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Article

An RNA Interference (RNAi) Toolkit and Its Utility for Functional Genetic Analysis of *Leishmania* (Viannia)

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Abstract: RNA interference (RNAi) is a powerful tool whose efficacy against a broad range of targets enables functional genetic tests individually or systematically. However, the RNAi pathway has been lost in evolution by a variety of eukaryotes including most *Leishmania* sp. RNAi was retained in species of the *Leishmania* subgenus *Viannia*, and here we describe the development, optimization, and application of RNAi tools to the study of *L.* (*Viannia*) *braziliensis* (*Lbr*). We developed vectors facilitating generation of long-hairpin or “stem-loop” (StL) RNAi knockdown constructs, using Gateway™ site-specific recombinase technology. A survey of applications of RNAi in *L. braziliensis* included genes interspersed within multigene tandem arrays such as *quinonoid dihydropteridine reductase* (*QDPR*), a potential target or modulator of antifolate sensitivity. Other tests include genes involved in cell differentiation and amastigote proliferation (*A600*), and essential genes of the intraflagellar transport (IFT) pathway. We tested a range of stem lengths targeting the *L. braziliensis* hypoxanthine-guanine phosphoribosyl transferase (HGPRT) and reporter firefly luciferase (LUC) genes and found that the efficacy of RNAi increased with stem length, and fell off greatly below about 128 nt. We used the StL length dependency to establish a useful ‘hypomorphic’ approach not possible with other gene ablation strategies, with shorter *IFT140* stems yielding viable cells with compromised flagellar morphology. We showed that co-selection for RNAi against adenine phosphoryl transferase (*APRT1*) using 4-aminopyrazolpyrimidine (APP) could increase the efficacy of RNAi against reporter constructs, a finding that may facilitate improvements in future work. Thus, for many genes, RNAi provides a useful tool for studying *Leishmania* gene function with some unique advantages.

Keywords: trypanosomatid protozoan parasite; *Leishmania braziliensis*; *Leishmania guyanensis*; virulence; gene knockdowns; site specific recombinase; *quinonoid* dihydropteridine reductase (QDPR); intraflagellar transport; hypomorphic mutations

1. Introduction

More than 1.7 billion people are at risk for the ‘neglected tropical disease’ leishmaniasis, with nearly 12 million exhibiting symptomatic disease and more than 50,000 deaths annually, and upwards of 100 million harboring asymptomatic infections [1–5]. *Leishmania* sp. have two distinct growth stages, the promastigote in the sand fly vector, and the intracellular amastigote residing within cellular endocytic pathways in the mammalian host. While the promastigote stage is readily cultured in the laboratory and amenable to molecular techniques, amastigotes require the use of macrophage infection systems or infections of animal models replicating key aspects of human disease.
Different species of *Leishmania* tend to associate with different clinical presentations ranging from localized mild cutaneous disease to more severe visceral or mucosal disease, with host factors also playing key roles [3,6]. *Leishmania* classified within the subgenus *Viannia* are widespread parasites of mammals within South and Central America [7,8]. *L. Viannia* sp. represent one of earliest diverging groups of *Leishmania*, with numerous differences from later-diverging subgenera including development within the insect hindgut, retention of the RNA interference (RNAi) pathway, and often the presence of RNA viruses [7,9,10]. Most importantly this extends to host responses and pathology, especially mucocutaneous disease presentations which most commonly arises from *L. braziliensis* (*Lbr*) infections and are uncommon in infections by species outside of *Viannia* [11].

In this work we focus on experimental applications of the RNAi pathway, an evolutionarily conserved post-transcriptional gene silencing mechanism in eukaryotes [12]. *Trypanosoma brucei*, a kinetoplastid parasite closely related to *Leishmania*, was the first trypanosomatid found to have a functional RNAi pathway [13]. As in other organisms, RNAi quickly became a key tool for diverse functional genetic analysis, ranging from individual to genome-wide applications [14–16]. In contrast, early studies showed that most species of the related trypanosomatid parasite *Leishmania* lack this pathway, in common with an evolutionarily widely separated group of other eukaryotic microbes [17,18]. These observations have raised many questions about the forces operating on microbes to retain or lose this otherwise universally conserved eukaryotic pathway, such as retention or loss of RNA viruses or retrotransposons [10,19].

While initially disappointing for application towards many *Leishmania* sp., phylogenetic mapping showed that RNAi had been retained in the lineage leading to the subgenus *Viannia*, before its loss in the lineages leading to the remaining species of ‘higher’ *Leishmania* such as *L. major*, *L. donovani* or *L. mexicana* [10]. The discovery of an active RNAi pathway in *Viannia* species raised the possibility that RNAi could be used as useful tool for genetic manipulation, as in other eukaryotes. In *Leishmania* classic gene knockouts by homologous recombination work efficiently, but are challenged by the need to disrupt at least two alleles, and/or the presence of multi-copy genes (although this is rapidly evolving with the implementation of CRISPR/Cas9 based tools [20]). The potential utility of RNAi in *L. braziliensis* emerged in several studies showing the impact of RNAi on cellular protein or RNA levels [10,21].

In this study we provide further evidence for the utility of the RNAi in *L. braziliensis*, surveying its activity against a spectrum of gene targets relevant to *Leishmania* biology or chemotherapy, such as flagellar, stage specific or metabolic genes. To accelerate these studies, we applied vectors facilitating the one step production of the dsRNA trigger hairpin-generating “stem-loop” (StL) constructs, using Gateway™ technology in a single step. A similar approach was described previously in African trypanosomes [22]. We explored several parameters relevant to the efficacy of RNAi, which in turn informed methods to generate hypomorphic loss of function mutations of otherwise lethal knockdowns, and co-selections for elevated RNAi efficacy by negative selection. As the great majority of *Leishmania* genes are shared across all species [23], RNAi performed within *Viannia* sp. will likely inform studies of *Leishmania* sp. more broadly outside of this subgenus.

