Trans-sialidase Stimulates Eat Me Response from Epithelial Cells

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Epithelial cell invasion by the protozoan parasite Trypanosoma cruzi is enhanced by the presence of an enzyme expressed on its cell surface during the trypomastigote life cycle stage. The enzyme, trans-sialidase (TS), is a member of one of the largest gene families expressed by the parasite and the role of its activity in mediating epithelial cell entry has not hitherto been understood. Here we show that the T. cruzi TS generates an eat me signal which is capable of enabling epithelial cell entry. We have utilized purified, recombinant, active (TcTS) and inactive (TcTS2V0) TS coated onto beads to challenge an epithelial cell line. We find that TS activity acts upon G protein coupled receptors present at the epithelial cell synapse with the coated bead, thereby enhancing cell entry. By so doing, we provide evidence that TS proteins bind glycans, mediate the formation of distinct synaptic domains and promote macropinocytic uptake of microparticles into a perinuclear compartment in a manner which may emulate entosis.

Key words: entosis, G-protein, host-parasite, lipid raft, trans-sialidase, Trypanosoma cruzi

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The protozoan parasite Trypanosoma cruzi is the aetiological agent of the tropical neglected Chagas disease, which is widespread in Latin America (1,2). In those infected, T. cruzi enters and proliferates amongst a variety of tissues; within epithelial and endothelial cells (3,4). Cell entry by T. cruzi is dramatically reduced by pertussis toxin (PT), which disrupts Gsi subunit signalling (5,6), and downstream intracellular events including reorganisation of the host cell’s cytoskeleton and recruitment of vesicles to the region of plasma membrane where the parasite attaches: the parasite synapse (7–11). While the exact identity and roles of G protein coupled receptor (GPCR) ligands in Chagas disease pathogenesis remains controversial, a role for GPCRs such as the bradykinin receptor B₂, cannabinoid receptor 1 and β₂ adrenergic receptor has been well established (12–14).

GPCRs are frequently resident in lipid rafts or microdomains which are concentrated at the parasite synapse during attachment and invasion of T. cruzi (11); their dispersal by cholesterol depletion with methyl β-cyclodextrin (MBCD) reduces entry of T. cruzi into host cells (15,16). Synaptic signalling engenders increased calcium ion concentration (17,18) resulting from cross-linkage of host cell receptors, and mediated via activation of cAMP (14,19) and phosphoinositol-3-kinase (PI3K) (20). This calcium ion flux enables mobilization of lysosomes (14) and early endosomes (20) (suggesting that T. cruzi may utilize both exocytic and endocytic pathways) to dock with the synapse providing for parasitophorous vacuole formation. Lysosome exocytosis directs acid sphingomyelinase to the parasite synapse promoting ceramide-rich microdomain formation and increased, localized endocytic activity (21,22). This endocytic component of cell entry was highlighted by cytochalasin D (cytD) inhibition of the early fusion of peripheral lysosomes with the plasma membrane at the parasite synapse (23) which required PI3K activation (20). Caveolin-dependent endocytosis also requires PI3K activation (24,25) and caveolin-1 (cav1), has been associated with T. cruzi entry of macrophages (16). Cav1 which is necessary for formation of caveolae is either sequestered at the cell surface forming invaginations (26) or recycled along microtubules under the control of multiple kinases which direct caveolin delivery (25).

Trans-sialidase (TS) is a T. cruzi trypomastigote surface enzyme which catalyses transfer of sialic acid from host to parasite surface. Seminal work established that a single amino acid, Tyr³⁴², is essential for activity (27) and that the enzyme can be entirely inactivated by point mutation at this site. Expression of TcTS itself, as well as other enzymatically inactive TS family members such as GP82, is strongly associated with virulence (28). While it is clear their roles as virulence determinants may arise from multiple functions, it is equally clear that they can act as important and specific mediators of infectivity, tropism and host cell invasion (29–31). Investigation into the role of TS during host cell entry by T. cruzi has described a reduction in entry upon inhibition of TS activity (32,33); however, reduced expression of some inactive members of the TS family has also been associated with reduced cell invasion (34,35). Moreover, TS activity has been implicated in roles including escape from the parasitophorous vacuole
TS promotes bead internalization by cells

To directly examine whether TS activity has a role in cell entry, or whether the observed ability to mediate cell entry arose from other structure-related activities of the TS protein, we compared the active recombinant enzyme with a nearly identical TS disabled by a point mutation in the active site. Four micron latex beads [selected as microparticles of similar volume to the T. cruzi trypomastigote – circa 30 fl (45)] were coated with purified active (TcTS) and inactive (TcTS2V0) recombinant enzyme, and their TS activity was then determined using a fluorometric assay (Figure S1). The number of beads internalized by MDCK II cells following 2 h of incubation was assayed by biotinylation of the cell surface and any attached beads. Unique, arbitrarily selected fields of cells were digitally captured and scored as internalized or attached. For each type of coated bead, two hundred cells were assayed in at least four separate experiments, and the number of beads which remained attached and internalized was compared for each protein (Figure 1). Time courses examining the kinetics with which beads coated in the active enzyme entered the cells (Figure S2) suggested that the proportion of beads taken up progressively increased over 8 h and a 2 h time point was selected for convenience. BSA and bradykinin coated beads and blank uncoated beads were used as controls.

Approximately, one third more beads coated in active enzyme (TcTS) were associated with MDCK II cells than beads coated in the inactive TcTS2V0. The proportion of beads internalized, however, was roughly similar and both were internalized in greater numbers than beads coated with bradykinin (Figure 1B), which appeared to be relatively poorly adherent but which was also able to mediate bead entry into cells. The proportion of internalized TcTS coated beads was also compared with that of BSA coated beads, however, no internalization of BSA coated beads was observed at all (Figure 1A and Table 1). Blank beads, preincubated with the culture media, were also included to control for any potential effects from serum proteins in the media that might be adsorbed onto the bead surface, however, these beads were entirely non-adherent to the cells under the conditions of these assays.

These results demonstrate that beads coated either in the active enzyme (TcTS), or the inactive TS (TcTS2V0), or the bradykinin peptide, in contrast to BSA, are able to direct uptake of microparticles by epithelial cells; but, that TS activity augments uptake relative to the inactive protein. Thus, while active and inactive TS proteins mediate proportionately similar internalization levels for the beads, the activity of the enzyme is a strong contributing factor to the sustained association between the cell and the coated bead, and hence facilitates considerably larger numbers of beads coated in active enzyme to enter epithelial cells.

