Regulation of the Extracellular Matrix Interactome by *Trypanosoma cruzi*

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**Abstract:** It has been shown that the invasive trypomastigote forms of *Trypanosoma cruzi* use and modulate components of the extracellular matrix (ECM) during the initial process of infection. Infective trypomastigotes up-regulate the expression of laminin γ-1 (LAMC1) and thrombospondin (THBS1) to facilitate the recruitment of trypomastigotes to enhance cellular infection. Silencing the expression of LAMC1 and THBS1 by stable RNAi dramatically reduces trypansom infection. *T. cruzi* gp83, a ligand that mediates the attachment of trypanosomes to cells to initiate infection, up-regulates LAMC1 expression to enhance cellular infection. Infective trypomastigotes interact with LAMC1 through galectin-3 (LGALS3), a human lectin, to enhance cellular infection. Silencing the expression of LGALS3 also reduces cellular infection. Some trypansom surface molecules also interact with the ECM to facilitate infection. Despite the role of the ECM in *T. cruzi* infection, almost nothing is known about the ECM interactome networks operating in the process of *T. cruzi* infection. In this mini review, we critically analyze and discuss the regulation of the ECM by *T. cruzi* and its gp83 ligand, and present the first elucidation of the human ECM interactome network, regulated by *T. cruzi* and its gp83 ligand, to facilitate cellular infection. The elucidation of the human ECM interactome regulated by *T. cruzi* is critically important to the understanding of the molecular pathogenesis of *T. cruzi* infection and developing novel approaches of intervention in Chagas’ disease.

**Keywords:** Extracellular matrix (ECM), *Trypanosoma cruzi*, cellular infection, ECM interactome.

REGULATION OF THE ECM BY *T. CRUZI* AND ITS GP83 LIGAND

The importance of the extracellular matrix (ECM) in *T. cruzi* infection has been highlighted [1, 2]. Our group and others have employed gene transcription microarray technology and have shown that certain host genes, including ECM genes, are significantly modulated by the parasite to facilitate the process of infection [3-8]. An integrative analysis of the observed gene expression profile at the transcriptome level, as well as protein-protein interaction (PPI) networks, can provide insights into the molecular mechanisms that contribute to the pathogenesis of Chagas’ disease. PPI network analysis has thus far not been exploited to elucidate the complex biological interactions occurring during the initial phases of *T. cruzi* infection. Our group studied the gene expression profile of human coronary artery smooth muscle cells (HCASM) modulated by *T. cruzi* trypomastigotes and the *T. cruzi* gp83 ligand [9-11] and here we report the interactome network operating in HCASM cells during the early process of *T. cruzi* invasion. The interactome data has shown that specific interactions between some ECM proteins are regulated by *T. cruzi* during early infection. THBS1, LAMC1, LGALS3 and fibronectin are part of the ECM interactome that *T. cruzi* regulates and uses in order to facilitate host cell trypanosome interactions leading to cellular infection.

We have demonstrated that *T. cruzi* up-regulates the levels of THBS1 expression in host cells during early infection. Stable RNA interference (RNAi) of host cell THBS1 knocks down THBS1 transcripts and protein expression causing inhibition of *T. cruzi* infection [4]. Exogenous THBS1 restored the level of infection of these THBS1 knock-down cells. Thus, host THBS1, regulated by the parasite, plays a crucial role in early infection. *T. cruzi* gp83, a ligand that *T. cruzi* uses to attach to host cells [9], increases the level of LAMC1 transcript and its expression in mammalian cells, leading to an increase in cellular infection. Stable RNA interference (RNAi) of host cell THBS1 knocks down LAMC1 transcript levels and protein expression in mammalian cells, causing a dramatic reduction in cellular infection by *T. cruzi* [3]. Furthermore, human LGALS3 binds to a trypomastigote surface mucin [12, 13] and to HCASM cells in a lectin-like manner [14] to significantly increase the cellular adhesion of trypomastigotes. Silencing LGALS3 expression in mammalian cells by antisense approach significantly reduces trypomastigote adhesion to cells.

*T. cruzi* infection causes extensive fibrosis and severe heart cardiomyopathy, which includes vasculopathy, leading to cardiac arrest that is frequently followed by death [15]. The fact that *T. cruzi* trypomastigotes modulate the expression of ECM proteins and subsequent PPI networks suggests that the altered ECM may contribute to the pathology manifested in Chagas’ disease. Here we present the first elucidation of the host cell ECM interactome network induced by *T. cruzi* and its gp83, a critical ligand used by the trypanosome to bind host cells, to facilitate cellular infection.

