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Diminazene resistance in Trypanosoma congolense is not caused by reduced transport capacity but associated with reduced mitochondrial membrane potential

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Abstract

Trypanosoma congolense is a principal agent causing livestock trypanosomiasis in Africa, costing developing economies billions of dollars and undermining food security. Only the diamidine diminazene and the phenanthridine isometamidium are regularly used, and resistance is widespread but poorly understood. We induced stable diminazene resistance in T. congolense strain IL3000 in vitro. There was no cross-resistance with the phenanthridine drugs, melaminophenyl arsenicals, oxaborole trypanocides, or with diamidine trypanocides, except the close analogs DB829 and DB75. Fluorescence microscopy showed that accumulation of DB75 was inhibited by folate. Uptake of [3H]-diminazene was slow with low affinity and partly but reciprocally inhibited by folate and by competing diamidines. Expression of T. congolense folate transporters in diminazene-resistant Trypanosoma brucei brucei significantly sensitized the cells to diminazene and DB829, but not to oxaborole AN7973. However, [3H]-diminazene transport studies, whole-genome sequencing, and RNA-seq found no major changes in diminazene uptake, folate transporter sequence, or expression. Instead, all resistant clones displayed a moderate reduction in the mitochondrial membrane potential \( \Psi_m \). We conclude that diminazene uptake in T. congolense proceed via multiple low affinity mechanisms including folate transporters; while resistance is associated with a reduction in \( \Psi_m \) it is unclear whether this is the primary cause of the resistance.

Keywords
diminazene aceturate, Isometamidium, nagana, oxaborole, resistance, Trypanosoma congolense
Animal African Trypanosomiasis (AAT), also called nagana, is an infection of livestock and wild animal species with African trypanosomes, principally *Trypanosoma congolense*, *Trypanosoma brucei* spp, *Trypanosoma vivax* and *Trypanosoma suis*, which is spread by the bite of infected tsetse flies. It has a very severe impact on livestock farming in sub-Saharan Africa, costing billions of dollars in lost agricultural production. AAT therefore affects issues such as food security, diet, and rural development, and the livelihood of rural African communities in endemic areas (Giordani et al., 2016). Although Human African Trypanosomiasis (HAT), or sleeping sickness, is in rapid decline in most of Africa due to active case finding and rapid treatment (Kennedy, 2019; Lehane et al., 2016), AAT remains a very serious problem throughout the continent. This is due in no small measure because just two old drugs have been the mainstay of AAT treatment for decades: diminazene aceturate (DA) and isometamidium chloride (ISM). Resistance to each has been reported from most countries in which the disease is prevalent (Giordani et al., 2016), and no new drugs have been introduced for over half a century. Yet, while there are many studies documenting treatment failures in the field, relatively little progress has been made in understanding the mechanisms of resistance in the clinically most relevant animal trypanosome species, *T. congolense* and *T. vivax*.

By contrast, much progress has been made in understanding resistance to the suite of drugs used to treat the various manifestations of HAT, caused by the *T. brucei* subspecies *T. b. gambiense* (in West and Central Africa) and *T. b. rhodesiense* (in East and Southern Africa). Most of the resistance mechanisms uncovered for HAT drugs involve aspects of drug uptake by the parasite. For instance, pentamidine (used for early stage gambiense HAT) and melarsoprol (now mostly used for late stage rhodesiense HAT) are internalized by two transport proteins in the trypanosome’s plasma membrane. The first of these is an aminopurine transporter called P2 (Carter & Fairlamb, 1993) encoded by the gene *TbAT1* (Mäser et al., 1999), the deletion of which has been shown to cause a loss of sensitivity to these drugs (Matovu et al., 2003). Subsequently, it was shown that a second and more important determinant for pentamidine-melarsoprol cross-resistance is the High Affinity Pentamidine Transporter (HAPT1) (Bridges et al., 2007; De Koning & Jarvis, 2001), which was more recently shown to be the aquaglyceroporin TbAQP2 (Baker et al., 2012; Munday et al., 2013). Rearrangements in the TbAQP2 locus were clearly linked to treatment failures, particularly with melarsoprol (De Koning, 2020; Graf et al., 2013). Similarly, resistance to ef-lornithine, used against late-stage gambiense HAT, is associated with the loss of an amino acid transporter, TbAAT6 (Alsford et al., 2012; Vincent et al., 2010), and resistance to suramin, the first-line drug against early-stage rhodesiense HAT, is the result of mutations that interfere with its uptake by receptor-mediated endocytosis (Alsford et al., 2012; Zoltner et al., 2016, 2020).

In *T. congolense*, resistance to isometamidium has also been attributed to reduced accumulation of the drug (Sutherland & Holmes, 1993; Sutherland et al., 1991, 1992; Tihon et al., 2017), possibly in part linked to reduced mitochondrial membrane potential (Eze et al., 2016; Wilkes et al., 1997), which diminishes sequestration of cationic drugs into the mitochondrion (Alkhaldi et al., 2016; Lanteri et al., 2008; Stewart et al., 2005)—the likely site of drug action. For DA, it has been shown in the *T. b. brucei* model that it is almost exclusively taken up by the P2/TbAT1 transporter (De Koning et al., 2004) as it is a very poor substrate for the HAPT1/TbAQP2 transporter (Munday et al., 2014; Teké et al., 2011). It was subsequently proposed that the same mechanism would apply in *T. congolense*, with an adenosine transporter designated TcoAT1 mediating DA uptake, and DA resistance being the result of specific single-nucleotide polymorphisms (SNPs) that could be used to screen for resistance (Delespaux et al., 2006; Delespaux & De Koning, 2013; Vitouley et al., 2011). However, expression of the putative TcoAT1 in a multi-drug resistant *T. b. brucei* clone, B48, showed that this gene encoded a P1-type broad specificity purine nucleoside transporter with no ability to transport diminazene (Munday et al., 2013), and genomic analysis has shown that there is no direct ortholog of the P2/TbAT1 transporter in the genome of *T. congolense*. Regrettably, this has left us still without any insights into the urgent problem of DA resistance in animal African trypanosomiasis.

One urgent question beyond that of the resistance mechanism is that of cross-resistance with isometamidium, which has been reported repeatedly from the field (Mamoudou et al., 2008; Sinyangwe et al., 2004), but inconsistently, as others report no cross-resistance and it used to be a rare occurrence (Giordani et al., 2016; Joshua et al., 1995; Sow et al., 2012). Clearly, it is possible that trypanosomes may have developed resistance to both drugs independently over the decades and are now resistant to both by separate mechanisms. Thus, it is currently unclear whether other diamidine drugs should be considered for novel drug development for AAT because of the risk of cross-resistance with DA. This adds to the many reasons why it is important to understand the mechanism of DA resistance in animal-infective trypanosomes, alongside the need to identify screening tools for AAT drug resistance to assess the real levels of resistance in the field.

Here, several of these issues are addressed in an attempt to develop a hypothesis for DA resistance in *T. congolense*, and we report that DA resistance is not linked to reduced drug accumulation in *T. congolense*, nor necessarily to cross-resistance with isometamidium. However, we present evidence of a lower mitochondrial membrane potential in the resistant clones, which is stable and somehow linked to DA resistance, although it is more likely to be the result of the adaptation to DA than the principal cause of the resistance in itself, as the degree of the mitochondrial membrane depolarization is limited, and the cells were not cross-resistant to most cationic drugs. Moreover, mitochondrial accumulation of the close DA analog DB75 did not appear to be diminished in the diminazene-resistant cells. Genomic analysis identified missense mutations in two vacuolar-type Ca\(^{2+}\)-ATPases that may have regulatory functions in mitochondrial metabolism. In sharp contrast to *T. brucei* spp, uptake of \(^{[3H]}\)-DA by *T. congolense* was low affinity and inefficient and involves, at least in part, folic acid transporters.
RESULTS

2.1 | In vitro induction of diminazene aceturate resistance in *T. congolense*

Four cultures of *T. congolense* TcoIL3000 were grown and passaged in parallel; a control without added drug and three separate, independent cultures in the presence of DA for adaptation to increasing concentrations, from which ultimately six clonal lines were obtained.

Resistance was very slow to develop, as documented in Figure S1. At the end of the adaptation, after 227 passages, the control culture tolerated 50 nM DA, whereas all the adapted cultures tolerated a maximum level of 800 nM DA, in what appeared to be at least two steps, with an initial plateau at 550 nM, where the strains were assessed for their level of DA resistance and cross-resistance pattern. At the 550 nM point, final clones were obtained from each of the parallel cultures, by limiting dilution. In this study we used one clone obtained from original line 4 (4C2), and two clones each from lines 5 (5C1, 5C2) and 6 (6C1, 6C3).

At the point of adaptation to 550 nM, the level of DA resistance and the extent of any cross-resistance with the diamidine drugs pentamidine and DB829, and with oxaboroles AN7973 and SCYX7158, were tested using an assay based on the viability dye Alamar blue (resazurin), in which live trypanosomes (but not dead ones) reduce blue and non-fluorescent resazurin to pink, fluorescent resorufin. This assay yields a fluorescent output that is proportional to the number of cells in the well (Gould et al., 2008). The results, summarized in Figure 1a, show that all the DA-Res strains display highly similar drug sensitivity patterns. Resistance to DA, as measured in this assay, was ~2.3–3-fold, and highly significant (Figure 1a for fold differences and p values, Table 1 for EC50 values). No cross-resistance with pentamidine was observed, but the level of resistance to DB829, a structurally closer analog of DA, mirrored the resistance profile of DA very closely, demonstrating that cross-resistance to diamidines is not inevitable, but observed only with close structural analogs. Importantly, there was no cross-resistance with isometamidium, nor with either of the oxaboroles tested, AN7973 and SCYX7158.

