Trypanosoma cruzi infections and associated pathology in urban-dwelling Virginia opossums (Didelphis virginiana)

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A R T I C L E   I N F O

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A B S T R A C T

Trypanosoma cruzi, a zoonotic protozoan parasite, infects a wide range of mammals. The southern United States has endemic sylvatic transmission cycles maintained by several species of wildlife and domestic dogs. We hypothesized that urban-dwelling opossums (Didelphis virginiana) in South Texas are infected with T. cruzi, and that tissue pathology would be associated with infection. In 2017, we collected blood, heart tissue and anal gland secretions from 100 wild opossums across three seasons that were trapped by animal control in South Texas. In addition, anal gland tissue and intercostal muscle were collected from 43 of the 100 opossums for which time allowed the extra tissue collection. All blood, tissue, and secretion samples were screened for T. cruzi DNA using qPCR with confirmation of positive status achieved through one or more additional PCR assays, including a qPCR to determine the parasite discrete typing unit (DTU). T. cruzi DNA was detected in at least one tissue of 15% of the opossums sampled: blood clot (9%), heart tissue (10%), anal gland secretions (12%), intercostal muscle (16.3%), and anal gland tissue (11.6%). Infection was detected in two or more different tissue types in nine of the opossums. The 35 tissues for which parasite DTU was determined was exclusively TcI. A DTU previously associated with locally-acquired human disease in the United States. T. cruzi-positive opossums were nearly 14 times more likely to exhibit significant heart lesions on histopathology (lymphoplasmacytic inflammation ± fibrosis) when compared to negative opossums (OR = 13.56, CI = 1.23–751.28, p-value = 0.03). Three triatomines were opportunistically collected from the study site, of which two were infected (66.7%), and bloodmeal analysis revealed canine, opossum, and human bloodmeals. Given the presence of parasite in opossum blood, unique potential for shedding of parasite in anal gland secretions, and evidence of vectors feeding on opossums, it is likely that opossums serve as wild reservoirs around urban dwellings in South Texas.

1. Introduction

Triatomines are hematophagous insects that transmit a hemoflagellate parasite, Trypanosoma cruzi, in their feces. With a variety of common names based on geographic location, including kissing bug, triatomines are distributed across the Americas, including the southern United States. Although transmission of T. cruzi is primarily vector borne, other forms of transmission exist, including oral and congenital transmission. Oral transmission has caused outbreaks in humans via contaminated food (Alarcon de Noya and Noya Gonzalez, 2015) and is thought to be a major transmission route for animals (Coura and junqueira, 2015). Infected humans and other mammals could potentially develop a deadly disease known as Chagas disease. The disease affects an estimated 8 million people worldwide (Montgomery et al., 2014). This parasite may cause cardiac abnormalities but may affect other organs as well. Six genetic variants, discrete typing units (DTUs), of the parasite exist: T. cruzi I-VI (TcI-TcVI) (Zingales et al., 2009). A unique bat DTU (TcBat) has been described as well (Pinto et al., 2012). DTUs may be exclusive to certain geographical areas and often overlap (Zingales et al., 2012). However, TcI and TcIV are the most common DTUs found in the United States (Curtis-Robles et al., 2018a). Infections of T. cruzi have been documented in animal hosts from different orders, including Chiroptera, Carnivora, Rodentia, Pilosa and others (Santos et al., 2019), with variation in the degree to which certain wild or domestic species serve as parasite reservoirs in different epidemiological settings. For example, in Argentina and Mexico, domestic cats and dogs have been identified as key reservoir hosts that maintain the domestic transmission cycle (Cardinal et al., 2007; Gurtler et al., 2007; Jimenez-Corolla et al., 2010). In the southern United States, T. cruzi is maintained in sylvatic cycles by several wildlife species,
including raccoons (Procyon lotor) and skunks (Mephitis mephitis) (Hodo and Hamer, 2017), with dogs as key hosts in peridomestic cycles (Curtis-Robles et al., 2017).