2. Materials and Methods

2.1. Leishmania Strains and Parasite Culture

*Leishmania braziliensis* (*Lbr*) M2903 (MHOM/BR/75/M2903) was obtained from D. McMathon-Pratt (Yale School of Public Health) and grown as promastigotes in Schneider’s Insect Medium (Sigma-Aldrich, St. Louis MO USA; cat. No. S9895) supplemented with 10% heat-inactivated fetal bovine serum (FBS), 2 mM L-glutamine, 500 units mL⁻¹ penicillin and 50 µg mL⁻¹ streptomycin (Gibco No. 5070). *L. braziliensis* M2903 SA (MHOM/BR/75/M2903) was obtained from S.C. Alfieri (Universidade de São Paulo, Brasil). The SA line had been adapted for growth and was serially maintained as amastigotes at 34 °C in modified UM-54 medium [24].
2.2. Leishmania Transfection

Stable transfection of *L. braziliensis* M2903 strain (MHOM/BR/75/M2903) was performed using the high-voltage procedure [25]. Parasites were grown to mid-log phase, pelleted at 1300 g, washed once with cytomix electroporation buffer (120 mM KCl, 0.15 mM CaCl$_2$, 10 mM K$_2$HPO$_4$, 25 mM HEPESS-KOH, pH 7.6, 2 mM EDTA and 5 mM MgCl$_2$) and resuspended in cytomix at a final concentration of 1 × 10$^8$ cells mL$^{-1}$. For transfection, 10 µg DNA was mixed with 500 µL of cells and electroporated twice in a 0.4 cm gap cuvette at 25 µF, 1400 V (3.75 kV cm$^{-1}$), waiting 10 sec between electroporations. Cells were then incubated at 26 °C for 24 hours in drug-free media and then plated on semisolid media containing the appropriate drug to select clonal lines. For selections using blasticidin deaminase (BSD gene), hygromycin phosphotransferase (HYG gene), streptothricin acetyltransferase (SAT gene) and the bleomycin-binding protein from *Streptoalloteichus hindustanus* (PHLEO gene) markers, parasites were plated on 10–20 µg mL$^{-1}$ blasticidin, 30–80 µg mL$^{-1}$ hygromycin B, 50–100 µg mL$^{-1}$ nourseothricin and 0.2–2 µg mL$^{-1}$ phleomycin, respectively (ranges reflect differences when using drugs singly or in combination). Colonies normally appeared by 14 days, at which point they were recovered, grown to stationary phase in 1 mL and passaged with the appropriate drugs. Plating efficiencies range from 60% to 95% for the un-transfected *L. braziliensis* M2903 strain; transfection efficiencies varied from 2 to 50 colonies per µg of SwaI-digested pIR vector controls.

*Lbr* expressing luciferase were generated by electroporation of SwaI-digested pIR1PHLEO-GFP65*-LUC (B6034; Supplementary Table S1) as described previously [10]. These parasites were then electroporated with SwaI-cut pIR2SAT-LUC-StL(A)-APRT-StL(b) (B6391; Supplementary Table S1) followed by selection with both phleomycin and nourseothricin. The presence of both constructs was confirmed by PCR tests. The LUC stem in this configuration is 508 nt [10].

2.3. Western-Blot Analysis

*Leishmania* promastigotes were collected and resuspended in phosphate-buffered saline (PBS) at 1 × 10$^8$ cells/mL. Cell extracts were prepared and Western blots performed after separation by SDS-polyacrylamide gel electrophoresis as previously described [26]. For HGPRT, primary antibodies were anti-*L. donovani* HGPRT and APRT antiserum [27] was used at a titer of 1:5000 and 1:1000, respectively. For normalization anti-*L. major* H2A [28] was used at a titer of 1:100,000, with goat anti-rabbit IgG as the secondary antibody (1:20,000, Licor Inc., Lincoln, NE USA.). Similar procedures were used for QDPR Western blot analysis with extracts from 1.6 × 10$^7$ cells. Gels were transferred to nitrocellulose membranes, which were blocked with a 5% skim milk solution and incubated with 1:500 dilution of rat anti-QDPR [29], a 1:1000 dilution of rabbit anti-PTR1 [30] or anti-*L. major* H2A as described above. IRDye™ anti-rat or rabbit goat immune globulin G were used as the secondary antisera at 1:10,000 dilution. Antibody binding to blots was detected and quantified using an Odyssey infra-red imaging system (Li-Cor).

2.4. Quininoid Dihydropteridine Reductase Assay

Parasites were harvested at log phase (4–6 × 10$^6$ cells/mL) and collected by centrifugation at 1250 × g for 10 min at 26 °C, washed twice with PBS, and resuspended at 2 × 10$^9$ cells mL$^{-1}$ in 10 mL of Tris-Cl, pH 7.0, with 1 mM EDTA and a mixture of protease inhibitors as described [31]. Cells were lysed by three rounds of freeze thawing and sonication, and the extracts clarified by centrifugation at 15,000 × g for 30 min at 4 °C. Protein concentrations were determined using Qubit Fluorometric Quantification (Invitrogen). *Quininoid* dihydropteridine reductase activity was measured at 25 °C as described [32] using *quinonoid* dihydrobiopterin generated continuously by horseradish peroxidase mediated oxidation of H$_4$-bioprotein. The standard reaction mixture contained 50 mM Tris-HCl, pH 7.2, 20 µg of horseradish peroxidase, 0.1 mM H$_2$O$_2$, 20 µM of H$_4$-bioprotein, 100 µM NADH, and purified QDPR or parasite lysates; all components were incubated for 3 min prior to initiation of reaction by addition of H$_4$B. The activity was measured by monitoring
NADH consumption at 340 nm ($\varepsilon_{340}$ for NADH is 6200 M$^{-1}$ cm$^{-1}$) in a Beckman DU-640 spectrophotometer.

