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Computation of polymer in-situ rheology

² using direct numerical simulation

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- 8 Abstract
- 9 In-situ rheology of synthetic polymers like hydrolyzed polyacrylamide, HPAM, cannot be directly
- 10 calculated from bulk rheology. The in-situ viscosity of polymers will be influenced by shear history and
- 11 local pore structure contraction / expansion. Transition from flux in porous media to bulk shear rate
- 12 requires empirical correlations. The conventional approach for in-situ rheology is to define apparent
- 13 viscosity vs. Darcy velocity and connecting Darcy velocity to bulk shear rate by using a correction factor,
- 14 which generally depends on both polymer and rock properties.
- 15 Estimating polymer in-situ rheology is important for estimation of displacement pressure gradients and
- 16 injectivity in the reservoir. In the following study, different models used to quantify the shear-to-flux
- 17 correction factor are compared by using direct numerical simulation in 3D real rock images. We found
- that the models developed based on capillary bundle approach are not able to predict accurately in-situ
- 19 rheology and microstructures of a rock sample can influence in-situ rheology. A novel approach is
- 20 suggested to provide a more accurate prediction of in-situ rheology by adjusting appropriately bulk power
- 21 index of the Carreau model before applying the correction factor. The pore scale simulations have
- 22 revealed that the key parameters of porous media influencing in-situ rheology are pore aspect ratio and
- 23 inaccessible pore volume.
- 24 Keywords: Pore scale modeling, apparent viscosity, polymer rheology µCT rock image, Direct Numerical
- 25 Simulation.

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