# CLONING AND FUNCTIONAL ANALYSIS OF TRANS-SPLICING FACTORS IN Trypanosoma brucei



1.

By

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A research thesis submitted to the Graduate School in partial fulfilment for the requirements of the Master of Science Degree

in Biochemistry of Egerton University.

# EGERTON UNIVERSITY

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### **DECLARATION AND RECOMMENDATION**

I declare that this thesis is my own original work and has not been presented before for the award of any degree.

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# DEDICATION

This work is dedicated to Lord God Almighty and my mother, Anne Awino.

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### ABSTRACT

Trypanosomes are parasitic protozoa that cause trypanosomosis. The disease is a threat to human population and a major impediment to livestock production and economic development in many countries in sub-Sahara Africa, where it is endemic. Presently, it is controlled by reduction of tsetse fly vector population, chemotherapy and chemoprophylaxis. However, none of these approaches is completely effective and thus the need for development of new approaches. The splicing machinery in trypanosomes presents a potential target for anti-trypanosome drug development. The development of new drugs is needed because of increasing incidents of resistance to available drugs. In trypanosomes, the mature messenger RNA is derived from independent pre-mRNA molecules in a process called trans-splicing. This is a variation from the more common cis-splicing which occurs in the mammalian hosts of trypanosomes, where the mature RNA is derived from one premRNA molecule. The process is important in regulation of gene expression in trypanosomes that is predominantly post-transcriptional. In this study, thirteen homologs of Trypanosoma brucei genes for trans-splicing and polyadenylation were identified in silico using Trypanosoma cruzi, yeast and/or human splicing and polyadenylation factors to query GeneDB, the repository of genome data for Trypanosoma brucei, Trypanosoma cruzi and Leishmania major. Degenerate PCR approach was used to clone the factors, which were subsequently sequenced. The amino acid sequences generated were used to query public protein databases and were also compared to homologous sequences from T. cruzi, L. major and Homo sapiens. Conserved RNA binding proteins domains and domains of proteins involved in multi-protein complex assemblies were identified. The kinetoplastid sequences were similar to each other, but were individually significantly different from human homologs. Sequence specific gene silencing of three factors (P14, CPSF 30 and U1-70k) using RNA interference (RNAi) technique was lethal suggesting the importance of the three factors in viability. Protein depletion was detected in the silencing of the P14 gene. Significant variations of the kinetoplastid sequences from human and importance of the factors in parasite viability suggest that some components of the trypanosome spliceosome are targets for the design of novel drugs.

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# LIST OF ABBREVIATIONS

AAT:	African Animal Trypanosomosis					
BCIP:	5-Bromo-4-chloro-3-indolyl phosphate					
BPS:	Branch Point Sequence					
CPSF:	Cleavage and Polyadenylation Splicing Factor					
CstF:	Cleavage Stimulating Factor					
DALYs:	Disability-Adjusted Life Years					
DDT:	Dichloro-diphenyl-trichloroethane					
DEAE:	Diethylaminoethyl					
DFID:	Department For International Development					
DFMO:	Difluoromethylornithine					
dsRNA:	Double Strand RNA					
HAT:	Human African Typanosomosis					
IPTG:	Isopropyl-β-D- thiogalactopyranoside					
LB:	Luria-Bertani					
miRNA:	microRNA					
NBT:	Nitro Blue Tetrazolium					
ORFs:	Open Reading Frames					
PAP:	polyA polymerase					
PCR:	Polymerase Chain Reaction					
PTGS:	Post-Transcriptional Gene Silencing					
Py:	Pyrimidine Tract					
RdRP:	RNA-dependent RNA Polymerase					
RISC:	RNA-induced Silencing Complex					
RNAi:	RNA Interference					
RRM:	RNA Recognition Motif					
SAP:	Spliceosome Associated Proteins					
SF:	Splicing Factor					
siRNA:	Small Interfering RNA					
SIT:	Sterile Insect Technique					
SL RNA:	Spliced Leader RNA					
SL RNP:	Spliced Leader Ribonucleoprotein					
snRNA:	Small Nuclear RNA					

snRNP:	Small Nuclear Ribonucleoprotein
SR:	Serine- Arginine
U2AF:	U2 Auxiliary Factor
WHO:	World Health Organization
X-Gal:	5-bromo-4-chloro-3-indoyl-β-D- galactopyranoside
ZPFM:	Zimmerman Post-Fusion Medium

# CHAPTER ONE INTRODUCTION

#### 1.1 Background information

African trypanosomes, members of genus *Trypanosoma*, cause diseases that affect man and domestic animals. The disease is known as sleeping sickness in human (or Human African Trypanosomosis, HAT) and nagana in cattle. Tsetse-transmitted trypanosomosis affects 36 countries in sub-Saharan Africa where more than 48 million cattle and 60 million people are at risk (WHO, 2002). It has been recognised as a cause of severe morbidity and mortality throughout sub-Saharan Africa and a major constrain to livestock production (Allsopp, 2001). Annual losses in cattle production alone are estimated at US \$1.2 billion (FAO, 2002). Despite its impact, trypanosomosis remains among the most neglected tropical diseases (Morel, 2003).

Trypanosoma brucei has three subspecies, two of which cause sleeping sickness. Trypanosoma brucei gambiense causes a chronic form of the disease and is found mainly in Western and Central Africa. Trypanosoma brucei rhodesiense causes a more acute form of sleeping sickness and occurs in Eastern and Central Africa while Trypanosoma brucei brucei does cause the disease in man. One of the control efforts of the disease is the use of drugs. In the recent years increased incidents of resistance to available drugs has limited their value. Tsetse control methods such as the use of trapping and bait technology, insecticide spraying and the Sterile Insect Technique (SIT) have also been applied with varying levels of success. The efforts to develop more efficient and sustainable methods and tools for the control of African trypanosomosis must therefore continue.

Drug development depends on the understanding of parasite biology. Parasites have developed mechanisms that enable them to survive and thrive in their hosts. Rational drug development is based on differences that allow the development of products that are selectively toxic to parasites. Trypanosomes have unique molecular mechanisms that are different from their human and animal hosts, which should be investigated and exploited for effective chemotherapeutic control.

In eukaryotic organisms, genes occur as coding regions (exons) interrupted by non-coding regions (introns) in genomes. They are decoded into precursor messenger ribonucleic acid (premRNA) by the process known as transcription. Synthesis of mRNA occurs in the spliceosome, which is a macromolecular complex consisting of small nuclear RNA (snRNA) and proteins (Jurica and Moore, 2003). The spliceosome contains various macromolecular interactions: RNA-RNA, RNA-protein and protein-protein interactions and the complex facilitates the process of splicing, which entails the removal of introns and joining of exons. The majority of trypanosome transcript lack introns and are transcribed as polycistronic units (Lücke *et al.*, 1997; Denker *et al.*, 2002; Liang *et al.*, 2003).

In trypanosomes, the mature mRNA is derived from independent pre-mRNA molecules in a process known as trans-splicing (Sutton and Boothroyd, 1986; Denker *et al.*, 2002; Garcia-Blanco, 2003). Trans-splicing in trypanosomes appears to be linked to polyadenylation, the addition of a poly-adenosine tail to the 3'-end of pre-mRNA (Clayton, 2002; Jurica and Moore, 2003). In mammals, however, trans-splicing of conventional pre-mRNAs appears to be exceedingly rare due to the presence of trans-acting inhibitors or lack of specific trans-activators (Garcia-Blanco, 2003). Proteins that are essential for trans-splicing but not for cissplicing have also been recorded to be absent in human, fly and plant genomes (Denker *et al.*, 2002). Cis-splicing involves formation of mature mRNA from a single pre-mRNA. The presence of trans-splicing-specific factors indicates that this process can be explored as a possible target for therapeutic intervention. Thus, in this study, insights to the *Trypanosoma brucei* trans-splicing and polyadenylation factors are sought.

### 1.2 The statement of the problem

Trypanosomosis is a major constrain to livestock farming in sub-Saharan Africa and limits the full potential of agricultural development in the 36 countries that hold the continent's greatest potential for expanded agricultural production (Swallow, 1999). Despite the annual loss of US \$1.2 billion in cattle production alone and an estimated US\$ 4.7 billion in agricultural gross domestic production (FAO, 2002), the disease remains among the most neglected in terms of drug development. Most available drugs are highly toxic and increasingly encounter parasite resistance. Since we may soon be confronted with complete absence of effective drugs, there is

an urgent need for research geared towards identification of unique biological processes that will facilitate clinical development of promising compounds. Investigation on trans-splicing and polyadenylation factors and their potential functions thus provides a foundation for appropriate target identification in efforts to develop anti-parasitic drugs. Improved chemotherapeutic control strategy will enhance the realization of Africa's optimum agricultural potential that would in turn support her economically disadvantaged inhabitants.

#### 1.3 General objective

To clone and undertake a functional study of factors involved in trans-splicing and polyadenylation in *Trypanosoma brucei*.

#### 1.3.1 Specific objectives

- i. To identify *Trypanosoma brucei* homologs of the *Trypanosoma cruzi*, *Leishmania major*, *Homo sapiens* and *Saccharomyces cerevisiae* splicing and polyadenylation factors in public database.
- ii. To amplify, clone and sequence genes encoding these factors.
- iii. To analyse the nucleotide and amino acid sequences of the cloned factors.
- iv. To express the open reading frames (ORFs) of selected factors in bacterial expression system.
- v. Undertake functional analysis of selected factors using RNA interference technique.

#### **1.4 Justification**

Kinetoplastid parasites such as trypanosomes have developed specific variations from common eukaryote mechanisms such as trans-splicing and translation (Denker *et al.*, 2002). The variations are determined at least partially by the protein components. Hence, it is reasonable to propose that trypanosome proteins can harbour some fundamental variations when compared to homologous proteins in the mammalian host. These variations are potential chemotherapeutic targets.

Trans-splicing is an essential step in the expression of all protein-coding genes in trypanosomes (Lücke et al., 1997; Sturm and Campbell, 1999; Palfi et al., 2000). The factors involved in

trans-splicing have also been shown to provide crucial communication links between splicing machinery and other processes such as transcription, capping and polyadenylation (Jurica and Moore, 2003). Analysis of factors involved in trypanosome trans-splicing will therefore provide insights into the machinery employed by them to carry out their functions. The investigations of trans-splicing and polyadenylation factors and their potential interactions will provide insights into the linkage between trans-splicing and polyadenylation, which appear to be closely coordinated. It also lays a foundation the discovery of potential drug target.

# CHAPTER TWO LITERATURE REVIEW

#### 2.1 Impact of trypanosomosis in Africa

African trypanosomosis affects man and his livestock. The disease limits Africas' agricultural potential since it adversely restricts livestock keeping and affects, directly or indirectly, other mans' economic activities. The economic deprivation is exacerbated by losses in milk production, tractive power, waste products that provide natural fuel and fertilizer and secondary products such as clothing and hides. This denies Africa a great economic benefit since affected regions are the continent's greatest potential for expanded agricultural production (Swallow, 1999).

#### 2.1.1 Human African trypanosomosis (HAT)

Human African trypanosomosis (HAT) also known as sleeping sickness occurs only in Africa. This devastating disease kills all those infected unless treated (Kioyi and Mattock, 2005). Although accurate statistics for HAT are not available, it is estimated that between 300,000 and 500,000 new infections occur per year, and more than 60 million people are at risk of contacting the disease (WHO, 2000; Fairlamb, 2003; Kubata *et al.*, 2005). An estimated 50,000 people die every year with 100% fatality rate in untreated patients (Table 1). However, these numbers are considered underestimates due to difficulties in diagnosis and remoteness in endemic regions (WHO, 2000).

HAT has been a major cause of depopulation of large tracts of Africa and mainly strikes the active adult population (WHO, 2002). This has led to abandonment of fertile lands in tsetse infested area by farmers due to fear of contracting the disease. It is estimated that the average rural inhabitant would lose 10 years of income, or \$615 (1986 U.S. dollars), because of premature death caused by the disease (WHO, 2002). With the scourge making a comeback, greater risks in human health are unavoidable particularly in regions experiencing large-scale migrations and inadequate or non-existent health systems, such as in southern Sudan, Democratic Republic of Congo and Angola. These regions also have suffered political instability, displacement of populations and war leading to extreme poverty in a large

proportion of the population. With these limitations and poor monitoring structures, the exact impact of the disease may be more severe than is estimated.

Statistic/Information	Statistical information in
	sub-Saharan Africa
Geographical location	36 countries
Population at risk	60 million
Number of deaths at 2001	50, 000
Global case fatality rate in 2002	N/A
Number of people infected in 2003	300,000-500,000
Global disease burden in 2001 (DALYs)	1.6 million

Table 1. Fact file: African trypanosomosis statistics

Abbreviations: DALYs, Disability-Adjusted Life Years; N/A, Not Applicable. Only 3-4 million people are under active surveillance.

Source: Fairlamb, 2003.

#### 2.1.2 African animal trypanosomosis (AAT)

African trypanosomosis remains a major constraint to livestock productivity and continues to impede intensification of crop-livestock systems across vast areas of the humid and sub-humid zones of Africa that hold the greatest potential for increased agricultural production. The main direct economic impact of trypanosomosis is on cattle through reduction of birth rates and increment of abortion and mortality rates. Annual losses here amount to an estimated 3 million deaths, mainly of young stock, with up to 25% mortality in pre-weaning calves (FAO, 2002). Mortality losses are compounded by lower reproduction, milk yields and weight gain. With 94% of the total African cattle population distributed at the fringes of the continental tsetse belt, it is estimated that direct losses amount to US\$ 1.2 billion yearly (FAO, 2002). The distorted cattle distribution also affects crop production. In the absence of cattle, there is no draught power for ploughing, less manure to use as fertilizer, less feeding of animals with crop residues and by-products (DFID, 2001; Kabayo, 2002). The productivity of the land therefore remains sub-optimal. Rearing of susceptible animals is uneconomical in areas with high tsetse challenge, but minimal mixed farming is possible in areas with low tsetse challenge. Other

valuable livestock such as camels suffer from the disease. However, little information on the economic impact of the disease in small ruminants is documented.

#### 2.2 Control of trypanosomosis

For more than one century, controlling trypanosomosis using a variety of approaches has been a major focus for research and development. These activities include strategies to control the tsetse fly vector and the parasite in livestock and man. These strategies have however recorded varied success.

#### 2.2.1 Vector control

Tsetse control has long been an important option for reducing the incidence of trypanosomosis. The goal is to significantly reduce the vector densities in areas populated by susceptible hosts. Initial attempts to control tsetse involved widespread bush clearing to destroy the flys' breeding habitats. This method is no longer employed since it poses a serious threat to biodiversity. Other methods used to control the vector population include spraying with insecticides, SIT, targets, traps and bait techniques (Aksoy *et al.*, 2001).

The discovery of dichloro-diphenyl-trichloroethane (DDT) and other persistent insecticides in 1945 paved way for ground spraying as the main line of defence against tsetse fly throughout Africa (Dransfield *et al.*, 1991; Allsopp, 2001). Other chemical compounds that have been used include chlorobenzene derivatives, halobenzene insecticides and synthetic pyrethrins. Aerial applications of insecticides using fixed wing aircraft were used with considerable success in savannah regions and from helicopters in forested areas (Aksoy *et al.*, 2001). Although quite successful, this method is environmentally damaging and has resulted in the spread of insecticide resistance in many vectors. The problem of reinvasion of previously cleared areas has also been exhibited (Schofield and Maudlin, 2001). Due to this shortfall in addition to high costs, there is minimal reliance to this approach but the dependence still exists.

Bait technology became widely used in the 20<sup>th</sup> century and is an attractive tactic for reducing tsetse fly population (Allsopp, 2001). It involves attracting tsetse with visual or olfactory baits to allow capture and killing. Improvement of this technique is the use of live baits (i.e. cattle) to

which insecticide has been applied by dipping or spraying or with pour-on formulations (Allsopp, 2001). It has become increasingly popular since it is suited to use by individual farmers or communities. Recent availability of synthetic pyrethroid insecticides, which are highly effective against insect pests but have a low mammalian toxicity, has boosted its use.

Additional environmentally acceptable methods for eradicating tsetse consist of an integrated campaign using insecticide-treated screens or traps. They have been made simpler into cloth screens impregnated with insecticides, called targets. The targets and traps developed have been shown to reduce fly population to low levels but only in some areas and species of tsetse (Allsopp, 2001). Moreover, to be efficacious, this control method demands regular target and trap maintenance and the active participation of the livestock keeping communities where traps are deployed (Allsopp, 2001).

Sterile insect technique (SIT) involves sustained systematic release of sterile male insects to fertilise with the wild females among the wild population, which are then unable to produce viable progeny. SIT does not require insecticides and is environmentally benign. It has been used successfully in Burkina Faso, Tanzania, Nigeria and most recently, in Zanzibar (Allsop and Phillemon-Motsu, 2000). However, the problem of reinvasion has been reported in the three countries except Zanzibar (WHO, 2004). While the low reproductive rate of tsetse makes this a highly desirable approach, it has been criticized due to the relatively large upfront costs that would be associated with its implementation (Aksoy, 2003). Unfortunately, the overall control of the vector is made more complex since tsetse flies have adapted to a wide range of habitats, from central African humid rain forests to the vast, semi-arid, open savannahs of Eastern Africa (Allsopp, 2001).

#### 2.2.2 Chemotherapy

Chemotherapy is one of the methods heavily relied on in trypanosomosis control. This method has the advantage that it can be used under any production system, in any ecological zone and farmers can individually apply trypanocides to their cattle. Various compounds have been developed for human and animal treatment. However, set against the spectacular advances in other areas of chemotherapy over the past, trypanosomosis in particular has fared extremely

badly. This is because of low profit margins expected by pharmaceutical companies since people most at risk are among the poorest in the world. The emergence of chemoresistance and toxicity highlights the limits of chemotherapy.

#### 2.2.2.1 Chemotherapy in human

Sleeping sickness evolves through clinically distinct stages namely early/stage 1 and late/stage 2. The early stage is a haemolymphatic stage in which parasites develop in the blood, lymph and peripheral organs. The parasites then spread to the central nervous system (late/encephalitic stage) where they cause serious neurological disorders. The type of treatment depends on the phase of the disease (Table 2). Pentamidine isethionate (Lomidine<sup>TM</sup>) is used only for the early phase of the disease since it does not cross the blood-brain barrier. It is active against *T. b. gambiense*, but is not used against *T. b. rhodesiense*, since primary resistance to it has been found in some areas. Other limitations include inactivity when dosed orally, slow elimination and toxicity. It binds negatively charged cellular components, such as phospholipids and nucleic acids and disrupts the structure of kinetoplast DNA (kDNA). Suramin (Germanin<sup>TM</sup>) is used for treating the early stages against *T. b. rhodesiense*. The mode of action is still not well understood but it is suggested to inhibit various glycolytic enzymes (Fairlamb and Bowman, 1980; Fairlamb, 2003). All of these drugs are expensive and have serious side effects.

The regimen for treatment of advanced stage (late phase/stage 2) of HAT where the central nervous system is affected involves the use of melarsoprol (Mel B, Arsobal<sup>TM</sup>). It inhibits trypanothione reductase, an enzyme essential for maintaining the correct thiol-redox balance for biochemical processes. Its use is limited since it causes encephalopathy in 5-10% of cases, half of which are fatal (Fairlamb, 2003). Other side effects include vomiting, abdominal colic, peripheral neuropathy, arthralgia and thrombophlebitis. A new drug effornithine (Difluoromethylornithine/DFMO, Ornidyl<sup>TM</sup>) has been developed that is effective in the treatment of both the early and the late stage of West African sleeping sickness. However, it is inactive against East African sleeping sickness due to innate tolerance of *T. b. rhodesiense* to the drug (Matovu *et al.*, 2001). It is currently mainly used as back-up drug for melarsoprol refractory *T. b. gambiense* cases, but recently also as first line treatment (EANETT, 2003). It is an ornithine decarboxylase inhibitor, which involves a 14-day treatment regime at the cost of

US\$ 250 per patient for the drug alone. Like melarsoprol, effornithine is expensive and requires weeks of hospitalisation. At present, only two drugs (DB 289, diamidine derivative and Megazol, nitro-imidazole) are under development (Legros *et al.*, 2002).

