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TRYPANOSOMA CRUZI GENOME
PLASTICITY AND EVOLUTION

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Cover artwork: Synteny map of two *Trypanosoma cruzi* genomes

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Trypanosoma cruzi genome plasticity and evolution
THESIS FOR DOCTORAL DEGREE (Ph.D.)

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A mi familia
The greatest and noblest pleasure which men can have in this world is to discover new truths; and the next is to shake off old prejudices

Frederick II of Prussia
ABSTRACT

Trypanosoma cruzi, a protozoan from the Kinetoplastidae family, is the etiologic agent of Chagas disease, a major public health problem affecting mostly the poorest areas of Latin America. Due to the complex nature of the parasite’s genome it has been impossible to produce a complete reference genome sequence, thus hampering the implementation of post-genomic approaches to unveil the mechanisms of generation of antigenic variation and the identification of new drug targets. My doctoral studies have focused on the application of combined genome sequencing and computational methods to produce a complete reference T. cruzi genome sequence and perform comparative analyses to better understand the mechanisms that allow T. cruzi to evade the mammalian host immune system and to briskly adapt to novel environments.

In paper I and II, different genome assembly strategies and second generation sequencing technologies were implemented to perform comparative analyses to identify elements of virulence between T. cruzi and two trypanosomatids that are non-pathogenic to humans: Trypanosoma cruzi marinkellei, a bat-restricted sub-species of the T. cruzi clade and the human avirulent species Trypanosoma rangeli. The studies reveal the expansion of T. cruzi-specific genomic traits specialised in the invasion of mammalian cells.

In paper III, using third-generation, PacBio sequencing data it was possible to assemble the complete reference genome sequence of a Trypanosoma cruzi isolate from the DTU-I clade. This breakthrough allowed us - for the first time - to explore in detail the genome architecture of the subtelomeric areas where many parasite virulence factors are encoded. One of the most interesting discoveries was the overrepresentation of interspersed retrotransposons and microsatellites in tandem gene arrays coding for surface molecules, hinting at a retrotransposon-driven mechanism of recombination for generating new sequence variants. Whole genome sequencing of 35 T. cruzi DTU-I isolates, collected from different locations in the American continent, made possible to identify and characterise the mechanisms of adaptability employed by the parasite.

Finally, paper IV analyses the mechanisms of genomic hybridisation in T. cruzi and the evolution over time of the hybrid offspring. The analysis revealed that during hybrid formation, the parasite integrates genetic material from each parental strains with the aid of retrotransposons and microsatellites, and the genome of these hybrid isolates moves quickly from a tetraploid to a diploid state. As a result, the hybrid strain has more genetic material,
mostly in the subtelomeres, providing the parasite with a pool of new surface molecule genes with the potential to possibly increase its fitness in a new environment.

In conclusion, the work presented here has advanced the understanding of parasite biology and provided a genomic resource to be exploited for the identification of drug targets and vaccine candidates.
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LIST OF ABBREVIATIONS

SNV       Single Nucleotide Variants
CNV       Copy Number Variant
Mbp       Megabasepair
InDel     Insertion/Deletion
SV        Structural Variant
NGS       Next Generation Sequencing
IPE       Illumina Paired End library
IMP       Illumina Mate Paired library
HLA       Human Leukocyte Antigen
RNA-Seq   RNA sequencing data
RAM       Random Access Memory
PCR       Polymerase Chain Reaction
dNTP       Deoxynucleotide triphosphate
HR        Homologous Recombination
NAHR      Non-allelic Homologous Recombination
NHEJ      Non-homologous End Joining
MMEJ      Microhomology-mediated End Joining
LD        Linkage Disequilibrium
MARK      Trypanosoma cruzi marinkellei
RANG      Trypanosoma rangeli
CHAPTER 1 - INTRODUCTION

1.1 A biologist’s field guide to genome sequencing technologies:

The chain terminator sequencing method described by Frederick Sanger in 1977¹ was a major breakthrough that allowed molecular biologists to better understand the information encoded by DNA. Further technological advances, such as PCR and the incorporation of fluorochromes, made it possible to read the nucleotide sequence of complete genes. A gene sequence database was soon available for the analysis of their molecular functions and potential interactions, but soon it became clear that to have a complete picture of all the molecular processes of an organism it was necessary to go “genome-wide”.

In the 1980’s, with the first automated sequencing machines, the idea of sequencing the entire human genome was proposed, and in 1988 the U.S congress funded the National Institutes of Health (NIH) and the Department of Energy (DoE) and in 1990 the initial research plan for the Human Genome Project (HGP) was released². The early stages of the HGP were dedicated to the development of new methods and technologies to analyse the genomic sections in a cheap and efficient way³.

In 1995, Haemophilus influenzae became the first organism to have its genome completely sequenced⁴. This was later followed by the genome of other majors organisms such as Saccharomyces cerevisiae (1996)⁵, Caenorhabditis elegans (1998)⁶, Drosophila melanogaster (2000)⁷, and Homo sapiens (2001)⁸⁹. The first draft genome for Trypanosoma cruzi was produced in 2005¹⁰. These genomes were sequenced using somewhat automated versions of the first generation Sanger sequencing machines that produced a low amount of sequencing data in short time but were very expensive to maintain and involved a lot of time consuming manual input¹¹. New technological advances in sequencing methods have reduced the operational costs while allowing a much higher level of production of data, these methods are usually referred as Next Generation Sequencing (NGS) technologies. The large amount of data produced by the second generation sequencing platforms have the shortcoming of having high error rates and producing significantly shorter sequence reads (35 - 150 bp) compared with their first generation counterparts (600 - 1000 bp)¹².
1.1.1 Second generation sequencing technologies: 454 and Illumina.

The first NGS platform was the 454 pyrosequencing®. Pyrosequencing is considered a variant of the sequencing by synthesis (SBS) method. Genomic DNA is randomly shredded into small fragments of the desired size, usually ranging between 150 - 550 bp, and attached to a small bead where the sequencing reaction takes place. The controlled addition of dNTPs produces a luminous signal that is detected by a charge-couple device (CCD) camera. In this way, it is possible to sequentially detect which dNTP has been incorporated in the reaction and thus determine the nucleotide sequence. The reads produced by the 454 pyrosequencing® platform ranged only between 250 - 450 bp but produced about 1 million reads per run which constituted a considerable advantage over the low yield Sanger sequencing machines.

The most widely used sequencing platform to date is the Illumina HiSeq®. Based on the cyclic reversible termination variant of the SBS method, the system works in a similar way as the Sanger sequencing technique. Following the random shredding of the genomic DNA, the fragments are organised into clusters and ligated to sequence adapters that serve as primers for a DNA polymerase reaction. This reaction is terminated once a labelled dideoxynucleotide triphosphate (dNTP) is incorporated into the sequence template. The dNTPs used in the reaction are labelled with a specific fluorochrome and these are recognised by a set of four lasers, each one with a specific wavelength per fluorochrome. Once the lasers have imaged the reaction, the labelled fluorochromes are washed away and a new sequencing cycle begins. In this way it is possible to determine the nucleotide sequence of each fragment.

The Illumina platforms produce short reads in the range of 75 - 300 bp with different sequencing data outputs ranging from 15 million (Illumina MiSeq®) to 6 billion (Illumina HiSeq X Ten®).

