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Finite element simulations of the bending of an IPMC sheet

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Background

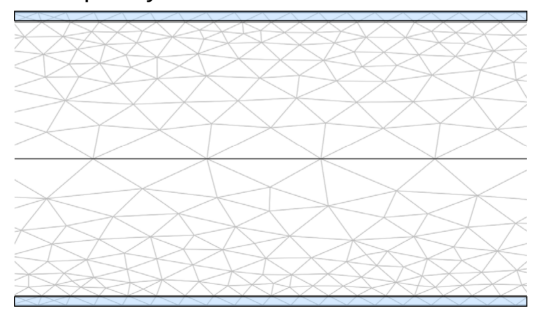


- IPMC ionic polymer-metal composites are highly porous polymer materials, such as Nafion, filled with some kind of ionic conductive liquid.
- The sheet of polymer material is coated with a thin metal layer usually platinum or gold.
- In electric field, freely movable ions inside the polymer migrate towards an electrode, causing expansion of material at the one end and contraction at the other end.
- FEM Finite Element Method
 - Used as a technic of modeling in this work

Introduction



- The goal: a time dependent FEM model for an IPMC
- Domains to be considered:
 - Mechanical
 - Electrostatic
 - Mass transfer
- The designed model is 2D
- Adding more complexity electrochemical reactions



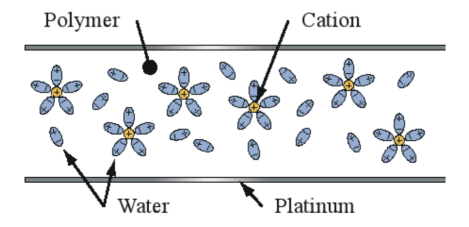
Simulations – cation migration



- Nernst-Planck equation:
 - Diffusion
 - Migration
 - Convection

$$\frac{\partial C_i}{\partial t} + \nabla \cdot \left(-D_i \nabla C_i - z_i \mu_i C_i \nabla \varphi \right) = -\vec{u}_i \cdot \nabla C_i$$

- Voltage causes migration of cations towards cathode
 - That's also the cause of the current in the circuit
- All anions are fixed to the polymer backbone



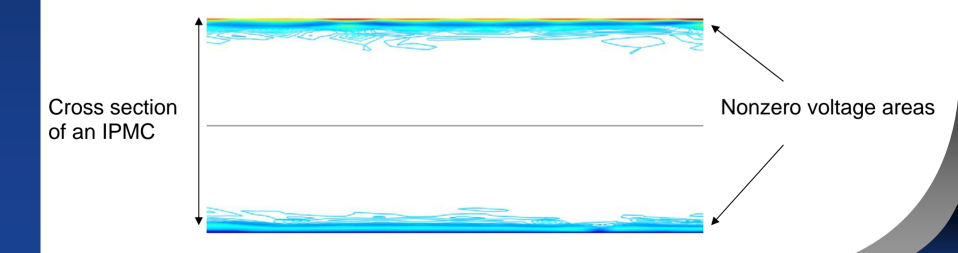
Simulations – cation migration (2)



- Charge imbalance inside the IPMC results in electric field
 - opposite to the applied field
- Gauss' law:

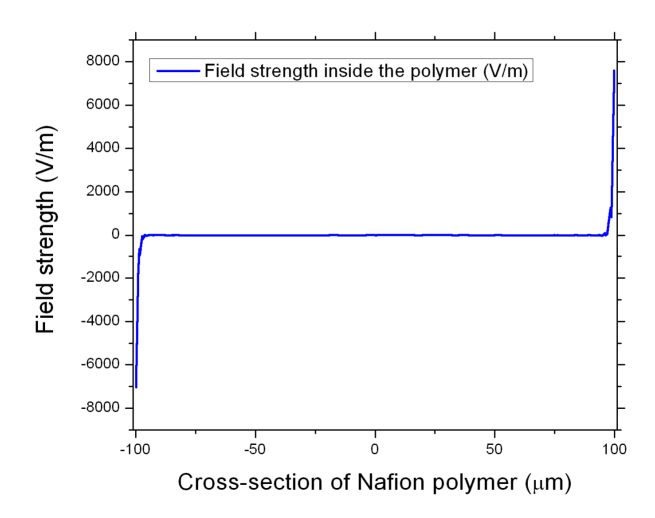
$$\nabla \vec{E} = \frac{F \cdot \rho}{\varepsilon}$$

 Steady state of cations – when the generated field cancels out applied field.