TcTS activity promotes entry in a PI-3 kinase dependent manner

In corollary experiments, we investigated the role of TS activity in attachment and internalization using sialic acid acceptor substrate lactitol and pre-treating MDCK II cells with sialidase. To facilitate analysis for these and subsequent biotinylation assays, rather than capturing large numbers of arbitrary fields as described for Figure 1, 200 cells with closely associated beads were located, captured and scored as internalized or attached. Attachment of TcTS coated beads was marginally, but significantly, reduced at concentrations of lactitol greater than 0.25 mM in comparison with TcTS2V0 coated beads, where no such reduction was observed (Figure 2A). Internalization of TcTS but not TcTS2V0 coated beads was also significantly inhibited by application of (1 mM) lactitol (Figure 2B). Since lactitol is an efficient acceptor substrate of TcTS and competitive inhibitor for sialylation of acceptor substrates on the epithelial cell surface, these results suggest that attachment and internalization are facilitated by sialylation of cell surface molecules.

Interestingly, MDCK II cells treated with sialidase to remove host cell sialic acid prior to incubation with TcTS...
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Figure 1: TcTS coated beads attach to and are internalized more effectively by epithelial cells than TcTS2V0, bradykinin and BSA coated beads. Attachment and internalization was assessed by biotinylation of the cell surface and attached beads. Beads shown in red and indicated with arrows are attached beads, where the scale bar = 10µm (A). Protein coated beads incubated with MDCK II cells for 2 h were scored using the biotinylation assay as the number of attached and internalized beads per 200 cells (B). **p = 0.002 and *p = 0.0455 (mean ± SEM) when compared to the control (N ≥ 5).

and TcTS2V0 coated beads actually afforded an increase in TcTS coated bead attachment (Figure 2C), but conversely showed a decrease in internalization (Figure 2D) which was not observed with the inactive enzyme. A result suggesting that while on sialidase treated cells, TcTS can bind receptors which are not influenced by sialylation; the enhanced internalization associated with the active TS is not simply a function of enhanced binding to sialylated surface proteins, or simply explained in terms of the glycan binding preferences of these proteins which are not sialic acid specific (Figure S3). Rather that enhanced internalization arises directly as a result of the trans-sialylation of specific receptors.

The release of intracellular calcium associated with T. cruzi entry into epithelial cells is inositol triphosphate (IP3) dependent (18) and can be prevented by treatment with PT (6), which interacts with heterotrimeric G protein subunit, Gαi. We considered whether this would also be the case for TS coated microparticle uptake. Our results demonstrate that attachment of TcTS coated beads was reduced by PT treatment whereas attachment mediated by TcTS2V0 and bradykinin was not significantly affected (Figure 2E). Correspondingly, internalization of TcTS coated beads was also partially inhibited (Figure 2F), as were bradykinin coated bead; whereas, TcTS2V0 remained unaffected. Thus, blockade of Gαi signalling...
Figure 2: Inhibition of TS, Goi or PI3K activity reduces bead internalization. TcTS and TcTS2V0 bead attachment was calculated per cell and determined post treatment with lactitol *p = 0.0349, **p = 0.0256 (A) and uptake was scored using biotinylation after incubation with lactitol *p = 0.0072 (B). Attachment of TcTS and TcTS2V0 coated beads was calculated per cell after pretreatment with sialidase *p = 0.0295 and p = 0.0061 (C) and internalization determined by biotinylation of the cell surface *p = 0.0161 (D). Attachment of TcTS, TcTS2V0 and bradykinin coated beads was also scored after treatment with PT *p = 0.0024 (E) and internalization *p = 0.0175, **p = 0.0596 (F) and wortmannin (WM) attachment and internalization *p = 0.0628, **p = 0.0308 (H). Where all p values are compared to the control (mean ± SEM, N ≥ 3).
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Table 1: Epithelial cell attachment and internalization of coated beads

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<th>Attached (%)</th>
<th>Internalized (%)</th>
<th>SD (%)</th>
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<tr>
<td>TcTS</td>
<td>47</td>
<td>53</td>
<td>14</td>
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<tr>
<td>TcTS2V0</td>
<td>57</td>
<td>43</td>
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<td>BSA</td>
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<tr>
<td>Bradykinin</td>
<td>62</td>
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appears to specifically inhibit the TS activity mediated augmentation of bead attachment and entry rather than the process of attachment and entry overall.

Significant variation between experiments was observed in the ability of coated beads to bind cells, indicative of the sensitivity of the bead binding interaction to environmental variation. Within each independently controlled experiment though, variation was less, allowing meaningful comparison between control and treated groups. Cell cytotoxicity was also assayed to show that any inhibition of internalization was due to G protein subunit inhibition and not to cell death (Figure S4A,B).

The recruitment of vesicles and the reorganisation of the cytoskeleton require the activation of PI3K (46–48), which can also occur via Gai signalling (49). In previous studies PI3K was inhibited by wortmannin (WM) before cells were challenged with T. cruzi resulting in decreased cell entry and suggesting that cell invasion by T. cruzi requires activation of this enzyme (50). With this in mind we treated cells with WM and assessed the effect on TcTS, TcTS2V0 and bradykinin bead attachment and uptake (Figure 2G,H). Treatment did not result in a significant reduction in attachment, but did lead to a reduction in TcTS and bradykinin (but not TcTS2V0) coated beads entering the cells. Taken together these results suggest that TcTS activity stimulates PI3K signalling during cell entry via Gai transactivation.

The internalized bead vacuole reaches a perinuclear position

We constructed an MDCK II cell line expressing both GFP-α-tubulin and cav1-RFP in order to consider the interaction between bead, microtubules and caveolin over time (Figure 3A). We observed a protracted association between cav1-RFP and TcTS coated beads; from initial recruitment to the bead synapase, through to vacuole formation and to a final destination in its perinuclear compartment. We found no increase in this association over time, however, where cav1 was initially recruited it appeared to remain in close proximity with the beads. It was apparent from this work, that at later time points beads were transported to a perinuclear position abutting the MTOC (Figure 3A). Investigation of the perinuclear compartment was undertaken with transmission electron microscopy of MDCK II cells incubated for 24h with TS coated beads (Figure 3B). These figures clearly show the presence of the centriole juxtaposed to the bead vacuole as well as highlighting a close association between the bead vacuole and the endoplasmic reticulum (ER).