To further investigate the limits of cross-resistance with other diamidines, a series of bis-benzofuran analogs (Bakunova et al., 2007) were tested (Table 1). These analogs were chosen to explore the effect of variable linker length between the benzofuramidine end groups, varying inter-amidine distance as well as the flexibility of the molecule. All the benzofuramidines displayed low micromolar activity (Table 1), and there was no cross-resistance with DA (Figure 1b). These results clearly show that the cross-resistance phenotype is
|                | EC<sub>50</sub> (Average ± SEM, µM) | IL3000          | 4C2     | 5C1     | 5C2     | 6C1     | 6C3     | Avg of DA-Res |
|----------------|----------------------------------|-----------------|---------|---------|---------|---------|---------|----------------|
| Diminazene     |                                  | 0.27 ± 0.03     | 0.65 ± 0.09 | 0.81 ± 0.09 | 0.79 ± 0.10 | 0.69 ± 0.09 | 0.63 ± 0.08 | 0.72 ± 0.04 |
| Pentamidine    |                                  | 1.01 ± 0.04     | 0.95 ± 0.06 | 0.94 ± 0.04 | 0.83 ± 0.04 | 1.01 ± 0.03 | 0.96 ± 0.03 | 0.94 ± 0.03 |
| DB829          |                                  | 1.07 ± 0.04     | 2.19 ± 0.16 | 3.27 ± 0.40 | 2.44 ± 0.27 | 2.60 ± 0.26 | 2.47 ± 0.20 | 2.59 ± 0.18 |
| Isometamidium  |                                  | 0.0098 ± 0.0007 | 0.0085 ± 0.0009 | 0.0074 ± 0.0014 | 0.0092 ± 0.0006 | 0.0068 ± 0.0008 | 0.0062 ± 0.0008 | 0.0076 ± 0.0005 |
| AN7973         |                                  | 0.12 ± 0.01     | 0.13 ± 0.02 | 0.14 ± 0.02 | 0.13 ± 0.02 | 0.13 ± 0.02 | 0.13 ± 0.02 | 0.13 ± 0.003 |
| SCYX-7158      |                                  | 0.52 ± 0.08     | 0.62 ± 0.13 | 0.53 ± 0.10 | 0.54 ± 0.12 | 0.57 ± 0.11 | 0.59 ± 0.10 | 0.57 ± 0.02 |
| Diminazene     |                                  | 0.33 ± 0.06     | 0.88 ± 0.03 | 0.80 ± 0.17 | 0.94 ± 0.06 | 0.95 ± 0.08 | 0.80 ± 0.12 | 0.88 ± 0.03 |
| Pentamidine    |                                  | 1.34 ± 0.21     | 1.23 ± 0.13 | 0.79 ± 0.06 | 1.05 ± 0.18 | 1.48 ± 0.51 | 1.32 ± 0.25 | 1.23 ± 0.13 |
| 55MB032        |                                  | 1.79 ± 0.13     | 2.22 ± 0.05 | 1.49 ± 0.06 | 1.44 ± 0.04 | 2.00 ± 0.11 | 2.15 ± 0.11 | 1.86 ± 0.17 |
| 45MB078        |                                  | 1.45 ± 0.06     | 1.99 ± 0.03 | 1.13 ± 0.05 | 1.16 ± 0.01 | 1.55 ± 0.27 | 1.81 ± 0.09 | 1.53 ± 0.17 |

(Continues)
very narrow and that many diamidines are still potentially useful for the treatment of DA-resistant AAT.

The DA-resistance of the five DA-resistant clones was then increased via further in vitro exposure until they could tolerate growth at 800 nM DA, and attempts to raise the resistance level further were not immediately successful; all subsequent experiments were done with the clones adapted to 800 nM DA. The in vitro growth rates of these final strains consistently appeared to be slightly slower than that of the drug-sensitive parental strain (Figure 2), although this reached statistical significance only for the 48 hr and 64 hr points of clones 5C2 and 6C1 ($p < .05$, unpaired student’s $t$-test). The EC$_{50}$ values, at ~2.2 nM, were similar for all clones but significantly higher than when they were measured before the cloning, at 550 nM DA medium concentration. This level of resistance, approximately ninefold compared with IL3000 WT (0.244 ± 0.004 nM in parallel experiment, $n = 4$, $p < .00001$, t-test, for all DA-RES strains versus IL3000 WT), was stable for at least 3 months of in vitro culturing in the absence of drug pressure (Figure 3a). Cross-resistance to DB829 was again observed in all clones (ca. fivefold), whereas sensitivity to pentamidine was identical in all clones including wildtype IL3000. Interestingly, sensitivity to AN7973 was slightly but significantly higher in most DA-Res clones (Figure 3d). Cross-resistance with phenanthridines isometamidium, and ethidium bromide, as well as with the melaminophenyl arsenical cymelarsan, was also tested for selected clonal lines. A slight loss of sensitivity to phenanthridines was consistently observed and reached significant (but lesser than twofold) levels in some clones. No change in cymelarsan sensitivity was observed (Figure 3e).
In order to confirm that the in vitro pattern of cross-resistance held up for in vivo infections, including for a field-derived DA-resistant strain, we infected groups of six mice that were treated i.p. with either DB829 (Figure 4a) or AN7973 (Figure 4b), using single doses of 5, 10, or 20 mg/kg body weight (two separate independent experiments with each drug, each experiment n = 6). For the resistant field isolate, we used strain KONT2/151 (Mamoudou et al., 2006). IL3000 was more virulent than KONT2/151, as is often observed with cultured trypanosome strains, and was sensitive to both DB829 and AN7973. DB829 delayed parasitemia and death with a dose of 10 mg/kg and was
Curative at 15 and 20 mg/kg, whereas AN7973 delayed parasitemia at 5 mg/kg and cured with 10 or 20 mg/kg. The DA-resistant field isolate was equally sensitive to the oxaborole but resistant to DB829, as even the highest dose, 20 mg/kg, merely delayed the onset of parasitemia.

### 2.3 Screen for potential diminazene transport inhibitors by DB75 fluorescence microscopy

In *T. brucei*, fluorescent diamidines such as DB75, a close structural analog of DA and DB829, have previously been used to monitor uptake and cellular distribution of such compounds (Stewart et al., 2005). In those experiments, the rate at which the kinetoplast (first) and the nucleus (second) became visible in the blue wavelengths ($\lambda = 405/435$ nm for excitation and emission, respectively) was much delayed in resistant strains, and in the presence of transport inhibitors (Stewart et al., 2005). Here, the technique was used to screen for potential transport substrates that competitively inhibit the uptake of the DB75 in animal trypanosome species. The utility of the technique was first confirmed in ex vivo *T. brucei* in mouse blood. Murine blood cells did not appear to significantly take up 10 µM DB75, ensuring a low background, but in *T. brucei*, parasite...
outlines would become visible after 4 min, followed by kinetoplasts at 5–6 min and finally nuclei were faintly stained by 8 min and bright by 15 min. These stages were all observed and scored in the presence of potential inhibitors. As expected, based on current *T. brucei* models (De Koning et al., 2004; Teka et al., 2011), the polyamines putrescine, spermine and spermidine (500 µM) had no discernible effect on DB75 staining, although the polyamine transporters of *Leishmania* spp and *T. cruzi* have been implicated in possible pentamidine uptake (Basselin et al., 1996; Diaz et al., 2014; Hasne & Ullman, 2005). By contrast, DA and the shorter pentamidine analog propamidine both dose-dependently delayed DB75 fluorescence (Table S1); both are known substrates of the TbAT1/P2 aminopurine transporter (De Koning & Jarvis, 1999, 2001) that is the sole uptake mechanism for DB75 in *T. brucei* (Ward et al., 2011).

Because of the involvement of the TbAT1/P2 purine transporter in DA uptake by *T. b. brucei*, we tested a range of purine and pyrimidine nucleosides and nucleobases for delaying DB75 cell entry in *T. congolense*, using the IL3000 WT cell line. Adenine and adenosine did not appear to slow DB75 fluorescence up to 1 mM, but were toxic to the cells at the concentrations of 200 µM and above. Other purines (inosine, guanosine, hypoxanthine, guanine, xanthine) or pyrimidines (uridine, cytidine, thymidine, uracil, cytosine) likewise failed to inhibit DB75 uptake; moreover, polyamines and amino acids (500 µM) were similarly ineffective in this assay (Table S1). Glycerol was also tested at up to 20 mM because some diamidines including pentamidine are similarly ineffective in this assay (Table S1). Glyceral was also tested at up to 20 mM because some diamidines including pentamidine are taken up by an aquaglyceroporin in *T. brucei* (Alghamdi et al., 2020; Baker et al., 2012), but the data indicated that glyceral also does not compete for the *T. congolense* DB75 transporter. However, the diamidines DA and, particularly, propamidine and pentamidine dose-dependently (100–500 µM) delayed DB75 fluorescence, and so did folic acid, from 200 to 500 µM (Figure 5). *T. vivax* was also assayed by this method and similar results were found, with no effect from a series of purines and pyrimidines, glycerol, and the cationic amino acids lysine and arginine, but dose-dependent inhibition by pentamidine and DA. For this parasite, there appeared to be some minor fluorescence delay in the presence of 500 µM folic acid or biopterin, but not at lower concentrations, and possibly in the presence of the polyamines spermidine and putrescine (Table S1).

### 2.4 Uptake of [³H]-Diminazene by DA-sensitive and -resistant *T. congolense* clones

In *T. brucei*, resistance to DA is linked to loss of the TbAT1/P2 transporter (De Koning et al., 2004; Graf et al., 2013; Matovu et al., 2003). While *T. congolense* does not have an equivalent transporter (Munday, Tagoe, et al., 2015), drug resistance in trypanosomatids has very often been associated with the loss of transporters at the plasma membrane (De Koning, 2020; Landfear, 2008; Munday, Settimo, et al., 2015) and we thus investigated whether uptake of [³H]-DA was altered in the resistant clones.