454 and Illumina, were the first platforms to produce high throughput sequencing data, and they made it necessary to further improve the algorithms required to process and analyse the new kind of data. Illumina is the largest supplier of sequencing machines to date, and part of the success has been the ability to produce equipment tailored for specific needs such as bench top sequencing (Illumina MiSeq®) to population-scale whole genome sequencing (Illumina HiSeq X Ten®), as well as the relatively low error rate (> 0.5 %). This has not been the case for the 454 pyrosequencing® platform, where the costs of the sequencing reagents and the relatively low throughput compared to Illumina forced the platform to be officially retired in early 2016.
1.1.2 Third generation sequencing: Long sequencing reads:

Second generation sequencing technologies provide researchers with large data sets that can be exploited for the analysis of models organisms with good reference genome sequences. Two examples of the application of second generation, short read sequencing in model organisms are the analysis of human variation in the 1000 Human Genomes Project\textsuperscript{16} and the ENCODE Project\textsuperscript{17}. The scenario is different for non-model organisms, as not all of them have been sequenced and the few that have are still not complete\textsuperscript{18}. The short length nature of the reads from second generation sequencing platforms makes it impossible to completely assemble highly repetitive regions (i.e: centromeres, telomeres, short tandem repeats, retrotransposons, etc) and high complexity regions (i.e: HLA loci).

Recently, a third generation of sequencing machines have been able to produce long sequence reads on the order of several kilobases. One of these platforms, from Pacific Biosciences (PacBio), is Single-Molecule Real-Time (SMRT) sequencing. PacBio SMRT sequencing fixes a DNA polymerases at the bottom of a micro well known as zero-mode waveguides\textsuperscript{19}. The polymerase allows a single molecule of the template DNA to be processed at a time. Labelled dNTPs are incorporated as the DNA molecule passes the DNA polymerase, while a laser excites a fluorochrome and a camera detects the emitted signal. Subsequently, the marked dNTP is detached by the polymerase to allow the incorporation of a new nucleotide until the entire molecule has been processed\textsuperscript{20}. Alternatively, the process can take place in a circular consensus sequencing (CCS) fashion, where the single DNA molecule is circularised and read several times in shorter segments and the consensus of these shorter segments is used to produce a longer read\textsuperscript{21}. The reads produced by the PacBio machines are on average 12.5 Kb in length, but can be as long as 50 Kb\textsuperscript{15}.

The length of these reads makes them suitable for reconstruction of complex regions in \textit{de novo} genome assembly projects\textsuperscript{22} or single molecule mRNA sequencing\textsuperscript{23}. Unfortunately, the error rate of these long reads can be as high as a 15% of the total length\textsuperscript{24}. These errors can be corrected using medium to high coverage (25 - 80 X) of PacBio data\textsuperscript{25} or high coverage (> 50 X) of Illumina data\textsuperscript{26}, but the process is highly demanding in terms of computational resources and time-consuming; a factor that should be taken into account in the experimental design of a given genome project. The production of PacBio data is also more expensive than Illumina data with an estimate of €885 per Gb for PacBio compared to €133 per Gb for Illumina\textsuperscript{15}, as of 2016.
1.2 Introduction to genome assembly:

Suppose for a moment that you receive a very important letter in the post and before you have the chance to read it, this document is thrown into a paper shredder. Twice. The task to assemble a genome sequence from scratch is similar to trying to put together the double-shredded letter with the hope that you can read it and that this message makes sense. The process to assemble sequencing data into a complete reference genome is called de novo genome assembly. To date, due to technological limitations, it is not possible for any sequencing method to read an entire genome at once. The fact that sequencing reads are usually shorter than the length of an entire genome sequence implicates the need for algorithms to order all the reads together into a genome as if they are a jigsaw puzzle. The problem of genome assembly has led to multiple innovations regarding the data produced and the algorithms to process the data.

Below, I give a brief description of the most popular sequencing data and assembly methods used to date.

1.2.1 Bestiarum vocabulum of sequencing data:

Different types of sequencing data that can be generated from second and third generation sequencing platforms. The generation of these data depend on the method used to build the sequencing library and the purpose of the sequencing experiment.

Libraries using long genomic fragments - usually between 50 and 100 Kb - can be produced from a highly purified DNA preparation, as required for the PacBio sequencing protocol or for the production of Bacterial Artificial Chromosomes (BAC), and Fosmids. The advantage provided by sequencing the latter relies on the distance information from the long inserts, which can be used for fragment ordering during a de novo assembly.

Mate paired libraries, also known as jumping libraries, are produced by fragmenting DNA into pieces between 1 and 20 Kb in length. The long pieces of DNA are submitted to a circularisation process where both extremes of the DNA molecule are joined together by biotinylation and later sheared into smaller fragments whose ends are sequenced. The sequencing data are used to resolve long repetitive regions in de novo genome assemblies.

The preparation of genome sequencing libraries with insert sizes between 180 - 1000 bp often starts with the enrichment of genomic material via cloning or amplification by PCR.
However, this enrichment step is optional and it will depend on the quantity of the starting genomic material, as some protocols require as little as 10 ng/μL \(^{30,31}\). Later, the genomic material is randomly sheared by means of physical (e.g: sonication), enzymatic (e.g: DNase I or transposases) or chemical (e.g: metal cations) methods to the desired size \(^{32}\). The resulting fragments are used as template for sequencing with second generation sequencing platforms. Two types of sequencing data can be produced with this kind of genomic library: singled and paired ends.

Singled end data results from the sequencing of a fragment of one of either of the ends of a genomic fragment. Singled end data is currently mostly used for transcript quantification in RNA-Seq experiments \(^{33}\).

Paired end data is the most widely used type of sequencing data and is produced by sequencing both ends of a given fragment. Besides the sequence information contained in the reads, the data also provides distance information between both ends of the sequenced fragment. This information is useful for *de novo* genome assemblies \(^{28}\) and the detection of genomic structural changes \(^{34}\).

1.2.2 Genome assembly algorithms:

The general idea of a *de novo* assembly experiment is to transform the millions of short sequencing reads into a single consensus that contains all the information encoded by the genome at a high level of accuracy. To achieve this, computer scientists have implemented different algorithms to be able to analyse the large amount of data in an organised way to determine the best possible reconstruction of a genome sequence \(^{35}\).

There are two categories of algorithms used to achieve this task: Overlap Layout Consensus (OLC) and de Bruijn Graph (DBG). The usefulness of these algorithms depends on the type of sequencing data produced and the available computational resources.

OLC-based assemblers work the best using long sequence reads such as those produced by the Sanger method and PacBio. Examples of this kind of genome assembler are the CELERA assembler used for the first versions of the *D. melanogaster* and *H. sapiens* genome assemblies, among others \(^{36}\). Given a set of relatively long reads, an OLC based algorithm will first hash the reads into smaller fragments of an arbitrary length - known as K-mers - and perform an all-against-all comparison of these to find overlaps. The overlaps are incorporated into a data structure called a graph, from where consensus sequences called contigs are
inferred. The nature of the algorithm, especially the all-against-all search step, makes these assemblers slow and impractical when working with large amounts of sequencing data\textsuperscript{35,37}, but recent advances have made it possible to assemble genomes to a high level of completion using OLC-based assemblers and long PacBio reads\textsuperscript{25}. Hybrid approaches have been developed to take advantage of high sequence coverage in combination with long sequences with limited success\textsuperscript{38}.