Drug	Species	Indication ½ life	Year of first use	Comments
Pentamidine isethionate	T.b.g	Stage 1 [9.4 hrs - IM] [6.4 hrs-IV]	1940	2 <sup>nd</sup> line drug when suramin therapy is contraindicated
Suramin Sodium	T.b.g T.b.r	Stage 1 [50 days]	Early 1920s	No significant clinical resistance has emerged
Melasoprol (Mel B)	T.b.g T.b.r	Stage 2 [35 hrs]	1949	Increased treatment failure
Eflornithine	T.b.g	Stage 2 [3 hrs]	1981	Difficult use
Nifurtimox¥	T.b.g T.b r?	Stage 2 [3.5 hrs]	1977	Not registered for HAT Case series only
				Toxic and action on T.b.r. poorly documented

Table 2. C	)verview	of human	trypananocidal	drugs
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T.b.g =Trypanosoma brucei gambiense, T.b.r =Trypanosoma brucei rhodesiense

IM = intramuscular, IV = intravenous

<sup>4</sup> = registered for treatment of *T. cruzi* infections and is used in combination with melarsoprol (EANETT, 2003). The table is modified from Legros *et al.* (2002).

### 2.2.2.2 Chemotherapy in animals

Diminazene aceturate (Berenil<sup>TM</sup>), isometamidium chloride (Samorin<sup>TM</sup>) and homidium bromide (Novidium<sup>TM</sup>) are the drugs widely used for the treatment of nagana or cattle trypanosomosis. These drugs are effective both for treatment and for prophylaxis. Diminazene aceturate binds to AT-rich regions of DNA and RNA duplexes exhibiting properties characteristic of intercalation and minor groove binding (Portugal, 1994). This unwinding inhibits type II topoisomerase. Isometamidium chloride acts by cleaving kDNA-topoisomerase

complexes (Wells *et al.*, 1995) while homidium bromide intercalates into the DNA thus inhibit DNA-primed polymerase. Due to their mode of action, they are mutagenic and tumorigenic and also cause necrosis and sloughing of extensive areas of unpigmented skin. Suramin, used in treatment of sleeping sickness, is also used against *Trypanosoma evansi* in camels. Cattle treated with these drugs may only be slaughtered and used for consumption, several months after treatment, when there is no residual drug left in the animal.

#### 2.2.3 Prophylactic strategies

Prophylaxis concerns all measures that can be undertaken to prevent the onset or transmission of a disease. In trypanosomosis, the prophylactic strategies involve actions taken against vector population and health care for both human and animal populations. Two management approaches are applied in human populations, these are, routine screening for the disease and treatment of the infected persons. In cattle, isometamidium chloride (Samorin<sup>TM</sup>) is the most commonly used prophylactic drug. Although extensively used in trypanosomosis control, chemoprophylaxis is expensive, and thus unsatisfactory long-term solution to the problem of African trypanosomosis.

#### 2.2.4 Vaccination

The possibility of developing a vaccine against trypanosomosis has been frustrated by the ability of the parasite to vary their surface antigens and thus evade the host immune response (Donelson, 2002). A conventional vaccine that primes animals' immune system against only one or a few antigens will therefore not be broadly effective against trypanosomosis. Lubega *et al.*, (2002) showed that tubulin rich preparation from *T. brucei* confers broad protection against African trypanosomes. This can be through internalisation of the antibodies by the parasite by a mechanism not yet established (Balaban *et al.*, 1995) though flagellar pocket is implicated (Gull, 2002). However, the compound may not be immunogenic enough since it is not exposed on the surface of trypanosomes.

#### 2.2.5 Trypanotolerant animals

The use of animals that are inherently tolerant to the effect of the disease and are able to remain relatively productive even when infected has been used as an approach to farming within the



11

trypanosomosis belt. It is used as a strategy for productive farming since it allows keeping of animals in the infested areas. In West Africa, these tolerant breeds include N'Dama (*Bos taurus*), Baoule, Laguna, Samba, Dahomey and Muturu cattle that have been bred in areas endemic to trypanosomosis and showed considerable resistance (Weits, 1970). In East Africa, the Orma Boran and Maasai zebu have been shown to possess a degree of natural resistance to trypanosomosis (Mwangi *et al.*, 1998). The main drawbacks of this method are that many farmers believe these breeds are less productive than others and few of such animals are available. They also constitute a reservoir for infection to other susceptible livestock.

#### 2.3 RNA splicing

#### 2.3.1 Spliceosome and splice sites

Most mRNAs encoded by nuclear genes are synthesized as precursor RNA molecules, in which the coding sequences (exons) are interrupted by intervening sequences (introns). The mRNA becomes functional when introns are excised and exons precisely joined together by a process called splicing. This splicing reaction is achieved by a dynamic RNA-protein complex called spliceosome (Jurica and Moore, 2003). It consists of five small nuclear RNAs (snRNAs) and more than fifty attached proteins (Ito *et al.*, 1999; Nilsen, 2000). The snRNAs are U1, U2, U4, U5 and U6. They have proteins attached thus exist as small nuclear ribonucleoproteins (snRNPs).

Precise removal of introns (without disturbing the reading frame) requires specific sequences that must be involved in the process. Splicing signals have been identified by examination of the boundaries of introns and exons from a large number of genes in a variety of eukaryotic cells (Padgett *et al.*, 1986; Nilsen, 2000). The signal identified for higher eukaryotes is:

### 5' AG|GUAAGU...... YNCURAC...... Y<sub>n</sub>NAG|G 3'

where the vertical line () denotes the boundary between the exon and the intron, Y indicates a pyrimidine, R a purine and N means any base.  $Y_n$  means region of about nine pyrimidines (Py). The red 'A' is the branch point adenosine that is involved in the splicing reaction. These include the three conserved sequence elements identified as the 5' and 3' splice sites and a site within

the intron near the 3' splice site known as the branch point sequence (BPS). Consensus sequences from plant introns show strong similarities to the animal consensus elements (Brown *et al.*, 2002). This identifies the sequence of a generic intron as:

Exon...GU......Intron.....AG...Exon

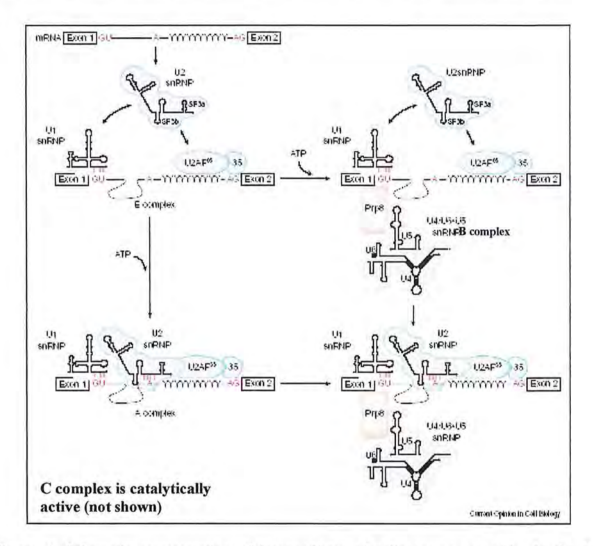
#### 2.3.2 Splicing mechanism

#### 2.3.2.1 Cis-splicing

The spliceosome is a multienzyme complex that consists of five snRNPs (U1, U2, U4, U5 and U6), non-snRNPs (helicases, DExD/H box proteins, serine-arginine, SR proteins) and conserved pre-mRNA sequence elements (5' splice site, 3' splice site, Py and BPS) (Jurica and Moore, 2003). The spliceosome assembly proceeds through a series of short-lived intermediate stages (Figure 1) dubbed E/CC, A, B and C (Jurica and Moore, 2003). In summary, the process is initiated upon binding of U1 snRNP to the 5' splice site, the mammalian branch point binding protein or splicing factor 1(mBBP/SF1) to the BPS and U2 auxiliary factor (U2AF) to the Py and 3' AG. This forms the early or commitment complex (E/CC). Associated protein factors such as U2AF are needed to promote the complex formation when the sequence at the BPS varies from the consensus. Subsequently, U2 snRNP binds to the BP to form A complex (a.k.a pre-spliceosome complex) in which ATP is consumed. The addition of U4/U6-U5 tri-snRNPs to pre-spliceosome results into complex B. The C complex is formed from massive rearrangement in which U6 replaces U1 at the 5' splice site, U6 and U2 interact, U5 bridges the splice sites and U1 and U4 becomes destabilized. This rearranged spliceosome is catalytically active and carriers out two trans-esterification reactions.

During the ATP dependent transition of complex E to pre-splicing complex A, a short helix results between U2 snRNP and the branch site, defining the adenosine that functions as the nucleophile in the first catalytic step. The U2 snRNP and U6 snRNP in the C complex leads to bulging of the adenosine exposing its nucleophilic 2' hydroxyl (OH) group that attacks the 5' splice site guanosine bound to U5 snRNP. The reaction results into cleavage of pre-mRNA at the 5' splice site, generating an exon RNA fragment that contains a 3' hydroxyl group and an intron-exon fragment with the intron in the form of a lariat due to a 5'-2' bond (Padgett *et al.*,

1986). The two RNAs are held together non-covalently. A further rearrangement in complex C brings the 3'OH of exon fragment into close proximity to 5' phosphate of intron-exon fragment enabling the second trans-esterification reaction. This results into 3' splice site excision, ligation of the exons and release of the free intron. This chemical process is cis-splicing (Figure 2).



**Figure 1.** Schematic representation of intermediates of spliceosome assembly. In the E complex, U1 snRNP binds 5'ss (red GU), and U2 snRNP loosely binds to the pre-mRNA near the 3'ss (red AG) in an ATP dependent process (A complex). U4/U6·U5 tri-snRNP binds to the 5'ss, in part through interaction between Prp8 and pre-mRNA forming B complex. Red vertical lines show RNA base-pairing interactions, and filled circles and squares depict snRNA 5' cap structures. Modified from Hastings and Krainer (2001).

Self-splicing introns have shown that RNA can catalyse transesterification reaction by a mechanism in which metal ions that interact with specific atoms in the RNA activate the attacking nucleophile and stabilize the leaving group (Sontheimer *et al.*, 1997; Nilsen, 2000; Gordon *et al.*, 2000). This, a form of autocatalysis is known as self-splicing.

Alternative mRNA splicing is the term used to describe the regulated process of differential inclusion or exclusion of regions of the pre-mRNA. It generates multiple mRNA isoforms (Breitbart *et al.*, 1987; Gascard *et al.*, 1998) hence proteins with different functions from a single gene. Multiple transcripts from a single gene can result from exon skipping, mutual exclusion of exons and retention of introns and/or selection of an alternative 5' or 3' site (Smith and Valcárcel, 2000; Maniatis and Tasic, 2002; Roberts and Smith, 2002; Lee and Irizarry, 2003; Reddy, 2004). It has been documented in plants (Reddy, 2004), Drosophila, human and most vertebrates (Smith and Valcárcel, 2000).

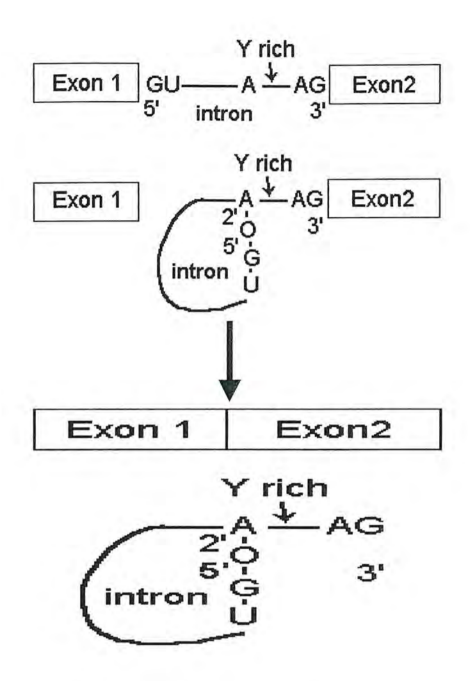


Figure 2. Two steps splicing (cis-splicing). Step one is the first trans-esterification reaction involving the 2' OH of the adenosine residue at the BPS and phosphate of the 5' splice site leading to lariat formation. Step two represents nucleophilic attack of 3'OH of the 5' splice site to the 5' phosphate of the second exon leading to exon ligation and intron excision.

#### 2.3.2.2 Trans-splicing in trypanosomes

Trypanosomes belong to a group of eukaryotic organisms in which the splicing mechanism involves independent pre-mRNAs, a process known as trans-splicing or intermolecular exonligation (Figure 3). Surprisingly, cis-splicing has been reported in poly (A) polymerase genes (PAP) genes in T. brucei and T. cruzi (Mair et al., 2000) against the almost two-decade-old tenets that trypanosomes exhibit only trans-splicing. Nonetheless, trans-splicing process is an essential step in the expression of all protein coding genes in trypanosomes that form polycistronic transcripts (Mandelboim et al., 2002). It involves interaction between 5' and 3' splice sites on separate transcripts and occurs in a variety of eukaryotic organisms including trypanosomes (Sutton and Boothroyd, 1986), euglena, nematodes, trematodes and chordates (Lücke et al., 1997; Mandelboim et al., 2002). In this process, a Y branched intermediate is formed as opposed to a lariat in cis-splicing. This occurs by addition of a short non-coding miniexon sequence derived from the splice leader (SL) RNA onto each protein-coding exon sequence present within polycistronic precursor transcripts (Sutton and Boothroyd, 1986; Lücke et al., 1997; Li et al., 2000). The splicing complex that carries out this process is known as trans-spliceosome. The SL sequence is derived from a large transcript called the SL RNA (Li et al., 2000; Landfear, 2003). The SL RNA is transcribed from arrays of tandemly repeated genes of 10-11 copies per haploid genome (Roberts et al., 1996) and is present in the cell in the form of a SL ribonucleoprotein, the SL RNP (Goncharov et al., 1999; Evans et al., 2001).

The SL RNA of all organisms has two domains (Mandelboim *et al.*, 2002). The SL sequence or miniexon, which in trypanosomes is thirty-five nucleotides long (Sutton and Boothroyd, 1986) followed by the intron of variable length and can be folded into a conserved secondary structure with three stem-loops (Mandelboin *et al.*, 2002). The primary sequence of SL RNA is not conserved among the trypanomastids, but the secondary structure is consistent (Sturm and Campbell, 1999) and fulfils a dual function of recruiting splicing co-factors and serving as a substrate for splicing.

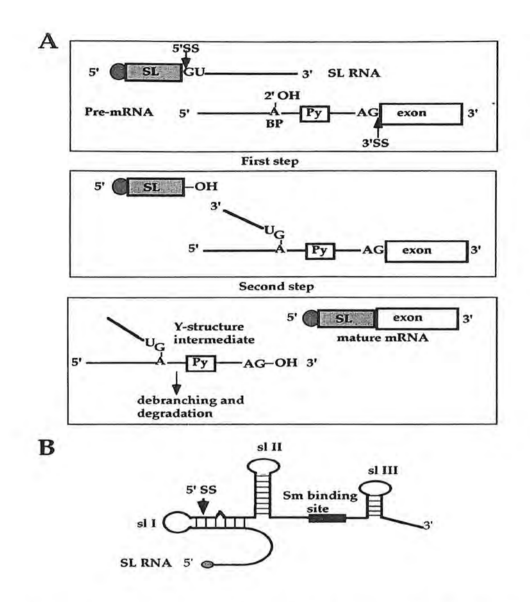


Figure 3. Mechanism of trans-splicing. (A) Schematic representation of trans-splicing. The 5' splice site GU on the SL RNA and the 3' splice site AG on the pre-mRNA are indicated. Y branched intermediate is shown. BP, branch point; Py, polypyrimidine tract. (B) Secondary structure of SL RNA. The three stem-loop structures (sl I, II, and III), the 5' splice site, and the Sm-binding site are indicated. The shadowed dot at the 5' end of SL RNA indicates the cap 4 structures. (Adapted from Liang *et al.*, 2003)

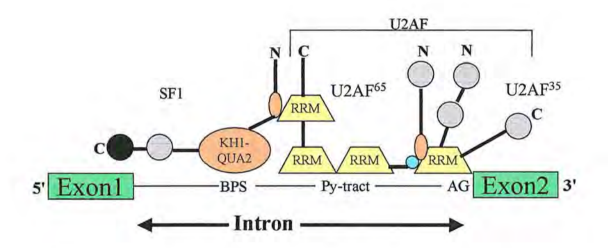
#### 2.4 Linkage between trans-splicing and polyadenylation

Factors involved in capping, splicing and polyadenylation interact with the carboxy-terminal domain of RNA polymerase II at an early stage of mRNA production in cis-splicing. These processes are therefore coupled to transcription (Jurica and Moore, 2003; Liang et al., 2003). This linkage involves specific sequence, such as the signal for polyadenylation (AAUAAA) in higher eukaryotes. Such sequences are absent in trypanomastid mRNAs (Liang et al., 2003). In trypanosomes, evidence suggests that maturation of polycistronic pre-mRNAs involves a temporal and mechanistic relationship between trans-splicing and 3'-end formation (Ito et al., 1999; Shepard et al., 2002; Hendriks et al., 2003; Liang et al., 2003). It has been shown that the pyrimidine rich sequences immediately upstream of the  $\alpha$ -tubulin trans-splice site are a major determinant for B-tubulin mRNA 3'-end formation in trypanosomes (Matthews et al., 1994). The Py thus plays a role in the coupling of these processes since its maturation leads to aberrant poly (A) site choice (Hug et al., 1994). This model is supported by the finding that the location of poly (A) site moves in concert with the 3' splice site AG in Leishmania (LeBowitz et al., 1993), a close relative of trypanosome. In Leishmania, the factor(s) responsible for cleaving the pre mRNA at the poly (A) site bind only a certain short distance away from the 3' splice site. The factors required for 3'-end cleavage and polyadenylation therefore associate with the premRNA after the machinery of the spliceosome has marked the 3' splice site region. The factors shared between 3'-end formation and trans-splicing couple the two processes. U1 snRNP is involved in coupling cis-splicing to polyadenylation. Similarly, an SL RNP specific protein may coordinate coupling of polyadenylation to trans-splicing in trypanosomes (Liang et al., 2003).

#### 2.5 Splicing, cleavage and polyadenylation factors

The spliceosome assembly proceeds through a series of short-lived intermediate stages dubbed E/CC, A, B and C (Jurica and Moore, 2003). The E complex consists of U1 snRNP base paired to the 5' splice site, U2 auxiliary factor heterodimer (U2AF) which binds to Py and 3' splice site, and SF1/mBBP. Associated protein factors such as U2AF are needed to promote the complex formation when the sequence at the BPS deviates from the consensus. The U2AF heterodimer consists of 65-Kda (U2AF<sup>65</sup>) and 35-Kda (U2AF<sup>35</sup>) subunits (Ito *et al.*, 1999; Shepard *et al.*, 2002) and is essential for viability in eukaryotes (Vàzquez *et al.*, 2003). U2AF<sup>65</sup>

contains an N-terminal arginine-serine-rich (RS) domain and three RNA recognition motifs (RRM) in mammals (Shepard *et al.*, 2002). It interacts directly with the Py and is involved in stabilizing the interaction of U2 snRNP with the BP (Gozani *et al.*, 1998). This activity requires the RS domain, which is thought to assist in the U2 snRNP-pre-mRNA duplex and the third RRM that interacts with SF3b155, a component of U2 snRNP (Shepard *et al.*, 2002). SF1 and U2AF<sup>65</sup> interact with each other facilitating cooperative recognition of the BP and Py tract that are adjacent to each other (Selenko *et al.*, 2003) (Figure 4). SF1 contains at least five distinct structural domains in which the amino-terminal region contacts the U2AF<sup>65</sup> (Guth and Valcárcel, 2000).



