IPSE, an Abundant Egg-Secreted Protein of the Carcinogenic Helminth Schistosoma haematobium, Promotes Proliferation of Bladder Cancer Cells and Angiogenesis

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Abstract

Background: *Schistosoma haematobium*, the helminth causing urogenital schistosomiasis, is a known bladder carcinogen. Despite the causal link between *S. haematobium* and bladder cancer, the underlying mechanisms are poorly understood. *S. haematobium* oviposition in the bladder is associated with angiogenesis and urothelial hyperplasia. These changes may be pre-carcinogenic events in the bladder. We hypothesized that the Interleukin-4-inducing principle of *Schistosoma mansoni* eggs (IPSE), an *S. haematobium* egg-secreted “infiltrin” protein that enters host cell nuclei to alter cellular activity, is sufficient to induce angiogenesis and urothelial hyperplasia.

Methods: Mouse bladders injected with *S. haematobium* eggs were analyzed via microscopy for angiogenesis and urothelial hyperplasia. Endothelial and urothelial cell lines were incubated with recombinant IPSE protein or an IPSE mutant protein that lacks the native nuclear localization sequence (NLS-) and proliferation measured using CFSE staining and real-time monitoring of cell growth. IPSE’s effects on urothelial cell cycle status was assayed through propidium iodide staining. Endothelial and urothelial cell uptake of fluorophore-labeled IPSE was measured.

Findings: Injection of *S. haematobium* eggs into the bladder triggers angiogenesis, enhances leakiness of bladder blood vessels, and drives urothelial hyperplasia. Wild type IPSE, but not NLS-, increases proliferation of endothelial and urothelial cells and skews urothelial cells towards S phase. Finally, IPSE is internalized by both endothelial and urothelial cells.

Interpretation: IPSE drives endothelial and urothelial proliferation, which may depend on internalization of the molecule. The urothelial effects of IPSE depend upon its NLS. Thus, IPSE is a candidate pro-carcinogenic molecule of *S. haematobium*.

Summary

*Schistosoma haematobium* acts as a bladder carcinogen through unclear mechanisms. The *S. haematobium* homolog of IPSE, a secreted schistosome egg immunomodulatory molecule, enhances angiogenesis and urothelial proliferation, hallmarks of pre-carcinogenesis, suggesting IPSE is a key pro-oncogenic molecule of *S. haematobium*.

Introduction

Urogenital schistosomiasis (UGS), primarily infection by the blood fluke, *Schistosoma haematobium*, affects the bladder and other pelvic organs. Egg deposition by the adult stage of the parasite in the bladder during infection is notable for its association with hematuria and bladder angiogenesis (1). This suggests that this parasitic worm triggers aberrant host endothelial responses. *S. haematobium* oviposition is also linked to urothelial alterations such as hyperplasia (1–6). It is unknown, however, if other factors produced by the *S. haematobium* adult worms, which live in the pelvic veins, also contribute to the bladder endothelial and urothelial changes of UGS. Both abnormal angiogenesis and epithelial
hyperplasia have been associated with pre-carcinogenic changes in endodermal organs. Indeed, UGS is categorized as a group 1 carcinogen, i.e., deemed to cause cancer in humans, by the International Agency for Research on Cancer (7). It is, however, unclear which components of *S. haematobium* eggs are pro-oncogenic.

One major protein secreted by the egg of *S. haematobium* is the ortholog of interleukin-4-inducing principle (IPSE) of the egg of the congener, *Schistosoma mansoni*, in which it was first discovered (8). *S. mansoni* IPSE features numerous host modulatory properties. As indicated by its name, IPSE leads to secretion of IL-4 from basophils and mast cells by engaging IgE bound to the high affinity IgE receptor on these two cell types. IPSE also contains a nuclear localization sequence which guides the protein into the nuclei of host cells and presumably alters cellular activity (9, 10). We have demonstrated that H03-H-IPSE, one of the major *S. haematobium* orthologs of IPSE (11), induces proliferation of mouse urothelial cells *in vitro* in a nuclear localization sequence-dependent manner (12). Furthermore, H06-H-IPSE, another major *S. haematobium* ortholog of IPSE, is internalized by both urothelial and neuronal cells (13), indicating that IPSE may be taken up by and influence diverse cell types. This led us to hypothesize that IPSE drives proliferation of human urothelial cells, skews them towards S-phase of the cell cycle, and also induces angiogenesis.

