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D-PEPTIDES DEVELOPED TO BE THERAPEUTICALLY ACTIVE AGAINST BETA-AMYLOID OLIGOMERS SHOW PROMISING PHARMACOKINETIC PROPERTIES

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Methods: Radioactively labelled peptide was administered via several administration routes and organs were harvested at different time points post injection. The amount of radioactive D-peptide in the organ homogenate was measured by liquid scintillation counting. Furthermore, binding to plasma proteins as well as brain membranes was determined, also using radioactively labelled peptide as indicator.

Results: Results show that all D-peptides indeed reach the brain where they may exhibit their therapeutic activity. Furthermore, the peptides show small elimination constants and long half-lives of more than a day in plasma as well as a high bioavailability after i.p., s.c. or p.o. administration.

Conclusions: Promising pharmacokinetic properties confirm that D-peptides may be very potent AD-therapeutic agents on their way to clinical studies.