2.5. Luciferase Assay

Logarithmic growth phase promastigotes (10$^6$) were suspended in 200 µL media containing 30 µg/ mL of luciferin (Biosynth AG, Staad, Switzerland) and added to a 96-well plate (Black plate, Corning Incorporated, NY, USA). The plate was imaged after 10 min using a Xenogen IVIS photomizer (Caliper LifeSciences), and luciferase activity quantitated as photons/sec (p/s).

2.6. Transmission Electron Microscopy

*Leishmania* promastigotes were harvested in logarithmic growth phase and fixed in 2% paraformaldehyde/2.5% glutaraldehyde (Polysciences Inc., Warrington, PA, USA) in 100 mM phosphate buffer, pH 7.2, for 1 h at room temperature. Samples were washed in phosphate buffer and postfixed in 1% osmium tetroxide (Polysciences Inc., Warrington, PA, USA) for 1 h, rinsed extensively in water, and block stained with 1% aqueous uranyl acetate (Ted Pella Inc., Redding, CA, USA) for 1 h. Following several rinses in water, samples were dehydrated in a graded series of ethanol solutions and embedded in Eponate 12 resin (Ted Pella Inc.). Sections of 95 nm were cut with a Leica Ultracut UCT ultramicrotome (Leica Microsystems Inc., Bannockburn, IL, USA), stained with uranyl acetate and lead citrate, and viewed on a JEOL 1200 EX transmission electron microscope (JEOL USA Inc., Peabody, MA, USA).

2.7. Construction of the Integrating pIR-GW Destination Vectors Facilitating Generation of StL Constructs Using Gateway Site Specific Recombinase

We first assembled a construct bearing a PEX11-MYC ‘loop’ fragment flanked by inverted Gateway® cassettes; these contain ccdB, a lethal gene that targets DNA gyrase, and CmR encoding chloramphenicol-resistance, flanked by two attR sequences necessary for site-specific recombination (attR1-ccdB-CmR-attR2; www.Invitrogen.com (accessed on 26 December 2022). For propagation, all constructs bearing the Gateway cassettes were propagated in *Escherichia coli* (E. coli) DB3.1 which contains a gyrA462 mutation conferring resistance to ccdB toxicity. First, the PEX11-MYC loop was excised from pIR1 SAT-GFP65-StL (B4733) [7,9,10] and inserted into pGEMT yielding pGEMT-stuffer (B5974). The Gateway cassette (attR1-ccdB-CmR-attR2) was amplified from the pDONR221 (Invitrogen) with primers S1 and S2, and inserted by blunt end ligation into the NheI site of pGEMT-stuffer (B5974), yielding pGEMT-stuffer -one GW (B6158). The second GW cassette was inserted by blunt end ligation into the pGEMT-stuffer -one GW AvrII site, in inverted orientation to the first cassette (with both in divergent orientation relative to the ccdB/CmR ORFs), yielding (B6218).

Plasmid pGEMT-stuffer—2 GW divergent contains a 3877 bp Sphi/HindIII fragment bearing the inverted Gateway cassettes and loop. This fragment was blunt-end ligated into the Smal site of pIR1SAT (B3451), yielding pIR1SAT-GW (B6223) (Supplemental Table S1). Similarly, the inverted Gateway/loop fragment inserted into the BglII (B) site of various pIR vectors by blunt end ligation, yielding the final vectors pIR1SAT-GW (B6223), pIRHYG-GW (B6544), pIR1PAC-GW (B6543), pIR1BSD-GW (B6542) pIR2HYG-GW (B6563) (Supplemental Table S1). The sequence of pIRHYG-GW is provided in Supplemental File S1.

2.8. Generation of Target Stem-Loop (StL) Constructs for RNAi

The molecular constructs used in this work and their synthesis are summarized in Table S2. Briefly, for StL constructs the ‘stem’ was obtained by PCR, inserted into the pCR8/GW/TOPO vector by TA cloning, (Invitrogen # K250020), and their orientation (same direction as attL2) confirmed by sequencing and restriction digestion. The stems were transferred from the pCR8/GW/TOPO donor vector to the pIR1-GW or pIR2-GW destination vectors described above; these contain sequences from the parasite small
subunit rRNA locus to enable integration into the genome, and inverted attR1-ccdB-CmR-attR2 cassettes. The gene of interest in pCR8/GW/TOPO was transferred to the pIR-GW vector with LR Clonase II (Thermo Fisher, Waltham MA USA) in an overnight reaction at room temperature. Reactions were terminated by incubating with proteinase K for 1 h at 37 °C (Gateway Technology with Clonase™ II manufacturer' protocol, Version A: 24 June 2004). Reactions were transformed into E. coli TOP10™ or DH5α (which select against both ccdB cassettes). All final stem-loop (StL) constructs named as StL expressers were confirmed by restriction enzyme digestion and DNA sequencing (Figure 1). Prior to transfection, constructs were digested with SwaI to expose the SSU rRNA segments mediating homologous integration.

**Figure 1.** Flowchart for generation of entry and StL constructs using Gateway site-specific recombinase technology followed by introduction into *Leishmania* to express dsRNA. A stem for each GOI (gene of interest) was amplified by PCR and then cloned into pCR8/GW/TOPO vector (Invitrogen) to generate an “entry” clone bearing a flanking attL1 and attL2 sites for recombination-based transfer. The pIR-GW destination vectors bear two opposing ccdB/CmR cassettes, each flanked by attR1 and attR2 sites. Following site-specific recombination by the Gateway LR reaction, the desired StL for each GOI was obtained and confirmed. For biological tests, each StL DNA was linearized with SwaI, electroporated into *Leishmania*, where it integrated into the small subunit ribosomal RNA locus (SSU). There it is transcribed by the strong rRNA pol I promoter to generate RNAs whose dsRNA regions are processed by the RNAi machinery.