Initial experiments with various concentrations of [³H]-DA, trying to standardize linear uptake conditions, were unsuccessful. Uptake was very slow, the label displayed high background from binding to the outside of the cell and a true linear phase of uptake could not be established with the requisite level of confidence and reproducibility. Therefore, accumulation of 0.1 µM [³H]-DA was monitored over 30 min in IL3000 and two of the derived DA-resistant clones, 4C2 and 6C3. This showed that even over half an hour there was no clear difference in the rate of accumulation of [³H]-DA in any of the three cell lines and that resistance is therefore very unlikely to be the result of induced changes in drug uptake rate (Figure 6a). Indeed, monitoring of DB75 uptake, quantified as incubation time required for staining of nucleus and kinetoplast to become detectable by fluorescence.
FIGURE 6  Investigation into the uptake of diminazene by T. congolense. (a) Uptake of 0.1 µM [3H]-DA by IL3000 and two derived DA-adapted clones. Uptake of DA was monitored for 30 min in in vitro grown cells of T. congolense IL3000, 4C2 and 6C3. There were no statistically significant differences between the three strains at any of the timepoints. (b) Efflux of [3H]-DA from T. congolense IL3000 and from clone 4C2, after loading with 0.1 µM [3H]-DA for 30 min, followed by a wash into fresh assay buffer. Samples were taken at the indicated time points after resuspension into fresh buffer. The data presented are the average of each time point of four independent experiments, each performed in triplicate. The two slopes, calculated by linear regression, were not significantly different from each other (p = .84, F-test), nor significantly different from zero (p = .14 and .46 for IL3000 and 4C2, respectively). (c) IL3000 parasites were isolated from mice and incubated with 0.1 µM [3H]-DA with and without 250 µM unlabeled DA. (d) IL3000 parasites from culture were incubated with 0.1 µM [3H]-DA label in the presence or absence of 100 µM DA, 100 µM pentamidine (PMD), 100 µM DB829 or 1 mM folic acid. (e) Uptake of 0.1 µM [3H]-DA by bloodstream forms of T. b. brucei clone B48 transfected with TbAT1 without inhibitor or in the presence of 250 µM DA or 1 mM adenosine (Ado). (f) Uptake of 0.1 µM [3H]-DA by T. congolense IL3000 from culture, in the presence and absence of 1 mM folic acid, expressed as % of uptake in the absence of folic acid at 30 min. The average and SEM of six independent experiments, each performed in triplicate, is shown. (g) T. congolense IL3000 from culture were incubated with 0.1 µM [3H]-pentamidine by T. congolense IL3000 from culture in the presence or absence of 1 mM pentamidine (PMD) or diminazene aceturate (DA). One representative experiment in triplicate is shown. (h) Uptake of 0.1 µM [3H]-folinic acid by T. congolense IL3000 from culture, and the effects of 1 mM folic acid or 1 mM DA. One representative experiment in triplicate is shown. All experiments were performed in triplicate and the average and SEM are shown. When error bars are not shown, they fall inside the symbol. Statistical significance was calculated using Student’s unpaired t-test: *p < .05; **p < .01; ***p < .001
microscopy, also showed no difference between IL3000 and DA-resistant clone 6C3, at either 1 or 10 µM DB75; lower concentrations did not produce detectable staining in either cell line.

Nor were there significant efflux from the cells, after preincubation with [3H]-DA for 30 min followed by incubation of up to 30 min in fresh medium without label or diminazene. Although the slope of both the IL3000 WT and the 4C2 clone trended downwards, in each of four such experiments the slopes were not significantly different from zero, nor different from each other (F-test, p > .05). Figure 6b shows the average of four experiments, which still shows a non-significant downward trend and no difference between the slope of the sensitive and resistant lines. In order to better understand why DA-resistance in *T. congolense*, in contrast to *T. b. brucei*, is not associated with loss of import or gain of efflux function, the mechanism by which DA is internalized by *T. congolense* was further investigated.

Uptake of 0.1 µM [3H]-DA was poorly saturable by even a large excess of unlabeled DA (Figure 6c), indicating that DA enters *T. congolense* by a low affinity and/or non-saturable process. Indeed, out of several potential inhibitors or substrates of the DA uptake, the process was consistently (but always partially) inhibited by 100 µM pentamidine, 1 mM folate, and 100 µM DB829 (Figure 6d). This is in sharp contrast to DA uptake in *T. brucei* which is completely saturable and 100% inhibited by a competitive substrate of the TbAT1 transporter (Figure 6e). The inhibition with folate was confirmed and quantified in a series of experiments with added time points, and found to be highly significant (Figure 6f). Moreover, in a separate experiment it was tested whether the inhibition by folate and pentamidine was additive and the data indicated that this was the case (Figure 6g), suggesting that in *T. congolense* DA is taken up by at least two separate low affinity transporters, one sensitive to folate and another that is sensitive to pentamidine. However, it must be noted that the inhibition by both compounds was modest in this experiment and that the joint administration still did not inhibit [3H]-DA uptake by even 50%. The complete lack of cross-resistance with pentamidine, despite high concentrations of pentamidine partially inhibiting DA uptake, is consistent with DA resistance not being the result of changes in its uptake rate. Indeed, the overlap in transport mechanisms of DA and pentamidine was even more clear from the inhibition of [3H]-pentamidine uptake by DA in wild-type IL3000 (Figure 6h). To further investigate whether (a) folate transporter(s) might contribute to DA uptake in *T. congolense*, we measured the uptake of [3H]-folate and found this to be partially inhibited by 1 mM DA (Figure 6i).

### 2.5 *T. congolense* folate transporters are engaged in diminazene uptake

Three genes encoding putative folate transporters of *T. congolense* were identified via homology searches in TriTrypDB in the original *T. congolense* IL3000 genome and amplified by PCR with Phusion proof-reading polymerase from IL3000 and from the DA-resistant isolates KONT2/133 and KONT2/151 (sequences, GeneIDs, and GenBank accession numbers in Table 2); all three strains are Savannah-type *T. congolense*. The genes were cloned into pGEM-T Easy for amplification in *E. coli* and for each gene at least four independent clones were sequenced. The gene sequences from the two KONT strains, both from Cameroon, were almost identical to each other, but displayed some SNPs relative to IL3000 (Table 3). For folate transporter 1 (TooFT1, 592 aa) there were two identical, conservative SNPs of IL3000 alanine to KONT serine (positions 49 and 331). For FT2 (640 aa), there were eight shared non-synonymous SNPs in the two KONT strains and one unique one in each, and for FT3, both KONT strains carried the same 15 amino acid changes relative to IL3000 (Fig. S2); a table with the percentage identity and similarity between these genes is included in Table S2. The genes were cloned into the pHD1336 expression vector and transfected into *T. b. brucei* strain B48. This clonal line lacks both the TbAT1/P2 transporter and a wild-type TbaQP2 gene, and is thus highly resistant to DA, the furamidines, pentamidine, and melaminophenyl arsenicals (Bridges et al., 2007; Munday et al., 2014; Ward et al., 2011). As the sensitivity to these drugs is completely restored upon re-introduction of the transporters (Alghamdi et al., 2020; Munday et al., 2014; Munday, Tagoe, et al., 2015) this is an ideal system to test potential diamidine transporters from other kinetoplastids. Clonal lines were obtained by limiting dilution; the correct integration of the expression constructs was verified by PCR.

Expression of the *T. congolense* folate transporters in *T. b. brucei* B48 confirmed that these transporters have some capacity to take up diamidines, but not the oxaborole AN7973 (Figure 7). Only FT2 of IL3000 did not significantly sensititize to any of the drugs tested, although its diamidine EC<sub>50</sub> values also trended to be lower than the control (B48 transfected with the "empty vector"). Sensitization to DA and DB829 was almost identical, but only the FT2 transporters of the KONT strains significantly sensitized to pentamidine. In all cases, the sensitization was modest, reaching ~50% with DA and DB829, and <40% for pentamidine, which is consistent with the transport experiments with TcL3000, in that the transporters appear to display significant and yet very limited diamidine uptake capacity. For comparison, the expression of TbAT1 in B48 cells, using the same expression vector and assays, resulted in ~200-fold sensitization to DA.

### Table 2 GenBank Accession Numbers of the *T. congolense* folate transporter genes

| FT1 | FT2 | FT3 |
|-----|-----|-----|
| KONT133 | MT782654 | MT782655 | MT782657 |
| KONT151 | MT782654 | MT782656 | MT782657 |

### Table 3 Number of non-homologous SNPs in *T. congolense* folate transporter genes, relative to IL3000

| FT1 | FT2 | FT3 |
|-----|-----|-----|
| KONT133 | 2 | 9 | 15 |
| KONT151 | 2 | 8 | 15 |
and pentamidine instead of twofold (Munday, Tagoe, et al., 2015). The data presented in Figure 7 do not suggest that the two KONT strains are DA resistant because of changes in DA uptake—or at least not by their folate transporters. With respect to DA and DB829, the IL3000 and KONT FT1 and FT3 performed identically, but the KONT FT2 clearly sensitized stronger than the IL3000 FT2.