The apparent perinuclear positioning of the vacuoles containing TS coated beads seemed analogous to previous work demonstrating repositioning of the host cell MTOC in close proximity to the T. cruzi vacuole (10). To confirm this repositioning effect, we measured the distance between MTOC and beads in fixed cells over a 9-h time course. We found a considerable decrease in the distance between the beads and the MTOC over time consistent with bead uptake and eventual positioning in close proximity to the MTOC (Figure S6). Microtubule polymerisation is integral to repositioning of the MTOC and has also been implicated in the invasion of T. cruzi where tubulin appears to accumulate in the region of the attached parasite (10). To investigate whether microtubule dynamics were also associated with bead entry, cells were treated with colchicine to depolymerize the microtubules and bead entry assessed by surface biotinylation. A modest reduction in bead uptake was observed in the 5 and 10 μM range for TcTS coated beads but not TcTS2V0, while bradykinin coated beads show a decrease in internalization for all colchicine concentrations (Figure 3C). This result suggests a microtubule-dependent component mechanism in the uptake of bradykinin and active TcTS coated beads which is absent for uptake of beads coated in the inactive enzyme. Visual confirmation of microtubule depolymerisation of MDCK II cells at these concentrations was performed by colchicine treatment of GFP-tubulin expressing MDCK II cells (Figure 3D).

Polymerisation of actin is required for effective internalization of TcTS and TcTS2V0 coated beads

Host cell invasion is often associated with cytoskeletal remodelling and T. cruzi has been proposed to invade epithelia by both tubulin and actin dependent mechanisms. Previous studies have described T. cruzi mediated depolymerisation of the cortical actin cytoskeleton in some cells (9) and induction of actin-rich pseudopodia (51,52). In this study, TcTS, TcTS2V0 and bradykinin coated beads were incubated with MDCK II cells before immunolabelling for β-actin. The resultant images show ruffling; with cup-like actin-rich projections closely associated with the beads, and in some cases surrounding them entirely (Figure 4A). Interestingly, this activity appeared to be independent of TS activity, as it was also observed when beads were coated with TcTS2V0 (Figure 4B). We inhibited microfilament polymerization with cytD prior to the addition of TcTS, TcTS2V0 and bradykinin coated beads. In each case this treatment almost entirely inhibited bead internalization even at low concentrations (Figure 4C), confirming a role for microfilament dynamics in microparticle uptake which did not relate to TS activity per se. The cytD treated cells were stained for β-actin after treatment to evaluate the extent of depolymerisation at each concentration (Figure 4D).
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Figure 3: Microtubule dynamics in perinuclear positioning of internalized microparticles. Beads coated with TcTS (DIC) were incubated with MDCK II cells stably expressing GFP-tubulin (green) and cav1-RFP (red) for 6 h. The arrows indicate the position of the bead, scale bar = 5 µm (A). MDCK II cells were incubated overnight with TcTS coated beads before preparation for transmission electron microscopy (B). The 4 µm white beads are shown here in close proximity to mitochondria (m), golgi (G) and ER, scale bar = 1 µm (i). The centrosome (arrow) is also visible in very close proximity to the bead, scale bar = 0.5 µm (ii). Close to the centriole the golgi is clearly visible, scale bar = 0.5 µm (iii). B-bead, PM-phagosomal membrane and N-nucleus. MDCK II cells were incubated with 0, 2, 20 µM of colchicine to depolymerize microtubules before incubating with TcTS, TcTS2V0 and bradykinin coated beads for 3 h. The number of internalized TcTS coated beads was determined by biotinylation (mean ± SEM, N = 3, **p = 0.008) (C). MDCK II cells expressing GFP-tubulin were imaged after application of the indicated concentrations of colchicine (D).

Cell cytotoxicity was ruled out as a cause for the decrease in internalization by LIVE/DEAD® assay where cell death remained at a basal rate despite addition of the drug (Figure 4E).

Cav1 is recruited to internalized beads and parasites but is not essential for internalization

A significant proportion of cell invasion by *T. cruzi* into epithelial cells is dependent on formation of a host cell plasma membrane-derived vacuole (20). To determine if the observed TcTS-mediated internalization of coated beads was similarly associated with plasma membrane markers we looked for evidence of association with a canonical marker from one of the major endocytic pathways – cav1. Immunofluorescence at short timepoints showed that many of the attached beads coated with both TcTS and TcTS2V0 were associated with increased concentration of cav1 (Figure 5A).
Previously, an association with cav1 has been described with *T. cruzi* invasion of macrophages (16) but its role in epithelial cell invasion has not been explored. We, therefore, considered whether live parasites attached to epithelial cells recruited caveolin to the parasite synapse. We found that attachment of trypomastigotes often resulted in a marked increase in cav1 at the parasite:cell synapse (Figure 5B), implying that attachment might promote cav1 translocation. We found this association irrespective of the parasite strain or epithelial cell line we used (not shown). We also noted that at a longer (6 h) time point, as many as half of the vacuoles formed remained associated with caveolin regardless of whether live or killed parasites, or beads were used (Figure 5C).

In light of its association with TcTS coated beads we considered whether cav1 might play a functional role in TcTS-mediated epithelial cell entry. We silenced cav1 expression in MDCK II cells, reducing expression levels by more than 90% (Figure 5D,E). The proportion of TcTS coated beads internalized by silenced cells showed no significant difference to unsilenced controls (Figure 5F). A result which suggests that although caveolin is a plasma membrane marker which is enriched at the bead synapse, it is present primarily as a bystander and that caveolin-dependent endocytosis is not a mechanism which promotes entry of TcTS coated beads into epithelial cells *per se*.

**Figure 4: Bead internalization is actin dependent.** MDCK II cells were incubated with TcTS (A) and TcTS2V0 (B) coated beads. The position of the beads is shown with DIC and arrows (A) and blue fluorescence (B). Immunolabelling with anti-β-actin (red) shows ruffling and increased concentration associated both kinds of coated beads after labelling with anti-mouse Alexa Fluor568. Scale bar = 5 µm. Biotinylation assay for bead internalization demonstrated that treatment with cytD completely prevents bead internalization after 3 h (C). Depolymerisation of microfilaments was confirmed by immunolabelling with anti-β-actin after treatment with 0, 5, 50 µM of cytD, scale bar = 5 µm (D). The level of cell cytotoxicity after addition of cytD was assayed by LIVE/DEAD<sup>®</sup> assay (E). Graphs show mean ± SEM (N = 3) where **p = 0.0007 and *p = 0.049 when compared with the untreated control.