### 3. Results

#### 3.1. Rapid Generation of ‘Stem-Loop’ Constructs as RNAi Triggers

To trigger the RNAi response, dsRNA generating ‘stem-loop’ constructs have been used in previous studies, often assembled in three steps with two ‘stem’ segments cloned in opposite orientations, separated by a short spacer/loop [10]. To accelerate this process, we utilized Gateway™ (Invitrogen) technology, incorporating precise, site-specific recombinase technology.
tion [22,33]. First we engineered a ‘destination’ expression vector, based on the Leishmania pIR vector series which achieves high levels of RNA expression following integration into the ribosomal small subunit RNA locus [10,34]. The final destination vector (pIR-GW generically) bears two Gateway recipient cassettes, arranged in inverted orientation and separated by a short segment destined to become the ‘loop’ inserted into one of the strong expression sites of pIR vectors (Figure 1). Each Gateway cassette contained a positive CmR and negative ccdB marker, flanked by attR1 and attR2 sites. Then, each stem to be tested was inserted into an ‘entry’ vector (pCR8/GW/TOPO), where it was flanked by donor attL1 and attL2 sites (Figure 1). Gateway LR reactions between the destination vector and entry DNAs were performed in vitro, and transformed into E.coli TOP10 or DH5α™ which selected against the presence of the target ccdB/CmR cassettes, which could be rapidly confirmed by loss of chloramphenicol resistance. Typically we obtained numerous recombinants of which more than 70% bore the expected configuration of the desired StL configuration, and one correct representative was selected for introduction into Leishmania. Prior to transfection, constructs were digested with SvaI to expose the SSU rRNA termini to direct integration into the SSU rRNA locus, where strong expression is driven from the rRNA promoter (Figure 1).

3.2. Testing the Effect of Stem Length on RNAi Activity Using L. braziliensis HGPRT-StL and Luciferase-StL Series Constructs

In T. brucei, stem lengths ranging in length between 100 and 1500 bp are effective in knocking down target genes [35,36], and stem lengths as short as 29–100 nt are active in other metazoan species [37,38]. To explore this in Leishmania, we varied the stem length in RNAi constructs targeting an integrated firefly luciferase reporter gene (LUC) as well as an endogenous cellular gene (HGPRT), assayed following transfection into WT Lbr. For HGPRT increasing the stem from 494 nt to 1005 nt reduced expression from 55% to 92% (Figure 2A). No strong differences were seen between similarly sized stems targeting the HGPRT ORF or 3′ untranslated region (Figure 2A).

For luciferase, we tested LUC StL constructs following introduction into a LbrM2903 transfectant stably expressing high levels of luciferase activity (Figure 2B). As with HGPRT, the longer luciferase stems resulted in greater reductions in LUC activity (Figure 2B). While stems of 158 bp or greater showed strong reductions (>93%), stems of 128 or 54 nt showed much smaller effects (35–32%; Figure 2B). Preliminary analysis did not suggest a strong association of the activity of the stems tested with siRNA abundance ‘hot spots’, small regions where siRNA levels greatly exceed the average across a gene, as seen with LUC or Leishmania Virus 1 (LRV1) in our prior study [19]. We did not observe synergy when two weak constructs were transfected simultaneously (not shown).

These data together with those shown below for IFT140 (Section 3.6) suggested that for the strongest effect stem lengths of >500 nt were preferable, with a significant drop-off below 128 nt. We did not see strong ‘positional’ effects of the location of the stems within the target gene, however the constructs tested tended to be progressively truncated from one side so it is possible these may have been overlooked. Importantly, our findings suggest differences in the dsRNA trigger length dependency in Leishmania relative to trypanosomes or other organisms. Further studies will be required to explore the basis for this.
Figure 2. The effect of stem length on RNAi activity. (A) Targeting the endogenous HGPRT. Various stem length HGPRT StL constructs were transfected into *L. braziliensis*, and HGPRT expression assessed in two colonies by Western blot with anti-HGPRT antisera and anti-H2A antisera for normalization (Supplementary Figure S1). The percent expression relative to WT *Lbr* is shown. (B) Targeting an introduced luciferase reporter. Various stem length LUC StL constructs were transfected into an *Lbr* line expressing LUC (Section 2.2). Luciferase activity was measured in triplicate and the percent reduction in activity calculated relative to the parental *Lbr*-LUC-expressing line.

3.3. StL-Mediated Specific RNAi of the Metabolic Target Quinonoid Dihydropteridine Reductase (QDPR) Interspersed within a Tandem Repeated Gene Array

We examined the efficacy of StL RNAi against the *L. braziliensis* *quinonoid* dihydropteridine reductase (QDPR) gene, encoding an enzymatic step of the reduced pteridine pathways involved in recycling oxidized $qH_2$-biopertin or $qH_2$-folates back to the active tetrahydro state in *Leishmania* [29]. In all *Leishmania*, QDPR genes are interspersed in a tandem array with two other genes, ORFq (q: a hypothetical protein) and $\beta_7$-proteasome ($\beta_7$: 20S proteasome $\beta_7$ subunit), the latter gene being essential where tested in other species including trypanosomes (Figure 3A; [29]). This rendered selective deletion of the interspersed QDPR targets considerably more challenging, but provided a suitable opportunity for testing the utility of RNAi in this context.

