania* and trypanosomes, as it supplies NADPH, an essential molecule for lipid synthesis and anti-oxidant defence, among other important cellular processes [247]. Therefore, it is easy to imagine that once inhibiting an enzyme from the oxidative branch, the depletion of the NADPH pool will be critical for these parasites, as observed when knocking down G6PDH [251]. In the case of 6PGDH, which catalyses the third step of PPP, its deletion also leads to the accumulation of 6-phosphogluconate, which inhibits phosphoglucose isomerase and consequently glycolysis. Moreover, the subsequent depletion in R5P production may also play a role [247].

We have demonstrated that RPIB isomerase is essential for *T. brucei* and *L. infantum* [261] and therefore, it would be worth exploring the metabolic implications of its inactivation/inhibition. At this point, we can only speculate why an enzyme involved in the non-oxidative branch of the PPP, in which interconversion reactions take place, is essential. In *Leishmania*, our *in vitro* data suggest that RPIB is critical for the development of intracellular amastigotes [261]. These forms have nutritional requirements that are more complex than those of the majority of prokaryotes and fungal pathogens. Specifically, *Leishmania* is one of the very few pathogens that infect macrophages and does not escape or subvert the
endocytic pathway, oddly dwelling in a mature phagolysosome [278]. This compartment may appear very hostile! However, it may be a permissive niche regarding nutrient availability. Yet, to benefit from this large nutrient supply, pathogens must employ mechanisms to counteract the host cell microbicidal machinery, and *Leishmania* has successfully mastered it [279].

A very dramatic modification in amastigotes concerns the global downregulation of nutrient transporters as a stringent metabolic response is activated [280]. The consequent overall decrease in nutrient uptake in amastigotes as they enter this metabolic stringency may render the parasites more dependent on RPIB for R5P synthesis. However, there are alternatives sources for R5P production that bypass RPIB. Some organisms, like yeast, have an alternative NADP-independent pathway for R5P synthesis which relies on the combined action of sedoheptulose-1,7-biphosphate (SBP), TKL and aldolase. *Leishmania* in opposition to trypanosomes lacks SBP sequence and *T. brucei* does not express TKL in bloodstream forms: could this render them more dependent on RPIB? It would be very interesting to investigate it in *T. cruzi*, as it encodes the putative SBP and expresses TKL in all developmental stages [247, 272, 276].

Another possibility is that the absence/inhibition of RPIB may lead to the accumulation of Ru5P, which may modulate the activity of the upstream PPP enzymes by negative feedback. This sort of regulation is absent in several key enzymes of central carbon metabolism in trypanosomatids [281-283], thus it is unlikely to operate.

Moreover, a study has demonstrated that human RPIA modulates hepatocarcinogenesis [284], and its knockdown leads to Ru5P accumulation, which is further converted into X5P by RPE. It has been reported that X5P can activate PP2A [285], which negatively regulates ERK signalling for cell proliferation. Whether this sort of regulation takes place in trypanosomatids is unknown. However, they encode for a RPE [266, 273, 286], PP2A [287] and MAP kinases [288]. This would be particularly interesting to investigate in *Leishmania* and *T. cruzi*, as RPE is not expressed in the mammalian stage of *T. brucei*.

These different parasites undergo several developmental stages in different hosts, experiencing environmental conditions that are dramatically different during their life cycle. Therefore, when looking at different parasites or different forms of the same parasite, it is not surprising that their metabolism also shifts and adapts, and if many similar traits can be found, certainly many particularities as well. A better understanding of their metabolic pathways and their nutritional needs is critical to enlighten us on how to exploit them as new venues towards novel antimicrobial strategies.

