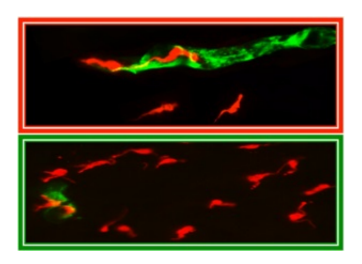


Nitric oxide protects against parasite invasion and brain inflammation by keeping the blood brain barrier intact

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Enhanced invasion of *Trypanosoma brucei* (red) into the brain of iNOS deficient (lower micrograph) is compared to normal (upper micrrograph) mice. In green: cerebral blood vessels. Credit: Olivera et al.

African trypanosomiasis is called 'sleeping sickness' because when the infection is untreated, trypanosome parasites will invade the brain and cause disruption of sleeping patterns and irreversible neurological damage. A study published on February 25th in *PLOS Pathogens* reports that in a mouse model of trypanosome disease, nitric oxide (NO) plays an unexpected role in preserving the integrity of the blood brain barrier (BBB), thereby reducing parasite invasion into the brain, and likely limiting neurological damage.



NO is generally thought to be a pro-inflammatory signal, promoting a strong immune response against pathogens. The resulting inflammation is a mixed blessing: on one hand, it helps to control potentially dangerous pathogens, but on the other, it can cause "collateral damage" to the inflamed tissue.

Martin Rottenberg and colleagues, from the Karolinska Institute in Stockholm, Sweden, are interested in how trypanosome <u>parasites</u> cause disease and in the host immune defense against them. In this study, they examine the role of NO in a <u>mouse model</u> of trypanosomiasis, with a focus on how the parasites manage to get through the so-called <u>blood</u> <u>brain barrier</u> (BBB), the border surrounding the mammalian <u>brain</u> that is normally impenetrable to foreign intruders as well as most host cells.

To their surprise, they found that when they infected mice that lack an enzyme called iNOS, which produces NO in response to an infection, with trypanosomes, the mutant mice had elevated numbers of parasites and host <u>immune cells</u> in the brain. The cause, they determined, was an increased permeability of the blood vessels that are part of the BBB.

When the researchers looked at iNOS levels in response to trypanosome infection in normal mice, they found that infection is followed by an increase in iNOS levels in immune cells called macrophages in and near the brain, which in turn produces more local NO. Examining the underlying mechanisms, the researchers found that a molecule called TNF mediates the increased permeability to proteins, leukocytes and parasites in the brain of mice lacking iNOS. NO inhibited both TNF production by macrophages and its inflammatory function.

TNF itself, they could show, acts through a molecule called MMP-9 whose function is known to affect cell motility and tissue invasion. Consistent with a key role of MMP-9, the researchers found that minocycline, an antibiotic drug related to tetracycline, can restore the



function of the BBB in mice lacking iNOS, and reduce the numbers of parasites and host inflammatory cells in their brains.

"In the present study", the researchers summarize, "we describe a protective, anti-inflammatory role of NO in the pathogenesis of an experimental neuroinflammatory disease, i.e. African trypanosomiasis. In particular, iNOS produced NO was critical for maintaining the integrity of the BBB, hampering continuous brain invasion of parasites and leukocytes. This was unexpected, since release of iNOS-derived NO into the brain often has been associated with toxicity both of the BBB and the nervous tissue."

More information: *PLOS Pathogens*, dx.plos.org/10.1371/journal.ppat.1005442

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