Simulations – cation migration (3)





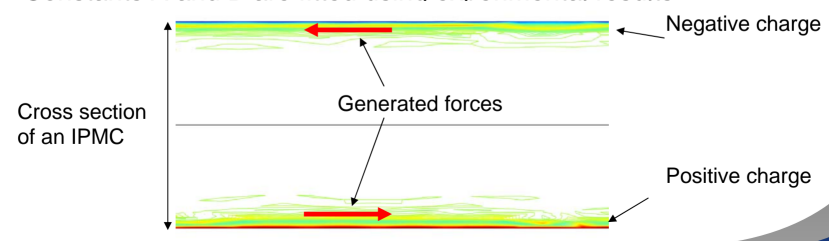
Actuation of an IPMC



- FEM model a bit different approach
 - Euler beam model is not used
 - Our model is dynamic, not static
 - Viscoelasticity is also considered
- Assumption charge imbalance creates longitudinal force inside the IPMC

$$\vec{F} = (A \cdot \rho + sgn(\rho) \cdot B \cdot \rho^2) \cdot \hat{x}$$

- We consider two terms charge density and square of the charge density
- Constants A and B are fitted using experimental results



Mathematics behind the bending



- Continuum mechanics equations are used in the model
- The stress is related to the force in each point of the polymer:

$$-\nabla \cdot \sigma = \vec{F}$$

The stress is related to strain:

$$\sigma = D\varepsilon$$

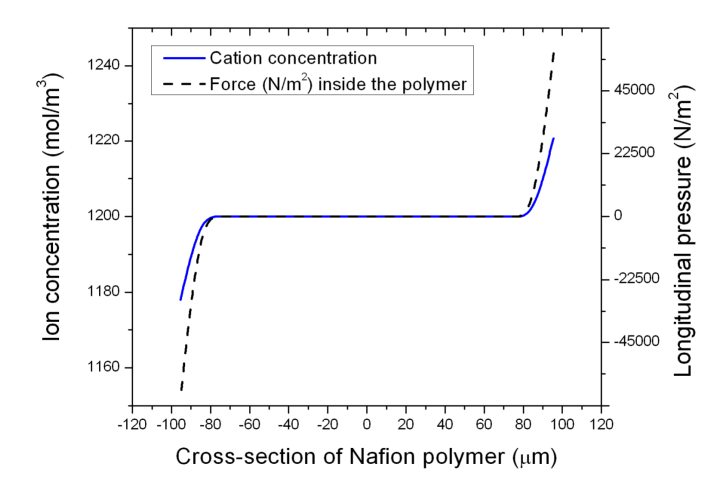
where ε and σ are column vectors with dimensions of 6

Finally, the strain is related to displacement:

$$\varepsilon_{i} = \frac{\partial u_{i}}{\partial x_{i}}, \varepsilon_{ij} = \frac{1}{2} \left(\frac{\partial u_{i}}{\partial x_{j}} - \frac{\partial u_{j}}{\partial x_{i}} \right)$$

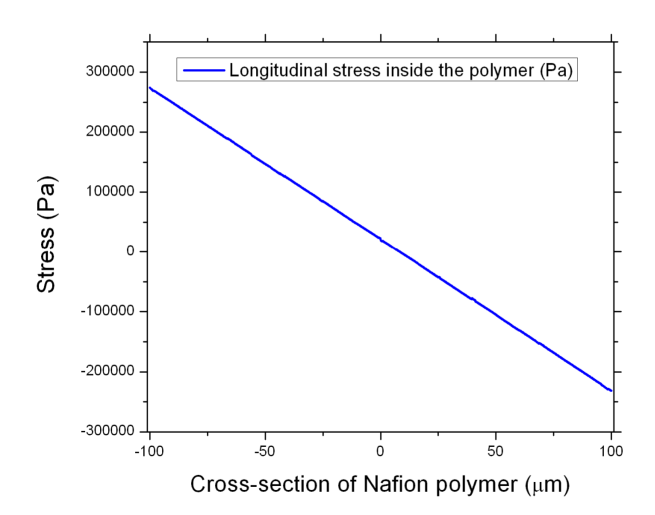
Simulations – cation migration, force





Simulations – generated stress





Transient analysis of bending



- Time dependent equations -> dynamic problem
- Motion of an IPMC sheet is described by:

$$\rho \frac{\partial^2 \vec{u}}{\partial t^2} - \nabla \cdot c \nabla \vec{u} = \vec{F}$$

Rayleigh damping model:

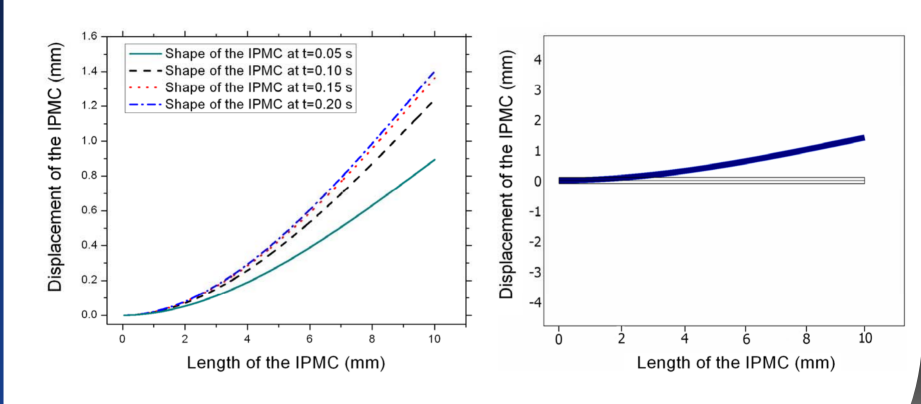
$$m\frac{d^{2}u}{dt^{2}} + \xi \frac{du}{dt} + ku = f(t) \qquad \qquad \xi = \underline{\alpha}m + \underline{\beta}k$$

 \bullet ξ is a damping parameter. m is mass and k is stiffness. By coupling those equations for the system of multiple degrees of freedom:

$$\rho \frac{\partial^{2} \vec{u}}{\partial t^{2}} - \nabla \cdot \left[c \nabla \vec{u} + c \underline{\beta} \nabla \frac{\partial \vec{u}}{\partial t} \right] + \underline{\alpha} \rho \frac{\partial \vec{u}}{\partial t} = \vec{F}$$

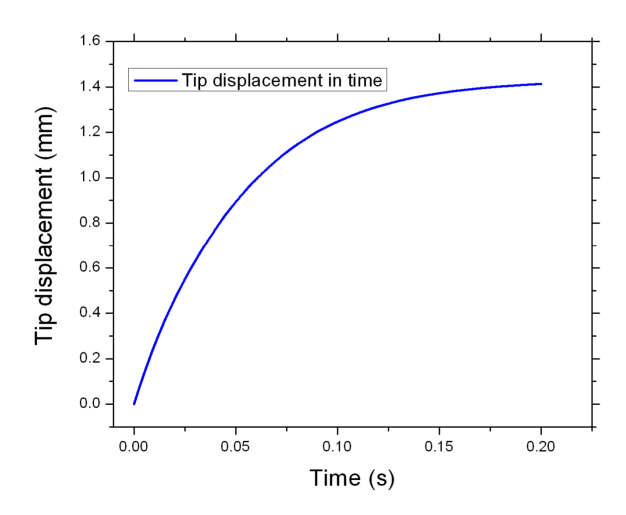
Simulation results





Tip displacement in time

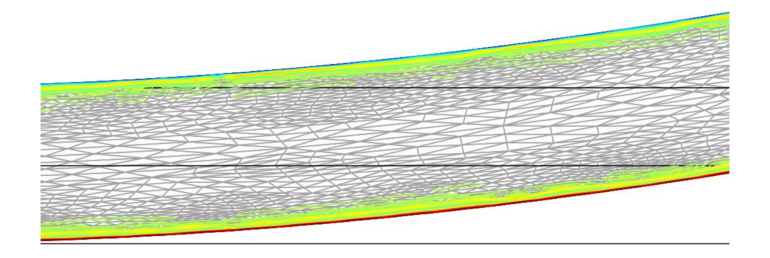




Conclusions



- We have a finite elment model, which...
 - describes the movement of ions inside the polymer
 - describes the electric field
 - describes the dynamic bending of an IPMC sheet