Here we demonstrate that *S. haematobium* eggs, in the absence of *S. haematobium* adult stage worms, are sufficient to initiate the bladder endothelial and urothelial alterations of urogenital schistosomiasis. Furthermore, we show *in vitro* that H03-H-IPSE is taken up by both endothelial and urothelial cells. H03-H-IPSE triggers angiogenic behavior in endothelial cells in culture and orchestrates urothelial proliferation and S-phase cell cycle skewing in a nuclear localization sequence-dependent fashion.

**Materials And Methods**

**Mice**

Female 6- to 7-wk-old BALB/c mice (Charles River Laboratories, Wilmington, MA, USA) were housed under 12 h light-dark cycles in temperature-controlled holding rooms with unlimited access to dry mouse chow and water. Newly received mice were acclimated to the animal facility for at least one week prior to experimental use.

**IPSE protein production and labeling**

Recombinant H03 H-IPSE, H06 H-IPSE, and an NLS mutant of H03 H-IPSE (the wild type NLS sequence SKRRKDY changed to SAAGAAY) were produced in HEK293-6E cells, and then purified via immobilized-metal affinity chromatography using an 8x His tag in the construct as described (11). H03 H-IPSE was conjugated to Alexa Fluor 488 using a Alexa Fluor 488 antibody labeling kit (Thermofisher Scientific, Waltham, MA) according to the manufacturer’s instructions; however, the pH was kept at 7.4 throughout the reaction to enrich for labeling of the terminal amine (pKa of 7.4). The efficiency of conjugation was confirmed by Nanodrop. The typical level of labeling was one mole of dye per mole of IPSE, which
suggested IPSE was only labeled on the terminal amine. Avoiding labelling of lysine or arginine residues in the protein backbone is critical, as each of the positively charged amino acids in the NLS is essential for nuclear translocation (9). Low labeling efficiency thus minimized the potential interference of the dye with IPSE’s functional domains.

**S. haematobium egg injection**

Bladder wall injections were performed as described previously (14). Female BALB/c mice were anesthetized with isoflurane, a midline lower abdominal incision was made, and the bladder was exteriorized. *S. haematobium* eggs (3,000 eggs in 50 µl) were injected into the wall of the bladder. Abdominal incisions were subsequently closed with 4–0 Vicryl sutures, and the surgical site was treated once with topical antibiotic ointment.

**Histology**

Bladder tissues were fixed in neutral buffered formalin, dehydrated, and embedded in paraffin. Five-micrometer sections were stained with hematoxylin and eosin. Histology was analyzed by a board-certified pathologist (JIO) in a blinded fashion.

**Ex vivo angiogenesis and microvascular leakage microscopy**

Microvascular leakage was assessed three weeks following *S. haematobium* egg or vehicle injection using an established protocol (15). Briefly, following FITC-lectin injection, 100 µL R50 Fluoro-Max red fluorescent microspheres, 0.048 µm in diameter (Thermo Scientific), were injected through the inferior vena cava. After 3–5 min in circulation, a sternotomy was performed, and the aorta was cannulated via the left ventricle with an 18-gauge angiocatheter and perfused with 1% paraformaldehyde for 3–5 min using a mini pump (Fisher Scientific). Grafts were harvested and mounted as described above. Microvascular permeability was assessed by using confocal microscopy to determine the extent of microsphere extravasation.

**Proliferation xCELLigence assay**

Cells were seeded at 5,000 cells per well in 200 µl of complete media in E-plates (ACEA Biosciences, San Diego, CA, USA) and grown overnight while monitored with an xCELLigence DP system (ACEA Biosciences) which monitors cellular events in real time by measuring electrical impedance across interdigitated gold micro-electrodes integrated on the bottom of tissue culture plates (16, 17). Cells were washed three times with PBS and cultured with 180 µl EGM-2 basal media (no growth factors or supplements) and incubated for a minimum of 6 h before further treatment. Treatments were prepared at 10 × concentrations and added to each well in a total volume of 20 µl. The xCELLigence DP recorded cell index readings every 15 min for 3 days after treatment. Cell index readings were normalized before
treatment and cell proliferation ratios were determined from four biological replicates and represent the relative numbers of cells compared to control cells. A two-way ANOVA with Holm–Sidak’s multiple comparisons test was used to compare IPSE treatment to medium-alone control, with $P \leq 0.05$ deemed significant.