In Latin America, peridomestic and urban wildlife like Didelphis sp. opossums have long been recognized as T. cruzi reservoirs where they bridge transmission cycles to domestic animals and humans (Herrera and Urdaneta-Morales, 1992; Roque et al., 2008). Furthermore, the lineages of T. cruzi discrete typing unit (DTU) Tcl are thought to have originated and evolved in opossums, and are consistently isolated from opossums (Yeo et al., 2005). Opossums have been identified as important reservoir hosts in some parts of the southern United States (Herrera and Urdaneta-Morales, 1992; Houk et al., 2010) but the pathological consequence of infection on animal health has not been thoroughly investigated. The Virginia opossum (Didelphis virginiana) has a wide geographic range spanning from Central to North America and is the only marsupial found in the United States (McManus, 1974). Opossums are primarily nocturnal animals that thrive in urban environments and peridomestic environments. They are opportunistically omnivores and insectivores, and could have a high risk of infection if they consume infected triatomines. Opossums and other peridomestic animals can attract triatomines to peridomestic environments thus increasing the risk of transmission to humans and other domestic animals (Ruiz-Pina and Cruz-Reyes, 2002). Being natural hosts of T. cruzi, opossums living in urban areas may pose a public health threat in areas where risk factors are already present (Yeo et al., 2005).

Texas is a hotspot of Chagas transmission, and autochthonous human, domestic animal, and wildlife cases have been reported (Curtis-Robles et al., 2016; Hodo et al., 2019; Nolan et al., 2018). The Rio Grande Valley (RGV) region of south Texas, along the United States-Mexico border, has at least four triatomine species and a higher than Grande Valley (RGV) region of south Texas, along the United States-Mexico border, has at least four triatomine species and a higher than

2. Methods

2.1. Study site

A large animal shelter located in the RGV of South Texas intakes over 30,000 animals a year including an average of 6750 opossums annually (from 2016 to 2018). Animals come to the shelter from multiple animal control agencies. Animal control agencies respond to resident complaints about nuisance opossums and trap or pick up trapped opossums which are then deposited at the animal shelter.

2.2. Opossum necropsy and tissue sampling

We conducted a repeated cross-sectional study of opossums from a South Texas shelter in the winter, spring, and summer of 2017. The opossums (n = 100) used in this study were euthanized for reasons unrelated to our study. City of animal origin (location) and season (winter, spring, summer) of collection were noted. Age was determined by shelter staff through weight approximation (< 2.2 kg = juvenile, > 2.2 kg = adult); based on age determination by weight (Petrides, 1949), only adults were used in this study. Necropsies were performed 20–60 min post-euthanasia. Heart tissue, blood, and anal gland secretions were collected from all 100 animals; in addition, anal gland tissue and intercostal muscle were collected from 43 animals for which time allowed the extra tissue collection. At the time of necropsy, anal gland secretions were collected by manual compression of the anal glands into a sterile vial and were maintained on ice. Using standard necropsy procedures, the heart, anal glands, and approximately 2–3 anterior ribs with attached muscle were completely excised from the body and placed into separate specimen bags. All cutting tools and forceps used were sterilized between subjects using a 50% Glorox® bleach solution, then rinsed with 70% ethanol, and then followed by flame sterilization using a handheld butane torch (Pro Chef’s Torch, BonJour). Approximately 8–12 ml of blood were collected from the thoracic cavity with a syringe and placed into sterile vials with no additives that were maintained on ice. Blood was centrifuged at 5488 RCF for 8 min to recover the blood clot. All samples were saved in −20 °C until processing.

After thawing, an incision was made along the coronary sulcus of the heart to expose the interior of the right ventricle followed by an incision towards the right atrium. An incision was then made from the apex to the left auricle to expose the interior of the left ventricle and atrium. A 1.5–2 cm section of tissue from each chamber was collected with half preserved in 10% neutral-buffered formalin for histopathology and half placed into a nuclease free tube with no additives for prompt molecular analysis. Anal glands and intercostal muscle excised from a rib section were evaluated with separate sections preserved in formalin and prepared for molecular analysis. Extra tissues were saved and stored in −20 °C.

