Effects of some natural leads on *Trypanosoma* and *Leishmania* strains

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Abstract. Well-known medical herbal compounds including apigenin, daidzein, phyllanthin and tyramine were assessed against *Trypanosoma* and *Leishmania* protozoans. Two strains of the bloodstream forms of *Trypanosoma brucei*: s427-WT and TbAT1-B48, and *Leishmania major* and *Leishmania mexicana* promastigotes were utilised. Among selected natural compounds, apigenin and daidzein displayed moderate activity against African trypanosomes with EC₅₀ 16 µM for wild-type sensitive control strain. Tyramine was not found to be very active for trypanosomes strains while all compounds were found to have trivial activity for the inhibition of *Leishmania mexicana* strains.

INTRODUCTION

A leading cause of numerous tropical diseases is protozoan parasites that are members of the Trypanosomatidae family. Approximately 65 million people of African countries are at risk to become infected with sleeping sickness which can be lethal if left untreated (WHO Report, 2017). Sleeping sickness is transmitted by the tsetse fly and caused by the *Trypanosoma brucei* subspecies *T. b. gambiense* and *T. b. rhodesiense* (Lindner et al., 2010). In 2009, after intensive control efforts, the number of cases reported dropped below 10 000 (9878) for the first time in 50 years. This downturn in number of cases has been sustained, with 2804 new cases reported in 2015, the lowest level since the start of systematic global data-collection 76 years ago (WHO, 2017).

Leishmaniasis refers to a collection of chronic diseases caused by protozoa of the genus *Leishmania* and is naturally transmitted by sandflies (Alvar et al., 2012; Stockdale et al., 2013). Pentavalent antimonials and pentamidine, used for the treatment of leishmaniasis, are toxic or have severe side effects, such as renal, cardiac and neural toxicity, risk of diabetes and shock (Tiuman et al., 2011). No vaccine is available for human leishmaniasis, while the drugs used against the parasite are very toxic and patients are frequently unresponsive to treatment (Koff et al., 1994; Kumar et al., 2014; Tiuman et al., 2011).

*Leishmania mexicana* infects many hosts (Berzunza et al., 2015) and has been detected in an assortment of wild and domestic mammals, while humans are opportunistic hosts (Courtenay et al., 2017; Martina et al., 2017). Cutaneous leishmaniasis caused by *L. mexicana* was first described in chicle collectors of the Yucatan peninsula in 1912 by Seidelin, therefore its name is “Chiclero’s Ulcer” (Seidelin, 1912).

The drugs currently being used for the treatment display a high level toxicity, inadequate efficacy and require a long
Moreover, it has been reported that there is development of drug resistance by the parasites (Croft et al., 2005; Kerboeuf et al., 2008; Paes et al., 2011). Thus, it is important to explore new drugs or compounds that might be allied with the conventional treatment. Natural sources such as plants, algae and micro-organisms constitute an important source of alternative drugs (Medeiros et al., 2011; Rondon et al., 2012; Wink, 2012; Gamboa-Leon et al., 2014; Sifaoui et al., 2014). It has been estimated that there are about 250,000 medical plant species in the world but on the other hand, the therapeutic potential of only about 6% of them has been biologically evaluated. Furthermore, only approximately 0.75% of the identified medical herbal compounds have been studied in clinical trials (Jameel et al., 2014). The foremost virtues of herbal medicine include their low cost, the low prevalence of severe adverse effects and superior efficacy.

In the current study we selected the well-known medical herbal compounds apigenin, daidzein, phyllanthin and tyramine. Because of the reported antileishmanial and various biological activities of these natural compounds (Wong et al., 2009; Ilangkovan et al., 2016), we decided to further investigate their activity against various Trypanosoma and Leishmania strains including the well-characterised and multi-drug resistant Trypanosoma brucei clonal line B48. The clonal line B48 is highly resistant to the two main classes of trypanocides, the melaminophenyl arsenicals and the diamidines, due to the loss the TbAT1/P2 and HAPT1 drug transporters (Bridges et al., 2007).