**TcTS coated bead attachment is inhibited by cholesterol depletion**

Cholesterol is an essential component of the lipid containing assemblies of the plasma membrane (53) and specific extraction of cholesterol from the plasma membrane by MBCD results in the dissipation of these assemblies. Previously, depletion of cholesterol has been shown to inhibit the ability of *T. cruzi* to invade cells (15), suggesting the integrity of these domains is important for cell entry by *T. cruzi*. To investigate the effect of removing cholesterol on attachment of TcTS and TcTS2V0 coated beads we considered whether the number of attached beads was reduced by treatment of the cells with MBCD. We found that attachment of both TcTS and TcTS2V0 coated beads to MDCK II cells was modestly, but significantly impaired by treatment with MBCD (Figure 6A), suggesting epithelial adhesion cell is favoured by concerted interaction with surface moieties tethered to cholesterol-dependent assemblies in the plasma membrane. Interestingly, the removal of serum from the media enhanced binding by beads coated in the active TcTS (perhaps by promoting contact with donor and acceptor substrates on the cell surface), but actually afforded reduced binding by beads coated in the inactive TcTS2V0.

**TcTS activity-independent entry is serum sensitive**

In testing the cholesterol dependence of internalization of both TcTS and TcTS2V0 coated beads, we aimed to...
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Figure 5: Caveolin-1 is recruited to bead and parasitophorous vacuoles in epithelial cells but reduced cav1 expression does not affect bead internalization. For MDCK II cells incubated with TcTS and TcTS2V0 coated beads and immunolabelled with anti-cav1 antibody, cav1 accumulates in close proximity to attached and internalized beads. Scale bar = 2 µm (A). Live parasites used to infect OK cells show considerable concentrations of cav1 at the parasite synapse during attachment (B). The scale bar = 5 µm. Most vacuoles of TcTS coated beads, live and dead parasites associate with cav1, but the association is strongest for live parasites (C). MDCK II cells were transfected with cav1 siRNA (Cav-) and non-targeting siRNA(NT) and separated by SDS-PAGE before transferring by western blot. The resultant membrane was labelled with anti-cav1 (green) and anti-β-actin (red) antibodies, untransfected cells (Un) were also run alongside and the membrane was imaged using an Odyssey® infrared imager (D). The transfection rate was calculated by immunolabelling cells with cav1 antibody and transfecting with glowing RNA (E), scale bar = 10 µm. Untransfected and transfected cells were biotinylated and imaged by confocal microscopy after incubation with TcTS coated beads for 2 h. (graphs show mean ± SEM, N ≥ 3) (F).

assess the uptake of the beads from MBCD treated cells compared with untreated cells. The use of MBCD requires that cells are assayed in the absence of sera supplements and we compared bead uptake in cells treated with 10, 20 and 30 mM of MBCD for an hour with that in cells simply exposed to sera free media for the same period and cells in normal media containing Foetal Calf Serum.

We observed a dramatic decrease in internalization of TcTS2V0 beads even without MBCD treatment in the absence of sera (>95% reduction). The absence of sera also reduced the internalization of TcTS beads but this effect was considerably less profound, with significant numbers of TcTS beads still internalized. TcTS and TcTS2V0 coated bead internalization was further inhibited by increases in MBCD concentration until internalization was completely blocked at 30 mM (Figure 6B). A live/dead cell assay showed that MBCD had cytotoxic effects only at 30 mM and above (Figure S6). These results confirm that internalization of TcTS and TcTS2V0 coated beads is cholesterol dependent, relying on the cholesterol containing assemblies of the plasma membrane. Strikingly

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though, the differential susceptibility to serum depletion we observed further uncurl a mechanism by which TS activity augments uptake of the beads coated in active enzyme, from the serum sensitive macroinocytotic mechanism by which TS proteins can direct cell entry in an activity independent manner.

The bead-cell synapse membrane shows increased lipid order
Actin dependent uptake from the plasma membrane and the presence of caveolin at the bead’s interface with the cell imply localized structuring of the plasma membrane. Lipid rafts can be defined as regions of the cell membrane which exhibit increased lipid order, being tightly packed and enriched with sphingolipids and cholesterol. Lipid rafts can often be detected at the synapses between cells (54) or between cells and pathogens (55) as cross-linkage of GPI linked glycoproteins or glycosphingolipid receptors is believed to be sufficient to serve as a driver for a localized increase in membrane order. Visualization of these regions can be achieved by use of the dichromatic dye Laurdan and localized quantification of membrane order in the cell using Laurdan on a widefield microscope can be achieved by computation of the generalized polarisation (GP) value (56). The GP is the normalized intensity ratio between the two emission channels where the maximum +1 is the most condensed and the minimum −1 is the most fluid. When these values are mapped onto micrographs, cell membrane which appears ‘warmly coloured’ represents regions of high lipid order, while regions that appear in the blue end of the spectrum are less well ordered. We used this dye to examine whether the regions of plasma membrane where TcTS and TcTS2V0 coated beads had attached showed evidence of increased lipid order indicative of the ability of such proteins to cross-link glycolipids and glycoproteins.

Images were collected separately at emissions of 435, 460 and 500 nm using a dichroic mirror block to obtain triple split images, the 435 and 460 nm images were combined as previously described (56,57) to produce a final GP image (Figure 7A). TcTS coated beads were found to lie in contact with regions of the cell surface membrane which appeared more ordered than the rest of the cell (yellow, orange and red pixels). Such regions were present throughout the cell surface but appeared to accumulate proximal to the beads. The same phenomenon was observed with the TcTS2V0 coated beads which also appeared to attach via more warmly coloured regions.

Using SimFCS software, the GP was calculated for each pixel in a 30 × 30 pixel square region of interest (ROI) encompassing the bead synapse and for an equivalent ROI from the same cell in close proximity (Figure 7A). Histograms were produced showing the spread of GP values, control ROIs exhibited more negative average GP value (−0.099) than the ROI in contact with a TcTS coated bead (+0.003) (Figure 7B). The same patterns are also shown when TcTS2V0 coated beads are attached to the cell membrane (Figure 7C) with a visible shift in average GP from −0.013 to +0.136 in contact with the bead. Comparison of multiple ROIs of membranes in contact with the protein coated beads show that the region of membrane in contact with both TcTS and TcTS2V0 coated beads is more ordered than the regions of membrane not in contact with either protein coated bead (Figure 7D). Importantly however, no discernible difference was observed in the increased GP values from active and inactive enzyme, suggesting that the ability to cross-link host surface proteins and induce raft formation at the bead synapse are similar. Lending support to this observation, we found only relatively minor differences between the glycan binding profiles of the active and inactive enzyme (Figure S3). Those significant differences that were observed were not all or nothing, but rather showed differential avidity for fucose, mannos, lacto-ß-biose I and sialic acid. Interestingly though, some of these glycans are associated with mucins which are GPI-linked. Our results are consistent with lipid raft formation at the bead synapse and suggest that attachment of both TcTS and TcTS2V0 coated beads (through cross-linking of glycosylated

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**Figure 6: Cholesterol depletion prevents internalization by both TcTS and TcTS2V0 coated beads.** Internalization of coated beads was assessed using the biotinylation assay. The level of attachment is shown with FCS and before and after treatment with MBCD for both TcTS and TcTS2V0 coated beads (A) (mean ± SEM) where *p < 0.05. The effect on protein coated bead internalization by 10, 20 and 30 mM of MBCD was also assayed and compared with untreated cells which had and had not been deprived of serum (B) (mean ± SEM, *p < 0.05).
moieties on host glycoproteins and glycolipids) lead to structuring of an epithelial cell synaptic membrane characterized by regions of high lipid order which do not form when cholesterol is depleted (Figure S7A,B).