2.3. Detection of parasite DNA in blood clot, anal gland secretions, and tissues

DNA was extracted from the blood clot and tissues using the E.Z.N.A. kit (Omega Bio- Tek, Norcross, GA). Prior to extraction, tissues were macerated and a 30 mg volume was used. For heart samples, all four chambers were prepared and macerated together to represent all parts of the heart. All extraction steps were conducted as per manufacturer instructions with the addition of a longer tissue lysis duration (18–24 h). To detect T. cruzi DNA in samples, the two-step process started with a multiplex Real-Time PCR to amplify a 166-bp segment of the T. cruzi 195-bp repetitive satellite DNA (Duffy et al., 2013). Next, any sample that screened positive (Ct value less than 40 with a sigmoidal amplification curve) was then subjected to discrete typing unit (DTU) determination using a second multiplex Real-Time PCR to amplify the spliced leader intergenic region (SL-IR) (Cura et al., 2015). Finally, any sample that screened positive but was negative on the SL-IR assay was then subjected to a third PCR using the T. cruzi 121/122 primers to amplify a 330bp region of kinetoplast DNA (Curtis-Robles et al., 2016; Virreira et al., 2003; Wincker et al., 1994). The proportion of samples testing positive on each PCR is presented, and samples that had positive results on two independent PCRs were considered to be infected. This requirement of positivity on multiple PCRs that amplify independent genetic regions was used to reduce the chance that positive samples may have resulted from amplicon contamination.
2.4. Histopathology

Histopathology evaluation was performed by a board-certified veterinary pathologist (CLH) on twelve opossums that were PCR-positive on at least one tissue type and on twelve randomly-selected opossums that were negative on all PCR tests. The formalin fixed tissues from these animals were routinely processed for histopathology and stained with hematoxylin and eosin. Inflammation was semiquantitatively scored (Inflammation Score = IS) for each heart chamber, intercostal muscle, and anal glands on a numeric scale as normal (0), minimal (1), mild (2), moderate (3), or severe-marked (4). Additionally, the presence of fibrosis, cardiomyocyte degeneration or necrosis, and the distribution (focal, multifocal, focally extensive) and location (interstititial, myocardial, epicardial) of lesions were recorded. An overall heart inflammation score for each animal was calculated by adding the inflammation scores from the left and right sides of the heart. For analysis, animals were dichotomized by heart pathology status (significant lesions present or absent), in which significant was defined as an overall heart inflammation score (≥3). As previously described, we chose the inflammation cutoff of three because it represented at least mild inflammation in one section and minimal in another, or at least moderate inflammation in any one heart chamber (Hodo et al., 2020).

2.5. Epidemiological and statistical analysis

Molecular prevalence in blood was calculated as the total number of opossums that had PCR-positive blood clot, over the total number of opossums enrolled in the study. Molecular prevalence in tissues and anal gland secretions was calculated as the total number of opossums with a PCR-positive tissue type, over the total number of opossums with that specific tissue type collected. This metric may be useful in estimating the prevalence of animals that have parasite localized in specific tissue; tissue tropism of the parasite could vary by parasite genetic strain (Vera-Cruz et al., 2003). Overall infection prevalence was calculated as the total number of opossums that were PCR-positive on any blood clot, anal gland secretion, or tissue type, over the total number of opossums enrolled in the study.

Statistical analysis was performed in RStudio version 1.1.423. (R Development Core Team, 2008). Bivariate analysis using Fisher’s exact test was used to evaluate the relationship between risk factors (sex, season, location) and PCR status (positive on at least one sample vs negative on all samples) and between presence of significant cardiac inflammation and PCR status. Risk factors were further investigated using a generalized linear model if \( P \leq 0.25 \) in the bivariate analysis (Ranganathan et al., 2017), followed by calculating odds ratios and 95% confidence intervals.

2.6. Opportunistic triatomine vector collection and processing

Ongoing education of shelter staff and distribution of educational materials were conducted to raise awareness of triatomine vectors and Chagas disease. Passive vector surveillance was conducted by engaging shelter staff in the safe collection of triatomines on the premises using methods detailed through our Texas A&M University Kissing Bug Citizen Science Program (Curtis-Robles et al., 2018a). Triatomines were identified morphologically to species, sexed, surface-sterilized, and dissected (Curtis-Robles et al., 2015; Lent and Wygodzinsky, 1979). At the time of insect dissection, the presence of host blood in the hindgut was noted and scored from 1 (no blood observed) to 5 (fully engorged bug). DNA was extracted (KingFisher Cell and Tissue DNA kit, Thermo Fisher Scientific, Waltham, MA) from the vector hindguts and tested for infection and DTU of T. cruzi using the molecular methods described above. In order to determine the vertebrate hosts upon which triatomines previously fed, a molecular bloodmeal analysis (BMA) was conducted using PCR amplification of host cytochrome B sequences using ‘herp’ and ‘BM’ primers and Sanger sequencing (Eton Bioscience Inc., San Diego, CA, USA) as previously detailed (Curtis-Robles et al., 2018c). Samples were considered positive for a human blood meal only when yielding human DNA sequences on two independent assays (Curtis-Robles et al., 2018c).