Cell lines

3B-11 cells were obtained from ATCC (Manassas, Virginia). ATCC uses morphology, karyotyping, and PCR based approaches to confirm the identity of human cell lines and to rule out both intra- and interspecies contamination. These include an assay to detect species specific variants of the cytochrome C oxidase I gene (COI analysis) to rule out inter-species contamination and short tandem repeat (STR) profiling to distinguish between individual human cell lines and rule out intra-species contamination. HCV-29 cells were obtained from Dra. Monica Botelho. These cells have undergone STR profiling using the following markers: amelogenin, D8S1179, D18S51, D21S11, FGA, TH01, and vWA. The 3B-11 cells and the HCV-29 cells were Mycoplasma-free as established using the Lookout Mycoplasma PCR detection kit (Sigma-Aldrich).

Tubule formation assay

Growth factor-reduced Matrigel (Corning, Corning, NY, USA) was plated into a 96-well μ-angiogenesis plate (ibidi, Planegg, Germany) at 10 µl/well, and incubated at 37 °C in 5% CO$_2$ in air for 60 min as described (18). 3B-11 cells were detached using Trypsin/EDTA and resuspended in DMEM (Gibco), and seeded at 30,000 cells/well in medium supplemented with 10 µM sulforaphane (SFPH, Sigma) (negative control), or 1.8 µg/mL or 3.6 µg/mL IPSE. The ibidi plate was incubated for 5 h in a humidified atmosphere of 5% CO$_2$ in air at 37 °C in a microscope stage top incubator (OKOLAB, Pozzuoli, Naples, Italy). At intervals, photomicrographs of cells and nascent and developed tubules were collected using a Leica DMi8 automated platform microscope under bright field at 2.5 × magnification, and LASX software (Leica).

Analysis of tubule formation

Automated angiogenesis assessment was performed on TFA 490² pixel images by ImageJ (NIH) with the phase-contrast Angiogenesis Analyzer plugin tool as described (19, 20). Settings used were as follows: 10 pixel minimum object size; 25 pixel minimum branch size; 2,500 pixel artefactual loop size; 25 pixel isolated element size threshold; 30 pixel master segment size threshold; with iteration number of 3. The four output metrics (mesh count, segment count, segment length, and junction count) were either plotted directly or as a percentage relative to the medium-alone blank treatment (treatment measure divided by medium-alone measure). A two-way ANOVA with Holm–Sidak’s multiple comparisons test was used to
compare IPSE treatment against medium-alone blank control for the four metrics with $P \leq 0.05$ deemed significant.

Combining the four metrics into a single evenly weighted variable was accomplished through the calculation of $Z$ standardized scores that were based on population values (21). The formula below generates the $Z$ score and represented the distance between the raw score and the population mean in units of the SD. Population values were estimated from 39 treatment replicates.

$$Z = \frac{\text{treatment metric value} - \text{metric population mean}}{\text{population standard deviation}}$$

The combined robust $Z$ score ($Z^*$) was generated for each replicate from the median $Z$ score of the four metrics. $Z^*$ scores were plotted, and IPSE treatments compared to medium-alone blank control using one-way ANOVA with Holm–Sidak's multiple comparisons test, $P \leq 0.05$ was considered to be statistically significant.

**Cell cycle analyses and CFSE assay**

The human bladder epithelium (urothelium) cell line HCV-29 was grown in T-75 tissue culture flasks in complete DMEM media (Gibco) under 5% CO$_2$ at 37 °C. For cell cycle assays, $1 \times 10^5$ urothelial cells were co-incubated with IPSE. Following 48 hours of culture, the cells were fixed and stained with propidium iodide for cell cycle analysis. For CFSE assays to assess cell proliferation, cells were stained with the CFSE dye prior to stimulation with IPSE and cultured for 48 hours. The CFSE dye was evaluated post-culture by flow cytometry using the FITC channel. The intensity of CFSE dye, which halves with each cell cycle, was used to track the generations of urothelial cells.