DBG-based assemblers work best with short reads from second generation sequence platforms such as Illumina and 454. The first assembler to implement a DBG approach was EULER\textsuperscript{39}. The DBG algorithm relies on the lower error rate in short reads to build a graph based on K-mers in a similar way to OLC-based assemblers. Overlaps are stored in the graph and the consensus assembly results from finding the most accurate Eulerian path in the graph\textsuperscript{37}. However, the algorithm tends to collapse repetitive sequences and to breaking the consensus when highly heterozygous sequences are encountered, which yields a fragmented assembly\textsuperscript{29}. Assemblers based on DBG are faster, but since the graph of K-mer overlaps is stored in the RAM of the computer, they can be computationally demanding. Despite these drawbacks, several implementations of DBG-based assemblers have been successfully applied to reconstruct the genomes of several organisms\textsuperscript{40,41}.

The final contigs, from both approaches, will contain the best representation of the genome with the given sequencing data and they can be ordered into scaffolds using long insert size mate paired libraries\textsuperscript{42,43}, fosmid pools\textsuperscript{27,44} and long PacBio reads\textsuperscript{45}. Additionally, the scaffolded genome assembly could be further improved by the incorporation of additional data such as optical and linkage maps\textsuperscript{46}.

1.2.3 A recipe for the perfect de novo assembly:

There are many factors involved in the successful reconstruction of a genome sequence, such as the presence of repetitive sequences, the level of heterozygosity, the ploidy of the organism, as well as the estimated genome size. Some studies have tried to benchmark the behaviour of different genome assemblers with different types of data from a set of species\textsuperscript{47,48}, and although these evaluations are an invaluable resource, they do not provide an universal protocol for the perfect assembly. In my opinion, each genome has its own optimal assembly protocol that uses the techniques and algorithms available at a given time.
Nevertheless, thanks to the benchmarking studies, there are some parameters that can be fine-tuned at the beginning of an assembly experiment. For example, in DBG-based assemblers, the first step should be the selection of the K-mer size for graph construction and for OLC-based assemblers, the minimum overlap length between two reads.

Most importantly, the generation of the adequate sequencing data is key for the success of the experiment. For an Illumina-only approach it is suggested to use at least three paired end libraries with insert sizes between 180 - 750 bp as well as at least three mate pair libraries with insert sizes between 2 - 15 Kb. With the recent introduction of third generation long reads, it is possible to sequence and de novo assemble small to medium genomes with a high level of success using these and other ordering data such as linkage and optical maps.

Ideally, a combination of different types of data is required, but this does not guarantee an easy reconstruction of the genome of an organism.

1.3 Methods for comparative genomic analyses:

The genome sequence has been compared to a “blueprint” for life, implying that it is a rigid structure encoding genes for all the biological functions of an organism and without much change beyond point mutation in coding sequences. Recent studies have shown that genomes can be highly dynamic and that their evolution involves a wide range of genomic variants and adaptive evolutionary processes. The generation of de novo mutations that allow for sequence diversity and adaptability is closely linked to the architecture of the genome: retrotransposons, simple repeats, gene clusters, etc.

The first comparative studies used protein and gene sequences to identify sequence changes that may produce be functional. At that time, when the number of sequences was small and their length was usually short, the comparisons were often performed by hand, but once the number of available gene sequences increased from tens to thousands, the first sequence alignment algorithms were implemented to perform comparative gene analyses. With the production of complete genome sequences and introduction of second and third generation sequencing platforms the field of comparative genomics was born, together with the difficulties of handling large amounts of sequencing data and mapping sequences with precision to their specific locations in genomes for comparative purposes.
In this section I will describe the basic methods used for comparative genomics using second generation sequencing data.

1.3.1 Alignment of sequencing reads:

The problem of correct placement of short reads in a genome is known as read mapping and there has been several implementations of methods for highly accurate mapping.

To be able to search the genome for matches with the reads in a rapid, yet accurate way, the genome sequence has to be indexed. A widely used indexing strategy is based on Burrows-Wheeler Transform (BWT), implemented in the BWA\textsuperscript{62} and Bowtie2\textsuperscript{63} aligners. In a BWT-based aligner, the reads are mapped in a seed-and-extend fashion, where segments of the short reads - referred as seeds - are aligned against the indexed genome and the placement is evaluated based on the maximum number of tolerated mismatches (i.e: diversity between individuals of the same species). This process continues until the correct position for the entire read has been found, or not found, in the genome. In the case of paired end reads, the process is extended to both reads in a pair, and if one of the reads cannot be mapped, the other one could still be included in the final alignment\textsuperscript{64}. Genomic regions rich in repetitive sequences constitute a considerable problem for read mapping and some tools have been developed to map reads in these regions correctly\textsuperscript{65}. One of these tools is Stampy\textsuperscript{66}, which uses a statistical model that takes into account not only a per-base match but also the surrounding area of the read to identify potential erroneous placements in repetitive regions. This method makes it possible to in many cases map reads in complex areas of a given genome and even to map sequencing data from other species with a sequence divergence up to 15\%\textsuperscript{65}.

To store the large amount of alignment information generated by these mappers, a new file format was created: The Sequence/Alignment Map or SAM\textsuperscript{67} and its binary version BAM. Currently, BAM files are the standard format to store read alignment data since they are smaller than the human readable SAM files, but due to the rapid increase in the amount of sequencing data and larger, population-scale genomic studies, new versions of the format, such as the CRAM format used in the 1000 Human Genomes Project, have been created to reduce the storage footprint of large genomic datasets\textsuperscript{16,68}. 


1.3.2 Genomic variant calling in a nutshell:

Genomic variant calling is usually achieved by mapping reads or assembled contigs against the complete reference genome sequence of a given organism 69.

The most widely used approach for variant calling is the mapping of reads from second or third generation sequencing platforms. Once the reads have been mapped to a reference sequence, the resulting mapping file has to be processed prior to variant calling. The first step is to sort all the alignments based on their specific coordinates in the genome. Subsequently, reads that are redundant, i.e: PCR and optical duplicates, are removed. Finally, labels to allow the identification of the sample should be added to the BAM file, e.g: sample name, sequencing platform, sample population, etc. Several tools have been produced to streamline the preprocessing of mapping files for variant calling. These include Picard Tools 70, SAMtools 67 and Sambamba 71. After these preprocessing steps, the mapping files are ready to be used for the identification of different types of genomic changes. Additional steps can be incorporated into the preprocessing, such as base quality recalibration, but these depend on the variant calling algorithm to be used and the reader is referred to the GATK Best Practices manual 72 for more details. After these preprocessing steps, the mapping files are ready to be used for the identification of different types of genomic changes.

Several tools have been developed to use mapped reads for genomic variation analysis and, based on their internal algorithm, they perform better in certain genomic regions than in others 73–75. In my experience, a set of high quality, reliable variant calls can be obtained by integrating the output from multiple methods. For instance, the approach taken in the 1000 Human Genomes Project used several variant callers and a consensus was created from all the results 76. In principle, variants are identified by searching for supported mismatches in the reads that can be linked to biology, such as sample diversity, rather than sequencing errors 69. To achieve this, the tool used uses mapping parameters to support a statistical model to weight the confidence of the identified variant 77. The statistical model usually relies on how many bases from the mapped reads support a given variant, the sequence quality of that base, the mapping quality of that base, among others 78. Once the genomic variants have been identified, it is possible to filter more complex situations and this will depend on the purpose of the experiment and the genomic characteristics of the studied organism 79.