### 2.6 The mitochondrial membrane potential is diminished in DA-resistant *T. congolense* clones

As DA resistance is apparently not linked to changes in DA uptake, it was hypothesized that an intracellular rather than a cell surface change must be responsible for the resistance phenotype. Moreover, DA is a DNA minor groove binder, and resistance arising due to mutations in a target protein is therefore not expected. Our observations with DB75 fluorescence microscopy showed that such drugs accumulate first in the kinetoplast, as reported for other species (Basselin et al., 2002; Stewart et al., 2005). This requires rapid uptake into the mitochondrion, which is driven by the mitochondrial membrane potential $\Psi_m$ as DA, DB75, and other diamidines are dications. We therefore investigated whether $\Psi_m$ was altered in the resistant cell lines. Fluorescence microscopy after DAPI staining confirmed that the kinetoplast was present and did not appear altered or damaged in the DA-resistant clones (Fig. S3).

It is already well established that (di)cationic trypanocides that accumulate in the trypanosome’s mitochondrion cause a reduction in $\Psi_m$, both through the very fact that they are cations, and through disruption of mitochondrial processes (e.g. Alkhaldi et al., 2016; Fueyo Gonzalez et al., 2017; Ibrahim et al., 2011; Lanteri et al., 2008). For instance, pentamidine has been shown to act as a cationic uncoupler of oxidative phosphorylation in isolated rat liver mitochondria (Moreno, 1996). However, the more important question here was whether, as (part of) the adaptation to DA, the cells had
permanently lowered their $\Psi_m$, as previously reported for isometamidium (Eze et al., 2016; Wilkes et al., 1997). The DA-Res parasites did present flow cytometry profiles that had shifted significantly to lower fluorescence and were broader than the control IL3000 cells (Figure 8a). For quantification, the peak of the IL3000 profile was set at 80 Arbitrary Units (AU). Taking the percentage of cells with fluorescence $>80$ AU as a measure for $\Psi_m$ (Fueyo Gonzalez et al., 2017; Ibrahim et al., 2011), a highly significant decrease was found in all four DA-Res cell lines tested, although the extent of the $\Psi_m$ decrease was not identical in all cell lines (Figure 8b).

### 2.7 Diminazene causes mitochondrial lesions

In order to further assess the T. congolense mitochondrion as the target for diminazene, and as a potential contributor to diminazene resistance, culture of IL3000 and of 6C3 were grown for 4h or 8 hr in the presence and absence of 1.5 $\mu$M DA. The cells were harvested at the indicated times and processed for transmission electron microscopy (TEM). DA caused substantial lesions in mitochondria of IL3000 at 4 hr (compare Figure 9a,b) and 8 hr (compare Figure 9e,f); large parts of the mitochondria of the DA-exposed cells became electrolucent (white arrowhead in Figure 9b), with a vacuolar appearance (alterations indicated by black arrows), with some with electrondense material within the mitochondrial lumen (arrowhead in Figure 9f). By contrast, the mitochondria in the control cells remained quite electron dense, and clearly membrane-delineated. For the 6C3 cells, the mitochondria looked unchanged after 4h of DA-exposure, with very intact, very electron dense kinetoplasts (where captured in the frame) and clear membranes (Figure 9c,d). After 8 hr of continued exposure to DA, intact mitochondrial structures could still be seen in most cells, but lesions had started to appear in some of them (Figure 9g,H; arrows), and a less structured kinetoplast (white arrow).

### 2.8 Genomic analysis of DA resistance

DA-resistant T. congolense were analyzed for SNPs and indels by Illumina paired-end whole-genome sequencing. Across the four samples (wild type and DA-resistant clones 4C2, 5C1, and 6C3), there were a total of 158,396 SNPs and 28,640 indels (see Files S1 and S2 for the statistics on SNPs and indels, respectively). Of the indels, 20,107 were insertions, while the remaining 8,533 were predicted deletions. Both datasets were filtered, firstly to remove mutations occurring in all four samples, as these are clearly not related to drug resistance.

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**Figure 8** Mitochondrial membrane potential in IL3000 and 4 diminazene-resistant clones determined by flow cytometry. (a) Sample profiles for TMRE fluorescence of 10,000 cells was obtained on a FACSCalibur flow cytometer, using the FL2-height detector. Sample graphs of the IL3000 DA-sensitive parental strain and the 6C1 DA-Res clone are given. (b) Average (and SEM, $n = 3$) percentage of cells with a fluorescence $>80$ A.U. in the analysis for IL3000 and 4 DA-Res clones. Significant differences with the sensitive control are indicated with red stars; difference between 5C1 and the other three DA-res cell lines are indicated with blue stars. *** $p < .001$
and subsequently to obtain lists of high, moderate, and low impact (as described in the methods) mutations in protein encoding regions (5,263 SNPs and 541 indels) (Table S3, worksheets "SNPs-HIGH-MODERATE-LOW" and "Indels-HIGH-MODERATE-LOW"). Once mutations that occurred in genes predicted to encode variant surface glycoproteins (VSGs), expression site-associated genes (ESAGs) and retrotransposon hot spot (RHS) proteins were removed, there were 19 high impact SNPs (Table S3, worksheet "SNPs-Hi-filtered") and 192
high impact indels (Table S3, worksheet “Indels-Hi-filtered”), in addition to 1,693 moderate impact SNPs (Table S3, worksheet “SNPs-Mod-filtered”) and 42 moderate impact indels (Table S3, worksheet “Indels-Mod-filtered”) predicted to result in non-synonymous mutations for the former, and in-frame deletions/insertions for the latter. Due to the previously described correlation between resistance to the related compound isometamidium chloride (a fusion molecule of ethidium and diminazene) and reduced mitochondrial membrane potential (Wilkes et al., 1997), as well as reduced potential observed in the DA-resistant cell lines, data were mined to identify genes that could be involved in maintaining or generating this membrane potential. Two copies of a vacuolar-type Ca\(^{2+}\)-ATPase (TcIL3000.A.H_000569200 and TcIL3000.A.H_000569400) harbored missense mutations in all three resistant lines (TcIL3000.A.H_000569200: A109T in clones 4C2 and...
No further candidates were found. Two missense mutations were found in folate transporter TcIL3000.A.H_000597400 (A132S and N131Y, both in clone 4C2 only, both heterozygous) and another two in folate transporter TcIL3000.A.H_000597700 (L122R in 4C2 and F371C in clone 6C3; both heterozygous). No mutations were found in the other folate or pteridine transporters. Missense mutations were also found in several copies of an amino acid transporter (TcIL3000.A.H_000388400, TcIL3000.A.H_000388500, & TcIL3000.A.H_000388600), a putative plasma-membrane choline transporter (TcIL3000.A.H_000774600) and two copies of a mitochondrial chaperonin HSP60 gene (TcIL3000.A.H_000771700 & TcIL3000.A.H_000772600) (Table S3, worksheet “SNPs-HIGH-MODERATE-LOW”).

2.9 | Transcriptomics analysis of diminazene resistance

To ascertain whether differential gene expression could explain increased levels of resistance to diminazene, three replicates of the wild-type parental line, as well as three replicates of each independent resistant clone, were subjected to transcriptomic analysis by Illumina paired-end sequencing. Data were processed as described in the methods, resulting in transcript abundances for 9,360 T. congolense genes (Table S3, worksheet “RNAseq_all”). Data were filtered to obtain differentially expressed genes relative to wild-type (q value ≤0.05), resulting in 274, 247, and 265 genes identified as being significantly differentially expressed in DA-resistant clones 4C2, 5C1, and 6C3, respectively, with 120 of these genes being common to all three clones (Figure 10). Genes were then filtered to remove VSGs, ESAGs, and RHS proteins, resulting in a final list of 61 genes, of which 15 were annotated as hypothetical (Table S3, worksheet “RNAseq_filtered”).

One of the most significant changes was downregulation of an array of H4 histone encoding genes (mean log2 fold change: −7.842) (Table S3, worksheet “RNAseq_DOWN”). Furthermore, there was significant downregulation of a putative protein involved in elongation of very long fatty acids (TcIL3000.A.H_000436100.1; mean log2 fold change: −6.526) and a putative ZIP Zinc transporter that is, however, indicated as a possible pseudogene (TcIL3000.A.H_000205900.1; mean log2 fold change: −4.099). Other downregulated genes of note included a cysteine peptidase, as well as four hypothetical proteins (TcIL3000.A.H_000205700.1, 000205600.1, 000429100.1 and 000429200.1) for which expression was entirely abolished in all resistant clones. None of the hypothetical proteins were found to contain a recognized domain using the SMART (Letunic & Bork, 2018), interpro (Hunter et al., 2009), and pfam (El- Gebali et al., 2019) tools. Not all of these may be equally relevant because the locus of five genes from TcIL3000.A.H_000205600.1 to TcIL3000.A.H_000206000.1 were all extremely downregulated and this includes a Histone H4, three hypothetical proteins that are not related to each other, and the Zip Zinc Transporter mentioned above.

There were only 13 significantly differentially expressed genes exhibiting a log2 fold change of >1.0 in all three resistant clones, and these were identified as primarily RNA helicases, in addition to copies of cysteine peptidase and several hypothetical proteins. As the data for the RNA helicases have different significance and expression values, it is unlikely that the identification of this gene family is due to multimapping as in the Histone H4 genes, in this case leading to a firmer conclusion that all individual genes identified are differentially expressed (Table S3, worksheet “RNAseq_UP”).

![Figure 10](image-url) Differential gene expression in three independent DA-resistant clones (4C2, 5C1 and 6C3) of T. congolense IL3000. For each DA-resistant clone, Log2 fold change was calculated compared with wild-type parasites, as well as significance (q-value), and plotted as a volcano plot. Differential gene expression was deemed significant if the gene exhibited a Log2 fold change of >1 or <=1 in all three clones, as well as a q-value of <0.05 (-Log10 q-value of 1.30103) (genes shown as asterisks). The most significantly upregulated (Helicase associated domain (HA2) putative: TcIL3000.A.H_000255300.1) and downregulated (Histone H4 putative: TcIL3000.A.H_000205600.1) genes are shown as circled asterisks to illustrate reproducibility across clones.
some cases, there was significant differential expression in only two of the DA-resistant clones, for example, a predicted long-chain fatty acid-CoA ligase (TcIL3000.A.H.00065500.1; mean q-value: .0356; mean Log₂ fold change: -0.297).