**Discussion**

Penetrating and replicating within mammalian epithelia and other non-phagocytic cell types is integral to the life-cycle of *T. cruzi*. As part of the arsenal evolved by the parasite to facilitate infection, trypomastigotes express several members of the *T. cruzi* TS family which have previously implicated in host cell invasion, including some, such as gp30 and gp82, which have been shown to induce intracellular Ca$^{2+}$ ion release (58,59). Here, we have demonstrated that while TS activity may not be necessary for cell entry, it does possess the ability to enhance entry into epithelial cells. This is in accordance with data previously presented demonstrating the reduction in *T. cruzi* internalisation after treatment with lactitol and after host cells were treated with sialidase to remove sialic...
acid (32,33). Furthermore, here, we show that purified recombinant TS alone can potentiate similar responses from the host cell as the whole parasite; and moreover, that the presence of a homogenous coating of inactive recombinant TS protein is sufficient to mediate efficient epithelial cell entry.

We observe a reduction of the distance between TS coated beads and the MTOC with time which is accompanied by an enrichment of tubulins around the bead vacuole. This correlates closely with the behaviour observed during formation of the parasitophorous vacuole of *T. cruzi*, where microtubule rearrangement and nucleation of microtubules from the vacuole was observed and in which GFP-tubulin expressing MDCK II cells also implicated the host centrosome in the internalization of *T. cruzi* by suggesting that it migrates towards the nascent parasitophorous vacuole (10). The final proximity of the bead vacuole to centrosome may imply that as with *T. cruzi*, the internalized bead is able to affect host cell microtubule dynamics and influence positioning of the MTOC. Alternatively, bead vacuoles may be actively transported by motors such as kinesins to a perinuclear location in a manner analogous to that observed for several other pathogens (60). *Toxoplasma gondii* infected cells also exhibit translocation of the membrane and assessed using generalized polarity a higher lipid order when compared to the surrounding with the host cell plasma membrane which exhibited (66) and has been linked to autophagosome formation (67). Microtubules (10). Similarly, in the absence of the host gamma and tyrosinated tubulins which polymerize into vacuole is able to act as an MTOC rapidly recruiting positioning of the MTOC. Alternatively, bead vacuoles may also act as an MTOC, enabling the formation of parasite synapse and membrane of the *T. cruzi* parasite and membrane of the *T. cruzi* parasite vacuole is able to act as an MTOC rapidly recruiting gamma and tyrosinated tubulins which polymerize into microtubules (10). Similarly, in the absence of the host centrosome, the parasitophorous vacuole of *T. gondii* can also act as an MTOC, enabling the formation of microtubules around the parasitophorous vacuole (62).

The vacuole formed by TcTS coated beads appears qualitatively different to that associated with apicomplexan parasites such as *T. gondii*. *Toxoplasma gondii* has recently, however, been shown to ligate sialic acid via microneme adhesive repeat (MAR) domains of some of their micronemal repeats like MIC1, which may be analogous to ligation of sialic acid by the TcTS (63,64). For *T. gondii* a sorting junction excludes certain components of the host plasma membrane during vacuole formation and in particular cav1 is specifically excluded from the parasitophorous vacuole. Nevertheless, the vacuoles of *T. gondii* and the TcTS coated beads are found in perinuclear positions in proximity to the centriole, mitochondria and golgi (62,65). An even more striking observation is the close association with ER which is reminiscent of the *T. cruzi* and *Leishmania* sp. vacuoles (66) and has been linked to autophagosome formation (67).

Both TcTS and TcTS2V0 coated beads formed synapses with the host cell plasma membrane which exhibited a higher lipid order when compared to the surrounding membrane and assessed using generalized polarity mapping for the lipophilic fluorophore, Laurdan. This implicates lipid rafts in the initial attachment of TS, suggesting that the receptors for TS are either located in microdomains or that they translocate there upon cross-linkage by the enzyme. The role of lipid rafts in epithelial cell invasion for other intracellular pathogens such as *Shigella flexneri* and *Salmonella typhimurium* is well established and requires the type II secretion system for cell entry where their primary entry molecules IpA/B/SipB (respectively) bind directly to cholesterol prior to engulfment (68). *Salmonella* adhesion to gut epithelia is mediated via the glycan fucose (69) and it is therefore interesting that we find that *T. cruzi* TS also binds strongly to fucose moieties irrespective of activity (Figure S3).

Lipid rafts are also termed signalling platforms due to the clustering of receptors and are essential for a range of pathogens. Previous support for lipid raft formation at the *T. cruzi* synapse comes from studies showing an accumulation of raft markers such as GM1 ganglioside, LDLr, AKT-PH-GFP and the cannabinoid receptor associated with invading *T. cruzi* (13,15,20,70). In this study, we observe considerable recruitment of the canonical lipid raft marker cav1 by both trypomastigotes and coated beads during internalization. We utilized RNA silencing of cav1 to investigate whether the observed accumulation at the bead synapse was integral to bead internalization. Although we were able to silence cav1 effectively in our epithelial cell cultures, we found no evidence that reducing cav1 expression led to inhibition of bead entry. It seems most likely therefore, that during TcTS-mediated cell entry, caveolin is recruited as a bystander to the micro-environment formed when TcTS binds to host cell receptors, but that cav1 plays little if any role in the uptake of TcTS coated beads per se. This result echoes previous experiments where trypomastigote invasion into fibroblasts from cav1 knockout mice appeared largely unaffected by the loss of the gene (71).