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This is consistent with the first gene network analysis induced by *T. cruzi* and its gp83 in human cells [16].

As reported previously, *T. cruzi* trypomastigotes are able to increase the levels of THBS1 expression in host cells during early infection. Thrombospondins constitute a set of unique multimatrix proteins that are synthesized, secreted and incorporated into the ECM by many cell types [17, 18]. Thrombospondins are unique members of the ECM; they have been described as ‘matricellular’ proteins. Most thrombospondin isoforms are expressed in myocytes, smooth muscle cells, fibroblasts and endothelial cells, all of which *T. cruzi* infects.

Little is known about the role that host THBS1 isoforms play in the process of microbial infections. It was suggested that *T. cruzi* might have receptors for thrombospondin [19]. Since nearly all cells that *T. cruzi* infects are surrounded by basement membranes (of which THBS1 is a key component), the ability of the parasite to effectively regulate and interact with THBS1 is also critically important for its passage through the host cell membrane.

The kinetics of change in gene transcript profiles of THBS1, THBS2, THBS3 and THBS4 was evaluated by real time PCR during the early process of infection of HCASM cells by *T. cruzi*. It was shown that THBS1 transcript levels increase approximately 6-fold at 60 min, followed by an increase of 2.7-fold at 120 min during infection of HCASM cells by *T. cruzi*. The transcript levels of THBS2, THBS3 and THBS4 do not change significantly during the same period of time of *T. cruzi* infection of HCASM cells [4].

TGF-β1 induced over-expression of THBS1 in human epithelial cells, causing a significant increase in the number of trypansomes bound to cells and parasite load as compared to cells transfected with vector alone or scrambled THBS1 antisense. Thus, there is a direct correlation between over expression of THBS1 in human cells with an increase of infection of these cells. *T. cruzi* infection of mammalian cells is inhibited by stable RNAi of host cell THBS1 [4]. Addition of THBS1 to these cells restores infection. Thus, host THBS1, regulated by the parasite, plays a crucial role in early infection. RNAi silencing of THBS1 does not alter the expression of other ECM proteins such as fibronectin and LAMC1 and/or LGALS3, nor their abilities to adhere to substrates, supporting the specificity of these studies. Therefore, these results indicate that THBS1 is required for the process of *T. cruzi* infection.

Since *T. cruzi* modulates the ECM by up-regulating THBS1 expression, the parasite is able to exploit THBS1 in addition to LAMC1 to navigate through the ECM and facilitate infection. THBS1 is required for the infection process of *T. cruzi* as evidenced by RNAi of that specific isoform. The *T. cruzi* up-regulation of host THBS1 expression to facilitate infection of human cells represents a new mechanism that may contribute to elucidating the pathogenesis of *T. cruzi* infection.

Similarly to THBS1, LAMC1 is also up regulated by *T. cruzi*, specifically by its gp83 surface glycoprotein. An additional *T. cruzi* trypomastigote surface glycoprotein, Tc-85 that binds to laminin, has also been implicated in infection [20]. LAMC1 is the most abundant laminin isoform in humans [17] and stable RNAi of host cell LAMC1 knocks down both LAMC1 mRNA transcript and protein levels in mammalian cells. As a result of silencing LAMC1, there is a dramatic reduction in cellular infection by *T. cruzi* [3]. Thus, host LAMC1, which is regulated by the parasite, also plays a crucial role in the early process of infection.

The fact that a *T. cruzi* trypomastigote ligand, gp83, increases LAMC1 transcript levels in HCASM cells correlates with the finding that laminin is deposited in the hearts of patients infected with Chagas’ disease [21]. This suggests that the regulation of LAMC1 in cardiomyocytes by gp83 may partially explain the cause of Chagas’ disease pathology.

Our findings indicating that *T. cruzi* gp83 ligand remodels the ECM by up-regulating the expression of LAMC1, together with the report that *T. cruzi* presents laminin receptors on its surface [20], suggests that the parasite exploits LAMC1 to navigate through the ECM and facilitate infection. Thus, the *T. cruzi* gp83 ligand is a virulence factor that modifies LAMC1 expression in the ECM to contribute to the pathogenesis of *T. cruzi* infection in human cardiomyocytes.