3 | DISCUSSION

In this manuscript, we have studied the phenomenon of diminazene resistance in T. congolense. This veterinary parasite does not have an equivalent of the T. brucei diminazene transporter TbAT1 (Munday et al., 2013) and thus it is clear that the T. brucei model for diminazene resistance, based on loss of TbAT1 function (De Koning et al., 2004; Graf et al., 2013), is not applicable to T. congolense.

We found that it was possible to induce a substantial level of diminazene resistance in T. congolense by slowly increasing the drug concentration in the culture medium. The level of DA resistance was entirely stable in all clones, after months of in vitro passage in the absence of drug pressure, and repeated cycles of stabilating and reculturing. Remarkably, all the independently generated DA-resistant clones displayed the same level of resistance, cross-resistance patterns, and growth rates, which could indicate that they had developed very similar adaptations.

One of the main aims of this study was to establish whether DA resistance in T. congolense necessarily leads to cross-resistance to other (potential) AAT drugs. Co-resistance with isometamidium chloride has been reported in the field (Afewerk et al., 2000; Ainanshe et al., 1992; Mulugeta et al., 1997), but it is not clear whether this constitutes genuine cross-resistance. This conclusion can be hard to reach in the field as cattle are often treated with both drugs and independently developed resistance to both drugs is therefore a real possibility.

However, there have been reports of single resistance to diminazene and to isometamidium, and more recently to both drugs (Geerts et al., 2001; Giordani et al., 2016; Sinyangwe et al., 2004). This clearly shows that multiple-resistance is not automatic, as it seems to be for pentamidine and melsarsoprol in T. brucei ssp. (Baker et al., 2013). Our observation that induced DA resistance in T. congolense did not change isometamidium sensitivity in any of the independent clones is certainly a strong confirmation of that hypothesis and to the best of our knowledge the first such evidence from a controlled laboratory study since experiments in 1963 by Frank Hawking with multiple T. congolense strains in guinea pigs and mice, which also found no evidence of cross-resistance between DA and phenanthridines (Hawking, 1963). We have also attempted to similarly induce isometamidium resistance in T. congolense, but this was not successful in vitro although it has been achieved in vivo (Tihon et al., 2017). Another important observation on the nature of DA resistance in T. congolense is that it was remarkably stable, which, considering the many reports of DA resistance from all over Africa, has implications for the future utility of the drug.

As mentioned earlier, it has been argued that the further development of diamidine drugs against AAT would be unwelcome and only result in more drug pressure on populations that already harbor DA resistance. The strong cross-resistance with furamidines such as DB75 and DB829, observed both in vitro and in vivo and with field isolates and in vitro-adapted clones, would certainly strengthen this argument. However, the complete lack of cross-resistance with pentamidine and the tested series of bis-benzofuramidines demonstrates that only the closest structural analogs display cross-resistance with DA, and that therefore not all diamidines should be ruled out from consideration for further development against AAT.

The caveat, however, would be that any new diamidine drug must be active, at a minimum, against all the major species causing nagana in sub-Saharan Africa, including DA-resistant strains of each of those species. Given the dearth of knowledge about DA-resistant T. vivax, in particular, this would still constitute a significant hurdle at the moment. In this context, the observation that there is no cross-resistance between DA and oxaboroles, promising new agents in development for HAT and AAT (Akama et al., 2018; Begolo et al., 2018; Jacobs et al., 2011), is highly significant.

In T. brucei, drug resistance is mostly linked to changes in drug accumulation, particularly through mutations in transporter genes (De Koning, 2020). In T. congolense, isometamidium resistance has also been linked to reduced accumulation (Mulugeta et al., 1997; Sutherland et al., 1991, 1992; Tihon et al., 2017), but no drug transporters have been identified in this species. We therefore investigated the uptake of [³H]-DA in wild-type IL3000 and found it to be very slow and, apparently, low affinity. This indicates that uptake of diminazene could be limiting to its efficacy, whereas in T. brucei ssp. uptake of diminazene, the furamidines, and pentamidine is very fast with high affinity (De Koning et al., 2004; De Koning & Jarvis, 2001; Tekah et al., 2011; Ward et al., 2011). We found no consistent indication that DA resistance in T. congolense is linked to reduced cellular accumulation of the drug, as the accumulation rate was similar in the parental and resistant clones.

Several compounds were identified as inhibitors of both [³H]-diminazene and fluorescent DB75 uptake in T. congolense, particularly pentamidine, propamidine, and folate. The latter observation indicates the involvement of folate transporters in diminazene uptake, although high concentrations were required for significant inhibition and even then diminazene uptake was only partially inhibited. Clearly, a so-far unidentified, folate-insensitive uptake mechanism is additionally involved in diminazene uptake. The notion of multiple low-affinity entry routes was further strengthened by the demonstration that inhibition by folate and pentamidine was additive, and yet still incomplete. To further investigate whether T. congolense folate transporters may contribute to diminazene uptake, all three then-known FTs were cloned and expressed in the diamidine- and melaminophenyl-resistant clone T. brucei B48 (Bridges et al., 2007). The expression of IL3000 transporters FT1 and FT3 appeared to induce a small but significant sensitization to diminazene, whereas in vivo resistance to the other two transporters, but the effect did not reach statistical significance. The folate transporters of two DA-resistant isolates
from Cameroon displayed several SNPs relative to the IL3000 sequences, but the levels of sensitization induced by those FTs were not significantly different from those of IL3000. However, FT2 from both resistant isolates significantly sensitized to all three diamidines including pentamidine, but FT2 from IL3000 did not sensitize to any of the diamidines tested. None of the FTs changed the EC_{50} value for AN7973. These observations show that the folate transporters have a minor capacity for diamidine transport, but also that SNPs observed in these genes are not the cause of DA resistance, either in the field or in our laboratory-adapted clones. The lack of genuine resistance mutations in the folate transporters is presumably linked to the observation that at least two of those transporters have this capacity, and that the FTs mediate only a proportion of DA uptake, considering the modest effect of 1 mM folate on [H]-diminazene uptake.

Based on the above reasoning, we must conclude that diminazene resistance in *T. congolense* is not principally the result of a reduced rate of cellular diminazene accumulation. The alternative cause of resistance would be in changes of the target. In the case of diminazene and other cationic drugs this target is the mitochondrion (Alkhaldi et al., 2016; Basselin et al., 2002; Fueyo Gonzalez et al., 2017; Lanteri et al., 2008), and the fluorescent DB75 has been shown to accumulate in the kinetoplast (Mathis et al., 2006, 2007; Stewart et al., 2005), binding to the kDNA or interfering with enzymes involved in the functioning and/or replication of the kinetoplast. We confirm the mitochondrial targeting of DA in *T. congolense* with TEM, showing mitochondrial lesions appearing after a 4-hr incubation with 1.5 µM DA (5 × EC_{50}), whereas no other cellular changes were apparent. While dyskinetoplastic *T. brucei* have been described, including as part of the adaptation to isometamidium (Eze et al., 2016), and *T. evansi* is considered a dyskinetoplastic *T. brucei* (Lun et al., 2010), dyskinetoplastic *T. congolense* have not been described, and our DA-resistant clones displayed normal kinetoplasts as observed by DAPI staining and TEM. Thus, unlike *T. brucei* spp (Dean et al., 2013; Eze et al., 2016), *T. congolense* does not appear to have the ability to become resistant to kinetoplast-targeting drugs by losing dependence on genes encoded by kDNA. *T. brucei* bloodstream forms have a unique mitochondrial function and lack the usual oxidative phosphorylation pathways, instead employing a trypanosome alternative oxidase (TAO) as the sole terminal oxidase (Chaudhuri et al., 1998; Ebiloma et al., 2018, 2019) and using the F_{1}F_{0}-ATPase to pump H^{+} from the mitochondrial matrix and so generate the necessary Ψm (Schnauber et al., 2005). Bloodstream *T. congolense* may have a similar energy metabolism and mitochondrion, seeing that they have similar sensitivity to inhibitors of TAO (Fueyo Gonzalez et al., 2017) and no sensitivity to cyanide (Bienen et al., 1991).

However, in order to reach and disable the kinetoplast and/or other mitochondrial targets, the drug must accumulate inside the mitochondrion—a process that is dependent on the mitochondrial membrane potential Ψm. We observed, in all four DA-resistant clones investigated, a statistically significant reduction in Ψm, which might serve to diminish the entry of diminazene into the mitochondrion, where the drug would otherwise be strongly accumulated. Although DA and structurally related diamidines bind to kDNA, it should be mentioned that the lesions observed with TEM were in the mitochondrial matrix rather than any observable alterations to the kinetoplast structure, leaving open the possibility of mitochondrial targets other than the kinetoplast. This is particularly important as the entry of the drug into *T. congolense* is low affinity and inefficient, as opposed to the energy-dependent, high affinity process in *T. brucei*. Diminazene uptake in *T. congolense* is thus very likely equilibrative rather than concentrative, implying that a failure of the drug to accumulate in the mitochondrion will eventually result in a reduced cellular uptake as well. A reduced (rate of) diminazene accumulation in the mitochondrion of the resistant cells would be consistent with the observed lack of diminazene-induced mitochondrial lesions at 4 hr and relatively few alterations even at 8 hr. However, we acknowledge that the reduced mitochondrial membrane potential is likely not the only, or main, factor contributing to DA resistance, particularly as we did not observe a reduced time-to-fluorescence for kDNA on incubation with DB75. Moreover, the lack of cross-resistance to pentamidine and isometamidium also indicates that a more specific adaptation is likely to be at least partly responsible for the narrow resistance phenotype. In light of these observations, it is likely that the reduced Ψm is the result of that adaptation, rather than the primary cause of the DA resistance.