**DEVELOPING TRYPANOCIDAL SPECIFIC INHIBITORS OF THE PPP**

Inhibitors against the trypanosomatids G6PDH and 6PGDH from the oxidative branch and RPIB from the non-oxidative branch have been described (Fig. 2). Indeed, the steroids dehydroepiandrosterone (DHEA) and epiandrosterone (EA) (Fig. 2A) were shown to inhibit the TbG6PDH in an uncompetitive fashion with *K*<sub>i</sub> values about 6-fold lower than those reported for the human enzyme and exhibit *in vitro* anti-parasitic activity against bloodstream forms *T. brucei* [251]. However, DHEA failed to decrease *T. cruzi* *in vitro* growth, but its brominated derivatives, which are more potent inhibitors, displayed EC<sub>50</sub> values in the low-mid micromolar range, similarly to benznidazole [289-291]. Interestingly, DHEA and EA did not inhibit *L.
mexicana G6PDH and had no impact on parasite in vitro growth rate of cultured promastigote parasites, meaning that the effect mediated by these drugs is specific to trypanosomes [253].

Some analogues/mimics of the high-energy intermediates (hydroxamate derivatives of D-erythronic acid, Fig. 2B) were found to be potent and selective inhibitors of Tb6PGDH [292], however, the best compounds showed no trypanocidal activity due to their poor membrane permeability. Therefore, phosphate prodrugs were developed to overcome this limitation, in particular, aryl phosphoramidate prodrugs of 2,3-O-isopropylidene-4-erythrohydroxamate (Fig. 2B) displayed both potent activity against Tb6PGDH and in vitro anti-parasitic activity [293, 294].

The analogous inhibitor of the isomerization intermediate, 4-phospho-D-erythrohydroxamic acid (4-PEH, Fig. 2C), was shown to be a good competitive inhibitor against RPIBs [264, 295], however it also strongly inhibits class A RPI enzyme found in humans [295]. 4-PEH was also shown to inhibit the TbRPIB, LrRPIB and LmRPIB, but only in the millimolar range [296], thus unlikely to show any potent anti-parasitic in vitro or in vivo activity.

Other possible future sources of parasite selective inhibitors may be those that are identified for similar enzymes activities in the glycolytic pathway, which bind similar substrates and have similar enzyme functions, such as phosphoglucose isomerase, fructose-bisphosphate aldolases and the triosephosphate isomerase. Thus, an inhibitor against these glycolytic enzymes could also be targeting enzymes of the PPP, which could be of added benefit.

CONCLUDING REMARKS

The PPP of trypanosomatids have attractive drug target candidates due to their essential role for parasites survival and/or infectivity, but also due to the lack or low homology with their human counterparts. Unfortunately, the target-based approach for drug discovery has not been as successful as initially expected over the past 10-15 years for several reasons. These usually concern poor membrane permeability or metabolic inactivation of the enzymatic inhibitors when tested against the whole parasites and or in animal model due to fast clearance. Others may include: 1) the enzyme may be abundantly expressed and/or present a fast turnover; 2) the compound can have higher affinity for other targets (off-target) that are present in the host and/or dispensable for the parasite; 3) in the case of Leishmania, the ability for stochastic DNA amplification may allow a higher expression of the molecular target; 4) protein moonlighting, in which the atypical function is the one detrimental for parasite survival. In the case of the mammalian stage of Leishmania, a drug has to cross the host cell, the phagolysosome and the parasite membranes, and if the enzyme is inside an organelle, another membrane has to be crossed. The fact that some of the targets of the PPP present dual localization, as they localize mainly to the cytosol, but also in part to the glycosomes raises particular concerns in terms of redundancy and/or accessibility. Indeed, it would be important to address whether the localization in the glycosomes is essential for the parasite, as its activity inside the organelle may be harder to target. Another critical issue awaiting further investigation is the enzymes druggability, what happens when the enzyme is inhibited and the substrate concentration increases, a competitive inhibitor would simply be outcompeted by the higher substrate concentration, alleviating the blockage and thus allowing the parasite to survive and possibly evolve resistance.