The identified genomic variants are stored in a format called Variant Calling Format (VCF). This format is structured such that a given genomic change can be localised quickly in the
genome as well as displaying all the characteristics of the variant. Additional information, such as allelic frequency and genotype are also included\textsuperscript{80}.

These genomic variants are stored in a format called Variant Calling Format (VCF). This format is structured in a way that a given genomic change can be localised quickly in the genome as well as displaying all its possible characteristic. Additional information of the variant in a given context such as allelic frequency and genotype are also included.

Several tools that provide a high level of accuracy have been developed. The Genome Analysis ToolKit (GATK) has been developed for organisms such as human or mouse\textsuperscript{77,81}, but it can be adapted to work with other species\textsuperscript{69}. A different approach has been used in FreeBayes, where variants are identified taking both haplotypes into account directly from the reads\textsuperscript{82}, and a recent re-implementation has been designed for the rapid analysis of genomic variation in human genomes\textsuperscript{83}. An alternative approach to the study of whole genome variation is based on sequence or genome assembly, where instead of comparing reads directly, they are first assembled into contigs\textsuperscript{84}. The advantage of the assembly methods is that it can detect variation in regions that are not well resolved in the reference sequence, such as the HLA locus in the human genome, and they can be much faster than methods that rely on read mapping\textsuperscript{85,86}. In the future, this approach will be useful for the study of genomic variation in organisms with a high degree of genomic complexity.
1.4 The biology of Trypanosoma cruzi:

Trypanosoma cruzi is a unicellular protozoan from the family Kinetoplastida, which contains other unicellular protists such as Leishmania sp., Trypanosoma brucei, Crithidia sp among others 87, and is the etiological agent of American trypanosomiasis, clinically known as Chagas disease 88. The disease and the parasite were first described by Carlos Chagas in 1907 while he was working as a medical doctor for a malaria eradication program in the Brazilian state of Minas Gerais 89. It is estimated that there are 6 - 7 million people infected with T. cruzi just in Latin America 90 but with many more at risk, particularly in the poorest regions 91.

In this section I will provide a very brief introduction to the biology of T. cruzi and its clinical features.

1.4.1 T. cruzi life cycle:

Trypanosoma cruzi has a complex, dual life cycle with four main stages 92,93. As an epimastigote, the parasite has the ability to survive in the insect intestine without causing any detectable pathology. In this stage, T. cruzi undergoes binary fission. When the parasite exits the gut in the vector feces, it transforms into metacyclic trypomastigotes. The vector takes a bloodmeal from the mammalian host by perforating the skin with its proboscide and locally anesthetizes the perforated area with its saliva. Once the vector finishes the meal, it deposits fecal material near the wound area. When the parasite exits the gut in the vector feces, it transforms into metacyclic trypomastigotes. After the anesthetizing effect expires the host auto-inoculates the metacyclic trypomastigotes through skin abrasions at the bite site or other mucosas. When the metacyclic trypomastigotes enter the bloodstream, they transform into the slender trypomastigote form that evades the host immune system and invades host cells, with a preference for myocytes 94. Once the parasite is inside the target cell, it transforms into a rounded form called amastigote, which proliferates quickly inside the host cell forming clusters which disrupt the cell membrane, followed by transformation of the parasites into trypomastigotes that can infect other cells or be ingested by another vector to continue the life cycle 95,96.
1.4.2 Mechanisms of cellular invasion and immune evasion by *T. cruzi*:

*Trypanosoma cruzi* establishes chronic infection by successful evasion of the innate and adaptive immune responses, a strategy shared with other kinetoplastids. However, instead of entering the phagocytic immune cells, as is the case for *Leishmania sp.*, *T. cruzi* invades other nucleated cells.97,98

The complement pathway and its serum components is the most important innate immune response challenge for *T. cruzi* 99,100. Antigenic molecules present on the surface of the parasite trigger the proteolytic cleavage of C3, resulting in C3b molecules that attach to the parasite surface antigens and promote the accumulation of the complement elements involved in pathogen cellular lysis 99,101. To avoid being lysed by the complement pathway, *T. cruzi* releases trans-sialidases and mucins that specifically attach to C3a and C3b and block them.

After *T. cruzi* penetrates the skin or mucosa, it invades non-professional phagocytic cells by attaching itself using surface molecules, such as trans-sialidases and mucins 102. Once
attached, *T. cruzi* can be internalised into the cell by two mechanisms: Ca\(^{++}\)-dependent lysosomal recruitment at the parasite entry site\(^{103}\), a pathway shown to be used by isolates from the TcI clade; or parasite infolding into the plasma membrane with followed by lysosomal fusion, a pathway favored by isolates from the TcII - TcVI clades\(^{94,104,105}\). To date, it is not known if the parasite exploits a specific host cell receptor to promote the internalisation\(^{104}\).

Lysosomes attached to the vacuole containing the parasite and the low pH environment - produced by oxidative radicals - facilitates the transition of the trypomastigote to the amastigote\(^{106,107}\). Inside the acidic phagolysosome, the parasite releases trans-sialidases that remove the sialic acid from lysosome-associated membrane proteins (LAMP1 and LAMP2), which appear to act as structural scaffolds for the phagolysosome\(^{108,109}\). Additionally, the parasite also releases pore-forming proteins that allow the parasite to escape to the cytosol, where the trypomastigote transforms into an amastigote and proliferates inside the cell\(^{110}\). After a variable number of replications, the amastigotes convert to trypomastigotes that disrupt the cellular membrane and are released to the bloodstream, ready to invade new cells.

The intracellular stage of the parasite protects itself from the host innate immune response by hijacking cellular signalling pathways and reshaping the cellular structure\(^{101}\), but the viability of *Trypanosoma cruzi* inside the host depends on its ability to evade the innate and adaptive immune systems. It is possible to detect different sequence variants of surface molecules involved in immune evasion between metacyclic trypomastigotes and amastigotes\(^{111}\).

This ability has been linked to specific gene families, most of them coding for surface molecules\(^{112}\). These molecules have been identified by studying the behaviour of the parasite in different culture conditions and other modified environments\(^{113,114}\). Two of the first gene families to be associated with virulence were trans-sialidases\(^{115}\) and mucins\(^{116}\).

Trans-sialidases were first described in *T. cruzi* in as a modified version of a sialidase enzyme involved in the incorporation of sialic acid on the parasite cellular surface\(^{117}\). Since *T. cruzi* is not able to synthesize sialic acid *de novo*, it is taken from the surface of host cells and transferred to acceptor molecules located on the parasite surface\(^{118}\).