**Endocytosis assays**

HCV-29 human derived urothelial cells [see (22)] were grown in MEM (Thermo Fisher Scientific, Waltham, MA) with 10% fetal bovine serum (Sigma-Aldrich, St. Louis, MO). For internalization assays, floating cells and adherent cells (released via 0.12% trypsin (Sigma-Aldrich, St. Louis, MO) without EDTA) were washed in fresh medium, and aliquoted into 24 well plates at 200,000 cells/mL in 1 mL. The cells were incubated with Alexa 488-labeled H03 at 1 µg/mL (H03 was conjugated to Alexa 488 using a kit from ThermoScientific, Waltham, MA) for 16 hours at 37° C. Cells were released via 0.12% trypsin without EDTA and washed 3 times with PBS (Sigma-Aldrich, St. Louis, MO). 0.4% trypan blue (ThermoScientific, Waltham, MA) was added to the cells (1:4) to quench extracellular Alexa 488 signal. The cells were analyzed by flow cytometry (Beckman Coulter, CytoFLEX) to isolate the intracellular Alexa 488 signal. Data were analyzed using FlowJo and GraphPad.

**Statistical analysis**

Statistical analyses were performed using GraphPad software. Except where noted otherwise, the Mann-Whitney U test and Student’s $t$ test were used to evaluate statistical significance for nonparametrically and parametrically distributed data, respectively. $P$ values of $< 0.05$ were defined as significant.
Results

S. haematobium eggs trigger urothelial hyperplasia of the bladder

We sought to determine whether *S. haematobium* eggs alone, in the absence of *S. haematobium* adult worms, were sufficient to induce urothelial hyperplasia in the bladder seen during urogenital schistosomiasis. To address the question, we injected *S. haematobium* eggs into the mouse bladder wall. Five weeks following egg injection, the urothelial lining of the bladder still exhibited significant hyperplasia (up to 12-cell thick urothelium compared to three cells characteristic of quiescent mouse bladder, Fig. 1). Thus, *S. haematobium* eggs were sufficient to trigger bladder urothelial hyperplasia.

S. haematobium eggs initiate proliferation of bladder blood vessels with increased permeability

Hematuria is a hallmark sign of urogenital schistosomiasis (UGS). By definition, hematuria involves leakage of erythrocytes from the lumen of blood vessels into surrounding bladder tissues and the bladder lumen. UGS is also associated with angiogenic changes in the bladder. We tested whether *S. haematobium* eggs were sufficient, without the presence of *S. haematobium* worms, to initiate angiogenesis and increase vascular permeability. Bladder wall injection of mice with *S. haematobium* eggs induced angiogenesis featuring dilated, leaky blood vessels (Fig. 2). We conclude that *S. haematobium* eggs suffice to orchestrate bladder angiogenesis and vascular leakiness.

IPSE is internalized by endothelial and urothelial cells

We next assayed the capacity of HCV-29 urothelial and 3B-11 endothelial cells to take up H03-H-IPSE. Using Alexa 488-labeled H03-H-IPSE, we noted that both cell types readily uptake the protein (Fig. 3).

IPSE induces tubule formation by endothelial cells

Having demonstrated that whole *S. haematobium* eggs induce angiogenesis *in vivo*, we next sought to determine if recombinant H03-H-IPSE protein alone could induce similar changes in endothelial cells *in vitro*. *In vitro* tubule formation by the mouse 3B-11 endothelial cell line was enhanced by addition of 1.8 µg/mL or 3.6 µg/mL H03-H-IPSE to cultures (Fig. 4), indicating that this protein is pro-angiogenic (p < 0.05).

IPSE mediates urothelial cell proliferation in a nuclear localization sequence-dependent manner

Given *S. haematobium* eggs stimulate urothelial hyperplasia in the bladder, we next assayed the ability of recombinant H03-H-IPSE protein to trigger the *in vitro* equivalent, proliferation of urothelial cells. Electrical impedance measurements of cellular proliferation using the xCELLigence platform revealed that
incubation of HCV-29 human urothelial cells with 43 pmol, but not 216 or 864 pmol (possibly cytotoxicity), of H03-H-IPSE induced proliferation of the urothelial cells (Fig. 5A).

The pro-proliferative effect of H03-H-IPSE on urothelial cells was also verified using CFSE staining. These assays demonstrated that H03-H-IPSE, up to the tested concentration of 65 pmol, drove proliferation of HCV-29 cells in a concentration-dependent fashion (Fig. 5B). H06-H-IPSE also induced proliferation of HCV-29 cells. Incubation of HCV-29 cells with a nuclear localization sequence mutant of H03-H-IPSE failed to enhance proliferation, suggesting that H03-H-IPSE's pro-proliferative influence depends upon the protein's ability to migrate into the nucleus of the host cell.