Lipid rafts and actin dynamics are closely associated, particularly in regard to endocytic processes (72) and actin has been identified as an important component of the *T. cruzi* cell invasion process. Actin dynamics are already known to play a critical role in parasite entry, early lysosomal fusions and parasite retention (23); moreover actin-rich structures and actin-like projections associated with trypomastigote entry have previously been described (51,73). Our results indicate that actin is vital for the internalization of beads and that polymerisation of actin in the region of the bead is observed in association with formation of pseudopod-like structures which may be a common requirement for uptake of microparticles. Membrane ruffling has been observed during the internalization of polystyrene beads coated in transfection reagents and in this case, upon entry, the beads associate with acidic endosomes and then with the LC3 autophagy marker (74), a marker which has also been previously associated with the *T. cruzi* vacuole (67) and entoptic vacuoles (40).
The evidence presented here regarding the mechanism of cell entry by coated beads comprises lipid raft formation, cholesterol and actin dependency for coated bead uptake, lack of association with clathrin, failure of cav1 silencing or microtubule depolymerisation to prevent internalization, serum sensitivity and finally, observation of actin pseudopodia and microfilament accumulation around the beads during uptake. Even in the absence of TS activity, TSs are able to mediate cell entry and our results indicate that this ability is acutely dependent on the presence of serum in the medium. We have not resolved the reasons for this dependency, however, it is consistent with the observed reliance of macropinocytosis on the presence of exogenous growth factors which are supplied in the serum (75). Taken together our evidence suggests that macropinocytosis is the mechanism of entry for TS coated microparticles and consistent with this it is also a mechanism that has also recently been reported for *T. cruzi* entry (76).

We have shown in this manuscript that immobilization of TS activity on microparticle surfaces serves to direct the activity to a synaptic region of the host cell plasma membrane where the activity results in activation of G proteins facilitating cell entry of the coated microparticles. Within the synapse formed, TS activity promotes redistribution of sialic acid by desialylation of donors and trans-sialylation of acceptors. Consistent with this, where activity is inhibited, or the cell surface is first sialidase treated, the effect of TS activity on bead uptake is reduced. Previous work has suggested roles for GPCRs; bradykinin-2 receptor, beta-adrenergic receptor-1 and cannabinoid receptor-1 (12–14). These proteins are heavily glycosylated and potentially may be activated as a result of sialic acid transfer resulting in the activation of G protein-dependent signalling events. Indeed the evidence which links the bradykinin-2 receptor to *T. cruzi* cell entry (12) coupled with recent work which shows that sialylation of this receptor stabilizes its retention at the cell surface and facilitates its dimerization thereby potentiating Gαi subunits of heterotrimeric G proteins by preventing the interaction between the G proteins and the GPCRs on the cell surface, negatively controlling the activation of cyclic AMP (79), and eliciting downstream effects on vesicle recruitment to the cell surface. This in turn may influence the composition of the synaptic membrane, potentially increasing microdomain formation through the action of the acid sphingomyelinase (21).

Several other *T. cruzi* molecules have been implicated in the production of an intracellular rise in calcium ions, which drives parasite invasion. Oligopeptidase B has the capacity to release an extracellular molecule that can induce an IP3 dependent mechanism of intracellular calcium release (19). The intracellular rise in calcium is detected by synaptotagmin VII which in turn regulates vesicle docking with the plasma membrane which aids parasite invasion (80). A distinct internalization pathway has also been described which is readily inhibited by the PI3-kinase inhibitor WM (20), and we have found that enhanced cell entry mediated by TS-activity may also be influenced by the activity of the PI3-kinase. The signalling pathways utilized by different *T. cruzi* strains can vary (81) and *T. cruzi* TSs are highly polymorphic in sequence and expression levels, so it is worth considering that those strains with higher TS activities will be more able to make better use of this pertussis-sensitive mechanism for enhancing cell entry and may be better at invading epithelial cells than strains with trypomastigotes that possess lower TS activities.

Epithelia represent natural barriers to infection and the cells which compose them are usually considered to refrain from phagocytosis of larger bodies such as parasites, bacteria or latex beads. They are though acknowledged to be able to take up unanchored neo- plastic cells by a process known as entosis. Moreover, up-regulation of endogenous surface sialidases such as Neu 3 is firmly associated with neoplasia (39). Here, we show that epithelia can also internalize microparticles coated with TS (and bradykinin) in a similar fashion. This process is enhanced (through the action of the TS) by the localized removal of sialic acid to as yet unknown acceptor substrates. This action appears to engender a pertussis and WVM sensitive *eat me* signal that results in enhanced macropinocytic uptake. Thus like entosis, this mechanism appears macropinocytic and enhanced by PI3 kinase activation (40). Whether this activation is sensitive to the removal of sialic acid has not been investigated here, however sialic acid-binding lectins have been shown to inhibit PI3K autophagy pathways (82) and autophagy plays a key role in the efficient invasion of cells by *T. cruzi* into parasitophorous vacuoles containing LC3 (67).

In summary, we observe here that TS mediated macropinocytic internalization of coated microparticles is modulated by G protein activation, suggesting that sialic acid transfer activates the GPCRs which regulate the process. We further postulate that TS mediated uptake by epithelia may have arisen by emulation of entotic *eat me* signals generated by endogenous sialidases and evolved as part of a surveillance system for epithelial neoplasms.

### Materials and methods

#### Drugs and reagents

Red, blue and blank sulphate-modified latex beads (4 µm) were purchased from Molecular Probes® (Invitrogen). The Sulfo-NHS-LC-biotin and BCA protein assay kit were purchased from Pierce. The inhibitors and cholesterol chelators; MBCD, WM, sialidase (*Vibrio cholerae*), lactitol, cytD, co and PT...
were purchased from Sigma-Aldrich. Antibiotics G418, hygromycin B and amphotericin B were purchased from Invitrogen.

**Antibodies**
The anti-Trypanosome Alpha Tubulin (TAT) antibody was a kind gift from Prof. K. Gull (Oxford University, UK) and rabbit polyclonal to cav1, caveolae marker and rabbit polyclonal to beta actin were obtained from Abcam. Monoclonal anti-polystyrene peroxidase conjugate clone His1 and monoclonal anti-iH-actin clone AC-15 (western blot experiments) were purchased from Sigma-Aldrich. IRDye 880RD goat anti-mouse IgG (H+L) and IRDye 800CW goat anti-rabbit IgG (H+L) were obtained from LI-COR Biosciences. Streptavidin conjugated Alexa Fluor 586 was purchased from Invitrogen while streptavidin conjugated Cy3 was purchased from Caltag Medsystems. Green-fluorescent Alexa Fluor 488 and red-fluorescent Alexa Fluor 568 conjugated anti-mouse IgG (H+L) and red-fluorescent Alexa Fluor 656 conjugated anti-rabbit IgG (H+L) were all purchased from Invitrogen.