LAMC1 also interacts with LGALS3, a human lectin which binds β-galactoside. This interaction is thought to play an important role in the early steps of *T. cruzi* infection of human cells [14]. LGALS3 is known to bind to the surface of HCASM cells, as well as to the surface of invasive *T. cruzi* trypomastigotes [14]. The binding of LGALS3 to the surface of both *T. cruzi* trypomastigotes and HCASM cells is almost completely inhibited by lactose (but not by sucrose), indicating that LGALS3 binds to the surface of both trypomastigotes and HCASM cells in a lectin-like manner [14]. LGALS3 increases by five-fold the adhesion of trypomastigotes to HCASM cells; this effect is specific since it is inhibited by lactose in a concentration-dependent and saturable manner. On cells which LGALS3 has been knocked down by antisense approach there was a significant reduction of bound trypanosomes compared to control cells. This effect of LGALS3 knock-down is reversed by exogenous LGALS3, indicating that cellular expression of LGALS3 is required for *T. cruzi* adhesion to human cells.

The fact that exogenous human LGALS3 specifically binds to the surface of HCASM cells and to *T. cruzi* suggests that LGALS3 bridges trypanosomes and cells, resulting in an enhancement of *T. cruzi* binding to human cells. The proposed mechanism by which this interaction occurs is through the association of LGALS3 molecules with *T. cruzi* 45 kDa mucin, 32 and 30 kDa surface proteins and with LAMC1 via its carbohydrate recognition domains [13,22,23]. The 45 kDa mucin found on the surface of invasive *T. cruzi* trypomastigotes specifically interacts with LGALS3 and plays an important role in trypomastigote adhesion [12, 13]. These studies suggest that this is a trypanosome trapping mechanism which enables the organisms to accumulate in the basement membrane prior to invasion, making LGALS3 a candidate molecule which enhances the pathogenesis of *T. cruzi*.

LGALS3 is also expressed in B cells from *T. cruzi*-infected mice [24] and is up-regulated by *T. cruzi* infection of mice [25]. The concentrations of LGALS3 that increase...
trypanosome adhesion to HCASM cells in vitro are similar to the concentrations of LGALS3 present in vivo [26]. Furthermore, LGALS3 levels increase approximately 300-fold during microbial infection in vivo [27]. These observations suggest that the parasite modulates the host and takes advantage of a host inflammatory molecule, LGALS3, to bind to host cells. These findings may contribute to determining a cause of the pathology manifested in Chagas’ disease.

Fibronectin is an adhesion ECM glycoprotein which has been shown to be involved in target cell invasion by trypanastigotes via the RGDGS peptide (fibronectin cell attachment site) binding to the trypomastigote surface leading to inhibition of trypomastigote internalization [28]. The trypomastigote ligand for fibronectin was purified by affinity chromatography, and identified as an 85 kDa protein which interacts with cells bearing fibronectin molecules, such as human monocytes, neutrophils and 3T3 fibroblasts [28]. Fibronectin also mediates the uptake of T. cruzi by human monocytes via RGDGS [29]. The involvement of fibronectin in T. cruzi cardiomycocyte invasion has been observed during T. cruzi infection both in vivo and in vitro. Pre-treatment of trypomastigastes with fibronectin before cardiomycocyte interaction reduced T. cruzi infection significantly. Further, enhancement of ECM components was detected in the myocardium during late acute infection with T. cruzi [30]. However, molecular genetic approaches have not been used to validate the role of fibronectin in the process of infection.

REGULATION OF THE ECM INTERACTOME BY T. CRUZI AND ITS GP83 LIGAND

The genetic architecture in the early T. cruzi infection process of human cells is unknown. To understand this aspect of infection, we conducted gene transcription microarray analysis followed by gene network construction of the host cell response in primary HCASM cells infected with T. cruzi or exposed to T. cruzi gp83. Using THBS1, LAMC1, and LGALS3 as the seed nodes for biological network construction, we built an interactome network of the early T. cruzi infection process which centered on the ECM. After seeding the initial three nodes, the network was expanded to one degree of direct biological interaction, resulting in base interaction networks with the seed node as a center point in each. In order to populate and build our interaction network, the Michigan Molecular Interactions (MiMI) Cytoscape plugin (version 3.2) was used. The MiMI interface uses a database which is itself constructed from merged data of well-known protein interaction databases, including BIND, DIP, HPRD, RefSeq, SwissProt, IPI and CCSB-HI1 [31].

The MiMI query engine was used to seed the THBS1, LAMC1, and LGAL3 initial nodes and their respective nearest neighbors to one degree of biological interactions. The networks were then merged for interconnections and the global interactome was visualized in Cytoscape (version 2.6.3) [32], thus, yielding a snapshot of the ECM interactome involved in early T. cruzi infection. This ECM-focused interactome contains 104 nodes representing protein coding genes connected by 218 edges representing biological interactions between nodes (Fig. 1).