**IPSE skews urothelial cell cycle status in a nuclear localization sequence-dependent fashion**

Our findings that H03-H-IPSE stimulated urothelial proliferation led us to examine whether this protein alters the cell cycle status of urothelial cells. Through propidium iodide staining of H03-H-IPSE-exposed HCV-29 urothelial cells, we ascertained that this protein indeed skews cells towards S-phase (Fig. 6). The ability of H03-H-IPSE to bias urothelial cells towards S-phase was nuclear localization sequence-dependent, since the nuclear localization sequence mutant form of H03-H-IPSE did not change the cell cycle status of HCV-29 cells.

**Discussion**

Schistosomiasis affects more than 200 million people worldwide (23). The majority of cases affect the urogenital tract, caused mostly by *S. haematobium*. As a blood fluke, *S. haematobium* adult worms live primarily in the pelvic veins, where worm pairs lay eggs that lodge in the pelvic organs. Eggs laid in the bladder induce granuloma formation, which is thought to facilitate passage of eggs into the urinary stream (24, 25). Eggs voided into fresh bodies of water hatch into miracidia, which infect intermediate snail hosts. Infected snails release cercariae, the larval stage which infects humans.

The complex life cycle of *S. haematobium* depends upon the parasite successfully negotiating host tissue and immune responses that may threaten its survival. For instance, hematuria is a cardinal sign of urogenital schistosomiasis, and represents a compromised urothelial and endothelial barrier which allows *S. haematobium* to continue reproducing. However, unchecked damage to the urothelium and bladder blood vessels results in hemorrhage and even host death, which is counterproductive to any parasite (26). Hence, it is possible that *S. haematobium* and humans have co-evolved survival strategies. The exuberant proliferative response of the urothelium and endothelium to *S. haematobium* eggs may be one such strategy, since this likely promotes bladder tissue repair.

Schistosomes may induce host tissue repair responses, in particular angiogenesis, to promote egg expulsion. Turner et al. reported that deposition of *S. mansoni* eggs in Peyers’ patches of the mouse small intestine is associated with vascular remodeling and an expanded venule network (27). In mice deficient in Peyers’ patches, egg excretion is lessened, leading to more eggs entrapped in tissues, and
consequently worsened host morbidity. We postulate that *S. haematobium* eggs similarly initiate angiogenesis in the bladder to facilitate their expulsion into the urinary stream.

Angiogenesis and urothelial proliferation benefit the *S. haematobium*-infected host in the short term but have been postulated to facilitate bladder carcinogenesis. *S. haematobium* is one of a handful of helminths that are known to be carcinogenic (28). Angiogenesis, such as that seen in bladders containing *S. haematobium* eggs, is a crucially important process for tumors, since their rapid growth puts them at risk of outstripping their blood supply. Expansion of local vascular beds may also enhance hematogenous spread of metastases. Urogenital schistosomiasis is associated with urothelial hyperplasia, a potentially pre-cancerous feature in the bladder.

Despite the prominence of urothelial and endothelial aberrations in urogenital schistosomiasis, our comprehension of the underpinnings of these pathological processes is limited. One mystery is whether *S. haematobium* eggs, without adult worms, are sufficient to activate urothelial hyperplasia and angiogenesis in the bladder. This is a significant matter because *S. haematobium* worms can live for years within their human hosts, whereas parasite eggs can remain in the bladder wall for decades and continue to drive chronic inflammation, even after successful treatment of infection. Understanding the contributions of *S. haematobium* worms to bladder pathogenesis is important to improving our knowledge base of chronic urogenital schistosomiasis. To that end, we observed that when *S. haematobium* eggs are injected into the bladder walls of mice, this elicits significant urothelial hyperplasia, angiogenesis, and vascular leakiness. These bladder changes are not seen when mice are injected with control vehicle. These findings intimate that *S. haematobium* worms play no or minimal role in the pathological changes of the bladder associated with urogenital schistosomiasis. Finally, these findings point to one or more *S. haematobium* egg-associated factors which mediate urothelial and endothelial changes.