**Recombinant proteins and activity analysis**
The expression plasmids containing polystyrene tagged TcTS and TcTS2V0 were kindly supplied by S. Schenkman and A.C.C. Frasch, respectively. The TcTS was a 70 kDa compound truncated at the C terminus to remove the SAPA repeat unit while maintaining the TS activity (B3) while TcTS2V0 has a TyrM2-His substitution (27). The DHBS E. coli strain was used to express the proteins which were induced using 1 µM IPTG and grown shaking for 3 h at 27°C. Cells were lysed in 1 x binding and wash buffer made as per the manufacturer’s instruction (Invitrogen) with lysozyme (1 mg/mL) and then sonicated. Purification was achieved by incubating Escherichia coli lysates with Dynabeads® His-tag (Invitrogen) as per the manufacturer’s instructions; eluates were kept at 4°C until use. Purification samples were separated using SDS–PAGE and the protein bands visualized using SYPRO® gel stain (Invitrogen). Latex beads were coated with TcTS and TcTS2V0 at concentrations determined by BCA assay or with bradykinin and BSA as determined by the manufacturer (Sigma). To coat the beads a 1% bead solution was made in PBS and resuspended and allowed to coat the beads overnight under agitation at 4°C. The beads were then centrifuged and washed in PBS twice to remove surplus protein and resuspended in PBS or sample loading buffer (SLB). Immunoblotting was performed on 10 µL of coated beads and an anti-His antibody was used to confirm identity of the purified protein.

TS activity was assayed as previously described by Schrader and colleagues (84) and adapting the protocol to a 12 well plate. Samples were screened using a BMG-labtech fluorostar plate reader at 350 nm (ex) and 460 nm (em).

**Cell culture and labelling**
MDCK II cells were maintained in DMEM supplemented with 10% Foetal Calf Serum, 5 mM L-glutamine, 50 µg/mL streptomycin and 50 units/mL penicillin. OK (Opossum Kidney) cells (HPACC) were maintained in IMEM and supplemented as described above. Cells expressing GFP-tubulin were a kind gift from Dr. Mette Mogensen (University of East Anglia, UK) and were supplemented with G418 (100 µg/mL).

Prior to experimental procedure, cells were seeded into 12 well plates containing 13 mm diameter coverslips to a density of 8 × 10^5 cells per mL and grown for approximately 20 h at 37°C. Cells were fixed with 3% PFA and permeablized with 1% NP40. The actin cytoskeleton was visualized using anti-α-actin antibody and cav1 vesicles with anti-cav1 antibody after blocking with 10% neutral goat serum.

**Attachment and internalization assays**
MDCK II cells were seeded as described above. 1.4 × 10^4 protein coated beads were then added to the cells and centrifuged for 1 min at 300 × g. The plate was returned to a 37°C incubator for 2 h before the media was removed and the cells were rinsed in PBS and biotinylated as previously described (85). For the inhibitor studies, cells were treated as previously described and incubated with beads for the following times: PT, 2 h (86); sialidase, 1 h (83); Cy3 and Co, 3 h (87). 20 nm of VM was incubated with cells for 30 min prior to addition of beads for 2 h, cells were incubated with MBCD and beads for 1 h and lactitol was added to the media with the beads at concentrations of 0.25, 0.5 and 1 µM and incubated with the cells for 3 h.

Biotinylation was assayed by confocal microscopy and the percentage internalization was assayed as the ratio of the number of beads in cells which were unbiotinylated (internalized) to biotinylated beads clearly associated with cells surface (attached) where the ratio was expressed as a percentage of total cell-associated (bound and internalized) beads. Attachment and internalization values were derived from scores of 200 cells with attached and/or internalized beads from digitally captured fields. Fields were captured either arbitrarily to avoid normalizing bias in comparisons of absolute numbers of beads attached or internalized (Figure 1) or were shot selected for association with beads in order to facilitate larger numbers of beads to be assessed in between group comparisons of the proportion of beads attached/internalized (other figures).

**MTOC/cav1 recruitment assay**
For the caveolin recruitment studies MDCK II cells already expressing GFP-tubulin were transfected with plasmid cav1-RFP using the lipid based transfection reagent Fugene6 (Roche Applied Sciences). A PcDNA3.1 cav1-RFP construct (a gift from Prof. T. Willemans) (88) were selected for by using a combination of G418 (1 mg/mL) and hygromycin B (200 µg/mL) and subjected to transfection with restriction enzyme fsp.

Cells expressing both fluorescent constructs were selected for by using anti-TAT antibody to label tubulin; cells were then DAPI stained and mounted ready for imaging.

**Invasion assays**
Tissue culture trypomastigotes (TCTs) from T. cruzi Silvio and 92101601P strains were obtained from the supernatant of infected Vero cell monolayers by centrifugation. Purified TCTs were killed by overnight treatment with amphotericin B (1 µg/mL).

MDCK II cells were seeded at a density of 1 × 10^5 and grown overnight and washed with DMEM before addition of TCTs resuspended in DMEM at an MOI of 50. Trypomastigotes were incubated with MDCK II cells for 1 h before rinsing uninvaded parasites with PBS and fixated with 4% paraformaldehyde for 10 min.

Dead TCTs were centrifuged onto MDCK II cells at 300 × g for 1 min and then incubated together for 6 h at 37°C prior to fixation (as above).

**Cav1 siRNA**
Cav1 (Genbank Acc. No: NM_001003296) was silenced using custom siRNA sequences: GCCGGAGAGATGAGCAGGA and GAATATCAGATAAACCCCG (Dharmacon). Non-targeting siRNA from the on-target plus control pool (Dharmacon) was used as a control. Cells were grown to 40% confluency before transfection using JetPrime transfection reagents (Polyplus). After 24 h transfection media was replaced with either fresh
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DMEM or DMEM plus TcTS coated beads. MDCK II lysates were prepared using RIPA buffer and subjected to western blot analysis using an Odyssey® infrared imager (LI-COR® Biosciences). Denaturation was done using Odyssey software by calculating the relative intensity of the loading control. TcTS coated beads were then incubated with transfected and untransfected cells for 2 h before biotinylation.

Fluorescence microscopy

The confocal microscopy was performed using a Zeiss LSM510 META, approximately 50 beads were assayed and each experiment was done in triplicate unless otherwise stated. Visualization of cavin1, β-actin and tubulin was achieved using a Zeiss Axioplan 2 microscope and a Zeiss AxioCamHR camera. All image analysis was done using Axiovision 4.7 software.

