Developmental Progression to Infectivity in *Trypanosoma brucei*
Triggered by an RNA-Binding Protein

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Developmental Progression to Infectivity in Trypanosoma brucei Triggered by an RNA-Binding Protein

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Unraveling the intricate interactions between Trypanosoma brucei, the protozoan parasite causing African trypanosomiasis, and the tsetse (Glossina) vector remains a challenge. Metacyclic trypanosomes, which inhabit the tsetse salivary glands, transmit the disease and are produced through a complex differentiation and unknown program. By overexpressing a single RNA-binding protein, TdRBP6, in cultured noninfectious trypanosomes, we recapitulated the developmental stages that have been observed in tsetse, including the generation of infective metacyclic forms expressing the variant surface glycoprotein. Thus, events leading to acquisition of infectivity in the insect vector are now accessible to laboratory investigation, providing an opening for new intervention strategies.

Reproducing the life cycle of infectious organisms in vitro offers unique opportunities for studying pathogenesis and for developing intervention strategies. The protozoan parasite Trypanosoma brucei is the causative agent of sleeping sickness in humans, and the related disease “nagana” in livestock, which remain a public health challenge in sub-Saharan Africa. Human African trypanosomiasis is fatal if left untreated, and the livestock disease profoundly impacts on the economic development of affected countries. The blood-feeding tsetse fly (Glossina subspecies) vector transmits the disease, and its range determines the distribution of the disease. Bloodstream forms (BSFs) are taken up with the bloodstream, and in the fly midgut stumpy forms (SFs) differentiate into procyclic forms that are no longer infectious to mammals (1). Several morphological and metabolic changes take place during this developmental stage, including repositioning of the kinetoplast (the mitochondrion genome) midway between the posterior pole and the nucleus, elaboration of mitochondrion cristae, silencing of variable surface glycoprotein (VSG) expression, VSG coat loss, and expression of procyclin on the cell surface (1). Procyclics then move from the midgut to the proventriculus, the terminal portion of the foregut, where they elongate and become epimastigotes, with the kinetoplast positioned next to the nucleus in the anterior position. Epimastigotes undergo asymmetric division to produce a long epimastigote, which probably degenerates, and a short epimastigote, which instead goes on to colonize the salivary glands where it attaches to the epithelium. Attached epimastigotes divide and ultimately differentiate into mature nondividing metacyclics that detach from the epithelium and are found in the lumen of the salivary glands. During metacyclogenesis, the mitochondrion regresses to the single tubule characteristic of BSFs, indicating that mitochondrial function is inhibited; the VSG coat, synthesized from VSG genes in metacyclic-type transcription units (metacyclic expression sites), reappears; endocytosis increases; and the cells reacquire infectivity to mammals (1). Although the intricate nature of trypanosome development in the fly has been recognized for more than a century (2), the molecular mechanisms are still mysterious, due in part to experimental challenges of studying parasites in the fly.

RNA-binding proteins (RBPs) are important regulators of trypanosome gene expression as mediators of posttranscriptional control (3).

A survey of the T. brucei transcriptome in infected tsetse tissues by high-throughput RNA sequencing (RNA-Seq) revealed that the mRNA abundance for RBP6 (Tb9273.2930, containing a single RNA-recognition motif) is increased by a factor of 13 (relative to midgut-derived procyclics) in trypanosomes from the proventriculus (fig. S1). Experimentally induced RBP6 overexpression in cultured noninfectious procyclics resulted in the appearance of developmental stages that have been previously described in tsetse; namely, long and short epimastigotes and metacyclic trypanosomes (Fig. 1A and figs. S2 to S4). The prevalence of metacyclics depended on RBP6 expression levels (fig. S5), and we also observed asymmetrically dividing long epimastigotes (fig. S6).

Metacyclics generated in vitro featured a kinetoplast at the cell’s posterior pole (Fig. 1B and fig. S4), an undulating membrane (Fig. 1C), and BSF-like motility and, like BSFs, they did not adhere to glass. Metacyclic-like cells also displayed a range of mitochondrial architectures from multibranched to tubular (Fig. 1C and fig. S7), an indication of mitochondrial regression (4), which is distinctive of BSFs. Relative to procyclics and epimastigotes, metacyclics generated in vitro had higher levels of callflagin (Fig. 1B and fig. S8), mirroring results from salivary-gland metacyclics (5). Compared to procyclics and epimastigotes, metacyclics also showed higher internalization of the fluid-phase marker dextran 10,000 (fig. S9)—an indication of active endocytosis, which is a characteristic feature of BSFs (4). Up to 20% of the metacyclic-like cells could be purified by DEAE-cellulose chromatography, which combined with the gradation in mitochondrial regression (fig. S7) and callflagin expression (fig. S8), suggested that the cultures contained metacyclics at different stages of differentiation.

**Fig. 1. T. brucei RBP6 induces metacyclogenesis.** (A) Timeline for the appearance of different cell types upon induction of RBP6 expression (three independent experiments; error bars, SD). (B) Immunofluorescent detection of callflagin (K, kinetoplast; N, nucleus) in a culture induced to express RBP6 for 10 days (DIC, differential interference contrast). (C) Live-cell imaging of the mitochondrion in a metacyclic cell with Mitotracker Deep Red FM. (D) Transmission electron microscopy confirms the presence of a variant surface glycoprotein (VSG) coat. Shown is a transverse section across the flagellar pocket (FP), which, like bloodstream form trypanosomes, contains fibrous material (MT, mitochondrion; FL, flagellum; PM, plasma membrane; SPMT, subpellicular microtubules).