We hypothesized that one of these egg-associated factors was the Interleukin-4 inducing Principle of Schistosoma mansoni Eggs (IPSE) (29). IPSE, also known as α-1 (8), features many host immunomodulatory functions. First, IPSE ligates Fcε receptor-bound IgE on the surface of mast cells and basophils to trigger IL-4 secretion (30–32). It also binds to immunoglobulins on the surface of B regulatory cells (Bregs) and thereby activates these cells (33). IPSE can also sequester chemokines; it was previously known as *S. mansoni* chemokine-binding protein (smCKBP) (34). Lastly, IPSE contains a nuclear localization sequence which directs the protein to host cell nuclei (9, 11), where it modulates transcription (10, 35).

Although IPSE has multiple immunomodulatory properties, our *in vitro* work indicates that H03-H-IPSE mediates its urothelial and endothelial effects via non-immune mechanisms, including through its nuclear localization sequence. The similar uptake of H03-H-IPSE by endothelial and urothelial cells hints that endocytosis occurs in either a non-specific fashion, or that these cell types share a receptor for H03-H-IPSE. We speculate that once IPSE is internalized by host cells, it may trigger pro-carcinogenic programs. Indeed, Roderfeld et al. showed that IPSE activated hepatocellular carcinoma-associated proto-
oncogenes, namely c-Jun and its associated signaling molecule, STAT3 (35). This supports a potential role for H03-H-IPSE in promoting bladder oncogenesis.

Another interesting issue regarding the potential causal link between IPSE and schistosomal bladder carcinogenesis is whether this molecule promotes the high rates of squamous cell carcinoma seen in association with *S. haematobium* infection (36–39). This remains an open question, since we did not have an immortalized schistosomiasis-associated squamous cell bladder carcinoma cell line available for testing. However, schistosomal bladder cancer can still be associated with carcinomas arising from the urothelium, the tissue of origin for the HCV-29 cells tested herein.

Superficially, targeting IPSE to prevent schistosomal bladder cancer appears to be a worthy prophylactic approach. However, it has been reported that immunization of *S. mansoni* mice with IPSE leads to larger granulomas with enhanced macrophage activity and a mixed type 1 and type 2 immune response (40). Since granulomas contribute to host tissue fibrosis, neutralization of IPSE during schistosomiasis may actually be undesirable.

This investigation has noteworthy limitations. Although we hypothesize that H03-H-IPSE is sufficient to drive urothelial and endothelial proliferation and related effects, we have not demonstrated that it is necessary. There may be additional *S. haematobium* egg-derived factors which contribute to urothelial hyperplasia and angiogenesis during urogenital schistosomiasis. Ideally, transgenic approaches would be used to show that H03-H-IPSE is critical in angiogenesis and urothelial hyperplasia *in vivo*. This approach could be in reach for future studies given that both germline transgenesis and genome editing of egg-expressed genes have now been reported in (41, 42)

In conclusion, we have provided data that support the hypothesis that *S. haematobium* eggs are sufficient to initiate bladder urothelial hyperplasia, angiogenesis, and vascular permeability associated with urogenital schistosomiasis. These changes may be orchestrated by IPSE. Our observations are particularly striking considering that IPSE can increase proliferation of transformed urothelial and endothelial cell lines that, at baseline, already exhibit significant proliferation. IPSE may indeed be a pro-oncogenic factor of *S. haematobium*.

**Declarations**

**Ethics approval and consent to participate**

All animal work was conducted according to relevant U.S. and international guidelines. Specifically, animal experimental work was reviewed and approved as protocol 14-03 by the Institutional Animal Care and Use Committee of the Biomedical Research Institute (Rockville, Maryland, USA). Our Institutional Animal Care and Use Committee guidelines comply with the U.S. Public Health Service Policy on Humane Care and Use of Laboratory Animals.
Consent for publication

All authors have given their consent for publication of this manuscript.

