

Modelling high energy density supercapacitors by molecular dynamics simulations

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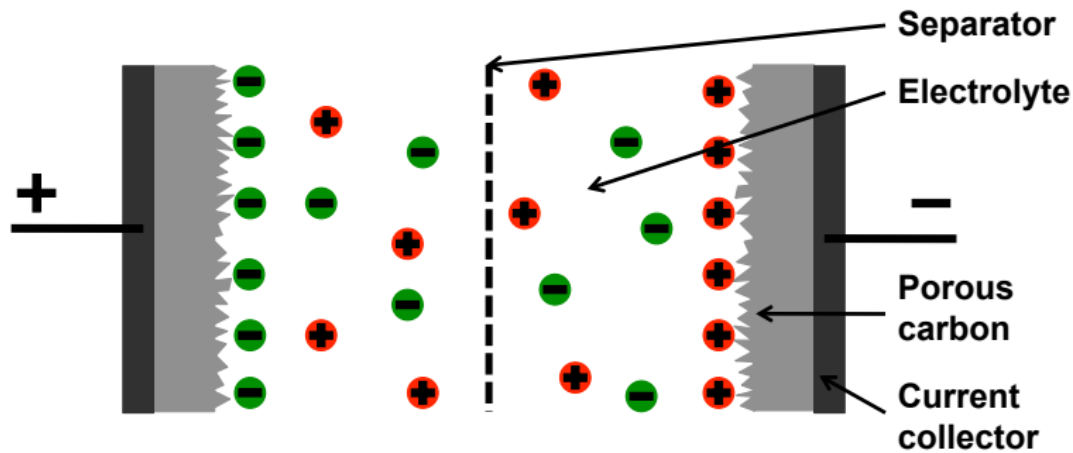
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Network on the electrochemical storage of energy (RS2E)
<http://www.energie-rs2e.com>

Forum Teratec - July 2nd, 2014

Outline

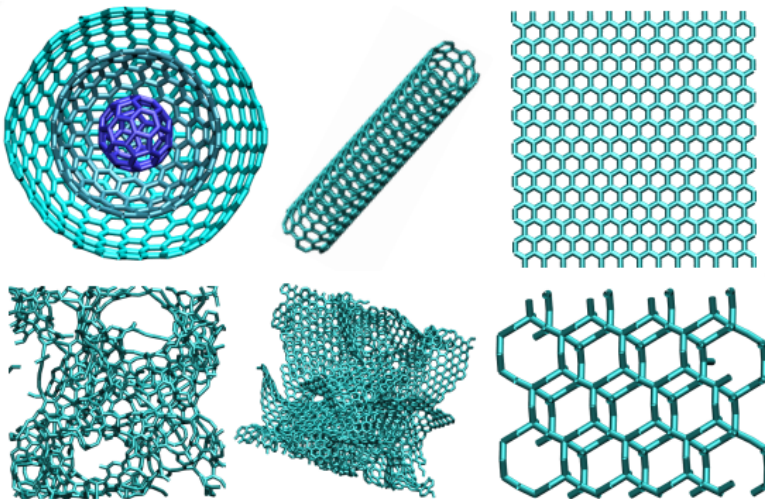
- 1 Supercapacitors
- 2 Molecular simulations
- 3 Molecular origin of supercapacitance
- 4 Conclusion

Supercapacitors: electricity storage devices



- Also called Electrochemical Double Layer Capacitors
- Charge stored through adsorption of ions at the **surface** of electrode
- No redox reaction in the bulk material → different from batteries
- $E = \frac{CU^2}{2}$ & $P = \frac{U}{4R}$

Electrode materials



- Commercial devices: carbon materials
- 1D, 2D, 3D materials
- Various porosities, surface area, chemical activation
- Efficiency measured with the **capacitance**

Simon & Gogotsi, *Nature Materials* 7, 845 (2008)

