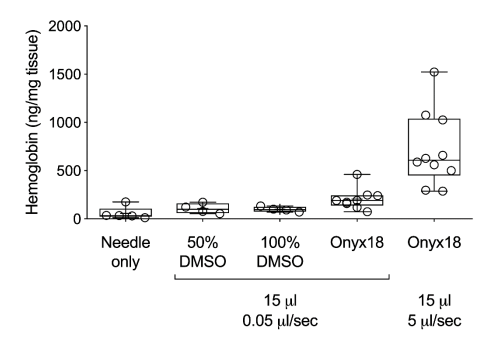
SUPPLEMENTAL MATERIAL



Supplemental Figure I: Hemoglobin concentrations after 15 μ I injections of 50% or 100% DMSO, and comparison to liquid polymer.

Supplemental Video 1: 3D reconstruction of images taken after mouse brains were cut in a cryostat after injection of 15 µl liquid polymer. Video 1A shows secondary hemorrhage in a non-anticoagulated mouse while Video 1B shows a mouse brain after warfarin anticoagulation. The liquid polymer is depicted in yellow, the secondary bleeding in red.

* Preclinical Checklist Preclinical Checklist: Prevention of bias is important for experimental cardiovascular research. This short checklist must be completed, and the answers should be clearly presented in the manuscript. The checklist will be used by reviewers and editors and it will be published. See <u>"Reporting Standard for Preclinical Studies of Stroke Therapy"</u> and <u>"Good Laboratory Practice: Preventing Introduction of Bias at the Bench"</u> for more information.	
This study invovles animal models: Yes	
Experimental groups and study timeline	
The experimental group(s) have been clearly defined in the article, including number of animals in each experimental arm of the study:	Yes
An account of the control group is provided, and number of animals in the control group has been reported. If no controls were used, the rationale has been stated:	Yes
An overall study timeline is provided:	Yes
Inclusion and exclusion criteria	
A priori inclusion and exclusion criteria for tested animals were defined and have been reported in the article:	Yes
Randomization	
Animals were randomly assigned to the experimental groups. If the work being submitted does not contain multiple experimental groups, or if random assignment was not used, adequate explanations have been provided:	Yes
Type and methods of randomization have been described:	Yes
Methods used for allocation concealment have been reported:	N/A
Blinding	
Blinding procedures have been described with regard to masking of group/treatment assignment from the experimenter. The rationale for nonblinding of the experimenter has been provided, if such was not feasible:	Yes
Blinding procedures have been described with regard to masking of group assignment during outcome assessment:	Yes
Sample size and power calculations	
Formal sample size and power calculations were conducted based on a priori determined outcome(s) and treatment effect, and the data have been reported. A formal size assessment was not conducted and a rationale has been provided:	No
Data reporting and statistical methods	
Number of animals in each group: randomized, tested, lost to follow-up, or died have been reported. If the experimentation involves repeated measurements, the number of animals assessed at each time point is provided, for all experimental groups:	Yes
Baseline data on assessed outcome(s) for all experimental groups have been reported:	N/A
Details on important adverse events and death of animals during the course of experimentation have been provided, for all experimental arms:	Yes
Statistical methods used have been reported:	Yes
Numeric data on outcomes have been provided in text, or in a tabular format with the main article or as supplementary tables, in addition to the figures:	Yes
Experimental details, ethics, and funding statements	
Details on experimentation including stroke model, formulation and dosage of therapeutic agent, site and route of administration, use of anesthesia and analgesia, temperature control during experimentation, and postprocedural monitoring have been described:	Yes

Different sex animals have been used. If not, the reason/justification is provided:

Statements on approval by ethics boards and ethical conduct of studies have been provided:	Yes
Statements on funding and conflicts of interests have been provided:	Yes

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