Availability of data and material

All data and material are available upon reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Author Contributions

Chinwike Terry Agbo: performed in vitro urothelial assays

Paul J. Brindley: designed xCELLigence assays, data analysis of blood tubule formation assays

Franco H. Falcone: designed IPSE expression constructs, generated recombinant IPSEs protein

Michael H. Hsieh: conceived and designed experiments, wrote manuscript, provided funding

Theodore S. Jardetzky: designed IPSE expression constructs

Mohammad Afzal Khan: performed ex vivo angiogenesis assays

Olivia K. Lamanna: performed endocytosis assays

Alex Loukas: data analysis of blood tube assays

Evaristus C. Mbanefo: performed in vitro urothelial assays

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Justin I. Odegaard: performed histological analyses
Luke F. Pennington: generated recombinant IPSE protein

Michael J. Smout: conducted data analysis of angiogenesis assays

Shannon E. Karinshak: performed blood tubule formation assays

Kimberly H. Thai: performed ex vivo angiogenesis assays

Yuanlong Zhao: performed in vitro urothelial assays

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**Figures**
Figure 1

Schistosoma haematobium eggs are sufficient to induce urothelial hyperplasia in the bladder. Mice underwent bladder wall injection with S. haematobium eggs. Five weeks later, their bladders were harvested, processed, sectioned, and stained with hematoxylin and eosin. Image shows the typical hyperplastic urothelium (more than 3 cell layers thick) resulting from egg injection.
Schistosoma haematobium eggs are sufficient to induce angiogenesis and increased vascular permeability in the bladder. Mice underwent bladder wall injection with S. haematobium eggs. Three weeks later mice were administered FITC-lectin to label blood vessels and Red Fluoro-Max microbeads (to measure vascular permeability) and their bladders harvested and examined by confocal microscopy. Left, middle, and right columns show FITC channel, red channel, and merged channels, respectively. Detection of red signal indicates leakage of Red Fluoro-Max microbeads out of blood vessels. Each row consists of representative images from a single mouse (n=3 controls, n=3 egg-injected).
H03-H-IPSE is internalized by endothelial and urothelial cells. Alexa 488-labeled H03-H-IPSE was incubated with HCV-29 urothelial and 3B-11 endothelial cells. Extracellular signal was quenched with trypan blue, and the remaining intracellular Alexa 488 signal measured by flow cytometry. Each symbol denotes an individual culture well replicate. Figure shows one of two representative experiments.
Figure 4

H03-H-IPSE drives increased tubule formation by 3B-11 endothelial cells. A, time-lapse movie showing spontaneous tubule formation by 3B-11 cells. B, time-lapse movie showing increased tubule formation by
3B-11 cells following incubation with H03-H-IPSE. C, Z-scores of spontaneous vs. IPSE-associated tubule formation by 3B-11 cells. 95% CI error bars shown, comparisons made against the control with one-way ANOVA and Dunnet multiple analysis correction. * p<0.05
H-IPSE enhances urothelial cell proliferation in a nuclear localization sequence-dependent fashion. A, H03-H-IPSE increases urothelial cell proliferation. HCV-29 human urothelial cells were incubated with 0, 43, 216, or 864 pmol of H03-H-IPSE and electrical impedance (cellular proliferation) measured in real-time using the xCELLigence platform. B, left panel, CFSE-labeled HCV-29 human urothelial cells were incubated with up to 65 pmol of H03-H-IPSE and analyzed for evidence of proliferation (decreased CFSE content per cell with each successive generation of cells). Each symbol denotes an individual culture well replicate (n=8 for each culture condition). B, middle panel, CFSE-labeled HCV-29 human urothelial cells were incubated with up to 65 pmol of H06-H-IPSE and analyzed for evidence of proliferation. Each symbol denotes an individual culture well replicate (n=8 for each culture condition). B, right panel, CFSE-labeled HCV-29 human urothelial cells were incubated with up to 65 pmol of a nuclear localization
mutant of H03-H-IPSE and analyzed for evidence of proliferation (decreased CFSE content per cell with each successive generation of cells). Each symbol denotes an individual culture well replicate.

Figure 6

H-IPSE polarizes urothelial cell cycle status in a nuclear localization sequence-dependent manner. Left panel, propidium iodide-labeled HCV-29 human urothelial cells were incubated with up to 65 pmol of H03-H-IPSE and analyzed for cell cycle status. Each symbol denotes an individual culture well replicate (n=8 for each culture condition). Middle panel, propidium iodide-labeled HCV-29 human urothelial cells were incubated with up to 65 pmol of H06-H-IPSE and analyzed for cell cycle status. Each symbol denotes an individual culture well replicate (n=8 for each culture condition). Right panel, CFSE-labeled HCV-29 human urothelial cells were incubated with up to 65 pmol of a nuclear localization mutant of H03-H-IPSE and analyzed for cell cycle status. Each symbol denotes an individual culture well replicate.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- Figure4B.mov
- Figure4A.mov