Electrolytes

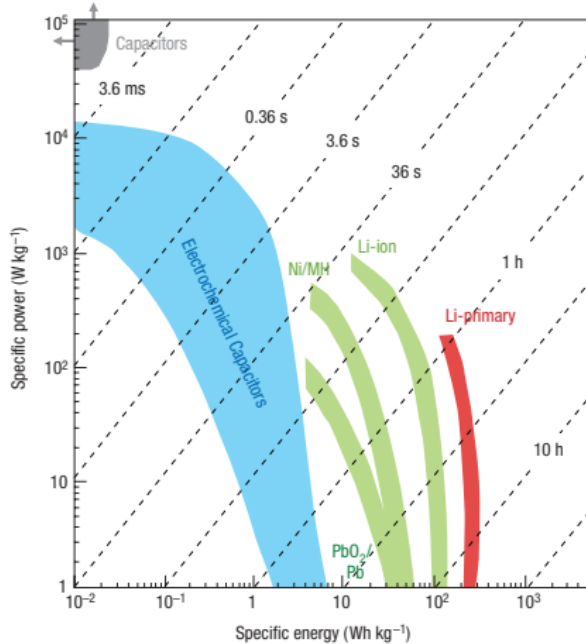
Need to find a good compromise between:

- Large electrochemical window
- High conductivity
- Good adsorption of ions on (porous) carbon materials
- Toxicity, safety...

Depending on the application:

- Ionic liquids
- Organic electrolytes (acetonitrile + organic salts)
- Aqueous electrolytes (inorganic salts)
- Solid electrolytes (ionic liquids + polymers)

Supercapacitors in the electricity storage landscape



Intermediate performances between conventional capacitors and Li-ion batteries.

- High specific power
- Correct specific energy
- Rapid charge/discharge (a few seconds)
- High cyclability (1 million cycles)

Applications



Alstom



Collaboration Alstom / Batscap



Tramway in Paris: braking energy is stored in a supercap (allows for a traction of more than 100 m)

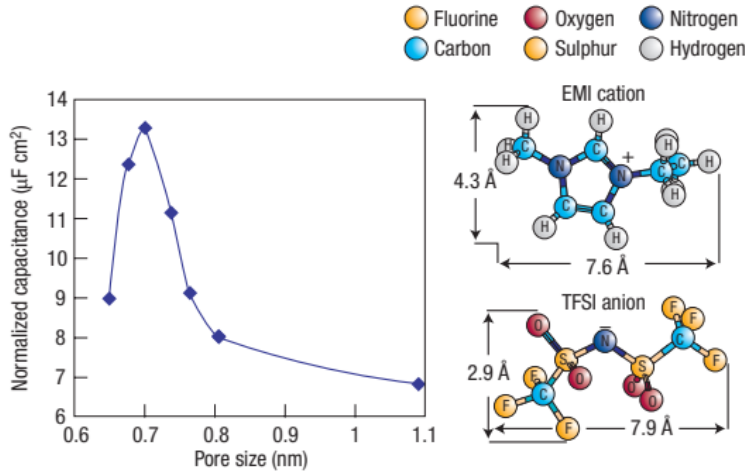
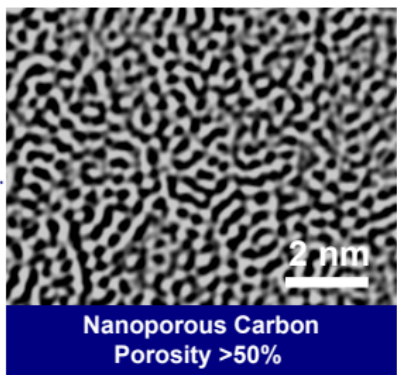
Applications



- Peugeot 308 e-HDI: 5V supercapacitor
- Stop-start system → lower fuel consumption (-15 %)
- Also in Formula One: KERS

Experimental discovery (Gogotsi & Simon)

Carbide Derived Carbon



- CDC: narrow pore size distribution
- Increase of the capacitance by +50 %!
- Optimum when pore size = ion size