**Figure 4**. Recognition of the 3' splice site. SF1 and the U2AF heterodimer recognize the branchpoint sequence (BPS), polypyrimidine tract (Py), and 3' splice site (AG) by forming a network of protein-RNA and protein-protein interactions mediated by conserved KH and RRM domain (Selenko *et al.*, 2003).

The A complex also known as the pre-spliceosome is the second step in spliceosome assembly. It is formed when U2AF recruits U2 snRNP to the BP, which displaces SF1 in a process in which ATP is consumed. Two multisubunit splicing factors, SF3a and SF3b, are components of U2 snRNP and are required in the binding to the BP. SF3a consists of three subunits (SF3a 60, SF3a 66 and SF3a 90) and SF3b consists of eight subunits (P14, P14b, SF3b 10, SF3b 49, SF3b 125, SF3b 130, SF3b 145, and SF3b 155). They anchor U2 snRNP tightly to the pre-mRNA. Addition of U2, U5 and U6 snRNPs to complex A as a preassembled tri-snRNP results in complex B. Subsequent formation of competent C complex involves the recruitment of additional protein factors along with significant structural rearrangement that destabilize the association of U1 and U4 snRNPs with the catalytic core (Jurica and Moore, 2003).

Vàzquez *et al.*, (2003) have characterized the *T. cruzi* U2AF<sup>35</sup>. It is 240 residues long, with four RRMs, one central and three at the C-terminal, and a C-terminal SR segment. In eukaryotes, U2AF<sup>35</sup> and U2AF<sup>65</sup> form an intimate heterodimeric complex in which the RRM of U2AF<sup>35</sup> and a central polyproline segment of U2AF<sup>65</sup> interact via reciprocal "tongue in groove" tryptophan residues (Vàzquez *et al.*, 2003). Guth and Valcárcel (2000) documented various factors that interact with U2AF<sup>65</sup> thus suggesting the importance of U2AF in viability in eukaryotes. It is also suspected to interact with poly (A) polymerase (PAP) thus linking transsplicing to polyadenylation. Poly (A) polymerase (PAP) is the enzyme that adds the poly adenosine (A) tail after the cleavage reaction. Palfi *et al.*, (2000) characterized seven proteins associated with snRNP core complex of *T. brucei*. The seven proteins were found to bind to conserved sequences in U1, U2, U4 and U5 snRNP resembling the Sm sequence in cis-spliceosomal snRNPs.

Members of the serine-arginine (SR) proteins have multiple functions in pre-mRNA splicing reaction. These include the removal of constitutively spliced introns and regulating alternative splicing, which they do both *in vivo* and *in vitro* (Graveley, 2000; Furuyama and Bruzik, 2002). They bridge between other splicing components where interactions are too weak for effective binding of the basal factors U1 and U2 as well as recruitment of other splicing factors. All SR proteins have a modular organization and contain an N-terminal RNA-binding domain that interacts with the pre-mRNA and a C-terminal RS domain that functions as a protein interaction domain (Philipps *et al.*, 2003). Ismaili *et al.*, (2000) characterized *T. brucei* SR domain-containing protein (TSR1IP) bearing homology to cis-spliceosomal U1-70 kDa proteins. They showed that it interacts with the 5' splice region of the SL RNA. However, its identity as a 70-kDa homologue is currently puzzling, since an U1 specific protein whose homology to the U1-70 kDa domain was recently identified (Palfi *et al.*, 2002).

Cleavage and polyadenylation specificity factor (CPSF) is required for both cleavage and polyadenylation reactions. It contains subunits of 160, 100, 70 and 30 kDa referred to as CPSF 160, CPSF 100, CPSF 70 and CPSF 30 respectively. CPSF 30 contains five CCHC zinc knuckles (Hendriks *et al.*, 2003). Both types of motifs have been implicated in binding nucleic acids. Its role is probably to cooperate with CPSF 160 in recognition of pre-RNA substrates and through interaction with poly (A) binding protein II (PAB II) to stabilize the polyadenylation complex (Zhao *et al.*, 1999). Hendriks *et al.* (2003) showed that CPSF 30 depletion by RNA interference is lethal and leads to disruption of RNA processing in *T. brucei*. The 70 kDa and 100 kDa subunits are closely related, with 23% identity and 49% similarity and their function is unknown (Zhao *et al.*, 1999).

Cleavage stimulating factor (CstF) binds downstream of the cleavage site at a GU-rich sequence. It associates with CPSF and is necessary for cleavage but not poly (A) addition. It contains 77, 64 and 50 kDa subunits with the CstF 77 being the central subunit bridging the other two and they are arranged in a linear fashion (Zhao *et al.*, 1999). CstF 64 contains a classical RNA-binding domain (RBD) close to its amino terminus (Evans *et al.*, 2001) while the 50 kDa subunit has a protein interaction domain and binds to RNA Pol II CTD.

Cleavage factors I and II (CF I, CF II) are responsible for actual cleavage reaction. Three polypeptides of 25, 59 and 68 kDa and a possible fourth one of 72 kDa copurify with CF I activity in mammals (Zhao *et al.*, 1999). It increases the CPSF-RNA complex stability, suggesting that this factor interacts with CPSF and contributes to the overall stability of 3'-end-processing complex. CF I contains RS domains similar to splicing factors. CF II has not been purified to homogeneity and its function not known (Zhao *et al.*, 1999).

RNA polymerase II, through its conserved carboxyl-terminal domain (CTD) of its largest subunit, is involved in the cleavage reaction through interaction with CPSF and CstF. Its Nterminal contains a catalytic domain with homology to nucleotidyltransferases (Zhao *et al.*, 1999). CPSF and PAP suffice for poly (A) addition to a pre-cleaved RNA substrate. However, rapid elongation and control of poly (A) tail length requires an additional factor PAB II (Zhao et al., 1999). The poly (A) binding protein in T. brucei has been identified and purified by Pitula et al. (1998).

More has been done on the understanding of SL RNA but little on trans-splicing and polyadenylation factors in trypanosomes. Few splicing factors have been characterized across the kinetoplastids. This has led to little information regarding the dynamic nature of the spliceosomal complex. Since the information available is highly disjointed, the exact process of spliceosome assembly is not available. Characterization and functional analysis of the factors involved is thus important. This will provide information on important factors and interactions that are essential for RNA maturation, hence identifying potential drug targets.

#### 2.6 RNA interference (RNAi)

RNA interference (RNAi) is a phenomenon leading to post-transcriptional gene silencing (PTGS) after endogenous production or artificial introduction in a cell double-stranded RNAs (dsRNAs) that are complimentary to known mRNAs, specifically destroying that particular mRNA, thereby diminishing or abolishing gene expression. RNAi has been linked to many previously described silencing phenomena such as PTGS in plants and quelling fungi (Dykxhoorn et al., 2003; Pauls and Esté 2004). It is an evolutionarily conserved mechanism of gene silencing that is thought to inhibit the replication and expression of selfish DNA elements and viruses (Caplen et al., 2002; Chi et al., 2003) hence maintains genome integrity. Whereas the transcription of the gene is normal in this process, the translation of the protein is prevented by selective degradation of its encoded mRNA. Although its protective role would be very important in plants and C. elegans (Milhavet et al., 2003) which lack an antibody-based immune system analogous to that found in animals, its role in mammalian cells remains unclear. It thus must have evolved as a cellular defence mechanism against foreign DNA and RNA regulating the expression of endogenous genes (Milhavet et al., 2003). This phenomenon has been used to study gene functions in vitro by injection of dsRNA that led to an efficient sequence-specific gene silencing in C. elegans (Fire et al., 1998).

#### 2.6.1 RNAi mechanism

Double-stranded RNA-mediated interference (RNAi) is a PTGS process in which doublestranded RNA (dsRNA) triggers degradation of homologous mRNA in the cytoplasm of a cell. Studies over the last few years have demonstrated that RNAi is mediated by the generation of 21- to 23-nucleotide dsRNA molecules, termed small interfering RNA (siRNA) (Dillin, 2003; Dykxhoorn et al., 2003). They have a characteristic structure, with 5'-phosphate/3'-hydroxyl ends and a 2-base 3' overhang on each strand of the duplex. Upon entry into the cell, dsRNA is cleaved into siRNAs by the action of RNase-III-type enzyme Dicer (Jones et al., 2004; Pauls and Esté, 2004). These siRNAs are incorporated into a protein-RNA complex, the RNAinduced silencing complex (RISC) (Dykxhoorn et al., 2003). There is a strict requirement for the 3' siRNA to be 5' phosphorylated to enter into RISC, and siRNA that lacks a 5' phosphate are rapidly phosphorylated by an endogenous kinase (Dykxhoorn et al., 2003). The duplex siRNA is unwound, leaving the antisense strand to guide RISC to the homologous target mRNA for endonucleolytic cleavage (Dykxhoorn et al., 2003; Pauls and Esté, 2004). The target mRNA is cleaved at a single site in the centre of the duplex region between the guide siRNA and the target mRNA, 10-nt from the 5' end of the siRNA. Mismatches greater than 1-2bp within the 21- to 23-nt siRNA effectively disrupts proper degradation of the target mRNA (Dillin, 2003). The synthesis of the protein encoded by the mRNA targeted by the siRNAs is thus prevented and that protein is selectively depleted from the cell thus RNA interference. The RNAi process is shown in Figure 5.

In worms, interaction between the siRNA and mRNA can lead to immediate cleavage by Dicer, liberating a new siRNA, and the degradation of mRNA by endo- and exonucleases (Dillin, 2003). Alternatively, the siRNA can serve as a primer for an RNA-dependent RNA polymerase (RdRP), creating many more siRNAs. The action of RdRP may explain the catalytic mechanism of RNAi, because only a few dsRNA molecules are required to degrade a much larger population of mRNAs.

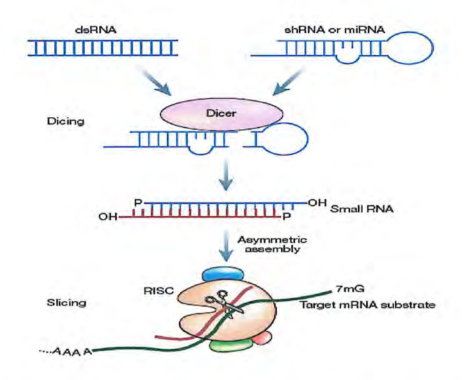


Figure 5. Overview of dsRNA-mediated mRNA degradation. Double stranded RNA is cleaved by dicer into 21- to 23-nt siRNAs, which is complexed with a large multiprotein complex, the RISC. RISC unwinds the siRNA to help target the appropriate mRNA (shown in green). The siRNA-mRNA hybrid is cleaved, releasing the siRNA, and the mRNA is degraded by endo and exonucleases.

Recent studies have identified the existence of endogenous siRNA molecules called microRNA (miRNA) (Jones *et al.*, 2004). These are transcribed in the nucleus as a large non-coding primary sequence that are processed by the RNase III enzyme Drosha to produce a hairpin RNA of ~70 nucleotides before being exported into the cytoplasm by exportin-5 (Jones *et al.*, 2004). Like siRNAi, pre-miRNA is then cleaved by the Dicer to produce a double stranded 21-to 23- nucleotide RNA duplex, which is incorporated into the RISC-like complex. However, in contrast to siRNA-mediated mRNA cleavage, miRNA is thought to block mRNA translation through imperfect complementary binding of the antisense sequence to the miRNA recognition elements (MREs) within the 3' untranslated region (UTR) (Jones *et al.*, 2004).

# 2.6.2 RNAi as a genetic tool

The discovery of the RNAi machinery provides a powerful tool for the study of gene function. Small fragments of double-stranded RNA (siRNA) can be introduced into cells to selectively degrade homologous endogenous mRNA enabling knock down of complementary genes. This technique has been used to investigate gene function in various organisms including *C. elegans* (Fire *et al.*, 1998), plants (Baulcombe, 1999), planaria (Sánchez-Alvarado and Mewmark, 1999), trypanosomes (Ngô *et al.*, 1998; Wirtz *et al.*, 1999; Shi *et al.*, 2000; LaCount *et al.*, 2002; Hendriks *et al.*, 2003; Alibu *et al.*, 2004), hydras (Lohmann *et al.*, 1999), *Drosophila* (Kinnerdell and Carthew, 1998; Misquitta *et al.*, 1999), mosquitoes (Caplen *et al.*, 2002) and mouse oocytes (Svoboda *et al.*, 2000).

## 2.6.3 RNAi and trypanosome genome studies

Characterisation of putative coding regions is a prerequisite for converting raw genomic sequence data into biologically relevant information (Gopal *et al.*, 2003). With the completion of trypanosome and leishmania genome projects (Berriman *et al.*, 2005; El-Sayed *et al.*, 2005a; Ivens *et al.*, 2005), a wealth of information is available at the repository genome database at GeneDB. Currently, GeneDB provides access to 32 genomes of various stages of sequencing curation pipeline as well as complete genomes with extensive manual curation. This includes sequences and associated annotation of bacteria, fungi, protozoa and arthropod (Arnaud Kerhornou and The Sanger Institute Pathogen Sequencing Unit, 2002; Hertz-Fowler *et al.*, 2004). The genome information will provide insight on parasitism, virulence, epidemiology (Gull, 2000) and parasite biology among other aspects. To exploit this information and the rapid accumulation of genome sequence in databases, techniques such as microarrays and RNAi (Donelson, 2002) that allow gene function studies are important.

The technique of RNA interference is useful for knocking down the expression of a specific mRNA in African trypanosomes and other organisms for the purpose of examining the function of its gene (DaRocha *et al.*, 2004). The technique circumvents the necessity of generating double knockouts, which is a requirement for such studies in diploid organisms. It works well in *T. brucei* (Ngô *et al.*, 1998) due to the presence of two predicted proteins representing potential Dicer candidates (El-Sayed *et al.*, 2005b). It can therefore be used to examine the

genes or a set of genes essential for trypanosome processes. Since this finding, more has been done on utilization of this technique in trypanosome gene functions studies (Wang et al., 2000; Morris et al., 2001; Hendriks et al., 2003), designing siRNAs (Redmond et al., 2003), development of cell lines (Chen et al., 2002), construction of plasmid vectors for RNAi delivery (Wirtz et al., 1999; Wang et al., 2000; Gull et al., 2002; Alibu et al., 2004; DaRocha et al., 2004; Motyka et al., 2004), in vivo analysis (Tschudi et al., 2003) and transfection methods. Redmond et al., 2003 have developed a web-based tool for designing RNAi targets in T. brucei. This T.brucei functional genomics project (TrypanoFAN) aims to utilise the information from the T.brucei genome project to produce a research resource and systematic collection of mutants by targeted gene inactivation using RNAi. These developments are important in the utilization of genome sequence for valuable functional assays.

#### 2.6.4 RNAi vectors for trypanosome gene research

The simplest method to achieve RNA interference in *T. brucei* is to use vectors with opposing bacteriophage T7 promoters, controlled by binding of the *tet* repressor to *tet* operators (Wang *et al.*, 2000; Morris *et al.*, 2001; LaCount *et al.*, 2002). The inducible RNAi vectors pZJM and  $p2T7^{Ti}$  are widely used for they allow for insertion of a sequence of interest between opposing T7 promoters under the control of tetracycline operator (Motyka *et al.*, 2004). This allows the transcription of the insert in both directions upon induction with tetracycline. Linearized vector allows integration by homologous recombination into ribosomal DNA (rDNA) spacer region, a transcriptionally inactive segment of the *T. brucei* genome (Morris *et al.*, 2001; LaCount *et al.*, 2002; Alibu *et al.*, 2004; Motyka *et al.*, 2004).

# 2.6.4.1 p2T7Ta blue vector

p2T7TA blue is a 6740bp vector derived from p2T7<sup>Ti</sup>B/GFP (LaCount *et al.*, 2002) and is shown in Figure 6. It has two promoters for bacteriophage T7 polymerase (shown as the big arrows facing each other). Each promoter has two binding sites for the bacterial *tet* repressor. The vector also has a hygromycin resistance gene, transcription of which is driven by a separate promoter. This gene allows for the selection of recombinants. Parasites express the T7 polymerase and the repressor so that transcription of the piece of DNA between the T7 promoters only happens when tetracycline is added to the medium.

Digestion with *Eam*1105I generates 3' overhanging T-ends, which allow PCR products with 3' A-ends to be cloned directly. Once inserted, opposing T7 promoters with downstream operators flank RNAi targets. Recombinant vectors are linearised at the *Not*I site and targeted to the rDNA locus (Wirtz *et al.*, 1999) in cells that express T7 RNA polymerase, *tet*-repressor and the selectable marker. Recombinants are selected with hygromycin B. Upon integration, the construct lies between rDNA spacer sequences. The operators bind *tet*-repressor in the absence of tetracycline so dsRNA is only synthesised when tetracycline is added to the growth medium. T7 terminators prevent transcription from extending into rDNA.

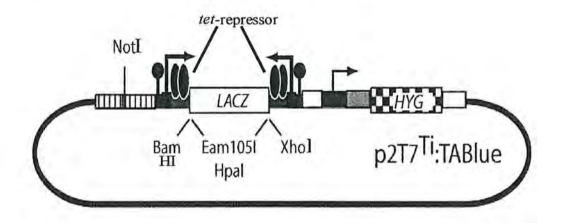


Figure 6. p2T7Ta blue vector. Two promoters for T7 polymerase are shown as big arrows facing each other, *Eam*11051 cloning site and hygromycin resistance gene for selection of transformants.

### 2.6.4.2 Stem-loop vectors

Stem-loop vectors such as pLEW79, pLEW100, pHD1146 and pHD1336 (Wirtz et al., 1999; Estévez et al., 2003) are also used (Appendix 3). They involve cloning of the insert twice in opposite direction on either sides of a 'stuffer' fragment. A representative map is shown in Figure 7. Within the transfected cells, a stem loop of complementary RNA is formed that can be 'Diced' as shown in Figure 8.

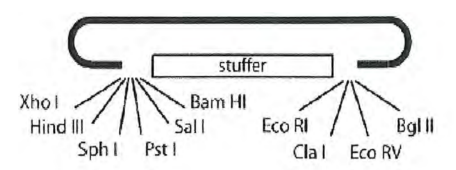


Figure 7. Vector for stem-loop cloning. The two multiple cloning sites for cloning of the insert are in opposite direction.

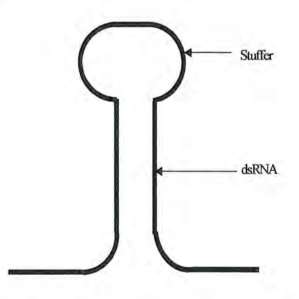


Figure 8. Double strand RNA (dsRNA) stem-loop. The dsRNA is formed in transfected cells. The siRNAs are formed from 'diced' dsRNA.

These vectors allow the cloning of the insert, which then undergoes homologous recombination in rDNA loci when linearized with *Not*I. The transcription occurs on induction with tetracycline, hence producing mRNA homologous to target the gene from which siRNA is produced.

# CHAPTER THREE MATERIALS AND METHODS

## 3.1 Data mining for T. brucei trans-splicing and polyadenylation homologs

In order to identify homologs of *T. brucei* genes for trans-splicing and polyadenylation, basic local alignment search tool (BLAST) searches (Altschul *et al.*, 1990; Altschul *et al.*, 1997) were done in GeneDB (Hertz-Fowler *et al.*, 2004), the repository of genome data for *T. brucei*, *T. cruzi* and *Leishmania major*. *T. cruzi*, human, and/or yeast splicing and polyadenylation factors were used to query the database. The accession numbers of nucleotide and protein sequences used in the search and *T. brucei* homologs generated are shown in Appendix 1. The search results were used to design primers, which were subsequently used to recover the genes from *T. brucei* by PCR.

#### 3.2 Trypanosome strain

The trypanosome strain used in this study was KETRI 3741 (MHOM/UG/72/KETRI 3741). It is a *T. b. rhodesiense* derived from a clone, KETRI 3666. These were grown in mice. KETRI 3666 is a derivative of KETRI 2537 that was isolated from human in 1972 in Busoga, Uganda. It was isolated by inoculation of whole blood from patient into monkey.