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The hallmark of metacyclogenesis is the re-acquisition of a VSG coat (1, 4). We detected the VSG coat by transmission electron microscopy (Fig. 1D) and the mRNA for several VSGs by reverse transcriptase–polymerase chain reaction (RT-PCR) and RNA-Seq (Fig. 2A and table S1). The VSG transcripts with highest expression were derived from monocistronic expression sites (Fig. 2B and fig. S10) with metacyclic-type Pol I promoters. Red rectangles, VSG genes; purple triangles, subtelomeric or telomeric repeats; yellow rectangles, ribosomal mobile element (RIME) sequences; gray rectangles, 70-base pair repeats; bent arrows, transcription start sites; red circles, Pol I promoters. (C) Alignment of the metacyclic Pol I promoters of expressed VSG genes to a previously characterized metacyclic promoter sequence of the MVAT4 gene (6). Bent arrow indicates the position of the putative transcription start site. Red and purple indicate >60% and <60% sequence conservation, respectively.

Fig. 2. T. brucei RBP6 induces the expression of monocistronic VSG genes. (A) RT-PCR detection of expressed VSG genes in cultures containing metacyclic cells. Ethidium bromide–stained agarose gel for samples processed with (+RT) or without (−RT) reverse transcriptase are shown. The identity of amplified products was confirmed by cloning and sequencing. (B) Diagrams of the genomic context of the VSG genes with metacyclic-type Pol I promoters. Red rectangles, VSG genes; purple triangles, subtelomeric or telomeric repeats; yellow rectangles, ribosomal mobile element (RIME) sequences; gray rectangles, 70–base pair repeats; bent arrows, transcription start sites; red circles, Pol I promoters. (C) Alignment of the metacyclic Pol I promoters of expressed VSG genes to a previously characterized metacyclic promoter sequence of the MVAT4 gene (6). Bent arrow indicates the position of the putative transcription start site. Red and purple indicate >60% and <60% sequence conservation, respectively.

The zebrafish regenerates its brain after injury and hence is a useful model organism to study the mechanisms enabling regenerative neurogenesis, which is poorly manifested in mammals. Yet the signaling mechanisms initiating such a regenerative response in fish are unknown. Using cerebroventricular microinjection of immunogenic particles and immunosuppression assays, we showed that inflammation is required and sufficient for enhancing the proliferation of neural progenitors and subsequent neurogenesis by activating injury-induced molecular programs that can be observed after traumatic brain injury. We also identified cysteinyl leukotriene signaling as an essential component of inflammation in the regenerative process of the adult zebrafish brain. Thus, our results demonstrate that in zebrafish, in contrast to mammals, inflammation is a positive regulator of neuronal regeneration in the central nervous system.

After a traumatic brain injury in the telencephalon, zebrafish can efficiently restore the tissue architecture and replace the lost neurons upon neurogenic activity of the ventricularly located radial glial cells (1–4). However, the signaling mechanisms involved in the activation of those stem cell niches after injury (reactive proliferation) and the production of new neurons (regenerative neurogenesis) from neurogenic progenitors are largely unknown.

Immune cell activation is among the first responses detected in the tissue after a severe central nervous system injury in both mammals and zebrafish (5–9). Various types of immune cells from the bloodstream (leukocytes) or resident in the brain tissue (microglia) accumulate in the injured tissue, secrete cytokines and chemokines to modulate the environment, and are responsible for the removal of cell debris and accumulated metabolites, as has been shown in mammals (10). Acute neuroinflammation in mammals has generally been considered to have a negative effect on neurogenesis and regeneration by promoting the formation of a glial scar and hampering the proliferation of the precursor cells and the migration, survival, maturation, and integration of the newborn neurons into the existing circuitry (11–15). In contrast, the inflammatory response seems to have different consequences in zebrafish, where adult neurogenesis can be beneficial, because it is a regenerative species (16). Here, we show that inflammation is required and sufficient for enhancing the proliferation of neural progenitors and subsequent neurogenesis by activating injury-induced molecular programs that can be observed after traumatic brain injury. We also identified cysteinyl leukotriene signaling as an essential component of inflammation in the regenerative process of the adult zebrafish brain. Thus, our results demonstrate that in zebrafish, in contrast to mammals, inflammation is a positive regulator of neuronal regeneration in the central nervous system.

References and Notes

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Supplementary Materials
www.sciencemag.org/cgi/content/full/338/6112/1352/DC1
Materials and Methods
Figs. S1 to S11
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References (7–10)
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Acute Inflammation Initiates the Regenerative Response in the Adult Zebrafish Brain

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The zebrafish regenerates its brain after injury and hence is a useful model organism to study the mechanisms enabling regenerative neurogenesis, which is poorly manifested in mammals. Yet the signaling mechanisms initiating such a regenerative response in fish are unknown. Using cerebroventricular microinjection of immunogenic particles and immunosuppression assays, we showed that inflammation is required and sufficient for enhancing the proliferation of neural progenitors and subsequent neurogenesis by activating injury-induced molecular programs that can be observed after traumatic brain injury. We also identified cysteinyl leukotriene signaling as an essential component of inflammation in the regenerative process of the adult zebrafish brain. Thus, our results demonstrate that in zebrafish, in contrast to mammals, inflammation is a positive regulator of neuronal regeneration in the central nervous system.