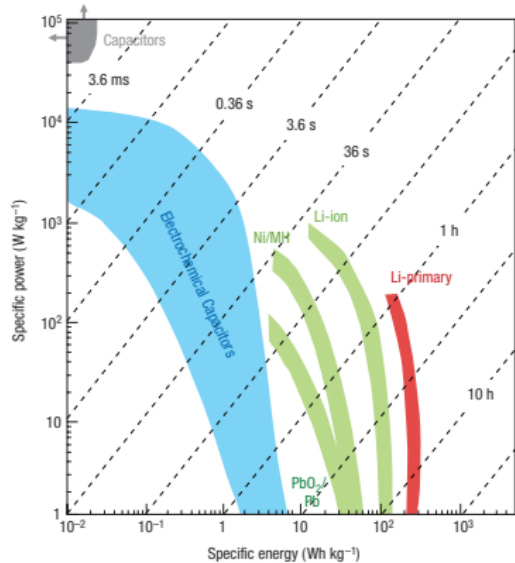
Chmiola *et al.*, *Science*, 313, 1760 (2006)

Largeot *et al.*, *J. Am. Chem. Soc.*, 130, 2730 (2008)

Challenges

Need for improvements:

- Increase of capacitance
→ design of new carbon materials
- Electrolytes with high potential window
- Electrolytes with high conductivity
at low temperature (-40°C)

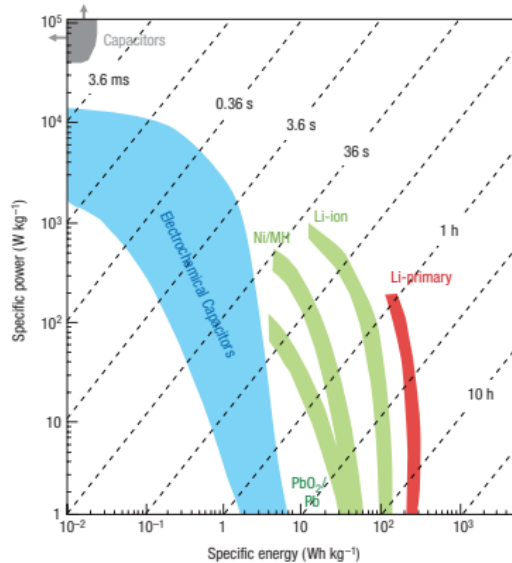


Much research work to be done in **materials** and **electrochemistry**, but...

Challenges

Need for improvements:

- Increase of capacitance
→ design of new carbon materials
- Electrolytes with high potential window
- Electrolytes with high conductivity at low temperature (-40°C)



Much research work to be done in **materials** and **electrochemistry**, but...

- Local structure of the liquid inside the nanopores is unknown
- Very difficult to probe it experimentally

→ **Simulation can help!**

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Simulation method: molecular dynamics

- System of N classical atoms/molecules interacting together
- Periodic boundary conditions
- Iterative integration of Newton's equation

$$m^i \frac{\partial^2 \vec{r}^i}{\partial t^2} = \sum_{j \neq i} \vec{F}^{j \rightarrow i} = -\frac{\partial V}{\partial \vec{r}^i}$$

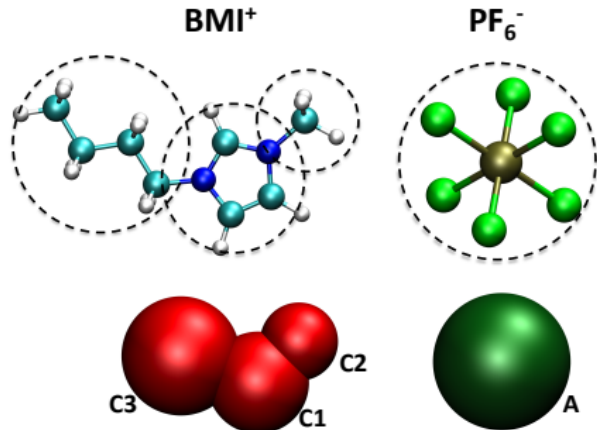
where V is the **interaction potential**