It has been estimated that the reference *T. cruzi* TcVI CL Brener strain contains about 1400 genes that code for trans-sialidase family members\(^{119}\). Based on a sequence clustering analysis, it was concluded from the sequence diversity of these genes that they split into four different groups with individual expression patterns and specific chromosomal locations\(^{120}\). Members of the trans-sialidases gene family are actively involved in the cell invasion...
process, where they participate in the cell adhesion process, thus facilitating the entry of the parasite into a specific host cell. Additionally, it has been observed that *T. cruzi* sheds trans-sialidases to the bloodstream as a way to modify the surface of target cells to facilitate invasion. Trans-sialidases play a fundamental role in parasite immune evasion by suppressing the triggering of the complement pathway. Early studies demonstrated that the removal of sialic acid attached to the parasite surface made it more vulnerable to the host complement system and phagocytosis. A specific group of trans-sialidases genes, described as Complement Regulatory Proteins (CRP), binds to the complement elements C3b and C4b and prevents them from recognizing the parasite. Trans-sialidases may also influence the regulation of T-cells by modifying the sialylation of the cell surface and preventing the activation of these cells against *T. cruzi*. It has also been shown that trans-sialidases can re-sialylate the surface of CD8+ T-cells, and thereby hampering the response of these cells to the parasite invasion. These characteristics, despite the variability, could make trans-sialidases suitable for immunotherapy development and possibly targets for testing new enzymatic inhibitors. More studies are required to understand the complete interaction with the host immune system and the mechanisms involved in the generation of sequence diversity in members of this gene family.

Mucins are heavy glycoproteins that can be found covering epithelial cells of the digestive and respiratory tract of mammals. They are also found in the surface membrane of *T. cruzi* and other protozoans as acceptors of sialic acid. In this way a negatively-charged sugar coat is formed as a means to evade the immune response of the host and the vector. This gene family, with approximately 900 gene copies, is one of the most diverse in the *T. cruzi* genome and the genes are variable in sequence and structure between isolates.

The TcMUC mucins are expressed in the mammalian host while TcSMUG mucins are expressed in the vector and both are post-translationally modified in a different way. Previous studies have shown that mucins have an important role in parasite immune evasion by stimulating dendritic cells via TLR2, and this activation has been found to delay the rapid response of other elements of the immune system, such as CD8+ T-cells. A novel gene family, Mucin-associated surface proteins, was identified after the first draft genome sequence was published. These genes comprise 6% of the CL Brener genome and are found near Mucin tandem gene arrays and they are expressed during the trypomastigote stage, which makes them possible vaccine candidates. The expression patterns of MASPs in the trypomastigote have been observed to be tissue-specific, and they seem to be more actively
Protozoan genes coding for specific molecules, such as mucins with glycosilphosphatidylinositol (GPI) anchors, are recognised by the innate system via Toll-like receptors (TLR) and promote the activation of innate effectors like macrophages and dendritic cells \(^{97,134}\). After the GPI-anchored molecule is recognised by TLR4 or TLR2, members of the mitogen-activated protein kinase (MAPK) and nuclear factor kB (NF-kB) are activated and trigger the release of IL-12, TNF and INF-\(\gamma\)\(^{129,134}\). Other receptors, such as TLR-9 can trigger the same response by recognition of nucleic acids \(^{135}\). *T. cruzi* use innate immune response pathways, specifically TLR-2 and TLR-4, to establish chronic infection. One example is the induction of protein phosphatase 2A by *T. cruzi*, which leads to the deactivation of MAPK and NF-kB molecules, thus promoting immune tolerance to secondary exposure \(^{136}\).

The slow initial response of the host cellular immunity is the direct result of the evasion of innate immunity by *T. cruzi* \(^{98,137}\). The adaptive immune response is triggered by cell death after the parasite has disrupted the cellular membrane, which results in a strong, CD8+-mediated immune response which targets the antigens exposed on the parasite surface at the late amastigote and early trypomastigote stages \(^{138}\). However, this parasitic antigenic coat is very variable and by the time the adaptive immune response has been mounted to target these antigens, a new antigenic combination of trans-sialidases and mucins is expressed on the parasite surface \(^{139}\).

### 1.4.1 The population structure of *T. cruzi*:

Early studies using a set of biochemical \(^{140}\) and molecular \(^{141}\) markers indicated that *T. cruzi* has an exclusively clonal population structure \(^{142}\). This clonal theory is based on using different sets of markers that show a clonal behaviour based on population genetic tests, such as high levels of linkage disequilibrium (LD), the presence of ‘near clades’ and overrepresentation of multilocus genotypes \(^{143}\). Analyses of genetic signatures in the kinetoplast minicircle (kDNA) indicated that these clonal groups had a single origin, thus reinforcing the clonal model for *T. cruzi* \(^{144}\). Later meta-analyses using additional markers such as isoezymes, Random Amplified Polymorphic DNA (RAPD), microsatellites, among others identified signatures of long-term clonal evolution and the clustering of the current strains into subgroups denominated as Discrete Typing Units (DTUs) \(^{145}\). Further
characterisation using a wide panel of markers in selected reference isolates established the subdivision of *T. cruzi* into six DTUs \(^{146,147}\). These clonal DTUs have been observed to have different patterns of virulence and tissue tropism in the mammalian host \(^{148}\).

However, these observations were challenged when genetic interchange to produce hybrid strains was demonstrated \(^{149,150}\). Analyses of different nuclear and mitochondrial markers have demonstrated that hybridisation events have occurred \(^{151}\) and that these hybridisation events have created three major groups \(^{10,152}\). Moreover, active genetic exchange in field isolates have also been demonstrated using nuclear and mitochondrial markers \(^{153}\). Population genetic studies of isolates derived from humans, indicated the existence of natural hybrids and strain variability linked to anthroponotic dispersal \(^{154}\). Furthermore, evidence of mitochondrial introgression has been observed in field and *in vitro* isolates \(^{155}\).

As a response to this challenging observations, the clonal model was further developed into the Predominant Clonal Evolution (PCE) model \(^{156,157}\), which states that despite the observation of genetic exchange in *Trypanosoma cruzi*, and other parasites, the exchange is rare and the main model of propagation is clonal, with random cases of selfing and parthenogenesis \(^{158}\). Thus far, the population genetics analyses of field isolates does not agree with this model \(^{159}\), and large studies have confirmed that there is more divergence within a single DTU than what would be expected under a clonal model \(^{160}\).

To date, the population structure of *T. cruzi* is far from completely understood.
CHAPTER 2 – PRESENT RESEARCH

2.1 Paper I and Paper II: Comparative genomics of Trypanosoma cruzi: Insights into mechanisms of virulence from other trypanosomatids.

The first draft genome assembly for Trypanosoma cruzi was produced using first-generation, Sanger sequencing data\textsuperscript{10} and was used to performed the first comparative analysis with the other kinetoplastid species Trypanosoma brucei and Leishmania major\textsuperscript{161}. This first assembly provided an adequate reconstruction of the core regions where the housekeeping genes are located, despite a high degree of polymorphism in the CL Brener reference strain, but the highly repetitive subtelomeric regions, where most of the surface molecule gene families are encoded, were only partially assembled and highly fragmented\textsuperscript{162,163}. In paper I and paper II we sequenced and assembled a less polymorphic TcI strain of T. cruzi from high sequence coverage of different types of second-generation reads, and compared this genome against two kinetoplastids that do not cause disease in the human host: Trypanosoma cruzi marinkellei (MARK) which is restricted to bats\textsuperscript{164} and Trypanosoma rangeli (RANG) which is highly prevalent in opossums but also common in humans\textsuperscript{165}.