We introduced a StL construct bearing a 588 nt QDPR stem into *L. braziliensis* parasites successfully. Relative to WT, QDPR protein was reduced more than 87.4% (Figure 3B) while QDPR activity was reduced by more than 88% individually in StL knockdowns (Figure 3C). The *Lbr* QDPR StL knockdowns grew normally in culture, suggesting that RNAi specifically targeted QDPR without significantly impacting the flanking essential proteasome subunit, although a partial reduction in expression cannot be ruled out. Thus RNAi can be used to successful probe the consequences of metabolic gene expression depletion in this most challenging context.

One functional consequence of QDPR ablation was shown by the increased sensitivity of the QDPR StL knockdown to the antifolate compound TQD (5, 6, 7, 8-Tetrahydro- 2, 4-quinazolinediamine) (Figure 3D) [39]. The QDPR knockdown parasite was 20-fold more sensitive, with an EC50 of 0.15 ± 0.015 vs. 3.49 ± 0.59 $\mu$M for WT ($p < 0.0006$). These data support the utility of RNAi knockdowns in *Lbr* as probes of pteridine metabolism and drug action.
Figure 3. RNAi of the tandemly dispersed gene QDPR. (A) QDPR gene organization in L. braziliensis. QDPR genes are tandemly repeated with ORFq (Q) and β7 proteasome (β7) up to 9 times in L. braziliensis genomes; only the ORFs are depicted, and the entire QDPR ORF was used as the tested stem. Proteasome subunits are known to be essential, while the requirement for ORFq has not been tested. (B) QDPR protein expression in QDPR-StL knockdowns. QDPR was detected with anti-QDPR antiserum and expression was calculated relative to that of H2A, from Western blots as described in the methods. (C) QDPR enzymatic activity in QDPR-StL knockdowns. QDPR enzymatic activity was assayed and normalized to total cellular protein. Data were from three independent experiments each performed in triplicate. * indicates \( p < 0.0235 \). (D) Growth inhibition by inhibitor TQD. WT and QDPR-StL transfected Lbr were inoculated into media containing 0–50 \( \mu \text{M} \) 5, 6, 7, 8-Tetrahydro-2, 4-quinazolinediamine (TQD) at 2 \( \times \) 10^5 cells/mL and allowed to grow until WT had reached late log phase, at which time parasite numbers were determined. The experiment was repeated three times, with results similar to that shown; the EC50s for the QDPR-StL (red) and WT (blue) lines were 0.15 ± 0.015 vs. 3.49 ± 0.59 \( \mu \text{M} \) (\( p < 0.0006 \)).

3.4. RNAi of an L. braziliensis Gene Important for Amastigote Replication

An important area of Leishmania biology is the study of genes that impact survival of the amastigote stage in the vertebrate host. Many (not all) Leishmania species can differentiate to amastigote-like forms in culture (axenic amastigotes), when grown at conditions resembling those within the parasitophorous vacuole, e.g., elevated temperature and low pH [40,41]. Here, we made use of a clonal derivative of Lbr M2903 (LbrM2903SA2) which had been adapted for growth as axenic amastigotes [24].

As a test we focused on the A600 locus, which in L. mexicana comprises four genes whose deletion had little impact on promastigotes, but precluded axenic amastigote replication in vitro [42]. The A600 copy number varies amongst species [42], with only two found in Lbr (A600-1 and A600-4) which show 88% nucleotide identity. We targeted these sequences simultaneously by a single StL construct using the A600-1 sequence, which shows long stretches of identity relative to A600-4 (118 nt, 51 nt and 45 nt).

We transfected the LbrA600-StL construct into Lbr SA2 promastigotes, obtaining many clonal transfectants all of which grew normally. Several were then inoculated into axenic amastigote growth medium, where they showed a severe growth defect, with 10-fold fewer amastigotes than WT (Figure 4), similar to the results obtained with L. mexicana A600 knockouts. These results establish the utility of RNAi knockdowns for the study L. braziliensis genes involved in amastigote differentiation and proliferation.
3.5. Targeting Essential Genes of the L. braziliensis Intraflagellar Transport (IFT) Pathway

The *Leishmania* flagellum plays key roles in the promastigote and amastigote stages, and previously we reported successful knockdowns of the paraflagellar rod proteins PFR1 and PFR2 which showed phenotypes comparable to complete deletions [10]. We extended these studies to a set of genes associated with intraflagellar transport (IFT), which mediates transport of cargo in both anterograde and retrograde directions and is required for proper flagellar assembly [43].

StL constructs targeting *Lbr IFT122, IFT140, IFT172*, representing both the anterograde and retrograde IFT, were transfected into WT *Lbr* (Table 1). However, we were unable to recover transfected colonies, despite multiple attempts and success targeting a nonessential gene, *LbrPFR1-StL* (Table 1). To establish that this result was dependent on the RNAi pathway, we introduced these same constructs into an RNAi-deficient mutant obtained by homozygous deletion of the *AGO1* gene (*Δago1*−), an Argonaute family protein encoding the key ‘slicer’ activity required for RNAi activity in *Lbr* [10,44]. Now, all *LbrIFT* StL constructs successfully yielded transfectants (Table 1), establishing that RNAi of the StL-derived dsRNA trigger was responsible for the lack of transfectants.

Table 1. RNAi of several IFT genes in *Leishmania braziliensis* is lethal.

<table>
<thead>
<tr>
<th>STL Construct</th>
<th>No. Transfectants with WT</th>
<th>No. Transfectants with Δ<em>ago1</em>−</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>PFR1-StL</em></td>
<td>243–360</td>
<td>360</td>
</tr>
<tr>
<td><em>IFT122-StL</em> (retrograde)</td>
<td>0</td>
<td>460</td>
</tr>
<tr>
<td><em>IFT140-StL</em> (retrograde)</td>
<td>0</td>
<td>265</td>
</tr>
<tr>
<td><em>IFT172-StL</em> (anterograde)</td>
<td>0</td>
<td>289</td>
</tr>
<tr>
<td>Negative control</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

These data are consistent with other studies in trypanosomes and *Leishmania* showing that IFT gene ablation displays an array of phenotypes, including essentiality [45–48].