#### 3.3 DNA preparation

KETRI 3741 *T. b. rhodesiense* strain was grown in mice and purified using anion exchange chromatography (Lanham and Godfrey, 1970). DNA extraction was conducted according to the methods of Van der Ploeg *et al.*, (1982) with few modifications. Briefly, the column-purified trypanosomes were harvested by centrifugation (3000 rpm, 10 min, 4°C) in a refrigerated microcentrifuge (Heraeus fresco Biofuge, Kendro Laboratory Products GmbH, Germany) and washed once with phosphate buffered saline (PBS) (Appendix 2). The cells were resuspended in 200µl of cell lysis solution (Appendix 2), transferred into 1.5 ml microcentrifuge tube and incubated at 37°C overnight. The lysate was phenol extracted by adding an equal volume of phenol/chloroform/isoamylalcohol (25/25/1, v/v/v) for 5 minutes (Sambrook *et al.*, 1989). This was centrifuged (5000 rpm, 3 min, 25°C) in the refrigerated microcentrifuge and aqueous upper

layer pipetted out leaving the interphase behind. The interphase was re-extracted and upper phases pooled. DNA was precipitated by adding 0.1 volume of 3M sodium acetate (pH 5.2) and 2 volumes of absolute ethanol. This was kept at -20°C overnight. The DNA was then pelleted (13,000 rpm, 10 min, 4°C) in the refrigerated microcentrifuge, washed with 1ml of 70% ethanol, air-dried and redissolved in 200µl of sterile water.

# 3.4 PCR amplification of genomic DNA

Individual amplifications were carried out in 25µl reaction volumes. The reaction mixture contained as final concentrations, 10mM Tris-HCl, pH 8.3, 1.5mM MgCl<sub>2</sub>, 200µM each of the four deoxynucleoside triphosphates (dNTPs), 100ng of each primer, 1µl DNA template, 1 unit of *Taq* polymerase (MBI Fermentas, Lithuania) and 15.3µl of nuclease free water. The reaction mixture was placed in a thermocycler (PTC-100<sup>TM</sup>, MJ Research, Inc. Watertown MA.) and incubated at 94°C for 1 min, the 35 cycles of 94°C for 30 sec (denaturation), 55°C for 30 sec (annealing), 72°C for 90 sec (extension) and a final extension at 72°C for 10 minutes. 10µl of the amplification product was electrophoresed through a 1% ethidium bromide stained agarose gel and viewed under ultraviolet (UV) illumination. The primers and amplified homologs are shown in Table 3.

SPLICING	PRIMER SEQUENCES
FACTORS	(5'-3' direction)
P14	FOR: GCGAATGTCCGATGAGCC
	REV: ATCCTCGGCCTTGACCT
CPSF 30	FOR: GCATGTTTACTGACAACGCTGCC
	REV: CTGCCTTCCCGTTGCATCACCACG
AF 35	FOR: GCATGTATCARGAYCGYTGCA
	REV: TTTAAGGGGGCATTCGCGAGAGATG
AF65	FOR: ATGGGGGGGTGATAGTCGCGGACAC
	REV: GCTACTCCACAAAAACACGGGGCT
CFII al	FOR: GCATGTCTTCTAATTGTCGCAG
	REV: GCTGGGGGCTTGTAAAGGTGG
U1 70k	FOR: GCATGGAGGCGTCCCACCGAGT
	REV: CGAGCATCTTCCCTTCTG
CST 50	FOR: GCATGTCTGGAGAAAACTTG
	REV: CGCCGTGGTCCATGCCGG
Zn 1	FOR: GCATGTGAGAGAGGGGGGGGGGGA C
	REV: CAAGGAAAGAAACATATGCAG
Zn 2	FOR: GTCTGCCCGCAGCTTTCGCGC
	REV: CAACGAAAGAAACATATGCAG
0001145	FOR: GCTGCGGCCGCATGTATATACATCTTTTCTCCTAT
SF3b145	REV: CCAGCGGCCGCCTATCAAAATTTTGTTGGTGCACG
GE2L125	FOR: CCTGCGGCCGCATGGAGGAGACGTACAGTCCCTT
SF3b125	REV: GCAGCGGCCGCCTATCTCGCGTATTTCTTGCGGG
	FOR: CCTGCGGCCGCATGACAATCGCTGCACAGGGGG
Sf3b49	REV: GCAGCGGCCGCCTATTAAGCGCTTGAGGCGTGT
SE2L10	FOR: CCTGCGGCCGCATGGATGTGCCTGGTGAGCTTCTC
SF3b10	REV: GCAGCGGCCGCCTATCAAACACACTCACTGGTCAT

## 3.5 Purification of amplified products.

The full-length amplification products were purified directly using QIAquick Gel Extraction Kit (Qiagen, GmbH Germany). Briefly, the DNA fragments were excised from the agarose gel using a clean sharp scalpel and weighed. Three volumes of yellow core buffer (QG) were added to 1 volume of the gel and incubated at 50°C until the gel was completely dissolved. One volume of isopropanol was then added to the sample and mixed. Sample was then applied to the QIAquick spin column and centrifuged at 13000 rpm for 1 minute. The flow through was discarded and QIAquick column placed back in the same collection tube. 0.5 ml of buffer QG was added to the QIAquick column and centrifuged at 13000 rpm for 1 min. To wash, 0.75 ml of buffer PE was added to the QIAquick column, allowed to stand for 5 min and centrifuged at 13000 rpm for 1 min. The flow through was discarded and the QIAquick column centrifuged for an additional 1 min at 13000 rpm. The QIAquick column was then placed in a clean 1.5 ml centrifuge tube and 30µl of nuclease free water added to the center of the QIAquick membrane. Column was allowed to stand for 1 min and centrifuged at 13000 rpm for 1 min to elute DNA. The centrifugations were done in a table-top microcentrifuge (eppedorf centrifuge 5415 C, GmbH & Co. Bremen, Germany).

## 3.6 Cloning of PCR products

Purified PCR products were cloned in pGEM-T Easy vector (pGEM-T<sup>®</sup> EASY vector Systems kit, Promega Corp., Madison, WI, U.S.A; map shown in Appendix 3) using a shotgun cloning strategy. Ligations were performed in a total volume of 10µl. Purified PCR product (3µl) was mixed with 50ng (1µl) pGEM-T Easy vector, 5µl 2x rapid ligation buffer and 1 µl of T4 DNA ligase. The reactions were mixed by pipetting and incubated at 4°C overnight. Control reactions containing pGEM-T Easy vector alone were also carried out. Transformation of DH5α high efficiency competent cells was carried out using the ligation mixture. 50µl of cells were mixed with the ligation reaction and incubated on ice for 20 min. Uptake of the DNA was facilitated by 1 min heat shock at 42°C, followed by 5 min on ice. 950µl of room temperature SOC medium (Appendix 2) was added and incubated for 3 hrs at 37°C. 50µl of IPTG (Isopropyl-β-D-thiogalactopyranoside; 100mM) and 40µl of X-Gal (5-bromo-4-chloro-3-indoyl-β-D-galactopyranoside; 50mg/ml) was spread on *Luria-Bertani* (LB)-ampicillin (125µg/ml) agar plates (Appendix 2). The transformed cells were pelleted by centrifugation at 3000 rpm for 10

min, resuspended in 200µl of SOC medium and 100µl plated on each of two plates. Plates were incubated overnight at 37°C. Recombinant clones were identified by blue: white screening of the colonies. Potential positives were screened by PCR using the primers used for genomic DNA amplification to check for insertion of the correct template sequence. A single colony of the positive clones was inoculated in 5ml LB medium (Appendix 2) containing ampicillin (125  $\mu$ g/ml) in a falcon tube and incubated overnight at 37°C with shaking.

#### 3.7 Purification of plasmid DNA

Plasmid DNA was purified using QlAprep® Spin Miniprep kit (Qiagen, GmbH Germany) according to manufacturers protocol. Briefly, overnight cultures were pelleted by centrifugation at 4000 rpm for 15 min on a refrigerated centrifuge (GEMCO refrigerated centrifuge, Sigma-Aldrich, St. Louis, MO). Pelleted bacterial cells were resuspended in 250µl buffer P1 containing RNase A, mixed and transferred into a 1.5ml microcentrifuge tube. Following this, 250µl of lysis buffer (P2) was added to the suspension and mixed gently by inverting the tube 4-6 times. The lysate was neutralized by addition of 350µl buffer N3 and mixed immediately by inversion until a cloudy precipitate was obtained. This was centrifuged for 10 min at 13000 rpm. The supernatant was applied to the QlAprep column by pippeting and centrifuged for 1 min at 13000 rpm. The flow through was discarded. The QlAprep spin column was washed by adding 0.5ml buffer PB, recentrifuged for 1 min and flow through discarded. Bound plasmid DNA was washed by adding 750µl of buffer PE and centrifuged for 1 min. The flow through was discarded and re-centrifuged for an additional 1 min to remove residual wash buffer. The QlAprep column was placed in a clean 1.5ml microcentrifuge tube and DNA eluted with 30µl of nuclease free water. DNA was stored at -20°C. The centrifugations were done in a table-top microcentrifuge (eppedorf centrifuge 5415 C, GmbH & Co. Bremen, Germany).

#### 3.8 Sequencing

Automated sequencing was undertaken for the purified plasmid with PCR inserts using dideoxy chain termination method (Sanger *et al.*, 1977) in an ABI 3100 sequencer (Applied Biosystems, Foster City, CA, USA) using appropriate fluorescent labelled terminators.

## 3.9 Expression and purification of recombinant protein

#### 3.9.1 Determination of open reading frames (ORFs)

The nucleotide sequences generated from the cloned products were used to query the *T. brucei* database GeneDB at http://www.genedb.org to evaluate similarity and therefore infer homology with sequences in the database. The sequences were translated using translation tool at Expert Protein Analysis systems (Expasy -http://us.expasy.org) to determine the ORFs. ORF specific primers containing restriction sites for *Not*I were designed to allow their restriction ligation into pET-28a expression vector (Appendix 3). All the factors except Zn2, SF3b10, SF3b49, SF3b125 and SF3b145 were ligated into pET-28a vector.

# 3.9.2 Amplification and purification of cloned inserts

Amplifications were carried out in 25µl reaction volumes. The reaction mixture contained as final concentration 10mM Tris-HCl, pH 8.3, 1.5mM MgCl<sub>2</sub>, 200µM each of the four dNTPs, 100ng of each primer, 0.1µl template, 1 unit of Taq polymerase (MBI Fermentas, Lithuania) and 16.3 µl of nuclease free water. The reaction mixture was placed in a thermocycler (PTC-100<sup>TM</sup>, MJ Research, Inc. Watertown MA.) and incubated at 94°C for 2 min, then 35 cycles of 94°C for 30 sec (denaturation), 55°C for 30 sec (annealing) and 72°C for 120 sec (extension) and a final extension at 72°C for 10 minutes. 20µl of the amplification product were electrophoresed through a 1% ethidium bromide stained agarose gel and viewed under ultraviolet (UV) illumination. The amplification products were purified using QlAquick Gel Extraction Kit (Qiagen, GmbH Germany) as above.

#### 3.9.3 Restriction digestion and ligation into pET-28a

A restriction digestion reaction in a volume of 20µl was prepared, with the reaction mixture containing 8.0µl of purified amplification product, 2.0µl of 10X buffer, 20 units of *Not*I (MBI Fermentas, Lithuania) and 9.8 µl of nuclease free water. The reaction mixture was incubated for 4 hrs at 37°C and was thereafter gel purified using QlAquick Gel Extraction Kit (Qiagen, GmbH Germany). A similar restriction digestion and purification reactions were carried out for 4.0µl of pET-28a vector using *Not*I (MBI Fermentas, Lithuania).

Ligation was conducted using Novagen pET-28 system (Biosciences, Inc, Darmstadt, Germany) according to manufacturers protocol. This vector allows the expression of a fusion protein tagged to six histidine amino acids (the His-tag) allowing easy purification on nickel columns. In brief, a reaction containing  $4\mu$ l of *Not*I linearized pET-28a vector,  $8\mu$ l of *Not*I digested insert,  $2\mu$ l of 10X buffer and 6 units of T4 DNA ligase (MBI Fermentas, Lithuania) was set up and incubated overnight at 16°C.  $5\mu$ l of the reaction mixture was used to transform DH5 $\alpha$  high efficiency competent cells, plated on LB-kanamycin ( $50\mu$ g/ml) plates and incubated overnight at  $37^{\circ}$ C. Colonies were PCR screened using insert specific primers. Positive colonies were grown on LB-kanamycin liquid medium ( $50\mu$ g/ml) with overnight shaking and plasmid purified using QlAprep Spin Miniprep kit (Qiagen, GmbH Germany) as above.

## 3.9.4 Expression and induction

Transformation of BL21 bacterial cells was carried out using 1µl of purified recombinant plasmid as described before (section 3.6). This was platted on LB-kanamycin plates ( $50\mu g/ml$ ) and incubated overnight at 37°C. A single positive colony was used to inoculate 10ml of LB-kanamycin medium and incubated at 37°C with shaking. 1ml of the culture was used to inoculate 100ml of LB-kanamycin medium and incubated at 37°C with shaking at 200 rpm. At OD<sub>600</sub> of 0.5 (Beckman DU 6480B spectrophotometer, Beckman Coulter, Inc. Fullerton, CA.), the culture was split into two equal parts. To one part, IPTG was added to a final concentration of 1mM (induced) and the other served as an uninduced control. Both were further incubated for 4 hrs. The bacterial cells were harvested by centrifugation at 4000 rpm for 15 min (GEMCO refrigerated centrifuge, Sigma).

## 3.9.5 Extraction of total cell proteins

Harvested cells were resuspended in 2ml of 1X PBS buffer.  $200\mu$ l of 4X SDS sample buffer (Appendix 2) was added and cells sonicated for 10 sec at 5-µm amplitude for 10 cycles using Soniprep 150 ultrasonic disintegrator (Sanoy, Integrated Services, TCP Inc.). Immediately the samples were heated at 85°C for 3 min. The prepared total cell protein was separated on 4 – 20% SDS PAGE gels.

# 3.9.6 Purification of recombinant proteins

The recombinant P14 and CPSF30 proteins expressed in BL21 bacterial cells were purified using Ni-NTA Spin Kit (Qiagen, GmbH Germany) according to the manufacturers protocol. Briefly, cells were harvested by centrifugation at 5000 rpm for 15 min (GEMCO refrigerated centrifuge, Sigma). The pellet was resuspended in 1 ml lysis buffer (Appendix 2) and incubated for 1 hr with agitation at room temperature. The lysate was thereafter centrifuged at 9500 rpm for 30 min at 25°C (GEMCO refrigerated centrifuge, Sigma). 20µl of the clear lysate was used as crude extract for SDS-PAGE analysis.

Ni-NTA spin column was equilibrated with 600µl of lysis buffer by centrifugation at 2000 rpm for 2 min in a microcentrifuge. Subsequent centrifugations were done at the same conditions. 300µl of lysate containing 6xHis-tagged recombinant protein was applied onto the Ni-NTA spin column, centrifuged and flow through collected. The column was washed three times with 600µl of wash buffer (Appendix 2) by centrifugation. The 6xHis-tagged recombinant was eluted twice with 150µl of elution buffer (Appendix 2) by centrifugation. The protein concentration was determined and protein used for antibody generation and SDS-PAGE analysis.

# 3.9.7 Determination of protein concentration

The purified protein was dialysed in 1X PBS buffer and concentrated. The protein concentration was determined using bicinchoninic acid (BCA\*) Protein Assay Reagent kit (Pierce, Rockford, Ill. USA). Briefly, working reagent was prepared by addition of 1 part of reagent B to 50 parts of reagent A and mixed. 0.1 ml each of standard, concentrated protein (serially diluted in 1X PBS buffer) and a blank (diluent/ 1X PBS) were pipetted into test tubes. 2 ml of working reagent was added to each tube and incubated for 30 min at 37°C. This reaction was cooled to room temperature and absorbance read at 562 nm on a Beckman DU 6480B spectrophotometer (Beckman Coulter, Inc. Fullerton, CA.). Concentration was then determined from a standard curve.

## 3.10. Antibody generation

Ten millilitres of pre-immune rabbit blood was obtained by bleeding aseptically from the central ear artery using a hypodermic syringe and needle. 1000µg of the purified recombinant P14 protein in Freund's complete adjuvant (Pierce and Warriner, Chester, UK) was injected intramuscularly into the right thigh muscle. After 2 weeks, a boost was given on the left thigh by injecting 700µg of the recombinant protein in Freund's incomplete adjuvant (Pierce and Warriner, Chester, UK). The rabbit was bled two weeks later and the blood stored at 4°C for two days. This was centrifuged at 13000 rpm for 10 min and serum pippeted out. The serum was stored at -20°C.

#### 3.11 Sequence analysis

Nucleotide sequences of cloned inserts were translated to protein using the translation tool at Swiss Bioinformatics Institute website - Expasy (http://us.expasy.org) (Bairoch, 1991). The generated amino acid sequences were used to query *T. brucei* database at GeneDB to determine E-values (the probability that the alignment is due to chance) at statistical significance threshold of 0.0001. The amino acid sequences were also compared with those of *Homo sapiens*, *Leishmania major* and *T. cruzi* via alignment with ClastalW (Thompson *et al.*, 1994; Altschul *et al.*, 1997) at http://www.ebi.ac.uk/clustalw/, biological sequence alignment editor – BioEdit (Tom Hall, *Ibis Therapeutics* Carlsbad CA.) and Needleman-Wunsch global alignment (NeedleN) (Needleman and Wunsch, 1970; Kruskal, 1983; Rice *et al.*, 2000). The amino acid sequences were used to query various public protein databases to identify conserved domains. These included the integrated resource of protein domains and functional sites, InterPro (Apweiler *et al.*, 2000; Mulder *et al.*, 2005; http://www.ebi.ac.uk/interpro/), prosite (Falquet *et al.*, 2002; Hulo *et al.*, 2004; http://au.expasy.org/tools/scanprosite/), MotifScan (Falquet *et al.*, 2002; http://myhits.isb-sib.ch/cgi-bin/motif\_scan) and Pfam (Sonnhammer *et al.*, 1998; Bateman *et al.*, 2004; http://pfam.wustl.edu/hmmsearch.shtml).

## 3.12 RNA interference

## 3.12.1 Plasmid constructs for RNAi

A 30µl restriction digestion reaction was set up consisting of 1µl of p2T7Ta vector, 3 µl of 10X buffer, 2µl Eam1051 restriction enzyme (MBI Fermentas, Lithuania) and 24 µl of nuclease free

water. This was incubated at 37°C for 4 hrs. All the reaction mixture was electrophoresed through a 1% ethidium bromide stained agarose gel and viewed under ultraviolet (UV) illumination. The band containing the linearized plasmid was purified using QlAquick Gel Extraction Kit (Qiagen, GmbH Germany) as above.

Primers (Table 4) were designed to amplify products 500bp or less. The amplification was carried out in a 25µl reaction volume. The reaction mixture was as in section 3.9.2 above. 20µl of the amplification product were electrophoresed through a 1% ethidium bromide stained agarose gel and purified using QlAquick Gel Extraction Kit (Qiagen, GmbH Germany). The linearized plasmid generates T-overhangs that allow T-A cloning of PCR products.

Purified amplification products were ligated into linearized p2T7Ta blue vector and incubated at 4°C overnight as described before (section 3.9.3). 5µl of the ligation reaction was used to transform competent DH5 $\alpha$  cells and plated on LB-ampicillin plates (100µg/ml) containing 150µl of IPTG (isopropylthio- $\beta$ -D-galactosidase; 100mM) and 40µl of X-Gal (5-bromo-4chloro-3-indoyl- $\beta$ -D-galactosidase; 50mg/ml) and incubated overnight at 37°C. White colonies were screened using insert specific primers. Positive clones were grown in LB-ampicillin medium at 37°C overnight. The plasmid was purified using QlAprep Spin Miniprep Kit (Qiagen, GmbH Germany) as described above.