In paper I we used high coverage of Illumina and 454 short reads and a hybrid assembly approach to reconstruct the genome sequence of T. cruzi TcI Sylvio X10/cl1 and MARK and performed comparative analyses. This strategy made use of the assemblers developed for early second-generation sequencing technologies with reads ranging between 50 - 75 nucleotides (nt) in length, such as the Velvet assembler\textsuperscript{166}; and the Celera assembler for the, relatively, long (~ 350 nt) 454 reads\textsuperscript{167}. Since genome assemblers perform better in different genomic regions and with different datasets\textsuperscript{168}, merging the individual assemblies produced a more contiguous and correct de novo draft sequence for these parasite genomes. This resulted in improved assemblies, producing the most contiguous T. cruzi genome sequence yet, when compared with previous attempts using only 454 short reads\textsuperscript{169}. This new assembly made it possible to perform intra-species comparative genomics for the first time. The initial analysis revealed that the two sub-species displayed a coding sequence divergence of nearly 7.5 \% and MARK was found to have a smaller genome compared to Sylvio X10/cl1. Both protozoans shared the same core genome components but the Sylvio X10/cl1 genome has more genes coding for surface protein families involved in infection and possible host specificity (See paper III).
In paper II high coverage of 454 paired end data was used to assemble the genome of *T. rangeli*. The lower repeat content of RANG made it possible to reconstruct the complete genome using the 454 short reads, which combined with the linking information produced long scaffolds. This would not have been possible for a more complex genome, where only this type of sequencing data is not enough to produce such contiguity \(^{167,168}\). We could assemble the complete genome of RANG into only 259 scaffolds with an average length of ?, with a mean scaffold size of 202 kilobases. The low repetitive content of RANG allowed the genome assembler to reconstruct the complete genome using the 454 short reads, which combined with the linking information produce long scaffolds; unlike more complex genomes where these sequencing data is not enough to produce such contiguity \(^{170,171}\). A drastic reduction in the copy numbers of surface molecule gene families was observed when compared with the *T. cruzi* CL Brener strain. The reduction of genes such as trans-sialidases and mucins could indicate an adaptive evolution process by genome reduction \(^{59,172}\), and it can be speculated that the absence of these genes could be involved in the inability of RANG to cause disease in mammals \(^{173}\). The levels of simple repeats in RANG were much lower than those observed in Sylvio X10/cl1 (See paper III), but the overall reduction of genome size seems to be almost entirely related to the contraction of repetitive gene families, which is in line with selection to reduce paralog content to redirect resources to other biochemical pathways that may confer an adaptive advantage \(^{174}\). On the other hand, the large numbers of members of these gene families in *T. cruzi* could indicate their importance for the the invasion of particular cells and tissues \(^{175}\).
Figure 2: Structure of *T. rangeli* telomeres showing reduction of surface molecule gene families in comparison with *T. cruzi* and *T. brucei*. (See paper II).

Subsequent recent comparative analysis carried out using Illumina reads (the methodology is described in paper III) from these species against the Sylvio X10/cl1 reference confirmed the stable nature of the core genome in these species, but a low mappability rate in the subtelomeric regions, which indicates that these regions have a more rapid rate of evolution than the core genome. Similar characteristics have been noticed in fungi, where genomic areas containing genes coding for virulence factors have a more rapid evolutionary rate compared to regions containing housekeeping genes, to the point of rendering them sample specific 176–178. Large genomic rearrangements were observed across the entire genome of these two protozoans when compared against the Sylvio X10/cl1 reference, including core regions, implying an active genomic restructuring process.

In the case of MARK, interspersed duplications were observed in chromosomal regions rich in mucins and retrotransposons while tandem duplications occurred in smaller regions within tandem arrays of trans-sialidases, mucins and MASPs. Multiple InDels with a slight bias towards short insertions, between one and five nucleotides were observed in the subtelomeric regions of MARK, accompanied by high levels of sequence diversity (7.24 SNV/Kb) in these areas compared with the core regions (1.1 SNV/Kb) when using Sylvio X10/cl1 as a reference; which together confirms that the surface proteins of MARK are diverged from those of *T. cruzi*, in response to the adaptation to different niches.
For RANG, deletions ranging from one to four kilobases in the subtelomeric regions were the most common genomic rearrangement. Interestingly, segmental duplications of regions containing trans-sialidase and mucin tandem arrays that were observed in MARK were absent in the same regions for RANG. Additionally, large genomic inversions in the subtelomeres and core regions were present exclusively for RANG in chr4 and chr16. Unlike the InDels observed in the MARK genome, it was not possible to detect an insertion or deletion bias in RANG, which could suggest a different mechanism of recombination in these regions compared to MARK or *T. cruzi*. Another characteristic of RANG was the lower sequence diversity observed in the subtelomeres compared with that of MARK (3.4 SNV/Kb).

**Figure 3**: Comparative analysis of *T. rangeli* and *T. cruzi marinkellei* against chromosome 2 of the *T. cruzi* Sylvio X10/cl1 reference strain. The large grey box represents a large (> 200 Kb) duplication in chromosome 2 for *T. cruzi marinkellei*. Green boxes represent retrotransposons, clear blue represent genes, red and dark blue segments represent SNV.

In conclusion, the expanded comparative analysis of these two trypanosome species with *T. cruzi* emphasised the specialised role of the subtelomeric gene families in the adaptive evolution of the parasite to new niches. These gene families can be expanded or eroded from the genome depending on their adaptive role and the extant paralogs are used to develop new specialised function tailored to invade cells and evade the immune system of a given host.
2.2 Paper III: Genome analysis of the *Trypanosoma cruzi* Tcl clade reveals mechanisms to generate antigenic diversity.

In this study, using high coverage of third generation sequencing data, it was possible to reconstruct the complete genome sequence of a *Trypanosoma cruzi* Tcl strain at the whole chromosome level, revealing the entire genomic architecture of the elusive subtelomeric regions of this protozoa for the first time. This new reference sequence and whole-genome data from 35 *T. cruzi* Tcl isolates and clones from different regions of the American continent were used to identify mechanisms involved in antigenic diversity generation in the parasite.

![Figure 4: Genomic diversity of Panamanian *T. cruzi* DTU-I isolates from patients. A 505 Kb segment from chromosome 3 shows the levels of sequence diversity in subtelomeric areas coding for trans-sialidases (far left) and core regions (centre).](image)

2.2.1 The *T. cruzi* genome architecture:

The genome of the *T. cruzi* Sylvio X10/cl1 reference isolate contains a large amount of retrotransposons (3.06 Mbp) and microsatellites (1.4 Mbp) interspersed throughout the genome, but with a significantly higher density in subtelomeric regions. Previous estimates of repetitive content - using second generation sequencing data - were much lower \(^ {169}\), most likely due to the inability of genome assemblers to reconstruct these complex areas using second generation data \(^ {29}\). The subtelomeric gene families are structured in tandem arrays consisting of two to three complete gene copies and a variable number of pseudogenes separated by microsatellite segments of variable length. Retrotransposons of the VIPER and R1 class were observed within or surrounding the tandem gene arrays. The presence of retrotransposons close to surface molecule genes and their pseudogenes generated the hypothesis that these elements are involved in the generation of new sequence variants. One
possible mechanism for this could be an RNA-guided DNA insertion mechanism\textsuperscript{179}, as has been reported in other eukaryotes\textsuperscript{180}. However, other types of data, such as RNA-Seq are required to confirm this.