MATERIALS AND METHODS

Materials
All tested pure compounds including apigenin, daidzein, phyllanthin and tyramine were purchased from Sigma (Steinheim, Germany). Stock solution in 100% DMSO for each compound was prepared and for the concentrations used in assay, the calculated amount of stock solution was taken and diluted with complete medium, ensuring that the final DMSO concentration did not exceed 1% in the final conditions. EC50 values were obtained using the Alamar blue assay and are given as averages in μM (±SEM), of 3 independent evaluations.

Cell culture

Trypanosoma brucei bloodstream forms (BSF) in-vitro
In this research, two strains of the bloodstream forms of Trypanosoma brucei were utilised. The first was the wild type strain of Trypanosoma brucei brucei (s427-WT) and the other was TbAT1-B48 that was acquired from the clone TbAT1-KO (Matovu et al., 2003) by exposure to pentamidine thus increasing resistance to pentamidine and melaminophenyl arsenicals. Consequently, these cells have neither TbAT1/P2 transporter nor the high affinity-pentamidine transporter genes (Bridges et al., 2007; Munday et al., 2014). Both strains were cultured in HMI-9 medium (pH 7.4) supplemented with 10% Fetal Calf Serum (FCS) and 14 μl/Litre of 13.4 M β-mercaptoethanol, as described by Hirumi and Hirumi (Hirumi et al., 1989). The medium was sterilized by filtration (0.22 μm, Millipore) inside a flow cabinet. The T. b. brucei cultures were incubated at 37 °C and 5% CO2 and passaged in vented flasks three times per week.

Leishmania major and Leishmania mexicana promastigotes
Leishmania major strain Friedlin (LmjF) and strains of Leishmania mexicana (MNYC/BZ/62/M379) were propagated in minimal essential medium (HO-MEM) with a pH value of 7.4 and 10% FCS using plastic flasks at 25 °C. The resultant cultures were then passed through fresh medium thrice per week.

Alamar Blue assay to determine the sensitivity to test compounds
Resazurin sodium salt (Alamar Blue) is commonly used as a cell metabolic function indicator. It is a non-fluorescent blue dye that is mixed with cell cultures containing various drug concentrations, in order to determine
the sensitivity of African trypanosomes or Leishmania cultures to the test compounds in vitro (Fumarola et al., 2004; Raz et al., 1997). In case there are no toxic effects caused by the drug, the color of the living cells changes to red and fluorescent from blue. Preparation of Alamar Blue involves dissolving 12.5 milligrams of resazurin sodium salt (Sigma) in 100 mL of phosphate-buffered saline (PBS) of pH 7.4, which was then filter-sterilized and stored in the dark at 4 °C.

**Drug sensitivity using Alamar Blue assay in T. b. brucei BSF**

For each test compound a solution of 200 µM in HMI-9 medium + 10% FCS is prepared using a 20 mM stock solution in DMSO; 200 µL of this is added to a first well of a 96-well plate. Of this, 100 µL is transferred to the next well, containing 100 µL of the same medium, achieving a 1:1 dilution, initiating a doubling dilution series across 2 rows of the plate; each experiment was positively controlled using pentamidine and the final well for each compound received 100 µL HMI-9 medium as negative control. To each well, 100 µL of a suspension of 2 × 10⁵ cells/ml was added, amounting to 1×10⁵ cells/ml as the final cell density. The plate is then incubated for 48 hours at 37 °C/5% CO₂ after which 20 µL of the Alamar Blue solution is added, followed by another incubation of the plate for 24 hours. A fluorimeter (FluoStar Optima) is used to read the fluorescence of the plate at the wavelengths of 590 nm for emission and 530 nm for excitation and the data were analysed using the GraphPad Prism 5 software package, fitting the fluorescence to a sigmoid curve with variable slope to determine the EC₅₀ value.

**RESULTS AND DISCUSSION**

In the present study, we demonstrated the effect of natural compounds on the growth of promastigotes of L. major, L. mexicana and different strains of Trypanosoma brucei brucei in vitro. The antitrypanosomal and leishmanicidal activities of natural compounds are summarized in Table 1.