3. Results

3.1. Demographic results

Of the 100 opossums sampled, 48 were female and 52 were male; exclusively adults were sampled. The number of opossums collected by season are as follows: winter (\( n = 42 \)), spring (\( n = 27 \)), summer (\( n = 31 \)). The majority of the opossums originated from McAllen, TX (92%) while the rest originated from Alamo, TX (6%) and Palmview, TX (2%). None of these factors (sex: p-value = 1, season: p-value = 0.44, location: p-value = 1) were associated with T. cruzi infection.

3.2. Molecular results and overall infection prevalence

Of 100 opossums, 15 (15%) had at least one sample that met the criteria for being called positive (positive on two independent PCR tests), including 9 animals that had two or more PCR positive tissues of which five were positive in all samples tested (Table 1). Based on the criterion of being positive on two independent PCR tests, the molecular prevalence in blood was 9%, molecular prevalence in heart tissue was 10%, and molecular prevalence in anal gland secretion was 12%. There were three additional samples that screened positive (CT value less than 40 in the initial PCR) yet were negative on all subsequent assays and were therefore considered negative in the prevalence estimates. These samples comprised two intercostal muscle (from animals with no other positive samples) and one anal gland tissue (from an animal that also had positive heart tissue), and in all cases had relatively high CT values (36.1, 97.9, and 38.7, respectively). If these samples were instead interpreted as positive, then the overall infection prevalence in opossums would be increased from 15% to 17% (17 of 100 animals). The DTU for all PCR positive blood clots, heart tissue samples, and five of 12 anal gland secretions was determined, and in all cases it was Tcl. The rest of the anal gland secretions (\( n = 7 \)) did not amplify on the SL-IR PCR that is used for DTU determination, yet were confirmed positive on the third PCR assay to amplify kinetoplast DNA. Intercostal muscle and anal glands were sampled from 43 opossums. The molecular prevalence was 16.3% (\( n = 7 \)) in intercostal muscle and 9.3% (\( n = 4 \)) in anal glands; all positive samples were infected with T. cruzi DTU Tcl. All of the samples in which DTU was determined (\( n = 35 \)) were exclusively Tcl.

3.3. Histology

Heart inflammation was evaluated via histopathology on twelve opossums that were positive on at least one tissue type (positive group) and twelve randomly-selected opossums that were negative on all PCR tests (negative group). Of the 12 hearts from the positive group, all 12 showed some level of inflammation (Table 1), including eight (66.7%) with bilateral lymphoplasmacytic inflammation and four with unilaterial lymphoplasmacytic inflammation. In contrast, of the 12 hearts from the negative group, eight showed some level of inflammation in which bilateral inflammation was present in only a single animal (8.3%) (Table 2). Overall, seven (58.3%) animals from the PCR-positive group had inflammation categorized as significant (overall heart inflammation score ≥ 3). In contrast, only one (8.3%) of the hearts from the negative group had significant inflammation (Table 1). Opossums from the positive group were nearly 14 times more likely to exhibit significant heart lesions when compared to opossums in the negative group (OR = 13.56, CI = 1.23–751.28, p-value = 0.03).

The section of heart from one of the PCR-positive opossum that had moderate inflammation in the left chamber also had myocardial fiber loss with replacement fibrosis (Fig. 1). Few eosinophils were observed
in one of the right chambers of a heart from a positive opossum (OP72). The heart from the negative opossum that had left sided moderate inflammation and the heart from the negative opossum that had right sided mild inflammation had numerous eosinophils, which are more commonly associated with other etiologies and not *T. cruzi* (Barr et al., 1991).