### 2.2.2 Mechanisms for antigenic diversity generation:

By studying the pattern of InDels and genomic rearrangements between TcI strains in the subtelomeric regions we identified two possible mechanisms to give rise to new sequence variants in the surface molecule repertoire. These mechanisms appear to be linked to the genomic architecture of the subtelomeres.

The analysis of InDels in the 35 DTU-I isolates revealed a strong bias towards one to three nucleotide insertions in genomic segments rich in microsatellites and retrotransposons, a bias that has been previously associated with recombination hotspots\textsuperscript{181,182} and gene conversion\textsuperscript{183,184}. These InDels were found within microsatellites containing tandem repeats with motifs such as n(A), n(AT) and n(AG) or within the LTR region of VIPER retrotransposons. The microsatellites and retrotransposons provide a source of homology that can be exploited by a double strand break (DSB) mechanism for recombination that in turn can generate sequence diversity\textsuperscript{185–187}. The microhomology provided by LTRs and microsatellites suggests two possible mechanisms: Non-Allelic Homologous Recombination (NAHR)\textsuperscript{188} and Microhomology-Mediated End Joining (MMEJ)\textsuperscript{189}. MMEJ has been observed in\textit{T. brucei} in the presence of sequence repeats\textsuperscript{190} and a recent study, which applied CRISPR-Cas9 to\textit{T. cruzi}, revealed signatures of DSB repair by MMEJ\textsuperscript{191}. On the other hand, NAHR has been associated with retrotransposons in higher eukaryotes\textsuperscript{192} and the generation of \textit{de novo} structural changes during meiosis\textsuperscript{193}, which has not been reported in\textit{T. cruzi} but in other protozoans\textsuperscript{194}.

Subsequently, we analysed genomic rearrangements in the 35 DTU-I genomes to identify actual recombination events in subtelomeric regions. In\textbf{ paper III} we have identified large structural variants such as deletions, tandem and interspersed duplications, genomic inversions larger than ten kilobases, as well as translocation-associated break ends occurring
in subtelomeric regions in *T. cruzi*. These types of large genomic events have been associated with NAHR\(^{195}\) but this is not the only possible mechanism, since NHEJ has also been linked to the generation of sequence translocations in AT-rich areas in humans, but at a much lower frequency than NAHR\(^{196}\). The breakpoints of the interchromosomal rearrangements were detected using an improved mapping strategy and a consensus of different methods to detect genomic rearrangements (See paper III). The breakpoints were frequently located within the coding sequences of surface molecules genes, short microsatellite segments and LTRs of VIPER retrotransposons.

Copy Number Variation (CNV) analysis in these isolates revealed a dynamic pattern of expansion and contraction of surface molecule gene family gene clusters. These were unique to each isolate regardless of their geographic origin or sample source, e.g: isolated from vectors or humans. Based on the results of the interchromosomal breakpoint mapping, it could be assumed that these expansions and contractions of specific gene families are the result of active intra- and inter-chromosomal reshuffling of subtelomeric segments for adaptive purposes, similar to what has been observed in certain fungi\(^{176}\), which may be retrotransposon-driven\(^{178,197}\).

**Figure 5**: Copy Number Variation (CNV) profile of chromosome 2 of a DTU-I isolate from Ecuador. The blue line represents the reference sequence. Each black dot depicts a coverage sliding window. The red line represents
the sample genome in comparison with the reference. The X-axis represents the position in the chromosome in base-pairs; the y-axis represents fold-change.

### 2.2.3 The population genetics of the *T. cruzi* TcI clade:

WGS data from 35 *T. cruzi* DTU-I isolates from different geographic locations of the American continent was used to study the genomic diversity of this clade. A PCA analysis using InDels revealed that these isolates mostly formed well defined clusters based on their geographic location, a pattern that may be a reflection of adaption to the specific environment and selective pressure, similar to that observed in many other parasitic species \(^{198,199}\). Linkage Disequilibrium (LD) was scanned genome-wide using the \(r^2\) statistic, which revealed that subtelomeric regions exhibit \(r^2\) values close to zero while core regions had values close to one \(^{200}\). The lower \(r^2\) values and the patterns of balancing selection observed in the subtelomeric regions indicate that gene the families in these areas evolve rapidly, while the core genome changes more slowly, implying a genomic duality similar to the one observed in certain fungi \(^{177,201}\).

**Figure 6:** TcI linkage disequilibrium \(r^2\) matrix for chromosome 16. X-axis represents the number of markers; y-axis represent the number of 10 Kb sliding windows. The \(r^2\) values close to 0 indicate recombination, while \(r^2\) values close to 1 indicate clonality.

Multiple diverse genotypes, reflected in the patterns of positive Tajima’s D, were observed in the subtelomeric gene families in parasites from different geographic locations. Samples isolated from vectors showed high levels of balancing selection in the subtelomeric regions whereas parasites isolated from mammalian hosts showed signatures of selective sweeps in the same regions. The balancing selection observed among Colombian clones derived from the same primary isolate, indicated that the parasite is constantly generating sequence diversity in the subtelomeres. The average \(F_{st}\) value for samples isolated from vectors was \(F_{st} = \sim 0.12\) whereas for human isolates it was \(F_{st} = \sim -0.05\) suggesting little to moderate genetic differentiation, respectively \(^{202}\).
Figure 7: Distribution of Tajima’s D in 10 Kb sliding windows for chromosome 10 in Panama samples, where a) represents isolates derived from human patients and b) represents isolates derived from vectors. Red areas show values with negative Tajima’s D values (i.e: selective sweeps) and cerulean areas indicate positive Tajima’s D values (i.e: balancing selection).

In conclusion, these data indicate that T. cruzi generates new antigenic variants via an active process of subtelomeric recombination - possibly mediated by MMEJ - while the core genome remains evolutionarily stable. Balancing selection in subtelomeric regions has resulted high levels of polymorphism for the surface protein gene families present in these areas. New alleles in these genes may confer an adaptive advantage since they can be exploited by the parasite for immune evasion or rapid adaptation to the immune system of a new host, as reported in other eukaryotes.
2.3 Paper IV: Comparative genomic analyses of *Trypanosoma cruzi* experimental hybrids reveal mechanisms of genetic exchange.

The identification of hybrid parasitic strains and genetic exchange is of great public health importance for the control of Chagas disease and the development of new drug targets \(^{205}\). Earlier studies using biochemical \(^{206}\) and single-locus molecular markers \(^{207}\) suggested that *Trypanosoma cruzi* has a clonal population structure. However, more recent studies using molecular markers and genome sequencing (ref) as well as the formation of hybrids *in vitro* have shown that genetic exchange occurs in *T. cruzi* \(^{150}\). These finding were later corroborated in multiple isolates \(^{151}\), which led to the restructuring of the *T. cruzi* population history \(^{146}\). In paper IV we extended the original analysis of in vitro hybrid strains using high coverage, whole genome sequencing of the parental and hybrid strains at different time-points.