A restriction digestion reaction in a volume of 50µl was set up. The reaction mixture contained 14µl of purified plasmid, 5µl of 10X buffer, 30µl of nuclease free water and 20 units of *Not*I (MBI Fermentas, Lithuania). The reaction mixture was incubated at 37°C for 4 hrs. Linearized plasmid DNA was precipitated by adding 0.1 volume of 3M sodium acetate (pH 5.2) and 2 volumes of absolute ethanol. This was kept at -20°C overnight. The DNA was then pelleted (13,000 rpm, 10 min, 4°C) in a refrigerated microcentrifuge (Heraeus fresco Biofuge, Kendro Laboratory Products GmbH, Germany), washed with 1 ml of 70% ethanol, air-dried and redissolved in 20µl of nuclease free water.

FACTORS	<b>PRIMER SEQUENCES</b> (5'-3' direction)
P14	FOR: GAGAAGATCTGCATGCCTTGTAACGGGGGATTCCC
	REV: CGGAATTCGTCGACATCCTCGGCCTTGACCTCCG
CPSF30	FOR: GAGAAGATCTGCATGCTGCAAGCATTGGTTTCGCGG
	REV: GGAATTCGTCGACTTGCATACCCGGACAGTTAGGTG
U170k	FOR: GAGAAGATCTGCATGCACCGAGTACGATAAGGACAC
	REV: CGGAATTCGTCGACAGCTTCCGCGGCACGCTCATAG

## 3.12.2 Trypanosome culture

Procyclic forms of *T. brucei* 1313-1333 cell line (Alibu *et al.*, 2004) was cultured in SDM79 (Brun and Schönenburger, 1979) supplemented with 10% fetal calf serum (FCS) at 27°C in the continuous presence of 0.5  $\mu$ g/ml and 15 $\mu$ g/ml of phleomycin and G418 respectively (Wang *et al.*, 2000; Allen *et al.*, 2003; Bakshi and Shapiro, 2004). The cells were cultured to a final density of between 5 x 10<sup>5</sup>/ml and 5 x 10<sup>6</sup>/ml with the cells number determined with a haematocrit counter.

#### 3.12.3 Transfection

The trypanosome cultures were centrifuged at 3000 rpm for 10 min (Koolspin up centrifuge, Burkard scientific, Uxbridge, Middx, UB8 2RT, UK) and supernatant filtered and stored in a new sterile tube. The cells were washed once in 5ml of Zimmerman post-fusion medium (ZPFM) electroporation medium (Appendix 2) and resuspended in 2.5ml of the same medium. 50µg of *Not*I linearized plasmid was added to 0.5ml of the 2 x 10<sup>7</sup> cells in 4 mm BTX cuvette (Harvard Apparatus, Holliston, Ma.) and mixed. Transfection was carried in a Gene Pulser<sup>TM</sup> electroporator (Bio-Rad, Hercules, Ca.) at 1.5kV in a resistance-timing mode of R2 (24 ohms). Immediately after transfection, the cells were transferred into 5ml of SDM-79 medium supplemented with G418 and phleomycin and incubated overnight at 27°C. Transfections were done in pairs for each gene. Circular plasmid DNA was used as control.

## 3.12.4 Selection of transformants

The cells were diluted to  $2 \times 10^5$ /ml using conditioned medium that was saved the previous day and selective antibiotic (hygromycin) added to a concentration of  $50\mu$ g/ml. The culture was plated on a 24 well plate in a four serial fold dilutions. The plates were put in a modular chamber and incubated. The transformants were cultured until a density of  $2 \times 10^6$ /ml was achieved, transferred into new wells and some stored in liquid nitrogen (-196°C). The cells were maintained in conditioned media with hygromycin.

#### 3.12.5 Induction of transformants

Induction of dsRNA was done by culturing the clonal population in the same medium with 100ng/ml of tetracycline. The cultures were monitored to detect morphological changes and the growth rate.

#### 3.13 Western blotting for P14 transformants

Mid-log phase cells (2 x 10<sup>6</sup>) were harvested, washed in 1x PBS buffer (Appendix 2) then resuspended in 50µl SDS-polyacrylamide sample buffer (Appendix 2). Control and transformants were not counted because they tended to clump together making counting virtually impossible. These were incubated at 95°C for 10 minutes. The samples were electrophoresed on 4-20% SDS-PAGE gel and transferred to nitrocellulose membrane (Bio-Rad Laboratories, Hercules, CA.) by electroblotting in transfer buffer (Appendix 2) at 168 mA for 2hrs. Nonspecific sites on the membranes were blocked by incubation in blocking buffer (Appendix 2) for 2hrs. The membrane was then incubated overnight in TBS buffer (Appendix 2) at intervals of 5 min followed by incubation for 1hr with alkaline phosphatase conjugated secondary antibody (goat anti-rabbit IgG, Sigma-Aldrich) diluted at 1:1000 (v/v). This was subsequently washed twice in wash buffer for 5 min each followed by a third wash in substrate buffer (Appendix 2) for 10 min. Bound antibodies were detected by incubation with Nitro Blue tetrazolium (NBT) (Sigma-Aldrich) and 5-Bromo-4-chloro-3-indolyl phosphate (BCIP) (Sigma-Aldrich) in the dark. The reaction was stopped by adding distilled deionised water.

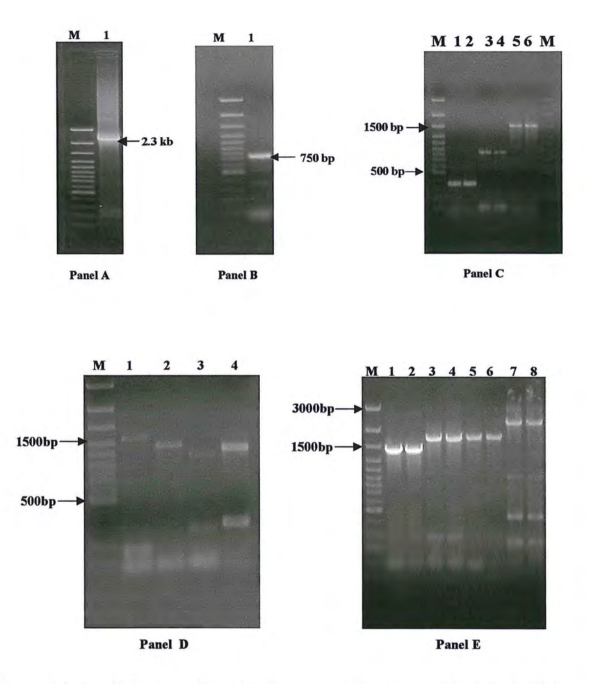
# CHAPTER FOUR RESULTS AND DISCUSSION

# 4.1 Data mining

A clean dataset of *T. brucei* nucleotide and protein sequences related to spliceosome was generated from sequences available at GeneDB. The searches had motifs also identified in Saccharomyces cerevisie, Caenorhabditis elegans, *T. cruzi*, *L. major* and *H. sapiens*. Combination of methods such as reciprocal BLAST hits, presence of protein domains and motifs, and phylogenetic trees exhibited orthologous relationship among these genes and those of better studied organisms such as yeast and human. The accession numbers of the data mining results are shown in Appendix 1.

## 4.2 Amplification of genomic DNA

Amplification of auxiliary factors AF<sup>35</sup> and AF<sup>65</sup> gave products of approximately 2500 and 750 bp respectively (Figure 9 Panel A and B). P14, cleavage and polyadenylation splicing factor 30 (CPSF 30) and cleavage stimulating factor 50 (CstF 50) amplification gave products of approximately 350, 800 and 1500 bp respectively (Figure 9 Panel C). PCR amplification of cleavage factor II (CFII a1), U1-70k, zinc finger 1 (Zn 1) and zinc finger 2 (Zn 2) gave products of between 1200 and 1300 bp (Figure 9 Panel D). Amplification products for splicing factors (SF) SF3b 145, SF3b 49, SF3b 125 and SF3b 10 gave products of between 1200 and 3000 bp (Figure 9 Panel E).



**Figure 9**: 1.0% ethidium bromide stained agarose gel showing amplification of splicing and polyadenylation factors from genomic DNA. Panel A: U2AF<sup>65</sup>; Panel B: U2AF<sup>35</sup>; Panel C: lanes 1-2, P14; lanes 3-4, CPSF 30; lanes 5-6, CstF 50; Panel D: lane 1, CFII-a1; lane 2, U1-70k; lane 3, Zn 1; lane 4, Zn 2; Panel E: lanes 1-2, SF3b 145; lanes 3-4, SF3b 49; lanes 5-6, SF3b 125; lane 7-8, SF3b 10. In all panels, M represents a 1 Kb ladder for panel A and E, and 100bp plus size ladder for B, C and D.

# 4.3 DNA sequencing

The genes associated with both E and A complex were sequenced. CFII-a1, Zn 1, Zn 2, U1-70k, AF<sup>35</sup> and AF<sup>65</sup> for the E complex while P14, SF3b 10, SF3b 49, SF3b 125 and SF3b 145 for the A complex. Cleavage and polyadenylation factors CFSF 30 and CstF 50 were also sequenced. The insert sizes from the sequencing and E-values from searches at geneDB using amino acid sequences from translated nucleotide sequences as query are shown in Table 5.

FACTORS	EXPECTED SIZE	SEQUENCING	E - value
	(from GeneDB)	(bp)	(at 0.0001)
	E Co	omplex	
CFII-al	1272	1273	3.7e-227
Zn 1	1047	943	6.8e-148
J1-70k*	1131	1103	2.2e-129
4F <sup>35</sup>	741	740	4.1e-111
AF <sup>65</sup>	2352	2312	9.8e-275
$Zn 2^{\Psi}$	1047	1039	8.4e-215
	А	Complex	
			-
P14*	507	355	1.0e-41
Sf3b 145	1434	1433	1.2e-197
Sf3b 49	1701	1688	4.1e-217
Sf3b 125	1707	1699	3.0e-280
Sf3b 10	2199	2200	0.00
	Cleavage and Pe	olyadenylation Factor	
CPSF 30*	879	833	8.0e-161

Table 5. Sequence from data mining, sequencing and E-values

\* Factors used for RNAi experiments

1566

CstF 50

 $^{\Psi}$  Nucleotide sequence was used instead of amino acid sequence due to short ORFs resulting from multiple stop codons, presumably resulting from sequencing errors

1550

2.7e-256

e = base 10

#### 4.4 Sequence analysis

The E-values of amino acid sequences (and nucleotide sequence for Zn 2) from the cloned factors were very low at statistical significance threshold of 0.0001 (Table 5). Comparison of amino acid sequences from the cloned genes with those from *H. sapiens*, *T. cruzi* and *L. major* are shown in Tables 6, 7 and 8. Percentage identity and similarity between cloned genes and those of *H. sapiens* were in the range of 15.4 - 31.1 and 22.8 - 49.3 respectively (Table 6). Human and *T. brucei* SF3b 10 homologs were incomparable since TbSF3b 10 had 732 amino acid residues while hSF3b 10 had only 86 amino acid residues. *T. cruzi* and *T. brucei* orthologs had percentage identity and similarity ranging between 52.8 - 83.0 and 61.7 - 88.8 respectively (Table 7). Results obtained for *T. brucei* and *L. major* orthologs varied between 22.8 - 66.5 and 32.3 - 76.7 for percentage identity and similarity respectively (Table 8).

Table 6. Comparison of the amino acid	sequences of cloned T. brucei factors and H. sapiens	
sequences.		

FACTORS	% IDENTITY	% SIMILARITY
SF3b125	22.3	31.7
SF3b49	19.9	29.5
SF3b145	15.4	22.8
P14	29.1	46.3
AF35	31.1	49.3
AF65	18.3	27.7
CstF50	22.4	38.3
CFII-a1	26.6	40.9
CPSF30	30.6	39.0

FACTORS	%IDENTITY	%SIMILARITY
SF3b125	69.3	78.9
SF3b49	71.7	78.6
SF3b145	59.1	70.9
SF3b10	53.9	68.5
P14	79.2	87.5
AF35	80.6	88.7
AF65	52.8	61.7
CstF50	67.6	77.3
CFII-a1	57.5	75.0
CPSF30	83.0	88.8

 Table 7. Comparison of the amino acid sequences of cloned T. brucei factors and T. cruzi sequences.

 Table 8. Comparison of the amino acid sequences of cloned T. brucei factors and L. major sequences.

FACTORS	%IDENTITY	%SIMILARITY
SF3b125	61.2	72.2
SF3b49	66.5	76.7
SF3b145	44.8	59.5
SF3b10	22.8	32.3
P14	40.0	50.3
AF35	63.7	72.2
AF65	30.8	43.4
CstF50	47.3	58.7
CFII-a1	36.3	54.4
CPSF30	47.4	59.4

## 4.5 Protein domains

# 4.5.1 U2 auxiliary factor 35 (AF35)

Genes homologous to *T. brucei* U2AF<sup>35</sup> for *T. cruzi*, *L. major* and *H. sapiens* had a central RNA recognition motif (RRM) flanked by two zinc finger motifs (Cx8Cx5Cx3H/CCCH). However, the human U2AF<sup>35</sup>, hU2AF<sup>35</sup> had an arginine-serine (RS) domain and a glycine tract at the C-terminus that were absent in the three kinetoplastids. Instead, *T. brucei* and *T. cruzi* accommodated a third and different zinc knuckles (CCHC/Cx2Cx4Hx4C) that is absent in *L. major*. The C-terminal of the kinetoplastids U2AF<sup>35</sup> had arginine and serine residues interrupted with other residues. Signalp domain was present in *T. brucei* and *T. cruzi* (Figure 10A). The amino acid alignment of the sequences exhibits significant similarity in the kinetoplatids (Figure 10B). The RRM domain characterized by two conserved sequence elements, RNP1 and RNP2 was conserved in the kinetoplastids and had helices. The tryptophan residue that interacts with the "groove" in U2AF<sup>65</sup> has been replaced with a lysine residue (green asterisk in Figure 10A) in all the kinetoplastids.

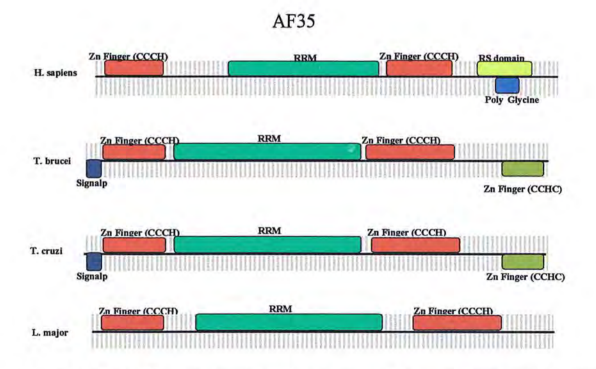


Figure 10A: U2AF<sup>35</sup> protein domains of cloned T. brucei, T. cruzi, L. major and H. sapiens

	β-1(RNP2)
Tb35	MYODRCI FFSKMGACRHGDGCIKVHVRPTTSPTVLFPFMYPNPAAI EH
Tc 3 5	MYODRCIFFSKMGACRHGDHCIKVHVRPTSSPTVLFPMMYPNPMALEH
Lm35	
Hs 3 5	MAEYLASIFGTEKDKVNCSFYFKIGACRHGDRCSRLHNKPTFSOTIALLNIYRNPONSSO
	te et et este este est est est est est e
	Helix α-A β-2 β-3(RNP1)
Tb 3 5	I QDREWNFHFERKYLRRHFEHFYKETWRTFME-LGRIAELRVVSNLGDHLLGNVYIKFED
Tc 3 5	I QDRQWDFHFDRKYLKRHFEHFYKETWRTFME-LGRIAELRVVSNLGDHLLGNVYIRFEE
Lm35	I KDREWNFELDKKYLKKHFEHFYKEVWRTFME-FGRI AELRVVSNLGDHLLGNVYIRFED
Hs 3 5	SADG-LRCAVSDVEMQEHYDEFFEEVFTEMEEKYGEVEEMNVCDNLGDHLVGNVYVKFRR
	·
	<u>Helix <math>\alpha</math>-B <math>\star</math> <math>\beta</math>-4 C C H</u>
Tb 3 5	SHDASRI VRELKAKKLNDI VLLPELSP VTNFAEAČCKEDLEGKČERGP QČNYLHI MKVSR
Tc 3 5	AADASHI ARELKAKKLNEI I LLPELSPVTNFADACCKEDLEGKCGRGAQCNYLHI I KVSR
Lm35	PQVATRI VKELRGKKLNDI I VLPELSPVTNFAEACCKEDLENRCQRGEQCNYLHI MKVSR
Hs 3 5	EEDAEKAVI DLNNRWFNGQPI HAELSPVT DFREACCRQYEMGECTRGGFCNFMHLKPI SR
	9 1 . 19 1 19 1 'essays's issell ''s so sailel iss
Tb 3 5	KL MEKLEKEQAKF WKKKEKHSSSSSSSRKRERSKERGGERSKERSKERGRDRQKSPRGYS
Tc 3 5	KL ME KL E KE QAK YWKKKEKHS RGS DRKRERS KDRGRERS RS PRP HP
Lm35	RLLEKLEKEQSKYWKKKERRHEHSSSDRKRSRSRSP
Hs 3 5	ELRRELYGRRRKKH <u>RSRSR</u> S <u>R</u> ERRS <u>R</u> S <u>RDRGRGGGGGGGG</u> GGGGGGGGGGGGGGGGGGGGGGGGG
	Polyelycine
	ССНС
Tb 3 5	SDMCHI CGKSGHI SRECPL-
Tc 3 5	S DL CHI CGKS GHI S RDCPL K
Lm3 5	ADTRW-
Hs 3 5	GGRERDRRS RDRERS GRF-
	RS domain

\*= Conserved amino acid; green box represent conserved RNA recognition amino acids

\* Lys (lysine) substitution for Trp (tryptophan).

Figure 10B. U2AF<sup>35</sup> amino acid alignment of T. brucei, T. cruzi, L. major and H. sapiens

# 4.5.2 U2 auxiliary factor 65 (AF65)

The three kinetoplastids lacked the N-terminal RS domain that is present in the  $hU2AF^{65}$  (Figure 11A). *T. brucei* however, had an arginine rich region at the N-terminal.  $hU2AF^{65}$  had three RRMs as compared to one in *T. brucei* and *L. major. T. cruzi* has two RRMs that are close. The amino acid alignment in Figure 11B is based on the signatures for RMM1, RRM2 and RMM3 from human  $U2AF^{65}$ .

**AF35** 

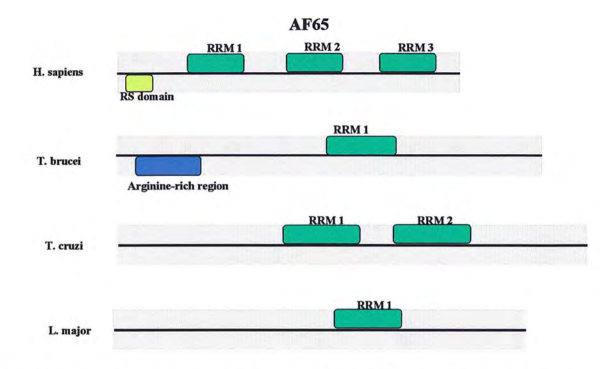
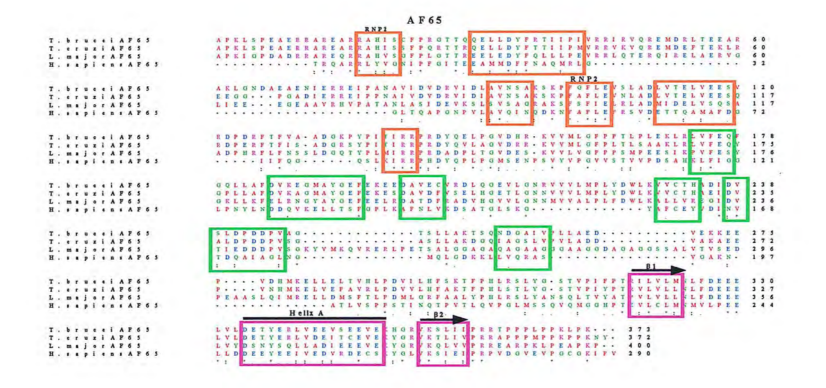


Figure 11A: U2AF<sup>65</sup> protein domains of cloned T. brucei, T. cruzi, L. major and H. sapiens



\*= Conserved amino acid, Orange= RRM1 green= RRM2 and violet= RRM3 boxes,  $\rightarrow$  =  $\beta$  sheets,  $\rightarrow$  = Helix Figure 11B. Structure-based multiple sequence alignment of U2AF<sup>65</sup> amino acid sequences of cloned *T. brucei*, *T. cruzi*, *L. major* and *H. sapiens*. RNP1 and RNP2 signature sequences in RRMs are indicated on top.