**Apigenin and daidzein**

Apigenin belongs to the flavone subclass of flavonoids (Kim, 2003) and is abundant in a variety of fruits, vegetables, and herbs. It frequently exists in food sources as a glycoside, which improves its solubility and bioavailability (Ross et al., 2002). An array of biological activities has been reported for apigenin, including the ability to inhibit proliferation and induce apoptosis in several cancer cell lines, as well as a capacity to inhibit angiogenesis (Zhou et al., 2017). Daidzein is a naturally occurring isoflavonic phytoestrogen, belonging to the non-steroidal estrogens, and is mainly derived from leguminous plants such as soybean and mung bean. Traditional Chinese medicine Gegen contains Daidzein as its main bioactive component is used often in the treatment of fever, acute dysentery, diarrhoea, diabetes, cardiac dysfunctions, liver injury etc (Knight et al., 1996).

The apigenin dimer ament of lavone, the only bioflavonoid investigated here, also showed some antileishmanial effect (IC₅₀, 6.0 µg/mL) (Tasdemir et al., 2006). In the current evaluation two different types of T. b. brucei were used including wild type (WT) and the highly drug resistant T. b. brucei clonal line B48. Apigenin and daidzein showed highly similar effects on both strains, as presented in Table 1. EC₅₀ values for these compounds were found between 13.4 to 16.9 µM and the results indicate that the TbAT1/P2 and HAPT1 drug transporters are not utilized by either compound. Considering the virtually identical anti-trypanosomal effects, the position of the 4-hydroxybenzene moiety on the main pharmacophore did not seem to be critical.

On the other hand, only apigenin showed some effect on Leishmania major promastigotes and neither compound was active against L. mexicana promastigotes at concentrations up to 100 µM.

**Phyllanthin**

Phyllanthin, found in many Phyllanthus species, has various biochemical and pharmacological properties, especially hepatoprotective effects. Indeed, we have previously reported that among the bioactive
constituents of *Phyllanthus amarus* Schum and Thonn (Euphorbiaceae), phyllanthin has been well examined for its biochemical and pharmacological properties, and particularly for its protective effects on the liver (Ilangkovan *et al.*, 2016). Phyllanthin has also been much investigated for its antitumor effects on various cancer cell lines (Krithika *et al.*, 2011). In the present research, phyllanthin was also examined for effects on *T. b. brucei* and *Leishmania mexicana*. Results are shown in Table 1 and it was observed that this natural compound has a similar if moderate effect on both strains of *T. b. brucei*, showing that phyllanthin is also not using the specified drug transporters and that compounds of this class would not display cross-resistance with diamidines and melaminophenyl arsenical drugs.

**Tyramine**

Tyramine is a substance generally found in nature, as it can be formed from the amino acid tyrosine by the action of the ubiquitous enzyme aromatic amino acid decarboxylase. Tyramine is a vasoactive amine that promotes blood pressure elevation, resulting in pain. Tyramine leads to cerebral vasoconstriction and subsequent rebound vasodilatation that causes a migraine attack in susceptible persons (Costa *et al.*, 2003). This compound was not found to be active against *Trypanosoma* and *Leishmania* strains and showed >100 µM EC50.

**CONCLUSION**

In this study, some new natural compounds were selected for determination of their activity against various *Trypanosoma brucei* strains and *Leishmania* species including multidrug resistant *T. brucei* clonal line B48. The results obtained herein revealed that the two related natural compounds, apigenin and daidzein displayed moderate activity against African trypanosomes and are not cross-resistant with current drugs for this infection. A systematic investigation of this scaffold is likely to result in the identification of compounds with substantially improved activity against kinetoplastid parasites, particularly the African trypanosomes, which are not only the etiological agents of human sleeping sickness but also of the livestock disease nagana, which causes agricultural losses in Africa that measure in the billions of dollar (Giordani *et al.*, 2016). For both conditions, new drugs are urgently needed.

**Conflicts of Interest**

The authors declare no conflict of interests.

**REFERENCES**