Genomic and transcriptomic analyses of the 4C2, 5C1, and 6C3 DA-resistant clones were performed to further investigate the causes of resistance. For each clonal line, several hundred genes were differentially regulated, relative to the parental clone IL3000. After filtering-out inconsistent and functionally irrelevant returns relatively few genes were significantly downregulated (including H4 histones, a cathepsin-like cysteine protease, and several hypothetical proteins with no known domains) or upregulated (mainly RNA helicases and a few hypothetical proteins). However, the highly stable level of resistance would be consistent with permanent mutations rather than the more transient changes in transcription. The genomic sequencing revealed 19 high impact SNPs and 192 high impact indels. SNPs were observed in several transporter genes, including amino acid transporters, some of the folate transporter genes and a putative choline transporter, as well as mitochondrial HSP60, but mostly in only 1 of the resistant clones. Most interestingly, however, in all three DA-resistant clones sequenced, missense mutations were observed in at least one of two copies of a vacuolar-type Ca^{2+}-ATPase.

The mutations in the vacuolar Ca^{2+}-ATPases could be relevant to a resistance mechanism that is linked to mitochondrial function, as trypanosome mitochondria are known stores of Ca^{2+} (Ramakrishnan & Docampo, 2018), which appears to be released to the cytosol upon membrane depolarization (Alkhaldi et al., 2016) through a Ca^{2+}/H^{+} exchanger (Ramakrishnan & Docampo, 2018), and the bioenergetics of trypanosomes is regulated by inositol 1,4,5-trisphosphate receptor-mediated Ca^{2+}-release from acidocalcisomes (Chiurillo et al., 2020; Huang, Bartlett, et al., 2013) and a mitochondrial Ca^{2+} uniporter
(Huang, Vercesi, et al., 2013). The *T. congolense* Ca^{2+}-ATPases are syntonic with *T. brucei* vacuolar Ca^{2+}/H^{+} antiporters PMC1 and PMC2, which are essential genes and localized to the acidocalcisomes (Huang & Docampo, 2015; Luo et al., 2004). RNAi knockdown of TbPMC1 expression increased cytosolic calcium levels but reduced the amount of Ca^{2+} in intracellular stores. Interestingly, diamidines including DB75 and DB820 have been shown to accumulate not just in the nucleus and kinetoplast of *T. brucei*, but also in the acidocalcisomes, as evidenced by a slow-onset yellow-orange fluorescence (Mathis et al., 2006). Acidocalcisomes have been described as drug targets and have essential regulatory roles in trypanosomes, particularly concerning ion distribution, pH, and Ca^{2+}-signaling (Docampo & Moreno, 2008). As such, the two missense mutations in the vacuolar-type Ca^{2+}-ATPase, each of which occurred in two independent clones, should certainly be followed up as potential contributing factors in *T. congolense* DA resistance, but this will clearly require extensive further validation. The very high level of DA-induced SNPs and indels is consistent with the DNA-targeting nature of this minor groove binder.

Altogether, we conclude that a clinically relevant level of DA-resistance in *T. congolense* can be induced in vitro and is stable, and not primarily due to reduced DA accumulation but instead associated with a reduced mitochondrial membrane potential, although this is not likely to be the main cause of the resistance phenotype. DA uptake is low affinity and partially mediated not only by the *T. congolense* folate transporters, but also by separate transporters that are sensitive to pentamidine. These results constitute the first systematic analysis of DA-transport and resistance mechanisms in *T. congolense* and provide some insights important for the development of new drugs against AAT.

4 | EXPERIMENTAL PROCEDURES

4.1 | In vitro culture of *T. congolense*

Cultures of bloodstream forms of Savannah-type strain IL3000 and the adapted clones derived from this strain were cultured exactly as described by Coustou et al. (2010), in a basal MEM-based medium supplemented with 20% goat serum 14 µl of β-mercaptoethanol, 800 µl of 200 mM glutamine solution, and 10 ml of 100× penicillin/streptomycin solution per liter of medium (pH = 7.3). *T. congolense* were cultured in 6- or 24-well plates at 34 ºC and 5% CO_{2}. For adaptation to DA, cultures were serially passaged in the highest concentration of DA tolerated, which was increased as the cells adapted, starting at 50 nM DA (Sigma). Thus one 24-well plate contained 1 row of “no drug” control culture of IL3000 bloodstream forms and three rows of independent cultures in the presence of DA, in three drug concentrations per passage, one above and one below the concentration under test.

4.2 | Resazurin assay with *T. congolense*

This drug susceptibility assay was performed essentially as described previously (Cerone et al., 2019) and is based on only live cells reducing the blue and non-fluorescent viability indicator dye resazurin sodium salt ("Alamar blue"; Sigma) to pink, fluorescent metabolite resorufin (Gould et al., 2008). The fluorescence was quantified using a FLUOstar Optima (BMG Labtech, Durham, NC, USA) at wavelength of 540 nm (excitation), 590 nm (emission), and the data plotted to a sigmoidal curve with variable slope using Prism 5.0 (GraphPad Software Inc., San Diego, CA, USA). The assay was set up in 96-well plates with the test drug added at doubling dilutions in culture medium, usually starting at 50 µM, over one or two rows of the plate, with the last well receiving 100 µl culture medium as a drug-free control, that is either 11 or 23 doubling dilutions. To each well, 100 µl of culture medium containing 5 x 10^5 bloodstream *T. congolense* cells was added, and the plate was incubated for 48 hr at 34 ºC/5% CO_{2}, followed by the addition of 20 µl of a 125 mg/ml solution of resazurin sodium salt in phosphate-buffered saline (PBS) and a further incubation period of 24 hr.

4.3 | In vivo experiments and ethics statement

Drug sensitivity testing in mice used the standardized single-dose procedure of Eisler et al. (2001). Groups of six NIH mice (Envigo) were used throughout, and infected i.p. with 10^7 *T. congolense* strain IL3000 or KONT2/151 in phosphate-buffered saline pH 7.4 (PBS). At 24 hr post infection (PI) a single dose of DB829, AN7973, or vehicle (DMSO, drug-free control) was administered i.p., at doses 10, 20, or 40 mg/kg bw for DB829 and 5, 10, or 20 mg/kg bw for AN7973. Parasitemia was monitored daily, scored by microscopic analysis of blood from a tail prick from each infected animal. Any mouse found to have high parasitemia during the experiment, or show any signs of suffering were euthanized by CO_{2} inhalation. All animal experimentation was performed at the University of Glasgow Joint Research Facility under the supervision of trained professionals; the facility is regularly inspected by a UK Home Office Inspector and adheres to all national and international regulations as stipulated by the UK Home Office for animal care and in accordance with the Animals (Scientific Procedures) Act 1986 as amended in 2012. The procedure had been expressly approved and licensed by the UK Home Office (project license PCF371688), and the experiment was performed by a trained animal technician under his personal Home Office license (PIL601/12386).

4.4 | Fluorescence microscopy screen for diamidine uptake inhibitors

NIH mice (Envigo) were infected with 100,000 trypanosomes of either *T. b. brucei* strain Lister 427 (i.p.), *T. congolense* strain IL3000 (i.p.) or *T. vivax* strain Y486 (Gibson, 2012) (i.v.) in PBS; each parasite species was injected in sterile saline solution. At peak parasitemia, the mouse was killed by CO_{2} inhalation and the blood collected by aortic bleed. This was kept on ice if not being used immediately for an experiment.
Approximately, 1 ml of whole blood was spun at 1,000 x g, and supernatant, buffy coat and a visible layer of parasites were removed to a new tube and spun again. The resulting pellet was washed with 1 ml of CBSS buffer (25 mM HEPES, 120 mM NaCl, 5.4 mM KCl, 0.55 mM CaCl2, 0.4 mM MgSO4, 5.6 mM Na2HPO4, 11.1 mM glucose; pH adjusted to 7.4 with NaOH), and the pellet resuspended in CBSS buffer at an appropriate concentration to visualize the parasites (usually 1 ml). The duration of viability of parasites was assessed at room temperature, 34ºC and 4ºC prior to completion of experiments. T. congolense parasites performed similarly when kept at 34ºC and room temperature during experiments; therefore, experiments were conducted at room temperature; T. vivax parasites were motile for longer and took up DB75 at a faster rate at 34ºC, so were kept at this temperature as much as possible during experiments (but not while under the microscope for observation).

For non-inhibited cells, 10 µM of DB75 was added and fluorescence observed using a Zeiss Axioscope fluorescence microscope, using the DAPI filter set (λex = 330 nm, λem = 400 nm) as well as brightfield. Images were obtained with Openlab imaging software (Improvision, Coventry, UK). For inhibited cells, the test inhibitor was added to the desired concentration (see Table S1) just before the addition of 10 µM DB75 and observation of fluorescence as before. Each potential inhibitor was tested at least twice on parasites from different mice, those which inhibited up to four times. Every experiment had cells with no inhibitor added visualized in parallel to control for slight variation in timing of appearance of fluorescence in kinetoplasts and nuclei between batches of ex vivo parasites.