# 4.5.3 Cleavage stimulating factor 50 (CstF 50)

Comparison of CstF 50 homologs showed that they have a tryptophan-aspartic acid (WD) and signalp domains except in *T. cruzi* where signalp domain was absent (Figure 12A). The WD domain signature is conserved among the species (Figure 12B). However, in the human homolog, aspartic acid (D) residue is replaced with glutamic acid (E) residue.

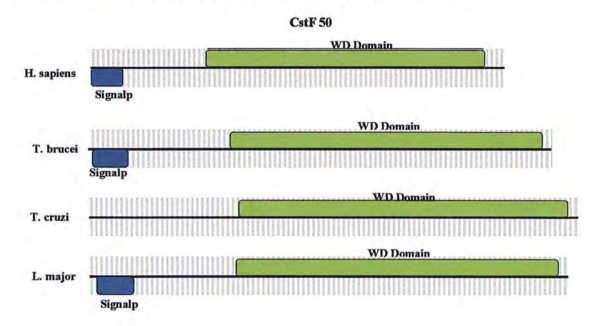


Figure 12A: Cleavage stimulating factor 50 (CstF 50) protein domains of cloned T. brucei, T. cruzi, L. major and H. sapiens

	Trp-Asp (WD) signature
T.bruceiCstF50	Trp-Asp (WD) signature
L.majorCstF50	VTSVVYSRTGNVVLTAGMDSTARLWDLRRL 30
T.cruziCstF50	SVKFSRTGNFILSAGMDSVARLWDLRR- 27
H.sapiensCstF50	SAIFSKNSKYILSSGKDSVAKLWEISTGRT 30
	* *** ** ** ** ** ** *

Figure 12B. CstF 50 tryptophan-aspartic acid (WD) signature

# 4.5.4 Cleavage factor II (CFII-al)

All the homologs of CFII-a1 had a pre-mRNA cleavage complex II protein Clpl domain. GTPase domain was absent only in *L. major*. However, *L. major* had a signalp domain, which was also present in *T. brucei*. ATP\_bind\_1 domain was only present in human protein and may be similar to AAA ATPase family domain in *T. cruzi*. P-loop domain was present only in *T. brucei* protein sequence (Figure 13).

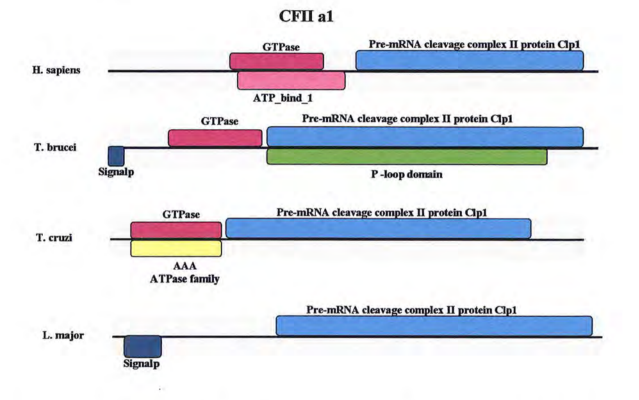


Figure 13: Cleavage factor II (CFII a1) protein domains of cloned T. brucei, T. cruzi, L. major and H. sapiens.

# 4.5.5 Splicing factor 125 (SF3b 125)

The four sequences had centrally located DEAD/D-box domain (Figure 14A), which is a conserved feature among the homologs (Figure 14B).

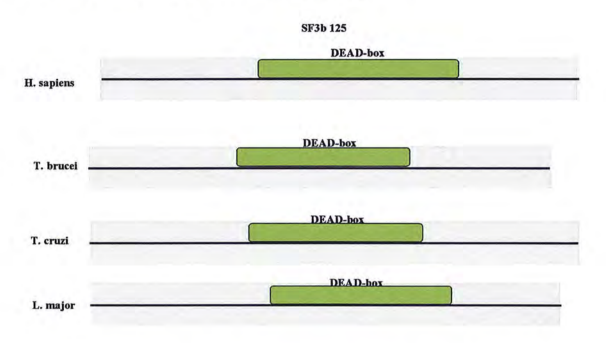


Figure 14A: SF3b 125 DEAD box domains of cloned T. brucei, T. cruzi, L. major and H. sapiens

T.bruceiSF3b125 T.cruziSF3b125 L.majorSF3b125 H.sapiensSF3b125 DEAD box signature NLHRVTY LVL DE ADR MLDMGF EP Q 24 NFF RVTY LVL DE ADR MLDMGF EP Q 24 NLL RVTY LVM DE ADR MLDMGF EP Q 24 NL QRVSY LVF DE ADR MFDMGF EY Q 24 \*: \*\* : \*\*\* : \*\*\*\*\* : \*\*\*\*\*

\* = Conserved amino acids

Figure 14B. SF3b 125 DEAD-box signature from cloned T. brucei, T. cruzi, L. major and H. sapiens.

# 4.5.6 Splicing factor 145 (SF3b 145)

The four species (*H. sapiens, T. brucei, T. cruzi* and *L. major*) had a proline rich (PSP) domain. However, the kinetoplastids had a signalp domain at the N-terminal while *H. sapiens* had a DNA binding splicing associated protein (SAP) domain in similar position. The human SF3b 145 had a DUF382 domain observably absent among the kinetoplastids (Figure 15A). The PSP domain signature was conserved amongst the four (Figure 15B).

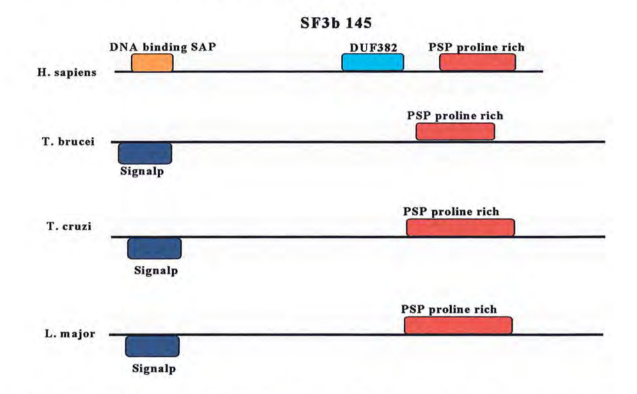


Figure 15A: Splicing factor (SF) 3b 145 domains of cloned T. brucei, T. cruzi, L. major and H. sapiens.

	PSP signature
T.bruceiSF30145	
T.cruziSF3b145	KNHHEPGHLSQRLRAALOM-GPHS-PPEWLYGMQAMRRLPPAYPTIKVPGLNAPIPP 55
L.majorSF3b145	KAHHTPGVLSKRLRQALGI-GPTA-PPPWLYSMQIMRRLPPAYPDLRIPGINAPIPA 55
H.sapiensSF3b145	LKEKKREDLSDELRISLEMPVGPNAHKVPPPWLIAMORYG-PPPSYPNLKIPGINSPIPE 59
	** ** ** ** ** ** *** ** ** ***

\* = Conserved amino acids

Fig 15B. Proline rich region (PSP) signature of SF3b 145

# 4.5.7 Splicing factor 10 (SF3b 10)

The hSF3b 10 sequence is small (86 amino acid residues) thus incomparable to the kinetoplastid homologs. The kinetoplastids had MutS III and V domains, DNA binding domain for DNA mismatch repair and ATPase domain for DNA mismatch repair (Figure 16A). Both *T. brucei* and *T. cruzi* had a signalp domain while *T. cruzi* and *L. major* had a P-loop containing nucleoside triphosphate hydrolases. The DNA mismatch repair MutS family signature was present in the three kinetoplastid sequences (Figure 16B).

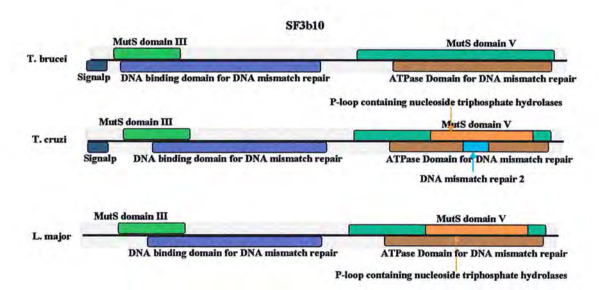


Figure 16A: SF3b 10 domains of cloned T. brucei, T. cruzi, L. major and H. sapiens

DNA mismatch repair MutS family signature	
ESDEGAARMLLVIDEFGKGTLSVDGA- 26	
EGGAGSGRSLLVLDEFGKGTLSLDGAA 27	
DGSRMAGRALVLVDEFGRGTSPEDGC- 26	
* * ******** **	

\* = Conserved amino acids

T.bruceiSF3b10 T.cruziSF3b10 L.majorSF3b10

Figure 16B: DNA mismatch repair MutS family signature

# 4.5.8 Splicing factor 49 (SF3b 49)

Homo sapiens SF3b 49 had two RRMs as compared to four RRMs in the kinetoplastids. Nterminal signalp domain as well as poly (A)-binding protein, h (PABP, h) and PABP domains were only present in the kinetoplatids (Figure 17).

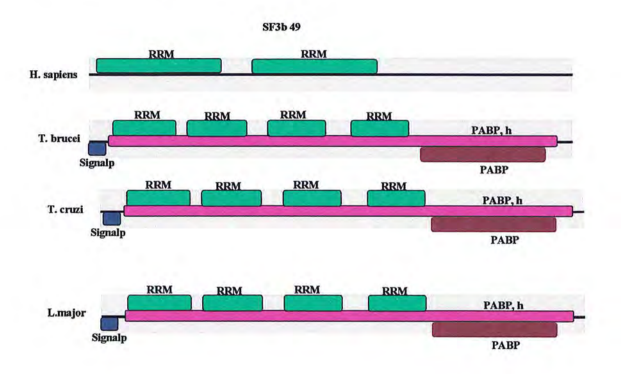


Figure 17: SF3b 49 domains of cloned T. brucei, T. cruzi, L. major and H. sapiens

# 4.5.9 P14

All the sequences (*H. sapiens*, *T. brucei*, *T. cruzi* and *L. major*) had a single RRM (Figure 18). *T. brucei* and *L. major* also had a signal pdomain that was absent in *H. sapiens* and *T. cruzi*.

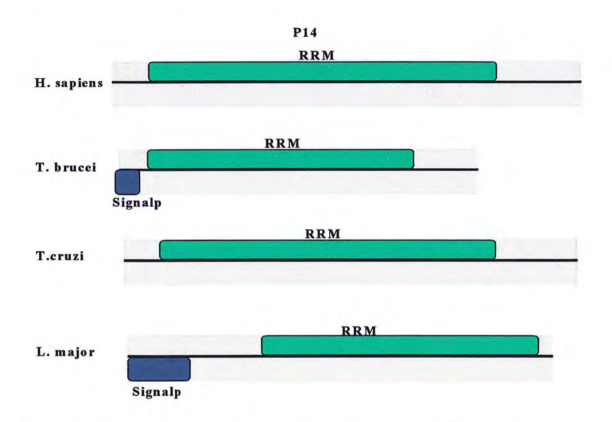


Figure 18: RRM domains in P14 of cloned T. brucei, T. cruzi, L. major and H. sapiens

# 4.5.10 Cleavage and polyadenylation factor 30 (CPSF 30)

CPSF 30 had zinc fingers of type CCCH and CCHC (zinc knuckle). CPSF 30 of *T. brucei*, *L. major* and *T. cruzi* had five CCCH zinc fingers and two zinc knuckles while *H. sapiens* had four CCCH zinc fingers and one zinc knuckle (Figure 19). Various deletions and additions are noticeable in the homologs. The *L. major* CPSF 30 is longer with several glycine residues at the C-terminal.

	CPSF30
	ССН
T. brucei CPSF30	METDNAARTSLAFEDTLPVEAPATAKESEICOPFOHGECENGAACPEEHV 50
T. eruzi CPSF30	MFTDSAARTSLAFEDALPDEPHATVKRR
L. majer CPSF30	MEVDDAAGTHEDEEDTLPKEOPRAEKKLEICODFORGRCRLGDACPORHI 50
H. sapiens CPSF30	MOETIAS VDHIKFDLEIAVEOOLGAOPLPFPGMDKSGAAVCEPFLKAACOKGGMCPFRHI 60
1.1.2.2.4.1.0.1.2.1.1.1.1.1.1.1	
	с с с н с с с н
T. brucei CPSF30	LS OF KS MRLEVCKHWP RGACVNGENCVYLHEYD DRYVP ACAPYORL GECS NPECFPOHVY 110
T. cruzi CPSF30	I S OF & T MR LEVC & HWLR GACVN GENCLYL HEYDDRYVPLCAFYORL GECTNPECPFOHVI 110
L. major CPSF30	ISAYRTVOTEVCEHWLEGACVNGDNCLYLHAYDNEYVFOCAFFEEVGECTNFECFFLHTE 110
H piens CPSF30	5 GEKTVVCEHWLEGLCEEGDOCEFLHEYDMTEMPECVEYSEFGECSNEECPFLHID 116
	с с с н с с с н
T. brucei CPSF30	OVEROPECAAYERGFCPLOPECELEHVFE. PPCVCYLTGFCPLOPECAS OHPVQQLYNEN 169
T. eruzi CPSF30	QVEROPECAAVREGFCFLGFECHLERVFE- PSCPFYMAGFCFLGFECTMGHFVOELYNEN 169
L. major CPSF30	PNESOPECAAVREGECELGEECELEBVERESACFYYLAGECELGEECELGEEIOERYDED 170
H. sapiens CPSF30	PESKIKDCPWYDEGFCKHGPLCEHEHTEE. VICVNYLVGFCPEGPSCKFMHPEFELPMOT 175
and the off of the second second	the second second second second second second second second
	ССНС
T. brucei CPSF30	DVSERLEORMLIER ADDPSENKNATCYR CFDPGHLSPNCPGMOSGLLERLLMALOE 225
T. eruzi CPSF30	SVSERLEQRMLIER VDDPTFNKNATCYR CFDPGHLSPNCPGVOSGLLERMLMAVOE 225
L. major CPSF30	AVSKRILARMIVER ADDPTFNRSATCYRAGCFDPGHLAPDCPGPOHSVLHKALGEIGE 228
H. sapiens CPSF30	TEOPPLPOOTOPPAKOSNNPPLORSSSLIOLTSONSSPNOORTPOVIGVMOS 227
27 08-00 0 06 06 0 000	the second se
	ССНС
T. brucei CPSF30	PG EQLYFQSDGRAARKCCFFCGEEGHEVRDCFKKQKPQQWGGHER 270
T. eruzi CPSF30	PG EQLYPQSDGRSARKCCFFCGDESHEVRDCPKKPKNHQWHT
L. major CPSF30	PGQFTGGLENSGHGGGGGGGSHERCFLCDQEGHTVKDCPMNTHETQHEAGGGGFGAFHGS 288
H picas CPSF30	QNSSAGNROPEPLEQVICYKCOEKGRYANECIKOHLAFLSGQ
	a second s
T. brucei CPSF30	GDATGRQ
T. eruzi CPSF30	A 271
L. major CPSF30	GGGAGVGNEERLEEGVAGGGGGGGSAGAGAGAGGGGGGGGGGEEEFGEDHY 339
H. sapiens CPSF30	

\* = Conserved amino acids

Figure 19: Zinc fingers of CPSF 30.

### 4.6 T. brucei U2 auxiliary factor 65 (TbU2AF65)

Analysis of TbU2AF<sup>65</sup> sequence and the well characterized hU2AF<sup>65</sup> showed conservation of some of the amino acid residues that could be associated with "groove" formation. Some were not conserved as shown in Figure 20 and Table 9 below. Of particular interest are two proline residues with six amino acid spacer. These are important in "groove" formation in hU2AF<sup>65</sup> hence highly conserved (Kielkopf et al., 2001). Sequence alignment between hU2AF<sup>65</sup> and TbU2AF<sup>65</sup> gave two proline residues at positions TbU2AF<sup>65</sup>Pro162 and TbU2AF<sup>65</sup>Pro169 with a six amino acid residue spacer. The other residues important in "groove" formation were substituted with amino acids of similar properties except in TbU2AF<sup>65</sup>Pro160 (Table 9). Sequence comparison between hU2AF<sup>65</sup> and the two Trypanosoma species homologs (TbU2AF<sup>65</sup> and TcU2AF<sup>65</sup>) and later between hU2AF<sup>65</sup> and the three kinetoplastid homologs (TbU2AF<sup>65</sup>, TcU2AF<sup>65</sup> and LmU2AF<sup>65</sup>) gave almost similar results. In both cases, the two proline residues with six amino acid spacer were found at positions TbU2AF65Pro213 and TbU2AF<sup>65</sup>Pro220. The representatives of hU2AF<sup>65</sup>Pro96 were proline residues which occurred at different positions, TbU2AF<sup>65</sup>Pro129 in the first comparison and TbU2AF<sup>65</sup>Pro174 in the second comparison. 83 residue spacer was noted in the alignment between the three kinetoplastids and human U2AF<sup>65</sup> homologs (TbU2AF<sup>65</sup> Pro129 and TbU2AF<sup>65</sup>Pro213), while 38 residue spacer was noted in the alignment between the Trypanosoma species and human U2AF<sup>65</sup> homologs (TbU2AF<sup>65</sup>Pro174 and TbU2AF<sup>65</sup> Pro213). No spacer was present in the case of alignment between human U2AF<sup>65</sup> and T. brucei U2AF<sup>65</sup> (Figure 20).

\* = Conserved proline residues

Figure 20. Alignment of amino acid residues of *H. sapiens* and *T. brucei* U2AF<sup>65</sup> for "groove" formation.

<b>Fable 9.</b> Residues in H	. sapiens and T. brucei U2AF <sup>65</sup> . The amino acid residues are implicated in
groove" formation bas	sed on sequence alignment between the T. cruzi (Tc), T. brucei (Tb), L.
major (Lm) and H. sa	piens (Hs). The columns under TbU2AF <sup>65</sup> represent T. brucei U2AF <sup>65</sup>

hU2AF <sup>65</sup>		TbU2AF <sup>65</sup>	
	Tb / Hs	Tb / Tc / Hs	Tb / Tc / Lm / Hs
Тгр92	Leu157	Val170	Pro125
Pro95	Pro160	Thr173	Asp128
Pro96	Leu161	Pro174	Pro129
Pro 97	Pro162	Pro213	Pro213
Phe99	Ala164	Val215	Val215
Pro 104	Pro169	Pro220	Pro220
Try107	Thr172	Pro223	Pro223

### 4.7 Purification of recombinant proteins in bacterial cells

### 4.7.1 CPSF 30 recombinant protein

Transformed BL21 *E. coli* cells expressed the CPSF 30 recombinant protein when induced with IPTG. The recombinant CPSF 30 protein purified on a Ni-NTA Spin Kit (Qiagen, GmbH Germany) gave a product of approximately 24 kDa when resolved on a 4 - 20% SDS-PAGE gel (Figure 21).