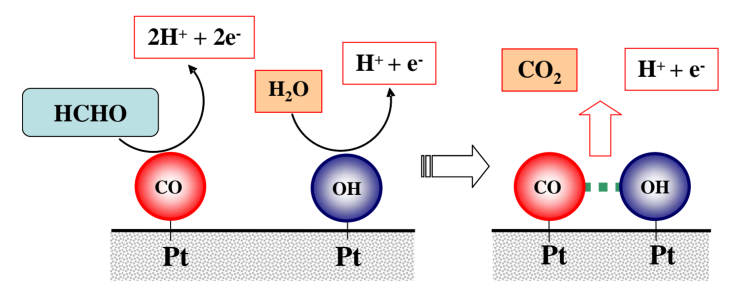


For next, we add some complexity – electrochemical oscillations

Self-oscillations



- Electrochemical oscillations common phenomena in nature
- Oscillations have also been experimentally studied
- Tests with IPMC:
 - immersed into acidic formaldehyde (HCHO) solution
 - applied constant voltage 0.75V
 - result: current oscillations and oscillating bending
- Cause of the oscillations are electrochemical reactions on Pt surface



Simulating self-oscillations



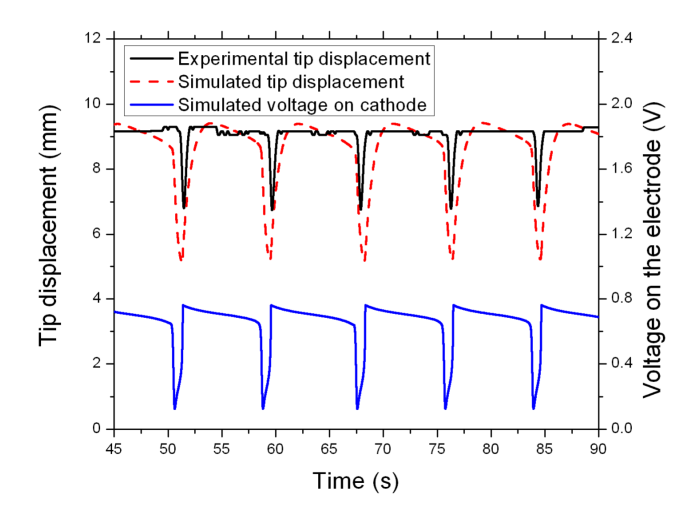
- A double layer were initialized near the platinum surface.
- On that suface, the following reaction is modeled:

$$HCHO + * \rightarrow Pt - CHO_{ads} + H^+ + e^-$$

- 4 dynamic variables must be modeled on the platinum surface
 - time dependence of adsorption coverage of CO
 - time dependence of adsoprtion coverage of OH
 - HCHO concentration change in double layer
 - time dependence of potential on cathode
- These equations were coupled to the basic model...

Self-oscillations – model and experiment

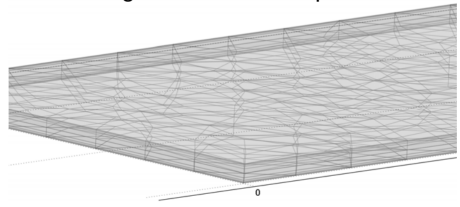




Conclusions and future plans



- 2D, basic FEM model was developed to describe time dependent actuation of an IPMC muscle
- In addition, more complexity was added by coupling a problem of electrochemistry.
- Also experimental and simulatios results were compared and quite good agreement were found
- For future:
 - Adding 3rd dimension to the model
 - Considering fluid viscosity and corresponding equations inside the polymer
 - Work on the simulating the oscillation equations



Questions



Thank you for your attention!