### 4.5 Transport assays

Transport assays for T. congolense bloodstream forms were performed exactly as described for T. b. brucei bloodstream forms (Wallace et al., 2002) and Leishmania major promastigotes (Al-Salabi et al., 2003). Depending on the assay, parasites were either purified from infected blood using a DEAE-cellulose column and a 6:4 ratio PSG buffer as described (Lanham & Godfrey, 1970), or collected from culture. Briefly, cells were washed into assay buffer (33 mM HEPES, 98 mM NaCl, 4.6 mM KCl, 0.55 mM CaCl₂, 0.07 mM MgSO₄, 5.8 mM Na₂HPO₄, 0.3 mM MgCl₂, 23 mM NaHCO₃, 14 mM glucose, pH 7.3) and adjusted to a density of 10⁵/ml just before use. About 100 µl of the cell suspension was added to 100 µl of radiolabeled substrate (³H-DA or ⁴H-pentamidine) in assay buffer, sometimes also containing a competitive inhibitor at 2 – 19 µM just before use. After a predetermined incubation time, the incubation was stopped by the addition of 1 ml ice-cold “stop solution” (1 mM/250 µM unlabeled permeant in assay buffer), and cells were separated from the extracellular radiolabel by centrifugation through the oil layer (1 min at 13,000 rpm in a microfuge). The cell pellets were harvested into scintillation tubes by cutting off the tip of the microfuge tube after flash-freezing in liquid nitrogen, incubated with 2% SDS for at least 1 hr, and overnight with scintillation fluid (Optiphase HiSafe III, Perkin-Elmer, Waltham, MA, USA) before being agitated overnight in scintillation fluid (Optiphase HiSafe III, Perkin-Elmer, Waltham, MA, USA). Tubes were then read using a Hidex 300 SL scintillation counter (Lablogic, Sheffield, UK). Raw disintegrations per minute (dpm) were corrected for background radiation and non-specific radiolabel association, by subtracting dpm counts from no-label controls and parallel determinations in the presence of saturating levels of non-labeled permeant, respectively. Radiolabeled ring-³H-DA was custom-made by PerkinElmer (CUST78468000MC; 60.7 Ci/mmol) and ⁴H-pentamidine isethionate was custom-made by Amersham (TRQ40084; 3.26 TBq/mmol). [3,5,7-³H]-folic acid (ART 0125; 56.8 Ci/mmol) was from American Radiolabeled Chemicals. Statistical analyses were performed with GraphPad Prism 8.1.

### 4.6 Cloning and expression of T. congolense folate transporters

Three folate transporters were identified in the original T. congolense IL3000 genome available in TriTrypDB (version 1 of the genome, released Oct 20 2010). The gene designated as Folate Transporter (FT) 1 (TcIL3000.10.7850) was annotated as a predicted pteridine transporter, and FT2 (TcIL3000.0.06950) and FT3 (TcIL3000.0.13340) were predicted folate transporters.

Subsequent to the completion of the cloning work described in this paper, the T. congolense IL3000 2019 genome was released in TriTrypDB (release Nov 6 2019); this contains several (additional) predicted genes all annotated as putative folate transporters. TcIL3000.A.H_000799700 aligns with our FT1 (GenBank accession numbers in Table 2 and Fig. S2). Three tandemly arranged, almost identical genes (97.8%–98.4% identity), TcIL3000.A.H_000597400, TcIL3000.A.H_000597500, and TcIL3000.A.H_000597700, align with our FT2 with 97.3%–99.8% identity by Clustal Omega analysis. And two genes annotated as being pseudogenes due to missing START and STOP codons, TcIL3000.A.H_000531600 (pseudogene) and TcIL3000.A.H_000558900: pseudogene, align with our FT3. The predicted CDS of TcIL3000.A.H_000531600 is the same as the IL3000 FT3, only having a predicted N-terminal extension with no START codon. The last 27 amino acids and STOP codon that are missing from the sequence given on the TriTrypDB gene page are actually present when the whole contig is examined. There are long stretches of TcIL3000.A.H_000558900 that are divergent (67.9% identical by Clustal Omega analysis). All are aligned in Figure S2.

Based on the TriTrypDB sequences obtained, primers were designed to amplify the three genes with either an Apal or HindIII 5′ restriction site and a 3′ XbaI restriction site. The Primer sequences were as follows: FT1_F 5′-ggggccATGTTGCGACGCGTCCG-3′; FT1_R 5′-cttagaCATATGTGGGCGCTCA-3′; FT2_F 5′-aagcttATGAGCGGCAAAGCGG-3′; FT2_R 5′-cttagaCATATGTGGGCGCTCA-3′; FT3_F 5′-ggggccATGTTGCGACGCGTCCG-3′; FT3_R 5′-cttagaCATATGTGGGCGCTCA-3′. Genomic DNA was extracted from parasites isolated from 1 ml of cardiac puncture blood.
from mice infected with each strain: IL3000, KONT2/133, and KONT2/151, using a Qiagen DNeasy blood and tissue kit, following the manufacturer’s instructions. Genes were amplified from the relevant gDNA using the primers given above and Phusion polymerase (New England Biolabs), following the manufacturer’s protocol. DNA fragments of approximately the correct size were extracted from 1% agarose gels and ligated into the pGEMT-Easy (Promega) subcloning vector. Positive colonies were identified by screening using the M13F/M13R sequencing primers on the vector and sequenced by Sanger sequencing (Source BioScience, Nottingham UK). The identified sequences were ligated into the vector pHDI1336 and transfected into T. b. brucei strain B48 (Bridges et al., 2007) using an Amaxa Nucleofector, program X-01, and TbBSF buffer as described previously (Burkard et al., 2007; Munday et al., 2013). Transfectants were cloned out, by limiting dilution, in standard HMI-11 medium (Hirumi & Hirumi, 1989) containing 5 µg/ml blasticidin S and clones screened for correct integration of the cassette by PCR. All clones and 5% CO₂

Drug sensitivity of the T. brucei cell lines was assessed using the standard resazurin assay, essentially as described for T. congolense, above and as described previously (Siheri et al., 2016). Briefly, the assay was performed in 96-well plates with 11 doubling test compounds dilutions, highest concentration 10 µM, and the 12th well containing cells in drug-free medium. Bloodstream form of 2 × 10⁴ trypanosomes were added to each well, and the plates were incubated for 48 hr at 37°C/5% CO₂ after which 20 µl of resazurin solution was added to each well and the plates incubated for another 24 hr.

4.7 Mitochondrial membrane potential (Ψm) assay

Changes in Ψm were determined by flow cytometry using the indicator dye tetramethylrhodamine ethyl ester (TMRE) as the probe, exactly as previously described (Cerone et al., 2019; Ibrahim et al., 2011). Briefly, bloodstream forms of 1 × 10⁷ T. congolense IL3000 and several IL3000-adapted DA-Res clonal lines in mid-log growth phase were centrifuged at 1,600 × g for 10 min at 4°C and washed in 1 ml pf PBS (pH 7.4) and finally resuspended in 1 ml PBS containing 200 nM TMRE. These cells were incubated for 30 min at room temperature and then placed on ice for a further 30 min prior to analysis on a Becton Dickinson FACSCalibur™ system (BD Biosciences) using a FL2-height detector, and CellQuest and FlowJo 10 software.

4.8 DAPI staining of nuclei and kinetoplasts

Approximately, 1 × 10⁶ cells were harvested in Eppendorf tube and centrifuged at 2,400 rpm for 10 min in a microfuge. The supernatant was discarded; the cells were resuspended in 1x PBS and washed by centrifugation (2,400 rpm for 5 min). The supernatant was discarded; the cells were resuspended in 50 µl 1x PBS pH 7.4, spread out on a microscope slide and allowed to dry for 30 min. The cells were then fixed by flooding the slide with 4% formaldehyde for 10 min, followed by washing twice in 1x PBS. The slides were allowed to air dry before placing a drop of Vectashield mounting medium containing DAPI (4′,6-diamidino-2-phenylindole dihydrochloride) (Vector Laboratories) on them. The slides were then covered with a cover slip and the edges sealed with nail varnish. The slides were viewed under a DeltaVision microscope (GE Healthcare) using softWoRx software and the images were processed using ImageJ software.

4.9 Transmission electron microscopy

T. congolense WT and diminazene-resistant clone 6C3 were prepared essentially as described previously for T. b. brucei (Cerone et al., 2019; Millán et al., 2017) at about 1 × 10⁶ cells/ml with or without diminazene at 1.5 µM in TCBSF3 medium. The samples were incubated at 34°C and 5% CO₂ for 4 or 8 hr, centrifuged at 1,100 × g for 10 min and washed in 1x PBS. The pellets were resuspended and fixed in 2.5% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.2) overnight at 4°C, then washed in 0.1 M cacodylate buffer (pH 7.2), and post-fixed in 1% OsO₄ for 1 hr on ice. After several washes in the same buffer, the samples were en bloc stained with 0.5% uranyl acetate in water for 30 min. Thereafter, samples were washed with water, dehydrated in an ascending acetone series, and embedded in epoxy resin. Ultrathin sections were collected and imaged on a JEOL 1200 Transmission electron microscope (JEOL, Japan). All images were analyzed and processed with FIJI software (Schindelin et al., 2012).

4.10 RNAseq

Total RNA isolations were performed on 1 × 10⁷ cells of a parental strain of wild-type T. congolense IL3000, as well as three replicates of each of the three independent DA-resistant clones (4C2, 5C1 and 6C3), using a commercial kit (RNaseq, Qiagen). Samples were enriched for mRNA by Poly-A selection, using the TruSeq stranded mRNA sample preparation kit (Illumina, catalog number 20020594) and subsequently quantified using Qubit high sensitivity reagents (Thermo). Average library size was determined using DNA high-sensitivity reagents for the Agilent bioanalyser 2100 (Agilent). For sequencing, 2 × 75 bp paired-end sequencing was carried out using an Illumina NextSeq 500 system (Illumina), resulting in an average of 12 million reads per sample. For RNA-seq data processing, reads contained in the resultant fastq files were aligned to the PacBio genome of the T. congolense strain IL3000 (Abbas et al., 2018) and gene abundances quantified, in both cases using Kallisto with default parameters (−b 100, −t 8) (Bray et al., 2016). Statistical analyses of differential expression between the wild-type parental strain and each of the three resistant clones.
were carried out using the R-based package Sleuth, with default parameters (for each transcript, a second fit was performed to a reduced model that presumes abundances are equal in the two conditions—wild-type and DA-resistant) as outlined in the manual (Bray et al., 2016), and figures were generated with Matlab (The MathWorks, Inc.). Genes exhibiting significant (q-value ≤0.05) differential expression between wild-type and resistant parasites were tabulated and filtered to remove VSGs, ESAGs and retrotransposon hot spot proteins, and subsequently sorted to obtain genes with significance based on q-value (i.e., p-value adjusted for multiple testing by false discovery rate). The full Sleuth output is available in Table S3, worksheet “RNAseq_all”.