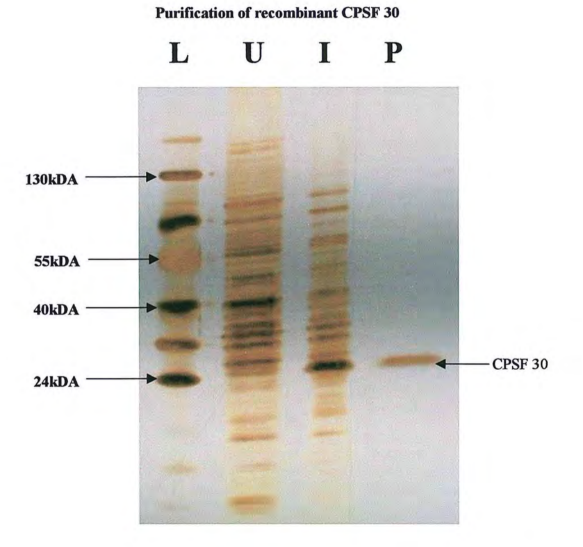
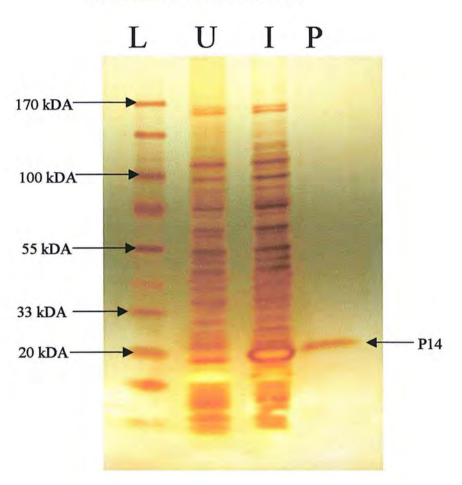


Figure 21. 4 - 20% SDS-PAGE gel showing purification of recombinant CPSF 30 protein. Lanes L: Protein ladder, U: uninduced recombinant, I: induced recombinant and P: purified recombinant CPSF 30.

#### 4.7.2 P14 recombinant protein

BL21 *E. coli* cells transformed with pET28a plasmid with P14 insert expressed the recombinant P14 protein when induced with IPTG. Purification of the recombinant P14 protein on a Ni-NTA Spin Kit (Qiagen, GmbH Germany) gave a product of approximately 20 kDa on separation on a 4 - 20% SDS-PAGE gel (Figure 22).



**Purification of recombinant P14** 

**Figure 22.** 4 - 20% SDS-PAGE gel showing purification of recombinant P14 protein. Lanes L: Ladder, U: uninduced recombinant, I: induced recombinant and P: purified recombinant P14.

### 4.8 Gene silencing of P14, U1-70k and CPSF 30 by RNAi

Difficulties were experienced in generation of clonal populations after induction due to clumping of the cells. However, growth arrest was evident among transfected cells in all the genes investigated (P14, U1-70k and CPSF 30). Unique morphological features were noticed in transgenic P14 and U1-70k cells but not in transgenic CPSF 30 cells (data not shown). The cells were abnormally large in size. In P14 transformants, the cells could not undergo complete division and remained held at the head in a group of about six cells just before death. Spherical cells like FAT cell stage (Ngô *et al.*, 1998) were noticed on induction for all the three factors.

#### 4.9 Western blot

Western blot on proteins derived from transfected, untransfected and control trypanosomes is shown in Figure 23. Detection of P14 protein in the untransfected trypanosome 1313-1333 cell line indicated that the gene encoding P14 was expressed. Similar detection was expected for control, however its absence could be due to "leakage" in the system (Alibu *et al.*, 2004). Depletion of P14 was observed in the transfected recombinants a phenomenon attributed to knock down of the P14 gene.

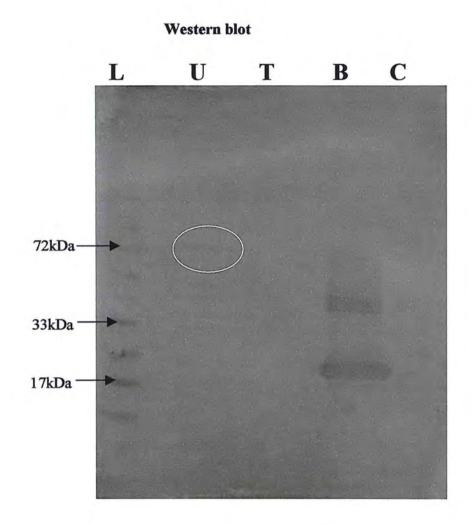


Figure 23. Results of western blot analysis of a trypanosome cell line transformed with P14 RNAi construct. L, protein ladder; U, 1313-1333 lysate; T, P14 RNAi transfected 1313-1333 cell; B, recombinant total bacterial lysate and C, control. The circle shows detection of P14 in untransfected 1313-1333 cell line lysate.

#### 4.10 DISCUSSION

#### 4.10.1 Sequence comparison

The amplification and sequencing results showed that *T. brucei* CFII-a1,  $AF^{35}$ , SF3b 145 and SF3b 10 were of similar sizes as those sequences from data mining at GeneDB, while U1-70k,  $AF^{65}$ , CstF 50, Zn 1, Zn 2, CPSF 30, P14, SF3b 49 and SF3b 125 were smaller in size. Comparison of the nucleotide and amino acid sequences of the cloned genes and those from data mining showed that the correct genes were recovered by the degenerate PCR amplification approach. This observation was further supported by the extremely low E–values (between 0.000 - 3.0e-280) at the stringent threshold limit of 0.0001.

Comparison of the *T. brucei* and human homologs showed the lowest percentage identity and similarity. This is because they are evolutionarily distinct and represents one of the earliest branches in eukaryotic lineage (Bringaud *et al.*, 1998; Stevens *et al.*, 1998; Verlinde *et al.*, 2001). Small U2 auxiliary factor (U2AF<sup>35</sup>) had high percentage identity and similarity across the four species in comparison to other factors. Moreover, the RNP1 and RNP2 motifs of RRM are conserved. This could be due to conserved intimate heterodimeric interaction of auxiliary factor ( $AF^{35}$  and  $AF^{65}$ ) in eukaryotes (Vázquez *et al.*, 2003). This could further be attributed to phylogenetic conservation of  $AF^{35}$  and residues of the U2AF<sup>65</sup> peptide that are critical for U2AF<sup>35</sup> binding (Kielkopf *et al.*, 2001). Similarly, CPSF 30 had an appreciably high percentage identity and similarity, presumably due to the conservation of the overall zinc finger motif structure and function (Hendriks *et al.*, 2003).

T. cruzi and L. major orthologs showed the highest percentage identity and similarity to T. brucei. Similar closeness was observed by El-Sayed et al. (2005b) at whole genome level. However, T. cruzi orthologs are more closely related to T. brucei than L. major. This closeness is in agreement with amino acid sequence alignment of a large sample of three-way cluster of orthologous genes (COGs) earlier observed by Haag et al. (1998) and El-Sayed et al. (2005b). The alignment revealed an identity of 57% between T. brucei and T. cruzi and 44% between L. major and the two other trypanosomes, reflecting phylogenetic relationships. Similarly, analysis of glucose transporter gene cluster (Bringaud et al., 1998) showed close evolutionary relationship between T. brucei and T. cruzi; members of the same genus. The difference between

T. brucei and T. cruzi is supported by the suggestion that among the monophyletic trypanosomatids, the Salivarian trypanosomes (also called African trypanosomes: subgenus Trypanozoon or Trypanosoma brucei group, T. congolense and T. vivax) emerged before T. cruzi (Bringaud et al., 1998). This variation could also be due to varied acquisition of an accelerated rate of evolutionary substitutions in Trypanosoma (Lake et al., 1988) and different rates of evolution (Stevens et al., 1998).

#### 4.10.2 Domains / motifs

The thirteen *T. brucei* trans-spliceosome genes studied showed domains that suggest their involvement in RNA splicing. The TbU2AF<sup>35</sup>, TbU2AF<sup>65</sup>, TbP14 and TbSF3b 49 have RRM domains involved in RNA recognition; a fundamental process in precise splice site and branch point recognition during RNA maturation. The TcU2AF<sup>35</sup> RRM domain has conserved residues (Thr 45, Leu 47 and Tyr 114) known to be directly involved in RNA recognition (Vázquez *et al.*, 2003). These residues are also conserved in TbU2AF<sup>35</sup>. However, Trp 134, the hallmark of the U2AF<sup>35</sup> RRM domain of eukaryotes (Vázquez *et al.*, 2003) is absent in TbU2AF<sup>35</sup>. This residue, which is necessary for the reciprocal "tongue in groove" heterodimerization with U2AF<sup>65</sup> is changed to Lys in the *T. brucei* ortholog. The *T. cruzi* ortholog has the same substitution (Vázquez *et al.*, 2003). This is a fundamental difference with the human homolog suggesting that the trypanosome gene products of U2AF<sup>35</sup> and U2AF<sup>65</sup> interact differently during spliceosome assembly. The third and different zinc knuckles (CCHC/Cx2Cx4Hx4C) in TbU2AF<sup>35</sup> and TcU2AF<sup>35</sup> is similar to the zinc finger domain found in a protein that binds the universal minicircle sequence of trypanomastids and is indicative of kinetoplastid DNA/RNA single strand binding protein (Tzfati *et al.*, 1995; Abu-Elneel *et al.*, 1999).

U2 auxiliary factor large subunit U2AF<sup>65</sup> interacts directly with the pyrimidine (Py) tract and branch point (BP) by the C-terminal RRM (Kielkopf *et al.*, 2001) and RS domain (Förch *et al.*, 2003) respectively. Its RRM also interacts with SF3b 155, a component of U2 snRNP (Shepard *et al.*, 2002). The protein is therefore thought to be involved in stabilization of the interaction of U2 snRNP with the BP through base-pairing interactions (Gozani *et al.*, 1998; Förch *et al.*, 2003). The hU2AF<sup>65</sup> N-terminal RS domain is missing in the three kinetoplastids. **TbU2AF**<sup>65</sup> however, has an arginine rich region at the N-terminal, which could be involved in direct

interaction with the BP and stabilization of the interaction of U2 snRNP with the BP as in hU2AF<sup>65</sup>. The RMM domains could be involved in interaction with pyrimidine (Py) tract and splicing factor 1/branch point binding protein (SF1/BBP) as suggested by Varani and Ramos (2003). TcU2AF<sup>65</sup> has two RRMs. This is suspected to be a split RRM when compared with the hU2AF<sup>65</sup>.

The cleavage stimulating factor 50 (CstF 50) sequences have WD domain (WD or betatransducin repeats) with a terminating Trp-Asp (W-D) dipeptide characteristic of the domain. The WD domain proteins form a large family with a high degree of diversity in sequence, multidomains and cellular functions (Yu et al., 2000). The sequence diversity occurs primarily in the two variable regions within the WD-repeat itself (Yu et al., 2000) thus substitution of aspartic acid with glutamic acid in human CstF 50. The TbCstF 50 could be involved in directing spliceosome complex assembly in which interactions between several proteins are involved. This is because the underlying common function of the domain is to coordinate multi-protein complex assemblies in signal transduction, transcription initiation complex assembly, chromatin assembly, RNA splicing, vascular trafficking, cell cycle control and apoptosis (Smith et al., 1999; Madrona and Wilson, 2004). The motif also provides an interface for protein-protein interactions (Zhao et al., 1999; Li and Roberts, 2001) either with other members of the WD family or with proteins carrying different motifs; most of the known proteins being members of multiprotein complexes (Yehuda et al., 1998). These interactions can occur simultaneously with several different proteins and their specificity is determined by sequences outside the repeats (Li and Roberts, 2001). Its interaction with RNA polymerase II and CPSF to stabilize the cleavage complex (Zhao et al., 1999) could be through the WD domain that interfaces for protein-protein interaction among different proteins (Zhao et al., 1999; Li and Roberts, 2001).

TbCFII-a1 has pre-mRNA cleavage complex II protein Clp1 domain that may be involved in ATP/GTP binding, P-loop (phosphate binding loop) with nucleoside triphosphate hydrolases and GTPase domain. These domains may be involved in nucleoside hydrolysis during cleavage and are similar to members of AAA+ family of ATPases. This family of ATPases forms dynamic oligomeric rings, and carries out diverse and important cellular functions, including those of helicases, unfoldases and ATPases (Orlova and Saibil, 2004), which have been recorded in the

splicing process. The function of the P-loop is to correctly position the triphosphate moiety of a bound nucleotide (Caruthers and McKay, 2002; Leipe et al., 2002).

SF3b 125 proteins have a DEAD box signature. The DEAD box represents the one letter code for the tetrapeptide, Asp-Glu-Ala-Asp and is a helicase domain characteristic of members of DExH/D box family domain (Will et al., 2002; Shi et al., 2004). The helicase 'superfamily' of proteins (RNA unwindases/ RNPases/ helicases) is characterized by a common general function of an ATP-dependent nucleic acid unwinding (de la Cruz et al., 1999). The 'superfamily' has been implicated in various aspects of RNA metabolism which include nuclear transcription, premRNA splicing, ribosome biogenesis, nucleocytoplasmic transport, translation, RNA decay and organellar gene expression (de la Cruz et al., 1999; Tanner and Linder, 2001; Cordin et al., 2004). TbSF3b 125 may therefore be involved in the ATP dependent A complex formation in which base pairing and ATP hydrolysis are involved. The domain could be specifically implicated in directing precise base pairing and correcting mismatch in the recruitment of U2 snRNP to the degenerate branch point. This process could occur through displacement of splicing factor 1/branch point binding protein (SF 1/BBP) as suggested by Fleckner et al. (1997) on the role of two DEAD box proteins, Prp5p and UAP56. The nucleic acid unwinding ability is very important in structural rearrangements and conformational changes during spliceosome assembly and correction of mismatches. The motif is specific to proteins that couple ATPbinding/hydrolysis and structural rearrangement (Fleckner et al., 1997; Xu et al., 2004) fitting well with formation of A complex, an ATP dependent process.

TbSF3b 145 exhibited a proline-rich domain (PSP) similar to homologs from *H. sapiens*, *T. cruzi* and *L. major*. It probably interacts with TbSF49 via its proline-rich domain since this domain is dispensable for the protein-protein interaction between human SF3b 145 and SF3b 49 (Igel *et al.*, 1998). *H. sapiens* SF3b 145 has a DNA binding SAP or SF found in ATP-dependent DNA helicase.

Interaction between TbSF3b 49 and TbSF3b 145 could be through RRM of TbSF3b 49 as observed in yeast homologs (Igel *et al.*, 1998). TbSF3b 49 may also bind the pre-mRNA via the RRMs. These interactions are for the stable recruitment of U2 snRNP to the degenerate BP, a

process that involves base pairing (Gozani *et al.*, 1998; Igel *et al.*, 1998). These inferences are supported by the facts that SF3b 49 can cross-link efficiently to RNA substrates in complexes A and B and also to bind both U2 snRNP and the pre-mRNA (Chiara *et al.*, 1996). The poly adenylate binding protein (PABP) and PABPh domains in TbSF3b 49 are thought to recognize the poly-A tail of mRNA and may be involved in the linkage of cleavage and polyadenylation.

The domains in TbSF3b 10 implicate the protein in the energy dependent mismatch repair during A complex formation. It has MutS (III and IV domains); a key protein of the *Escherichia coli* DNA mismatches repair system that recognizes mispaired and unpaired bases and has intrinsic ATPase activity (Lamers *et al.*, 2004). The ATPase domain in TbSF3b 10 could be associated with ATP binding activity that induces a state in which MutS slides away from the mismatch to allow new molecules to bind the mismatch (Lamers *et al.*, 2004) or discrimination between homoduplex and heteroduplex DNA (Schofield *et al.*, 2001). Alternatively, MutS domain can act as a motor protein that uses the ATPase activity to translocate along the DNA in search of a signal for strand discrimination (Blackwell *et al.*, 1998). In *T. cruzi* and *L. major*, a P-loop domain with nucleoside triphosphate hydrolase could be associated with ATP binding and hydrolysis. The binding and hydrolysis cause ATP-dependent conformational change that allows recruitment of other proteins (Alani *et al.*, 2003). These features concur with the structural rearrangements and energy consumption associated with spliceosome assembly (Schwer and Guthrie, 1992; Chan *et al.*, 2003; Xu *et al.*, 2004).

TbCPSF 30 has five zinc finger (type CCCH) motifs and two zinc knuckles (CCHC) as described by Hendriks and colleagues, 2003. TbCPSF 30 may be involved in both cleavage and polyadenylation. These involve RNA binding and protein-protein interactions through the motifs. These motifs typically function as interaction modules and bind to a wide variety of compounds such as nucleic acids (Hendriks *et al.*, 2003), proteins and small molecules (Krishna *et al.*, 2003). They are also structurally diverse and are present among proteins that perform a broad range of functions in various cellular processes, such as replication and repair, transcription and translation, metabolism and signalling, cell proliferation and apoptosis (Krishna *et al.*, 2003). Zhao *et al.* (1999) reported that CPSF as well as poly (A) polymerase (PAP) remains bound to

the cleaved RNA and elongate the poly A tail in the presence of poly (A)-binding protein II (PAB II). Therefore, TbCPSF 30 could be involved in the transcription.

Protein synthesis occurs in the cytoplasm, but many proteins are required in the nucleus and have to be imported. The splicing process which occurs in the nucleus requires recruitment of spliceosome complex proteins. Marchetti *et al.* (2000) demonstrated the presence of an energy dependent nuclear import system in trypanosomes. The signalp domain found in most of the trans-splicing factors could be involved in directing the transportation of these proteins across the trypanosome nucleus membrane. The nuclear import process depends on nuclear localization signals (NLS) present only in nuclear proteins and can be either signal sequences or signal patches (Görlich, 1998; Moore, 1998). The signalp domain could therefore be a signal sequence or patch that directs importation into the nucleus, by nuclear import receptors. Each type of receptor protein is specialized for the transport of a group of nuclear proteins sharing structurally similar nuclear localization signals (Smith and Raikhel, 1999).

## 4.10.3 T. brucei U2 auxiliary factor heterodimer (TbU2AF)

The U2AF heterodimer consists of 65-kDa (U2AF<sup>65</sup>) and 35-kDa (U2AF<sup>35</sup>) subunits (Ito *et al.*, 1999; Shepard *et al.*, 2002) that form an intimate heterodimeric complex in which the RRM of U2AF<sup>35</sup> and a central polyproline segment of U2AF<sup>65</sup> interact via reciprocal "tongue in groove" tryptophan residues (Vàzquez *et al.*, 2003). Three-dimensional structure of this interaction is mallet-shaped (Figure 24). In the "tongue in groove" interaction, the U2AF<sup>65</sup> peptide forms a type II proline helix that wraps tightly around the globular U2AF<sup>35</sup> domain at the head, while one long U2AF<sup>35</sup>  $\alpha$  helix escapes the molecular embrace to form the handle (Kielkopf *et al.*, 2001). The proline residues (forming the "groove") enclose the hU2AF<sup>35</sup> tryptophan 134 (hU2AF<sup>35</sup> Trp134) in a tight molecular embrace of aromatic/aliphatic interactions (Kielkopf *et al.*, 2001). In *T. brucei*, the non-polar and aromatic U2AF<sup>35</sup> Trp134 observed in human is replaced with a polar and basic U2AF<sup>35</sup> Lys122. However, the residues Thr 45, Leu 47 and Tyr 114 of U2AF<sup>35</sup> RRM known to be directly involved in RNA recognition are conserved.