Increased THBS1 expression significantly modulates the interactome cross talk between cells. This change in network topology potentially favors parasite invasion and infection of host cells. As such, THBS1 interacts with several proteins ranging from adhesion receptors (CD36 and CD47) to structural proteins (COL7A1) and zymogens (PLG) as shown in Fig. (1). CD47 is a receptor for the C-terminal domain of THBS1 and this interaction may be important in membrane transport and signal transduction. In addition, CD47 is involved in the intracellular calcium increase which occurs when the cell adheres to the ECM. An increase in cytosolic Ca++ in T. cruzi trypomastigotes was detected at the single cell level after association of the parasites with myoblasts. Ca++ mobilization in the host cells was also detected upon contact with trypomastigotes. Confirmatively, pretreatment of the parasites with the Ca++ chelators decreased the trypomastigotes’ association to myoblasts indicating that calcium mobilization is required for cell invasion [33].

The structural protein COL7A1, or collagen 7 alpha 1, is typically found in the basement membrane and also associates with THBS1. Once COL7A1 interacts with THBS1, it makes the ECM conducive to parasite motility and cellular invasion with the help of proteins from the laminin family of genes (Fig. 1). Plasminogen (PLG) links LAMC1 and THBS1. PLG (a zymogen), is cleaved into plasmin (a serine protease), and angiotatin (an angiogenesis inhibitor). Plasmin is known to cleave fibronectin, THBS1, and LAMC1, thus taking part in ECM modifications.

T. cruzi must navigate through the basal lamina, which contains LAMC1. The T. cruzi gp83 ligand modifies LAMC1 expression in the ECM and contributes to the pathogenesis of T. cruzi infection in human heart cells. The fact that LAMC1’s network is connected to THBS1 through PLG but also COL7A1 and MMP2 suggests that this network could facilitate parasite mobilization. Matrix metallo proteinase 2 (MMP2) is a type IV collagenase that participates in the rearrangement of the ECM, which could facilitate parasite mobilization. In the subnetwork of LAMC1, there is an indirect connection (through LAMA1) to Dystroglycan (DAG1). DAG1 is a dystrophin-associated glycol-protein responsible for transmembrane linkage between the ECM and host cell cytoskeleton. The extracellular form of DAG1 can bind to merosin alpha-2-laminin in the ECM [34]. If T. cruzi can alter the dystroglycan complex, it could consequently manipulate or weaken the host cell cytoskeleton without first gaining entry into the cell. This could yet be another mechanism by which T. cruzi invades host cells. In reference to the ECM network reviewed, LAMC1 also has a second-degree interaction (one node in between the genes) with LGALS3 through myocin (MYOC). MYOC is a secreted protein believed to have a role in cytoskeletal function, specifically vesicular transport and extracellular matrix conformation [35]. Unlike other intracellular pathogens which avoid contact with host cell lysosomes, T. cruzi requires the low-pH environment of lysosomes to initiate egress from the vacuole and delivery to the host cell cytoplasm where replication takes place [36]. Therefore the control of vesicular trafficking by T. cruzi improves the rate of trypomastigote entry and amastigote replication in the host cell. The fact that LAMC1 is connected to LGALS3 through MYOC in the ECM interactome suggests the importance of LGALS3 in T. cruzi’s manipulation of host ECM.
Increased LGALS3 expression in the ECM promotes the adhesion of \textit{T. cruzi} to host cells and subsequent infection [14]. In addition, LGALS3 has numerous ECM interacting partners [37] including collagen IV, hensin, laminins, fibronectin, tenasin and elastin. LGALS3 regulates adhesion of these ECM proteins to a variety of host cells. Matrix metalloproteases, which are more active in \textit{T. cruzi} infected mice, regulate LGAL3 function [38]. When metalloproteases are activated in the ECM, they can cleave LGALS3 and negatively regulate its function. Consequently, increased activation of matrix metalloproteases 1 and 9 (MMP-1 and MMP-9) is associated with ECM destruction and myocarditis in \textit{T. cruzi} infection.

Multiple types of collagen interact with the 3 central seed nodes, THBS1, LGALS3 and LAMC1. A glimpse of the importance of collagen in early \textit{T. cruzi} infection was reported by Velge \textit{et al.} [39]. Their results suggest that a \textit{T. cruzi} surface glycoprotein, made up of an 80-85 kDa subunit and a 58/68 kDa subunit, binds host cell fibronectin and collagen. In addition, this interaction may occur during the initial phase of parasite-cell recognition [39].

Thus, the elucidation of human ECM interactome regulated by \textit{T. cruzi} is of paramount importance to the understanding of the molecular pathogenesis of \textit{T. cruzi} infection and developing novel strategies of intervention in Chagas’ disease.

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REFERENCES