The table shows the number of colonies (per plate) obtained after transfection of various StL constructs into WT or RNAi-deficient Δ*ago1*− *Lbr*. *PFR1-StL* was used as a positive control [10].

![Figure 4. Impact of LbrA600-StL amastigote growth in *L. braziliensis*. WT or A600-StL transfectants grew normally as promastigotes; the figure shows parasites inoculated at a density of 5 × 10⁵ /mL and the cell numbers determined daily for 1 week. WT (■) was repeated three times along with 3 different A600-StL knockdown clonal lines (#1–3; triangles). Across 2 experiments with WT (3 replicas each) and 7 different A600-StL lines the difference was highly significant (*p* < 0.0001).](image-url)
3.6. Systematic Generation of Hypomorphic Mutants by Exploiting the Stem Length-Dependency of RNAi in L. braziliensis

While plus/minus tests of gene essentiality are useful they provide little information about the role of the encoded protein within the cell. When inducible systems are available, observations following shutoff and prior to cell death can be informative, however presently this can only be achieved using conditional degradation domains in *Leishmania* [49]. The studies of the stem length-dependency of the efficacy of RNAi described in Section 3.2 above suggested that it could be possible to engineer partial loss of function mutants through successively reducing the stem length, until a point was reached where viable cells could be obtained, which ideally might show informative defects. We tested this with the *IFT140* gene, which is essential in trypanosomes and probably *Lbr* (Table 1). Curiously, viable deletion mutants with different phenotypes were possible in *L. major* and *L. donovani* [47,48] suggesting there is considerable variation amongst *Leishmania* species for the consequences of IFT ablation.

Transfection of a 941 nt *IFT140* StL construct yielded no transfectants in WT *Lbr* (Table 1), however as the stem length was reduced to 562 nt and further to 131 nt, increasing numbers of transfectants could be recovered (Figure 5A). Notably, the 562 nt stem yielded about 10% as many transfectants, and the cells recovered from these colonies grew slowly in culture (Figure 5A). In contrast, transfectants recovered with constructs with stems shorter than 562 nt were recovered at normal frequencies, grew like WT, and showed no obvious differences in flagellar length or motility.

![Figure 5](image-url)

**Figure 5.** Systematic identification of viable hypomorphic RNAi mutants. (A) Map of the *IFT140* stems tested, number of colonies obtained after transfection, and the growth properties (when viable).
(B) Transmission EM of WT Lbr showing normal kinetoplast (K) and flagellum (F). (C) Transmission EM of hypomorphic LbrIFT140-StL (562 nt stem) cells, where defects in the parasite flagellum are evident (F) along with accumulation of vesicles within the flagellar pocket (P); not shown is that these cells are more rounded.

This suggested that the 562 StL IFT140 transfectant was a candidate hypomorph, as it displayed a shortened flagellum and weak motility (not shown). Transmission EM of one clone showed that these knockdown cells exhibited modified shape and accumulated vesicles in the flagellar pocket similar to the phenotypes in other trypanosomes, such as T. brucei where ablation of IFT140 was initiated by conditional RNAi prior to cell death [45], or in L. mexicana IFT140 knockouts [48] (Figure 5C). These data suggest that the stem-length dependency of RNAi in L. braziliensis will prove a useful tool in generating informative, hypomorphic mutants of otherwise essential genes, facilitating inquiries into their cellular targets and/or mechanism of action.

3.7. A Negative Selection System for Enhancement of RNAi Activity in L. braziliensis

It is often helpful to have screens or selections to monitor the efficacy of RNAi under various circumstances; for example, in screening for genes acting within the RNAi pathway, or when RNAi exhibits considerable clonal variability, as in some fungi [50–52]. GFP and luciferase based screens are commonly used, which both perform well in Leishmania [10], but here we sought a selectable system which could facilitate other applications.

We tested the 4-aminopyrazolopyrimidine/adenine phosphoryl transferase 1 (APP/APRT1) system in L. braziliensis. APRT converts APP into a toxic metabolite inhibiting Leishmania growth, and thus cells with decreasing APRT levels show increasing resistance to APP. Importantly, APRT1 is not essential in most media due to the alternative salvage route through adenosine aminohydrolase [53].

First, we introduced an APRT1-StL construct into WT Lbr, where transfectants were readily obtained and grew normally. Western blot analysis confirmed significant reduction to undetectable level in APRT1 expression (Figure 6A). Unlike WT parasites whose growth was completely inhibited by 500 µM APP, LbrAPRT1-StL transfectants were able to grow at the highest concentration tested, albeit at reduced growth rates (Figure 6B,C). Other studies showed that the APP EC50s were 50–100 µM for WT and 500–1000 µM for APRT1-StL (not shown). Thus, RNAi knockdown of APRT1 leads to APP resistance as expected.

We developed a dual stem-loop reporter parasite that simultaneously expresses an APRT1-StL along with a LUC-StL with a 508 nt stem. This was obtained by transfection an Lbr line expressing luciferase with a second construct bearing both APRT1-StL and LUC-StL cassettes (Section 2.2). This reporter line allows manipulations of overall cellular RNAi activity to be tracked through simple luciferase assays. We chose a mid-sized stem as we wanted to provide some range for detection of elevated RNAi activity before saturation.

To illustrate its utility, APRT1-StL+LUC-StL/LUC parasites were plated on increasing concentrations of APP, which would select for low APRT1 expression which could potentially arise through alterations in overall cellular RNAi activity. Colonies were picked and parasites were grown thereafter in 50% of the selective concentration of APP used in plating.