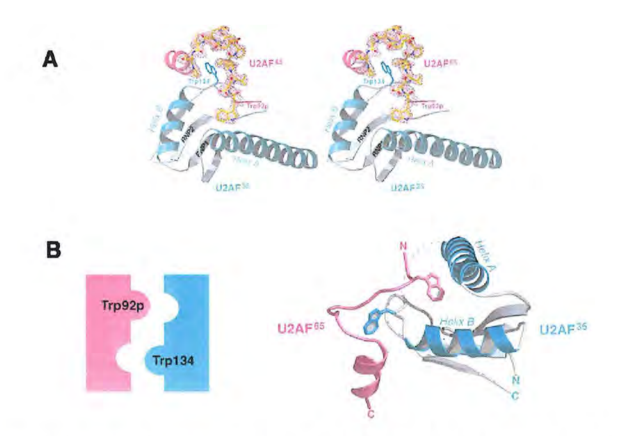


Figure 24. Schematic representation of human U2AF heterodimer. (A) Stereoview of the  $hU2AF^{65}$  proline-rich loop enveloping  $hU2AF^{35}Trp134$ . The  $hU2AF^{65}$  peptide is shown in pink, or color-coded for atom type (yellow, carbon; red, oxygen; and blue, nitrogen) and the U2AF<sup>35</sup> domain is shown in light blue. (B) A schematic representation of the reciprocal "tongue in groove" and tryptophan binding sites. (Adopted from Kielkopf *et al.*, 2001)

The residues important in "groove" formation in hU2AF<sup>65</sup> are Trp92, Pro95, Pro96, Pro97, Phe99, Pro104, and Try107 with a mandatory six residue spacer between the conserved hU2AF<sup>65</sup>Pro97 and hU2AF<sup>65</sup>Pro104 (Kielkopf *et al.*, 2001). Comparison of hU2AF<sup>65</sup> and TbU2AF<sup>65</sup> showed two sets of proline residue pairs with a six residue spacer at TbU2AF<sup>65</sup>Pro162 and TbU2AF<sup>65</sup>Pro169, and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro220. Conservation of these proline residues is suspected to be high. The role played by hU2AF<sup>65</sup>Pro97 and hU2AF<sup>65</sup>Pro104 might be undertaken by either TbU2AF<sup>65</sup>Pro162 and TbU2AF<sup>65</sup>Pro169 or TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro214 might be undertaken by either TbU2AF<sup>65</sup>Pro162 and TbU2AF<sup>65</sup>Pro169 or TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro220 in TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro169 or TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro220 in TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro169 or TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro220 in TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro220 in TbU2AF<sup>65</sup>Pro213 and TbU2AF<sup>65</sup>Pro220 in TbU2AF<sup></sup>

human U2AF<sup>65</sup> homologs, the amino acid residue at similar position to hU2AF<sup>65</sup>Pro95 in *T. brucei* is an aspartic acid residue (TbU2AF<sup>65</sup>Asp128). Interestingly, at position 127 of TbU2AF<sup>65</sup>, there is a proline residue (TbU2AF<sup>65</sup>Pro127). The TbU2AF<sup>65</sup>Pro127 could be implicated in a similar role played by hU2AF<sup>65</sup> Pro95. No inference could be drawn on the 83 and 38 residue spacers unless the proline residues implicated in interaction with TbU2AF<sup>35</sup> are determined. Despite these variations, the residues in TbU2AF<sup>65</sup> could be involved in formation of the TbU2AF<sup>65</sup> "groove" that encloses TbU2AF<sup>35</sup>Lys122 in a tight molecular aliphatic interaction similar to hU2AF heterodimer interaction suggested by Kielkopf *et al.* (2001).

The variations between hU2AF<sup>65</sup> and TbU2AF<sup>65</sup> could be due to the fact that the U2AF<sup>35</sup> residue actively involved in interaction with residues of U2AF<sup>65</sup> polyproline "groove" has switched from hU2AF<sup>35</sup>Trp134 to TbU2AF<sup>35</sup>Lys122. These changes may be implicated in positioning amino acid residues in appropriate positions allowing the formation of a "groove" and stable interactions with the TbU2AF<sup>35</sup>Lys122. If these variations are experimentally proven by mutational analysis to assess the residues that are important in molecular recognition hence interaction, it can demonstrate an example of host and parasite specific protein-protein interaction in the spliceosome complex. Such fundamental differences could be exploited as drug targets by designing specific disruptors.

#### 4.10.4 Gene knock down

Cell death observed in cell lines expressing recombinant CPSF 30, U1-70k and P14 suggests the importance of these factors in trypanosome viability. TbCPSF 30 depletion was lethal and no morphological changes were evident similar to findings by Hendriks *et al.* (2003). However, Hendriks and co-workers recorded complete cell death after 5-6 days as compared to 10 days observed in this study. This could be attributed to the fact that a different cell line was used.

TbP14 depletion resulting into incomplete division in cells and death infers the importance of this factor in viability. This may be due to the fact that P14 is positioned within the inner cage of the SF3b structure (Golas *et al.*, 2003). The depletion could result into significant disruption of SF3b consequently affecting the trans-spliceosome complex and trans-splicing process. The abnormal morphological shapes in transgenic P14 and U1-70k cells could be due to disruption of

expression of genes that are involved in maintenance of shape. Abnormal shapes have been documented in suppression of clathrin heavy chain (Allen *et al.*, 2003) in which enlargement of the flagella pocket resulted into a 'BigEye' phenotype. The appearance of FAT cells (Ngô *et al.*, 1998; Inoue *et al.*, 2002) suggests effects on expression of  $\alpha$ -tubulin. This can be due to disruption of processing of  $\alpha$ -tubulin pre-mRNA. There were difficulties in monitoring growth kinetics since the cells formed clumps. The clumping is characteristic of dying trypanosomes. However, the degree of clumping was unusual. The reasons for the extensive clumping were unknown.

The RNAi experiments highlighted the importance of trans-spliceosome and polyadenylation factors in trypanosomes. However, the levels of RNA and proteins should be determined to conclusively determine the effects of specific silencing. This was not achieved in this study because clumping of cells precluded accurate monitoring of cell numbers. Investigations on protein-protein interactions in trypanosome trans-spliceosome could be done using tandem affinity purification (TAP) tagging (Rigaut *et al.*, 1999; Puig *et al.*, 2001; Knuesel *et al.*, 2003; Gould *et al.*, 2004). This could offer insights into the nature of interacting pairs of proteins. Small molecules that disrupt such interactions are potential drugs.

### **CHAPTER FIVE**

### CONCLUSIONS AND RECOMMENDATIONS

Trypanosomosis is a widespread constraint on human health, livestock production and mixed farming in tropical Africa. The disease has denied Africa her agricultural potential and causes considerable mortality and morbidity in endemic foci. The main control strategies applied (vector control and chemotherapy) have varied limitations. In recent years, increased incidents of resistance to available drugs have limited their value. Continued use of the few available drugs may result in multiple resistance that could make chemotherapy increasingly ineffectual, though it remains the mainstay for control of parasitic diseases (Verlinde et al., 2001). It currently protects more cattle than all other techniques combined (FAO, 2002), can be used under any production system, in any ecological zone and allows farmers to individually apply trypanocides to their cattle. The limited number of available drugs is a consequence of market economy principles: since people most at risk from tropical diseases are among the poorest in the world, there is currently little perceived financial incentive for pharmaceutical companies to invest in development of new drugs. Trypanosomosis has therefore remained among the most neglected tropical diseases (Morel, 2003) in terms of drug development. Re-emergence of the disease (Legros et al., 2002) will worsen the delicate situation. This justifies the investigation of trypanosome specific process such as regulation of gene expression as a potential chemotherapeutic target.

Gene regulation is vital to cell viability, growth and development in all eukaryotic organisms. One aspect of gene regulation is the requirement for pre-mRNA processing through splicing into mature RNA species. This process is particularly important in regulation of gene expression in trypanosomes whose genes are organized in polycistronic transcription units. This occurs via trans-splicing, specific to trypanosomes but absent in human and mammalian hosts.

The identification and functional analysis of the trans-splicing and polyadenylation factors in *T. brucei* is an important contribution to understanding the trans-spliceosome as a potential drug target. The low percentage identity and similarity between the *T. brucei* trans-splicing and polyadenylation factors and those of human and suspected difference in protein-protein interactions, defines the variations in the process of RNA maturation. Additionally, the long

evolutionary distance between trypanosomatids and their mammalian hosts (Verlinde *et al.*, 2001) endows the trans-splicing and polyadenylation factors with distinct properties. For optimum utilization of these findings, I would recommend further studies aimed at generating exhaustive information that would be exploited in development of disruptors specific to parasite trans-splicing process. RNAi technology could be used as a tool to analyse the genes for validation as potential drug targets during such studies. Interacting factors whose silencing is lethal to the parasite should be adequately characterized and amino acid residues involved in molecular recognition determined. This should include auxiliary factor (AF<sup>35</sup> and AF<sup>65</sup>), SF3b 145, SF3b 49, CPSF 30, U170k and P14 as well as other factors that could be important in viability. Successful undertaking of the above recommendations would improve chemotherapeutic control not only to trypanosomosis, but also to other diseases caused by parasites that exhibit trans-splicing such as leishmania, *T. cruzi*, trematode infections caused by schistosomes and nematode infections caused by filaria. This would enhance realization of Africa's optimum agricultural potential that would in turn support her economically disadvantaged inhabitants.

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## **APPENDIX 1**

### Accession numbers of spliceosome factors sequences

Spliceosome nucleotide and proteins were used and generated in data mining. The accession numbers are given for *T. brucei*, *L. major*, *T. cruzi* and *Homo sapiens* in that order. Those for *Saccharomyces cerevisiae* and *Caenorhabditis elegans* are given the initials sc and ce. (The accession numbers are temporary)

Data Mining		
Factors	Accession number	
	E complex	
Zn1	TB05.28F8.690	
	LmjF35.1040	
	Tc00.1047053507305.40	
Zn2	TB05.28F8.690	
	LmjF35.1040	
	Tc00.1047053507305.40	
U170k	Tb10.70.4700	
	LmjF34.0495	
	Tc00.1047053504105.160	
CFII-a1	Tc00.1047053507027.59	
	LmjF30.2410	
	Tb06.4F7.820	
	Q92989	
	sc YOR250C	
AF35	Tb10.70.4300	
	Tc00.1047053510943.60	
	LmjF03.0190	
	AAH01923	
AF65	Tc00.1047053510265.40	
	LmjF03.0520	
	Tb10.70.3950	
	CAA45409	
	ce AAM44400	
	sc AAA64215	

Factors	Accession number
	A complex
P14	AAH15463
	Tb10.6k15.3190
	LmjF36.3040
	Tc00.1047053506885.70
	sc YIR005W
SF3b10	AAH00198
	ТЬ03.26Ј7.490
	LmjF29.1710
	Tc00.1047053509617.30
SF3b49	AAH04273
	Tb09.211.0930
	LmjF35.5040
	Tc00.1047053506885.70
	sc YOR319W
SF3b125	Tb10.70.0140
	LmjF36.2130
	Tc00.1047053510187.290
7791 1 45	AAH08208
SF3b145	Tb06.4M18.90
	LmjF30.0570
	Tc00.1047053504071.50
	AAH14125
	sc YMR240C
	Cleavage and polyadenylation factors
CstF50	Tb10.61.0570
	LmjF33.0280
	Tc00.1047053508899.30
	AAP36871
CPSF30	Tb11.01.4600
	LmjF09.0720
	Tc00.1047053510219.30
	EAL23878 sc YPR107C

## **APPENIX 2**

# **Buffers and solutions**

## **DNA** extraction

### **Cell lysis solution**

Tris-Cl (pH8.0)	10mM
NaCl	100mM
EDTA	100mM
SDS	0.01%
Proteinase K	100ng/ml

## Phosphate buffered saline (PBS) pH 7.3

137 mM
2.7 mM
4.3 mM
1.4 mM

### **Bacterial culture media**

20g
5g
0.5g
10ml
5ml
20ml

## LB\* medium

Tryptone	10g
Yeast extract	5g
NaCl	10g

LB* agar	
Tryptone	10g
Yeast extract`	5g
NaCl	10g
Agar	15g

 $^{\dagger}$  Filter Sterilize through a 0.2  $\mu m$  filter

\* Autoclave

# Protein extraction and purification buffers

4X SDS Sample Buffer	
Tris-HCl, (pH6.8)	250mM
Sodium dodecyl sulphate (SDS)	8 %
β-mercaptoethanol	4 %
Glycerol	40 %
Bromophenol blue	0.02 %

Lysis buffer (pH 8.0)	
Urea	8M
NaH <sub>2</sub> PO <sub>4</sub>	0.1 <b>M</b>
Tris-Cl	0.01M
NaCl	0.3M

Wash buffer (pH 6.3)	
Urea	8M
NaH <sub>2</sub> PO <sub>4</sub>	0.1 <b>M</b>
Tris-Cl	0.01M

Elution buffer (pH 4.5)	
Urea	8M
NaH <sub>2</sub> PO <sub>4</sub>	0.1M
Tris-Cl	0.01M

### Transfection medium

(pH 7.0)
132 mM
8 mM
8 mM
1.5 mM
1.5 mM
90μΜ

### Western blot

**Transfer Buffer (pH 8.3)** 48mM Tris-Cl 29mM Glycine 20% methanol 0.037% SDS

# Washing Buffer (pH 7.4)

25mM Tris-Cl 137mM NaCl 3mM KCl 0.3% Tween-20

### Blocking Buffer (pH 7.4)

25mM Tris-Cl 137mM NaCl 3mM KCl 1% BSA /1% Blocking reagent (Boehringer Mannheim, GmbH, Germany)

Nitro Blue tetrazolium (NBT) (Sigma) Stock Solution 1tablet of NBT dissolved in 1ml distilled deionized water

5-Bromo-4-chloro-3-indolyl phosphate (BCIP) (Sigma) stock solution 1 tablet of BCIP in 100% N, N- dimethylformamide (Sigma).

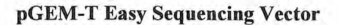
Substrate Buffer (pH 9.5) 100mM Tris-Cl 100mM NaCl 5mM MgCl<sub>2</sub>

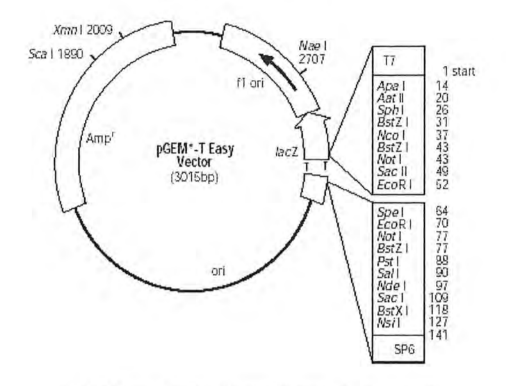
Substrate solution 330µl NBT 33µlBCIP 10 ml of substrate buffer (pH 9.5)

TBS Buffer (pH 7.4) 25mM Tris-Cl 137mM NaCl 3mM KCl

# **APPENDIX 3**

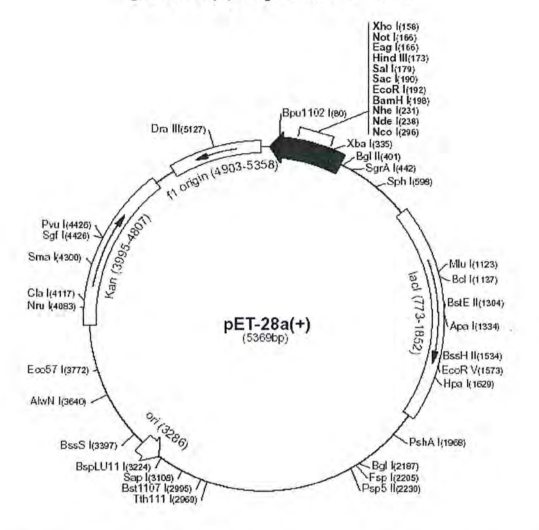
# Vector maps





pGEM-T Easy Vector circle map (Promega Biotechnology)

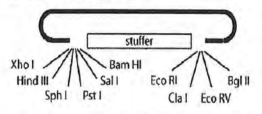
## pET-28a(+) Expression Vectors



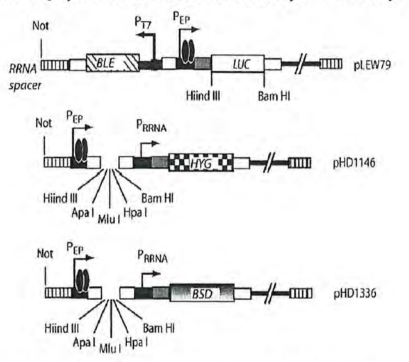
The pET-28a (+) vector map. It carries an N-terminal His•Tag/thrombin/T7•Tag configuration plus an optional C-terminal His•Tag sequence. Unique sites are shown on the circle map. The cloning/expression region of the coding strand transcribed by T7 RNA polymerase is shown above.

# **RNAi VECTORS**

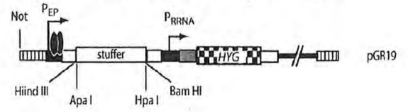
A. Vector for stem-loop cloning



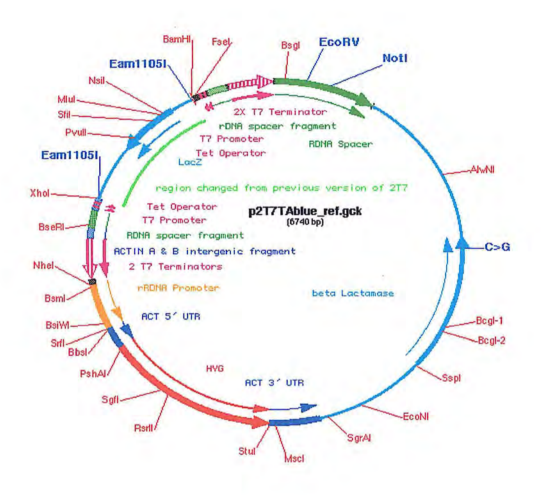
B. Inducible polymerase I vectors for insertion of ready-made stem-loops



## C. Inducible polymerase I vector for direct cloning of stem-loops

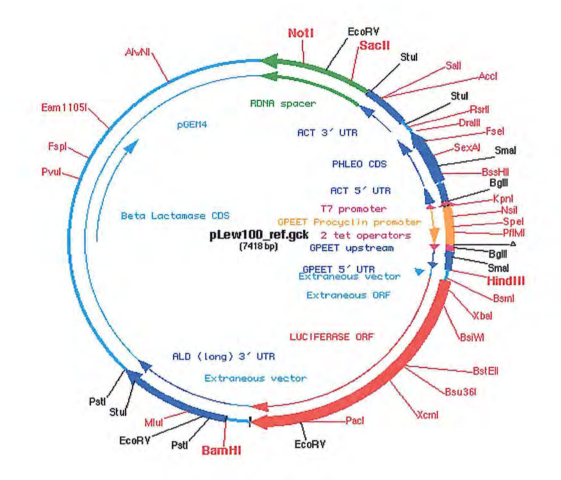


### p2T7TAblue RNAi vector

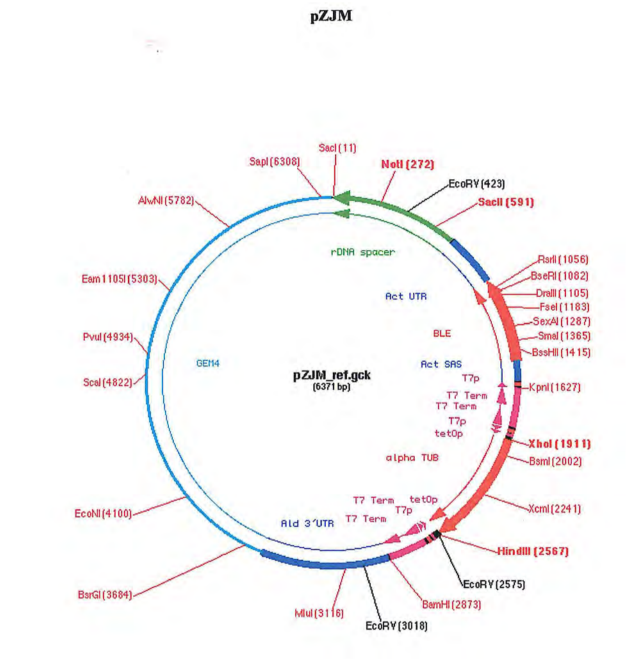


Source: http://tryps.rockefeller.edu/Seqs/Graphics/construct\_images.pdf

**pLEW 100** 



Source: http://tryps.rockefeller.edu/Seqs/Graphics/construct\_images.pdf; Wirtz et al., 1999



Source: http://tryps.rockefeller.edu/Seqs/Graphics/construct\_images.pdf