### 2.3.1 Genome analysis of the parent strains:

Two IPE libraries and a single IMP library with an average insert size of eight kilobases from both parental strains were sequenced at a total coverage depth of 140 X per each sample. These data were used to create *de novo* assemblies of the two parental strains for comparative purposes. Despite the high sequence coverage and the long-insert library, the repetitive nature of the *T. cruzi* genome made it impossible to completely reconstruct these genomes. As mentioned earlier, it is clear that either high coverage of single molecule long reads, or multiple data sets are needed to decipher the repetitive regions of the *T. cruzi* genome \(^{171,208,209}\). We were able to completely assemble the core regions, where most of the housekeeping genes are located, and to evaluate the conservation of core gene synteny between the parent strains and our Tc1 reference sequence, as previously described \(^{210,211}\). The synteny was found to be well conserved, which is in agreement with previous analyses of Tc1 strains. Since the parental assemblies did not cover the entire genomes, we decided to proceed with the comparative analysis using the sequencing reads and taking advantage of the newly assembled and closely related Sylvio X10/1 reference genome.
Figure 8: Synteny in the core regions between the parental strains PI-0 and PII-0 and the Sylvio X10/cl1 reference genome. Each coloured block represents a syntenic block in each sample. It can be noted that some blocks have been expanded in the parental strains (clear green and plum) or missing (dark blue).

A systematic analysis of genomic variation in the parental strains showed that the vast majority of the sequence and structural changes were limited to the subtelomeric regions, which is in agreement with the analyses of other T. cruzi TcI strains (See paper III). Multiple genome rearrangements were detected and the most common were sequence break-ends similar to the unbalanced translocations observed in higher eukaryotes. Copy number variation (CNV) analyses showed that the subtelomeric regions, and certain loci in the core regions were expanded compared with the Sylvio X10/1 reference strain, and these expansions accounted for the higher repetitive content of the parents and this made it even more difficult to assemble the parent strains de novo. This observation also agrees with the variability of the genome size in T. cruzi field isolates from the same clade as well as differences in the karyotype of isolates from the same clade.

Interestingly, the subtelomeric gene families in the parental strains were found to have lost gene copies after 800 generations of growth in culture. The eroded regions contained genes involved in cell invasion and immune system evasion such as trans-sialidases and mucins. This may indicate that T. cruzi evolves in culture by selecting for the removal of genes that are not necessary in this environment, to allocate resources to other pathways. This mechanism has been widely observed in other species. This observation has implications for future evolutionary studies of the T. cruzi surface molecule repertoire and requires further analysis and experimental follow-ups.
Figure 9: CNV evolution of a parental strain over time. PI-0 is the sample at the initial state, and PI-120 are isolates after 800 generations. Each black dot depicts a coverage sliding window. The red line represents the sample genome in comparison with the reference. X-axis represents the position in the chromosome in base-pairs; y-axis represents fold-change.
2.3.2 Genome analysis of the hybrid strains:

Strain-specific genomic variants, such as SNPs and InDels, were identified in the parental clones to make it possible to identify parent-specific genetic material in the hybrid offspring. We compared the reads from the hybrid genomes against the *T. cruzi* Sylvio X10/c1 reference and detected parent-specific signatures distributed throughout the genome as well as new mutations. Parent-specific genomic blocks in the hybrid isolates were surrounded by short InDels, principally insertions ranging in size between one and five nucleotides, found within long stretches of simple repeats in intergenic regions and within the LTR segments of VIPER retrotransposons. As expected, the hybrid strains had a higher level of polymorphism in both housekeeping and surface molecule genes compared to the parental strains, due to the presence of both parental haplotypes for much of the genome. It is tempting to speculate that these genomic signatures are the result of microsatellite instability associated with defective DNA mismatch repair \(^\text{219}\) and recombination associated with cell division \(^\text{220–222}\). While the meiotic machinery has been described in *T. cruzi* and other protozoan parasites \(^\text{194}\) and, although the presence of gametes has been reported in other kinetoplastids \(^\text{223}\), there is no experimental evidence of gametes in *T. cruzi*. It is therefore still more likely that the recombination occurs during mitosis.

![Figure 10](image)

**Figure 10**: a) Patterns of parent-specific genomic material in a hybrid. b) CNV in a trans-sialidase tandem array of a hybrid strain (red lines) compared with the reference (blue line). c) InDels (red and blue bars) around breakpoints (grey bars) in the hybrid offspring. Red blocks show genes and green blocks show retrotransposons.
By analysing the sequence coverage distribution and the pattern of structural changes in the hybrid offspring, it was possible to identify, almost entirely subtelomeric, regions that were expanded or eroded after the hybridisation event. These sequence expansions increased the genome sizes of the hybrids between 34.1% and 48.6 % compared to the parents, which seems to be why hybrid strains, such as TcVI CL Brener have larger subtelomeres, compared to non-hybrid strains such as TcI Sylvio X10/1. It is currently unclear whether the expansion occurs through a 4n hybrid intermediate or through a different mechanism. The extensive expansion of surface molecule genes and the genomic rearrangements observed in these samples are very similar to the ones observed in pathogenic fungi, where they have been associated with increased virulence. It is possible to hypothesise that the combination of new surface molecule genes from both parents provides the hybrid strains with a pool of new sequences as substrates for the MMEJ-like recombination mechanism, as observed in other TcI strains, to create new antigenic variants in order to increase the potential for forming viable hybrid offspring (See paper III). After 800 generations of growth in culture, the hybrid genomes showed a similar pattern of genome erosion as in the parent strains.
CHAPTER 3 – FUTURE PERSPECTIVES

Unlike other protozoan parasites, such \textit{Plasmodium falciparum} and \textit{Trypanosoma brucei}, the lack of a complete reference sequence has hampered the implementation of post-genomic studies in \textit{Trypanosoma cruzi}.

The present work describes the complete genome assembly and analysis of a \textit{T. cruzi} TcI strain, furnishing the parasitology research community with a valuable resource to better understand the biological aspects of Chagas disease. The new reference sequence will serve as the scaffold for the implementation of large scale population genomic studies, gene expression analysis in different stages of the parasite life cycle, the characterisation of transcription dynamics of the parasite virulence factors, just to mention a few.

In this work, using this genome sequence and an integrative genomic data analysis approach, it was possible to characterise a retrotransposon-driven mechanism that could be involved in the generation of antigenic variation in \textit{T. cruzi} which could potentially be targeted with chemotherapeutic agents. However, this is just a small step in the application of post-genomic methods quest for new therapeutic interventions for Chagas disease and new analytical strategies should be implemented.

There is an urgent need for the discovery of new, more effective drug targets and vaccine candidates for Chagas disease and other parasitic malaises, but the identification of those, with the current methodologies, is an expensive and time-consuming process\textsuperscript{224}. Genomic approaches have opened a new way to analyse complete gene families of protozoan genomes and their potential interaction with the host, providing a new alternative to identify drug targets and vaccine candidates more efficiently and cost effective.

\textit{Omics} datasets - composed by genomics, transcriptomics, proteomics, among others - provide an attractive source of biological information for different pathogenic organisms stored in public databases, such as the NCBI Short Reads Archive (SRA)\textsuperscript{225}. Some initiatives are already planning to use large public and private genomic datasets to identify links between genes and diseases in a personalised way\textsuperscript{226}. These large datasets could be mined with the latest data analytics\textsuperscript{227,228} for the identification of vaccine candidates, new drug targets and the synergy between these molecules and the human host, in a cheaper, faster and effective way.
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