Minimal to mild inflammation was observed in intercostal muscles from two animals in the positive group and four animals in the negative group. *Sarcocystis* sp. cysts (sarcocysts) were observed in the intercostal muscle of four positive animals, in one case associated with mild inflammation. Minimal to moderate inflammation was observed in 75% of the anal gland tissues from the positive group (Table 1) and all anal gland tissues of the negative group (Table 2).

### 3.4. Triatamine vector results

A total of 3 triatomines were collected by shelter staff, including one prior to the start of the study and two across the study period, and all were identified as adult female *Triatoma gerstaeckeri* (Table 3). All three bugs showed engorgement: two were assigned a blood meal score (BMS) of 4 and one was assigned a BMS of 5, indicative of recent meals. Two (66.7%) triatomines were PCR positive for *T. cruzi*; both were infected with DTU TcI. Bloodmeal analysis of one PCR-positive triatomine showed the presence of domestic canine (*Canis lupus familiaris*) DNA, and the other PCR-positive bug showed opossum (*Didelphis virginiana*) DNA. Blood meal analysis in the PCR-negative bug revealed human DNA.

### Table 1

Molecular and histopathology results (inflammation scores; IS) in Virginia opossums (*Didelphis virginiana*) of South Texas that tested PCR-positive on at least one tissue. All PCR-positive tissues were determined to be DTU TcI unless notes in footnote.

| ID  | Sex | Blood Clot Secretion | Heart Tissue IS | Significant Heart Inflammation | Intercostal Muscle IS | Anal Gland Tissue IS | Number of PCR Positive Tissues |
|-----|-----|----------------------|-----------------|-------------------------------|-----------------------|----------------------|---------------------------------|
| OP03 F | Negative | Positive | Negative b b b | N/A | N/A | N/A | 1/3 |
| OP05 F | Negative | Positive | Negative b b b | N/A | N/A | N/A | 1/3 |
| OP06 F | Negative | Positive | Negative b b b | N/A | N/A | N/A | 2/3 |
| OP07 M | Positive | Positive | Positive 3 2 Yes | N/A | N/A | N/A | 3/3 |
| OP09 F | Negative | Positive | Negative 0 1 No | N/A | N/A | N/A | 1/3 |
| OP17 F | Positive | Positive | Positive 2 2 Yes | N/A | N/A | N/A | 3/3 |
| OP34 M | Negative | Positive | Negative 1 0 No | N/A | N/A | N/A | 1/3 |
| OP63 F | Positive | Negative | Positive 0 2 No | Positive 0 Negative 2 3/5 |
| OP69 F | Positive | Positive | Positive 2 2 Yes | Positive 0 Positive 3 5/5 |
| OP72 M | Positive | Positive | Positive 2 3 Yes | Positive 0 Positive 2 5/5 |
| OP81 M | Positive | Positive | Positive 2 2 Yes | Positive 0 Positive 1 5/5 |
| OP92 M | Negative | Positive | Positive 1 1 No | Negative 0 Negative 2 1/5 |
| OP94 M | Positive | Positive | Positive 2 3 Yes | Positive 0 Negative 0 4/5 |
| OP94 M | Positive | Positive | Positive 2 2 Yes | Positive 2 Positive 1 4/5 |

N/A = Not available.

* No DTU detected, confirmed positive using 121/122 PCR
* No histology performed.
* Sarcocysts present.
* Eosinophils present.

### Table 2

Histopathology results (inflammation scores; IS) in Virginia opossums (*Didelphis virginiana*) of South Texas that tested PCR negative.

| ID  | Sex | Heart IS | Significant Heart Inflammation | Intercostal Muscle IS | Anal Gland Tissue IS |
|-----|-----|----------|--------------------------------|-----------------------|----------------------|
| OP01 M | 0 1 No | N/A | N/A |
| OP11 F | 0 0 No | N/A | N/A |
| OP23 M | 0 1 No | N/A | N/A |
| OP38 M | 0 0 No | N/A | N/A |
| OP58 M | 0 1 No | 1 1 |
| OP60 F | 3* 1 Yes | 1 3 |
| OP66 F | 0 1 No | 0 2 |
| OP74 M | 2 0 No | 0 2 |
| OP88 M | 1 0 No | 0 2 |
| OP95 M | 0 2* No | 1 2 |
| OP99 M | 0 0 No | 0 2 |
| OP100 M | 0 0 No | 1 1 |

N/A = Not available.