In the absence of APP selection, luciferase expression was reduced 122 fold in the APRT1-StL+LUC-StL/LUC parasites (Figure 7), slightly less than the 200–300 fold reduction seen previously [10] but greater than the 30–50 fold seen in Figure 2B with similarly sized stems. These experimental series were performed months or even years apart, and we suspect small differences in vector design or clonal variation may be a contributing factor. Since each series of experiments was internally controlled, we consider these differences of negligible significance in this context.

In the APRT1-StL+LUC-StL/LUC parasites subjected to APP selection, luciferase activity decreased an additional 2.9 to 6.8 fold, as the APP concentrations were increased from 125 to 1000 µM (Figure 7). Overall relative to WT parasites, the efficacy of LUC silencing increased from 122- to nearly 830-fold. Thus, in one step we were readily able to identify
parasite showing elevated levels of RNAi activity against a luciferase reporter. This proof of principle experiment establishes a useful tool for both monitoring and manipulating RNAi activity amongst clones or mutants that may prove of great utility in future studies.

Figure 6. RNAi of the APRT1 gene yields increased APP resistance. (A) Transfection of APRT1-StL leads to loss of APRT1 expression. The figure shows a Western blot of promastigotes tested against anti-APRT1 antisera. Lane 1. L. braziliensis WT; lane 2, L. donovani WT; lanes 3–7, Lbr APRT1-StL clones 1 to 5. (B,C) Growth inhibition by APP in the indicated concentrations is shown. (B): WT Lbr; (C): Lbr APRT1-StL knockdown clone 1. The APRT1-StL line is significantly more resistant to APP than WT at the concentrations shown (n = 4; p < 0.0001).
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Figure 7. A negative selectable system to identify enhanced RNAi activity in L. braziliensis. Double StLs (LUC and APRT1) expressed in a single construct were transfected in the Lbr LUC reporter line and selected on M199 plates containing various concentrations of APP. Colonies were readily obtained and grown in APP, and luciferase activity was measured from 10^6 parasites as described in the methods. The fold reduction of colonies selected in varying concentrations of APP relative to no drug are shown in brackets above the plotted luciferase activity. The average and standard deviations calculated from at least 10 transfectant colonies is shown. *** indicates a \( p < 0.001 \) for comparisons with no APP selection.

4. Discussion

RNA interference has proven a valuable tool for the study of gene regulation in many eukaryotes including African trypanosomes. While lost in many Leishmania sp., those of the subgenus Viannia retained a functional pathway, opening up its use as a tool for genetic analysis. In this work we describe a useful Gateway™ site specific recombination based system for rapidly and efficiently generating stem-loop constructs suitable for RNAi tests, and applied it towards a spectrum of Leishmania genes of interest to illustrate its range and potential. For perspective, Table 2 summarizes the genes and the outcomes obtained in this or previous studies.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Gene ID</th>
<th>Results</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>LbrAGO1</td>
<td>LbrM.11.0360</td>
<td>Reduced RNAi activity</td>
<td>[10]</td>
</tr>
<tr>
<td>LbrLPG1</td>
<td>LbrM.25.0010</td>
<td>Little mRNA change; remains LPG+</td>
<td>[10]</td>
</tr>
<tr>
<td>LbrLPG2</td>
<td>LbrM.20.2700</td>
<td>3-fold lower mRNA; remains LPG+</td>
<td>[10]</td>
</tr>
<tr>
<td>LbrLPG3</td>
<td>LbrM.29.0780</td>
<td>3-fold lower mRNA; remains LPG+</td>
<td>[10]</td>
</tr>
<tr>
<td>LbrHGPRT</td>
<td>LbrM.21.0990</td>
<td>Reduced protein</td>
<td>This work</td>
</tr>
<tr>
<td>LbrPFR1</td>
<td>LbrM.31.0160</td>
<td>Reduced mRNA; abnormal swimming</td>
<td>[10]</td>
</tr>
<tr>
<td>LbrPFR2</td>
<td>LbrM.16.1480</td>
<td>Reduced mRNA; abnormal swimming</td>
<td>[10]</td>
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Table 2. Cont.

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<th>Gene ID</th>
<th>Results</th>
<th>Reference</th>
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<tr>
<td><em>Leishmania</em> RNA virus 1</td>
<td>LgyM4147</td>
<td>LRV1 elimination</td>
<td>[19]</td>
</tr>
<tr>
<td></td>
<td>LbrLEM2700</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>LbrLEM2780</td>
<td></td>
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<td></td>
<td>LbrLEM3874</td>
<td></td>
<td></td>
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<tr>
<td><em>Lbr</em> δ Amastin</td>
<td>Lbr.M20.0160</td>
<td>Reduced mRNA; impaired viability of intracellular amastigotes</td>
<td>[21]</td>
</tr>
<tr>
<td><em>(gene family)</em></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><em>Lgy</em>VTC4</td>
<td>Lgy4147 VTC4</td>
<td>Reduced mRNA; polyphosphate levels decreased; little impact on growth</td>
<td>[54]</td>
</tr>
<tr>
<td><em>Lbr</em>APRT1</td>
<td>LbrM.26.0130</td>
<td>Resistant to APP</td>
<td>This work</td>
</tr>
<tr>
<td><em>Lbr</em>QDPR</td>
<td>LbrM.20.3970</td>
<td>Altered drug sensitivity</td>
<td>This work</td>
</tr>
<tr>
<td><em>Lbr</em>A600</td>
<td>LbrM.20.3230</td>
<td>Growth defect as amastigotes</td>
<td>This work</td>
</tr>
<tr>
<td><em>LbrIFT140</em> (retrograde)</td>
<td>LbrM.32.0380</td>
<td>Unable to recover transfectants of WT Lbr</td>
<td>This work</td>
</tr>
<tr>
<td><em>LbrIFT122</em> (retrograde)</td>
<td>LbrM.35.1120</td>
<td>Unable to recover transfectants from WT Lbr</td>
<td>This work</td>
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<tr>
<td><em>LbrIFT172</em> (anterograde)</td>
<td>LbrM.21.1210</td>
<td>Unable to recover transfectants from WT Lbr</td>
<td>This work</td>
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<td><em>Lbr</em> BBS1, BBS3, BBS4</td>
<td>LbrM.34.4180</td>
<td>No effect on growth or morphology</td>
<td>This work</td>
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<tr>
<td></td>
<td>LbrM.16.1430</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>LbrM.35.2520</td>
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</table>