ACKNOWLEDGEMENTS
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TABLES & FIGURES
Figure 1. Pentose phosphate pathway in trypanosomatids. The enzymes of the oxidative and non-oxidative branch are depicted in dark and light green, respectively: G6PDH, glucose-6-phosphate dehydrogenase; 6PGL, 6-phosphogluconolactonase; 6PGDH, 6-phosphogluconate dehydrogenase; RPIB, ribose-5-phosphate isomerase B; RPE, ribulose-5-phosphate epimerase; TKT, transketolase; TAL, transaldolase. Enzymes from glycolysis (ALD, aldolase) and riboneogenesis (SBP, sedoheptulose biphosphatase) are represented in red and purple, respectively. The oxidative PPP, non-oxidative PPP and riboneogenesis metabolic flow are depicted with dark green, light green and purple lines. Metabolites in red are generated in glycolysis. 6PG, 6-phosphogluconate; 6PGL, 6-phosphogluconolactone; DHAP, dihydroxyacetone phosphate; E4P, erythrose-4-phosphate; F6P, fructose-6-phosphate; G3P, glyceraldehyde-3-phosphate; G6P, glucose-6-phosphate; R5P, ribose-5-phosphate; Ru5P, ribulose-5-phosphate; S7P, sedoheptulose-7-phosphate; S1,7BP, sedoheptulose-1,7-biphosphate; X5P, xylulose-5-phosphate. The dashed purple line highlights riboneogenesis, which has not been formally demonstrated to operate in trypanosomes. The circles next to some enzymes indicate genetic validation by RNAi and/or gene knockdown has been performed in at least one of the three parasites (red), dual localisation between the cytosol and the glycosomes (yellow), expressed only in the insect-stage of T. brucei but not in bloodstream forms (light blue) or not encoded by Leishmania parasites (dark blue). Modified and updated from [243].
Figure 2. Available inhibitors of enzymes from the pentose phosphate pathway in trypanosomatids. A-B) On the left are represented the PPP enzymes and the reactions they catalyse and on the right the available inhibitors or classes of inhibitors. The enzymes of the oxidative and non-oxidative branch are depicted in dark and light green, respectively: G6PDH, glucose-6-phosphate dehydrogenase; 6PGDH, 6-phosphogluconate dehydrogenase; RPIB, ribose-5-phosphate isomerase B. Metabolites: 6PG, 6-phosphogluconate; 6PGL, 6-phosphogluconalactone; G6P, glucose-6-phosphate; R5P, ribose-5-phosphate; Ru5P, ribulose-5-phosphate. Inhibitors: DHEA, dehydroepiandrosterone; EA, epiandrosterone; 4PEH, 4-phospho-D-erythronate. For more detailed information [251, 264, 289-296].
### Table 1. Essentiality data and crystal structure availability for PPP enzymes in T. brucei, T. cruzi and Leishmania spp.

<table>
<thead>
<tr>
<th>Protein</th>
<th>Essentiality data</th>
<th>Crystal structure</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T. brucei</td>
<td>T. cruzi</td>
</tr>
<tr>
<td>Phosphogluconolactonase</td>
<td>Not essential [297]</td>
<td>-</td>
</tr>
<tr>
<td>Phosphogluconate dehydrogenase</td>
<td>Essential [301]</td>
<td>Not essential [297]</td>
</tr>
<tr>
<td>Ribose 5-phosphate isomerase</td>
<td>Growth defect [263, 297]</td>
<td>-</td>
</tr>
<tr>
<td>Ribulose 5-phosphate epimerase</td>
<td>Not essential [297]</td>
<td>-</td>
</tr>
<tr>
<td>Transketolase</td>
<td>Not essential [303]</td>
<td>-</td>
</tr>
<tr>
<td>Transaldolase</td>
<td>Not essential [276]</td>
<td>-</td>
</tr>
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* Blood stream forms. The references in brackets correspond to the cited in the text of the manuscript. - no data

### Graphical Abstract