

## It's complicated: Benefits and toxicity of antiprion antibodies in the brain

January 28 2016







A 3-D surface representation of the prion protein. The areas rendered in color illustrate the interaction with the two neurotoxic antibodies, POM1 and ICSM18 Credit: Reimann et al.

Immunotherapy to ameliorate neurodegeneration by targeting brain protein aggregates with antibodies is an area of intense investigation. A study published on January 28th in *PLOS Pathogens* examines seemingly contradictory earlier results of targeting the prion protein and proposes a cautionary way forward to further test related therapeutic approaches.

Damaging aggregation of proteins in the brain is a hallmark of several neurodegenerative diseases, including Alzheimer and Parkinson diseases. In prion diseases such as Creutzfeld-Jakob disease, aggregates of abnormally folded versions of the PrP protein cause the death of brain cells. The use of anti-PrP antibodies to target such aggregates and mediate their destruction by the immune system has been proposed as a therapy, with the caveat that such antibodies could themselves be toxic.

Prompted by the publication of seemingly contradictory reports on the potential neurotoxicity of antibodies against the <u>prion protein</u> in mice, Adriano Aguzzi from the University of Zurich, Switzerland, and colleagues set out to rigorously test several antibodies side-by-side. The researchers examined whether toxicity was influenced by where and how the antibodies were delivered into the brain, the antibody dose, which area on the PrP protein the antibodies bind to, and the genetic background of the mice.

For several of the antibodies, they found, toxicity depended strictly on the administered dose, results that could explain some of the seemingly



different earlier results. Other factors tested, including which target regions (or epitopes) on the PrP protein the antibodies bound to, also seemed to affect the toxicity of at least some antibodies.

Because the researchers observed toxicity for most of the antibodies that bind to the so-called globular region of PrP, they conclude that such antibodies might not be suitable for immunotherapy. In contrast, since no toxicity was found in tests of antibodies against the flexible tail region of PrP, they suggest that those might be more promising candidates.

"In summary", they say, "these data illustrate that the efficacy profile (i.e. the curative effectiveness versus the potential <u>toxicity</u>) of anti-prion <u>antibodies</u> is complex and depends both on intrinsic factors such as, crucially, the nature of the engaged epitope, and extrinsic factors such as the route of administration. Detailed analyses and mapping of the involved epitopes and—most importantly—appropriate dose-escalation studies in vivo are prerequisite not only for preparing clinical trials in humans, but also to avoid the reporting of contradictory, confusing, and potentially misleading results."

**More information:** Reimann RR, Sonati T, Hornemann S, Herrmann US, Arand M, Hawke S, et al. (2016) Differential Toxicity of Antibodies to the Prion Protein. *PLoS Pathog* 12(1): e1005401. DOI: 10.1371/journal.ppat.1005401

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