Of the 22 targets listed, 6 (27%) showed changes in survival, growth and/or morphology when knocked down (PFR1, PFR2, IFT140, IFT122, IFT172, VTC4). When other criteria were evaluated phenotypes were detected in a further 9 (41%; APRT1, A600, QDPR, AGO1, δ-amastin, and 4 LRV1s). In contrast, no phenotypes were detected with the methods applied with 7 target genes (HGPRT1, LPG1, LPG2, LPG3, BBS1, BBS3 or BBS4). Our findings can be compared with the more extensive studies from African trypanosomes, where about 1/3 of genes targeted in chromosomal surveys or genome wide studies show changes in survival, growth or morphology [16,55]. As in trypanosomes, genes involved in the synthesis of abundant surface glycoconjugates such as *Leishmania* lipophosphoglycan LPG1-3 showed little phenotype, while other genes such as those implicated in the parasite flagellum (PFR or IFT) yielded readily detected phenotypes. Importantly phenotypes were obtained in several repetitive gene families, including *Lbr* A600, PFRs, δ-amastins and QDPR, the latter having additional complexity as an ‘interspersed’ repetitive gene family. Similarly we have been able to target cytoplasmic RNA viruses effectively such as the LRV1 totiviruses of *Lbr* and *Lgy*. These data suggest that RNAi offers another strong option for functional genomics in *Leishmania*.

The specificity of RNAi knockdowns can be assessed in several ways. First, the dependency of RNAi on a functional Argonaute (AGO1) can be used to support conclusions about the essentiality of a given RNAi target, as illustrated by studies with several essential IFT genes (Table 1). Secondly, the highly expressed integrated SSU:IR-StL constructs used here typically yield very high levels of siRNAs and occasionally dsRNA, both of which can have off target effects [19]. The specificity for the target gene may be assessed by reintroduction of a ‘recoded’ RNAi-resistant target gene [21], or by selection for rare spontaneous excision of the integrated SSU:IR-StL construct from the rRNA gene array [56], both of which should restore the WT phenotype.

While most studies summarized in Table 2 have been carried out with *L. braziliensis*, RNAi was also effective in *L. guyanensis* (VTC4, LRV1). However, in previous work we showed in both LUC and LRV1 studies that the efficacy of RNAi was significantly less than in *Lbr* [10,19]. The factors responsible have not been studied, and in our previous
studies we noted that even with strong RNAi induction target RNA levels sometimes did not decline [10]. Thus, the 'penetrance' of RNAi efficacy can vary widely amongst specific genes, and/or species or strains, something that should be considered in future studies. Fortunately, the ease by which RNAi constructs may be generated using site-specific recombinase technology (Figure 1) allows this to be explored with minimal effort.

Interestingly, with the selectable APRT1/APP system we were able to increase the strength of RNAi nearly 7 fold in Lbr (Figure 7), suggesting it may be possible in the future to develop lines where the efficacy of RNAi is enhanced. The feasibility of this was shown in mammalian cells where increased expression of Argonaute-2 enhanced RNAi activity [27].

While the introduction of CRISPR/Cas9 technology provides an attractive alternative to RNAi for gene ablation, there are some useful applications of RNAi as well. Inducible RNAi systems are readily reversible, and the stem length dependency shown here offers the possibility of developing 'graded' RNAi responses to yield stable hypomorphic lines. This was illustrated with the Lbr IFT140 gene, where shorter stems led to viable cells, with a key one exhibiting reduced transfection efficiency, slower growth and flagellar defects (Figure 5), allowing further exploration of the impact of RNAi on flagellar biology or cell physiology. While the data provided here serve as a guide, it is likely that the optimal stem length (and possibly position) will have to be evaluated empirically for any given gene and/or phenotype. Nonetheless, the ability to systematically generate stable hypomorphic mutants adds another dimension to the utility of RNAi in *Leishmania*.

### Supplementary Materials

The following supporting information can be downloaded at: [https://www.mdpi.com/article/10.3390/genes14010093/s1](https://www.mdpi.com/article/10.3390/genes14010093/s1), Table S1: Molecular constructs; Table S2: Oligonucleotide primers; Figure S1: Western blot of HGPRT-StL transfectants. This figure shows the Western blot whose quantitation is shown in Figure 2A; File S1: Sequence of the pIR1HYG-GW vector.

### Author Contributions

Study conception: L.-F.L. and S.M.B.; Experimental design, performance, data analysis: L.-F.L., K.L.O., S.J., J.E.M., E.A.B. and S.M.B.; Manuscript preparation: S.M.B. and L.-F.L.; Supervision and funding acquisition: S.M.B. All authors have read and agreed to the published version of the manuscript.

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### Institutional Review Board Statement

Not applicable (no vertebrate animals).

### Informed Consent Statement

Not applicable.

### Data Availability Statement

All data are included in the text or Supplementary Materials.

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### Conflicts of Interest

The authors declare no conflict of interest.

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