4.11 Whole-genome analysis

For whole-genome analysis, DNA was extracted from in vitro cultures (3 × 10⁶ cells) of the T. congolense IL3000 wild type line, as well as the three DA-resistant lines (4C2, 5C1, and 6C3), using the QIAGen DNeasy kit. Libraries were prepared using the QIAseq FX DNA library kit (QIAGen, cat: 180475). The libraries were quantified using Qubit high sensitivity reagents (Thermo), and average library size was determined using DNA high sensitivity reagents for the Agilent bioanalyzer 2100. The four samples were subsequently paired-end (2 × 75 bp) sequenced using Illumina NextSeq500 to an average of 15 million reads per sample. SNP and indel analysis were carried out on the resulting data using the GATK pipeline (McKenna et al., 2010). Reads were first trimmed using cutadapt (parameters: −q 30, 30 −minimum-length = 70 −pair-filter = any) (Martin, 2011) and subsequently aligned to the PacBio T. congolense IL3000 assembly (Abbas et al., 2018) using STAR (parameters: −outSAMtype BAM unsorted −readNameSeparator_−readFilesCommand zcat −runThreadN 8) (Dobin et al., 2013). The resulting bam files were further processed with Picard (functions: AddOrReplaceGroups & MarkDuplicates) (Mckenna et al., 2010) and GATK (functions: SplitNcigarReads, HaplotypeCaller, VariantFiltering, SelectVariants) to generate separate files for SNPs and indels. Further filtering was carried out on these files using the Snpeff tool (parameter: SnpSift.jar filter “(FILTER = ‘PASS’)”) (Cingolani, Platts, et al., 2012). Final datasets for SNPs and indels were created using the Snpeff command (Snpeff.jar), and data were extracted into tabular form using the SnpSift command (Cingolani, Patel, et al., 2012) [extractFields—CHROM POS ID REF ALT FILTER “ANN[*.GENE] “ANN[*.GENEID] “ANN[*.FEATURE] “ANN[*.EFFECT] “ANN[*.IMPACT] “ANN[*.CODON] “ANN[*.AA] “ANN[*.ALLELE] “GEN[*.GT]”). The resulting data were filtered firstly to remove SNPs occurring in both wild-type and resistant lines, and subsequently to select for SNPs and indels predicted to have a high (a variant that is assumed to result in a highly disruptive impact on the protein such as truncation or loss of function), moderate (non-disruptive variant that is assumed to impact protein effectiveness such as non-synonymous SNPs or in frame deletions) or low (Mostly harmless variants such as synonymous SNPs) impact, in both cases using Microsoft Excel.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest regarding the publication of this research.

AUTHOR CONTRIBUTIONS

Tim G. Rowan, Liam J. Morrison, Michael P. Barrett, Rose Peter, Harry P. De Koning performed conception or design of the study. Lauren V. Carruthers, Jane C. Munday, Godwin U. Ebiloma, Pieter Steketee, Siddharth Jayaraman, Gustavo D. Campagnaro, Marzuq A. Ungogo, Leandro Lemgruber, Anne-Marie Donachie performed acquisition, analysis or interpretation of the data. Liam J. Morrison, Harry P. De Koning wrote the manuscript.

DATA AVAILABILITY STATEMENT

Sequence data have been deposited in the European Nucleotide Archive (accession number PRJEB39051). All new sequence information on cloned folate transporters was deposited with GenBank and accession numbers are included in the manuscript text (Table 2). All other relevant data are included in the manuscript or its Supporting Information.

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REFERENCES

Abbas, A.H., Silva Pereira, S., D’Archivio, S., Wickstead, B., Morrison, L.J., Hall, N. et al. (2018) The structure of a conserved telomeric region associated with variant antigen loci in the blood parasite Trypanosoma congolense. Genome Biology and Evolution, 10, 2458–2473. https://doi.org/10.1093/gbe/evy186
Afewer, Y., Clausen, P.-H., Abebe, G., Tilahun, G. & Mehltz, D. (2000) Multiple-drug resistant Trypanosoma congolense populations in village cattle of Metekel district, north-west Ethiopia. Acta Tropica, 76, 231–238. https://doi.org/10.1016/s0001-706x(00)00108-x
Ainanshe, O., Jennings, F. & Holmes, P. (1992) Isolation of drug-resistant lines of T. congolense from the Lower Shabelle Region of southern Somalia. Tropical Animal Health and Production, 24, 65–73. https://doi.org/10.1007/bf02356946
Basselin, M., Lawrence, F. & Robert-Gero, M. (1996) Pentamidine uptake by Trypanosoma brucei. Biochemical and Biophysical Research Communications, 225, 312-316.

Bienen, E.J., Webster, P. & Fish, W.R. (1991) Trypanosoma congolense: Development of genetic tools. Molecular and Infectious Disease, 15, 153-160.

Begolo, D., Vincent, I.M., Giordani, F., Pöhner, I., Witty, M.J., Rowan, T.G. & Jarvis, S.M. (2002) Resistance to pentamidine and melarsoprol in African trypanosomes. Molecular and Biochemical Parasitology, 115, 113-122.

Bray, N.L., Pimentel, H., Melsted, P. & Pachter, L. (2016) Near-optimal alignment of long and short reads. Nature, 535, 338-343.

Bakunova, S.M., Bakunov, S.A., Wenzler, T., Barszcz, T., Werbovetz, K.A., Al- Atabani, A.A., Martinek, J., Panicucci, B., Dardonville, C., Zíková, A. & Al-Salabi, M.I., Wallace, L.J. & De Koning, H.P. (2003) A molecular pharmacology of bisbenzofuran cations. Molecular Pharmacology, 63, 587-593.

Carruthers ET al. (2014) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 140, 616-622.

Carruthers ET al. (2015) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 148, 137-144.

Carruthers ET al. (2016) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 166, 54-60.

Carruthers ET al. (2017) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 176, 27-35.

Carruthers ET al. (2018) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 190, 114-120.

Carruthers ET al. (2019) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 208, 179-186.

Carruthers ET al. (2020) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 209, 107992.

Carruthers ET al. (2021) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 217, 107958.

Carruthers ET al. (2022) Uptake of pentamidine and melarsoprol in Trypanosoma brucei. Experimental Parasitology, 218, 107964.
for Parasitology: Drugs and Drug Resistance, 7, 350–361. https://doi.org/10.1016/j.jipddr.2017.10.002

Vickerman, K. (1969) The fine structure of Trypanosoma congolense in its bloodstream phase. Journal of Protozoology, 16, 54–69. https://doi.org/10.1111/j.1550-7408.1969.tb02333.x

Vincent, I.M., Creek, D., Watson, D.G., Kamleh, M.A., Woods, D.J., Wong, P.E. et al. (2010) A molecular mechanism for efollorithine resistance in African trypanosomes. PLoS Path, 6, e1001204. https://doi.org/10.1371/journal.ppat.1001204

Vitooley, H.S., Mungube, E.O., Allegge-Cudjoe, E., Diall, O., Bocoum, Z., Diarra, B. et al. (2011) Improved PCR-RFLP for the detection of diminazene resistance in Trypanosoma congolense under field conditions using filter papers for sample storage. PloS Neglected Tropical Diseases, 5, e1223. https://doi.org/10.1371/journal.pntd.0001223

Wallace, L.J., Candlish, D. & De Koning, H.P. (2002) Different substrate recognition motifs of human and trypanosome nucleobase transporters selective uptake of purine antimetabolites. Journal of Biological Chemistry, 277, 26149–26156. https://doi.org/10.1074/jbc.m20283200

Ward, C.P., Wong, P.E., Burchmore, R.J., De Koning, H.P. & Barrett, M.P. (2011) Trypanocidal furamidine analogues: influence of pyridine nitrogens on trypanocidal activity, transport kinetics, and resistance patterns. Antimicrobial Agents and Chemotherapy, 55, 2352-2361. https://doi.org/10.1128/aac.01551-10

Wilkes, J.M., Mulugeta, W., Wells, C. & Peregrine, A.S. (1997) Modulation of mitochondrial electrical potential: a candidate mechanism for drug resistance in African trypanosomes. Biochemical Journal, 326, 755–761. https://doi.org/10.1042/bj3260755

Zoltner, M., Campagnaro, G.D., Taleva, G., Burrell, A., Cerone, M., Leung, K.-F. et al. (2020) Suramin exposure alters cellular metabolism and mitochondrial energy production in African trypanosomes. Journal of Biological Chemistry, 295, 8331-8347. https://doi.org/10.1074/jbc.RA120.012355

Zoltner, M., Horn, D., De Koning, H.P. & Field, M.C. (2016) Exploiting the Achilles’ heel of membrane trafficking in trypanosomes. Current Opinion in Microbiology, 34, 97-103. https://doi.org/10.1016/j.mib.2016.08.005

SUPPORTING INFORMATION
Additional Supporting Information may be found online in the Supporting Information section.

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