Laurdan imaging

Membrane order was evaluated using the fluorescent probe 6-dodecanoyl-2-dimethylaminonapthalene (Laurdan) (Roche). Laurdan was solubilized in dimethylsulfoxide (DMSO) to a stock concentration of 1 mM and kept at room temperature in the dark. MDCK II cells were grown overnight in chamber coverglass systems (Nunc), and protein coated beads were added and centrifuged for 1 min at 300 × g to promote bead-cell contact and transferred to 4°C for 15 min. Unattached beads were then removed by rinsing with ice cold PBS and Laurdan added at 10 μM. Cells were then incubated in the dark with gentle rocking for 15 min, to allow incorporation of the probe. Cells were then rinsed again with PBS and imaged immediately.

For the cholesterol depletion experiment cells were incubated with 10 μM of MBCD prior to addition of beads. GP average of the cell membrane in each GP image was analysed using SimFCS software.

Transmission electron microscopy

Cells were grown on 10 cm plates until sub-confluent, whereupon protein coated beads were added and incubated with the cells overnight at 37°C. Cells then were washed in PBS and fixed with glutaraldehyde (2.5%). These cells were collected by scraping, and were pelleted at 86 × g. The pellets were then stored in 2.5% glutaraldehyde until processing for electron microscopy. At that point residual phosphate was removed by rinsing with 0.1 M cacodylate buffer. The pellets were then postfixed for 1 h at room temperature with 1% osmium tetroxide, 0.8% potassium ferricyanide and 5 μM calcium chloride before being further rinsed in cacodylate buffer. The cells were then dehydrated using increasing concentrations of graded acetone before embedding the pellet in Epon. Ultrathin sections were stained with uranyl acetate and lead citrate. Observations were made using a Zeiss EM-900 transmission electron microscope.

Imaging of TS coated beads (Glycan assay)

We utilized biotinylated polyacrylamido (PAA) glycans, which were purchased from Lectinex; http://www. lectinex.com/ and were dissolved in 10 μL PBS to a concentration of 1 mg/mL. One hundred microlitres of pre-coated bead suspension was aliquoted into 12 microcentrifuge tubes per protein. Each of the 10 synthesized glycans was added to the beads (10 μg) and mixed for 15 min at 4°C. Beads were then spun for 5 min at 160 × g twice and washed with PBS to remove excess glycan. The streptavidin conjugate (Cy3) was added directly to each bead suspension (0.5 μL) and incubated for 10 min at 4°C. Beads were washed with PBS, centrifuged at 160 × g for 10 min and 10 μL of the suspension was added to slides. Beads were imaged immediately using a Axioplan 2 (Zeiss) microscope and Axiovision 4.5 software. The glycans were numbered and keyed by another investigator so that these experiments were done blind. Beads were given a score of 0, 1 or 2, and at least 50 beads were assayed. The scores were added together and the relevance was calculated by Chi-squared analysis (see Material and Methods).

Live/dead cell assay

Cell death was determined by the cell LIVE/DEAD assay (molecular probes, Invitrogen) as per manufacturer instruction. Imaging of the cytoskeleton and LIVE/DEAD assay was achieved using an AxioPlan 2 microscope (Zeiss), 63× objective and a AxioCamMR Camera (Zeiss).

Statistical analysis

Statistical analysis was conducted using Prism software (GraphPad Software Inc.) using an unpaired Student’s t-test; p < 0.05 was considered statistically significant. Unless otherwise stated graphs show the mean of triplicate experiments with ± SEM.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Figure S1: Active trans-sialidase can be purified using dynabeads technology. TcTS and TcTS2V0 was purified using His-tag DynabeadsTM, fractions; Lysate (L), Pellet (P), Flow through (F), washes 1–4 (W) and the eluate (E) were separated by SDS–PAGE and the resultant gel was stained with SYPRO® Ruby protein gel stain (A). About 1.4 × 10^4 μl of sulphate modified latex beads were coated with 10, 15, 20, 25, 30, 35 μg of TcTS and were then separated from the beads by SDS–PAGE and analysed by western blot using anti-his antibody (B). Trans-sialidase activity was assayed by fluorometric assay where MuGal (4-methylumbelliferyl-beta-D-galactoside) acts as the acceptor substrate for the sialic acid from the 3′SL (3′sialyl lactosel) donor, i.e., gives activity with increasing amounts of TcTS. The activity per 20 μg of purified protein was measured in addition to protein coated beads and compared with Trypanosoma cruzi trypomastigote lysate (mean ± SEM, N = 2) (D).

Figure S2: The percentage of internalized TcTS coated beads increases with time. The percentage of internalized TcTS coated beads was plotted over an 8 h time period.

Figure S3: The TcTS and TcTS2V0 glycans binding profile. Beads coated with TcTS and TcTS2V0 were incubated with 10 synthetic glycans (key) which were then detected by microscopy using streptavidinCy3 (red) (A). The beads were then scored according to the amount of fluorescence observed and the mean score is shown in graph B, where * depicts a significant difference according to Chi-square analysis.

Figure S4: Pertussis toxin does no lead to excessive cell death. MDCK II cells were labelled using a LIVE/DEAD™ cell assay kit after treatment with pertussis toxin (PT) and immediately imaged using a Zeiss Axiosplan 2 microscope showing live cells in green and dead cells in red (A) scale bars are 5 μm. One hundred cells were scored dead or alive in triplicate experiments and shown in (B) (mean ± SEM, N = 3).

Figure S5: The MTOC migrates towards internalized TcTS coated beads. The mean distance of the MTOC from the centre of the bead was measured at a range of time points (mean ± SEM, N = 3; B).

Figure S6: MBCD does not lead to excessive cell death at concentrations below 20 μM. MDCK II cells treated with a range of MBCD concentrations were labelled using a LIVE/DEAD™ cell assay kit...
and immediately imaged using a Zeiss Axioscope 2 microscope showing live cells in green and dead cells in red (A) scale bars are 5 µm. One hundred cells were scored dead or alive in triplicate experiments and shown in (B) (mean ± SEM, N = 3).

Figure S7: MBCD treatment dissipates the lipid order present at the bead-cell interface. Cells were treated with the cholesterol depleting reagent, MBCD, prior to treatment with Laurdan and incubation with TcTS and TcTS2V0 coated beads. The resultant images demonstrated that for both sets of protein coated beads the cells appeared a lot bluer and in the region in contact with beads the cells no longer exhibited the yellow/orange characteristic (A). The cell surface membrane in general was a lot bluer in accordance with a loss of lipid order but this was not seen in some of the vesicles which retained their order suggesting that cholesterol is lost from the cell surface membrane. The corresponding histograms demonstrate a negative shift in the average GP to −0.098 in the presence of TcTS coated beads and to −0.019 for TcTS2V0 (B).

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