* Eosinophils present.

in one of the right chambers of a heart from a positive opossum (OP72). The heart from the negative opossum that had left sided moderate inflammation and the heart from the negative opossum that had right sided mild inflammation had numerous eosinophils, which are more commonly associated with other etiologies and not *T. cruzi* (Barr et al., 1991).

Minimal to mild inflammation was observed in intercostal muscles from two animals in the positive group and four animals in the negative group. *Sarcocystis* sp. cysts (sarcocysts) were observed in the intercostal muscle of four positive animals, in one case associated with mild inflammation. Minimal to moderate inflammation was observed in 75% of the anal gland tissues from the positive group (Table 1) and all anal gland tissues of the negative group (Table 2).

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A total of 3 triatomines were collected by shelter staff, including one prior to the start of the study and two across the study period, and all were identified as adult female *Triatoma gerstaeckeri* (Table 3). All three bugs showed engorgement: two were assigned a blood meal score (BMS) of 4 and one was assigned a BMS of 5, indicative of recent meals. Two (66.7%) triatomines were PCR positive for *T. cruzi*; both were infected with DTU TcI. Bloodmeal analysis of one PCR-positive triatomine showed the presence of domestic canine (*Canis lupus familiaris*) DNA, and the other PCR-positive bug showed opossum (*Didelphis virginiana*) DNA. Blood meal analysis in the PCR-negative bug revealed human DNA.
Different strains of *T. cruzi* show differential tissue tropism; for example, TcI is commonly found in heart, skeletal muscles, and intestine, and less commonly in the brain (Cruz et al., 2015). In previous studies on *Didelphis marsupialis* infected with various strains of *T. cruzi* (Carreira et al., 1996), mild inflammation and few amastigotes were observed in skeletal muscle, indicating variation in *T. cruzi* strains and coinfection with multiple strains may impact pathology outcomes (Roellig et al., 2009).

Almost all of the anal glands in the positive and negative group showed lymphoplasmacytic inflammation with few eosinophils. The high incidence of inflammation in anal gland tissue could be due to *T. cruzi* or could be normal or incidental for opossums (Brown, 1972). Lymphoplasmacytic inflammation could be a result of other pathogens like *Sarcocystis sp.* and *Besnoitia darlingi* (Barr et al., 1991). *T. cruzi* is often multifocally distributed in tissues which may affect results if the regions sampled for PCR versus histology differ. Additionally, parasite DNA may not be detectable due to limitations that occur during the DNA extraction process, such as issues at the enzyme digestion step due to excess tissue (James et al., 2002). Although measures were taken to prevent fecal contamination, it possible that there may have been PCR inhibitors (Monteiro et al., 1997) from feces around the anal glands affecting the detection of parasite DNA.

Three triatomines were opportunistically collected in outdoor kennels at the shelter, including two *T. cruzi*-infected bugs which harbored TcI - the same DTU that infected the opossums at the shelter. The three bugs had bloodmeals from an opossum, dog, and human. At the shelter, dogs and opossums are housed in outdoor environments. Although the bloodmeal analysis procedure cannot inform where the insects fed on these hosts, their high levels of engorgement suggest the bloodmeals were recently acquired and may have been obtained at the shelter. Alternatively, less likely, insects may have fed in the surrounding environment before dispersing to the shelter. Although vectors were found on site at the shelter, the infected opossums in our study likely acquired infection prior to arrival at shelter because opossums do not remain on site at the shelter for long, and triatomines are widespread in the RGV (Curtis-Robles et al., 2018b).

We show opossums are infected with *T. cruzi*, sometimes associated with pathology in multiple organs, in the RGV where ongoing human and canine transmission occurs (Curtis-Robles et al., 2017; Nolan et al., 2018). The high prevalence of *T. cruzi* infection in anal gland secretions of infected animals may signal a potential source of infection for humans and animals. Future research will determine the relative importance of opossums as reservoirs of *T. cruzi* in the United States and the threat they pose to domestic animals and public health.

### Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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