- Numerical resolution
- Trajectory of the atoms over several nanoseconds
- Determination of **structural**, **thermodynamic** and **transport** properties

Electrolyte: coarse-grained model

BMI⁺: 1-butyl-3-methylimidazolium
→ 3 sites

PF₆⁻: hexafluorophosphate
→ 1 site



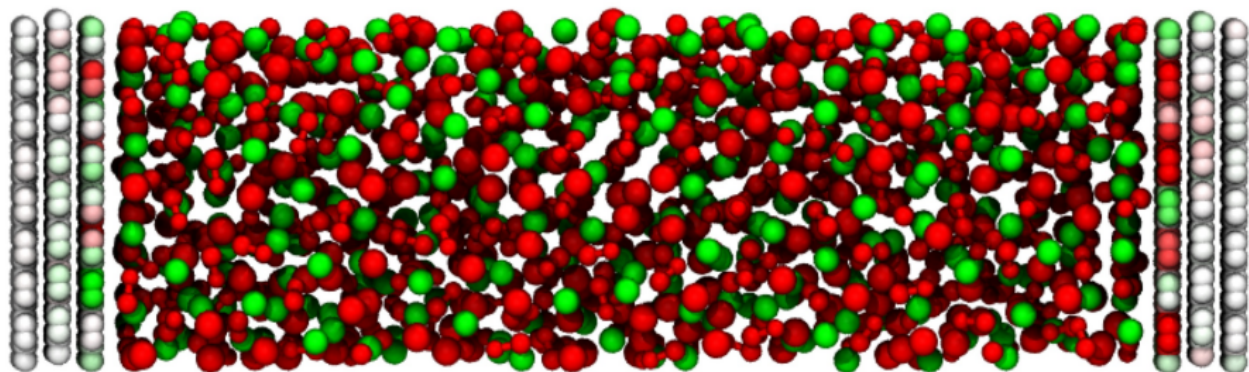
Interaction potential:

$$V = \sum_{i,j>i} \left[4\epsilon_{ij} \left(\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^6 \right) + \frac{q_i q_j}{r_{ij}} \right]$$

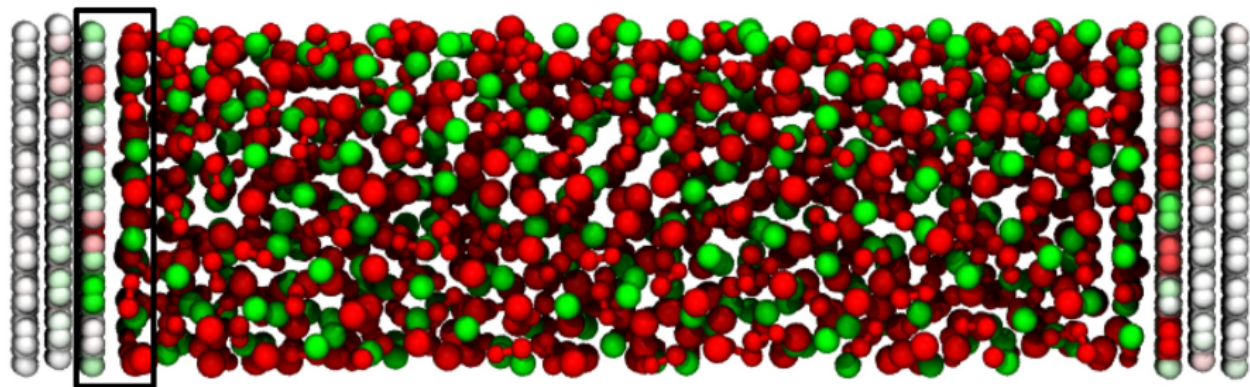
Roy & Maroncelli, *J. Phys. Chem. B* **114**, 12629 (2010)

Merlet, Salanne & Rotenberg, *J. Phys. Chem. C* **116**, 7687 (2012)

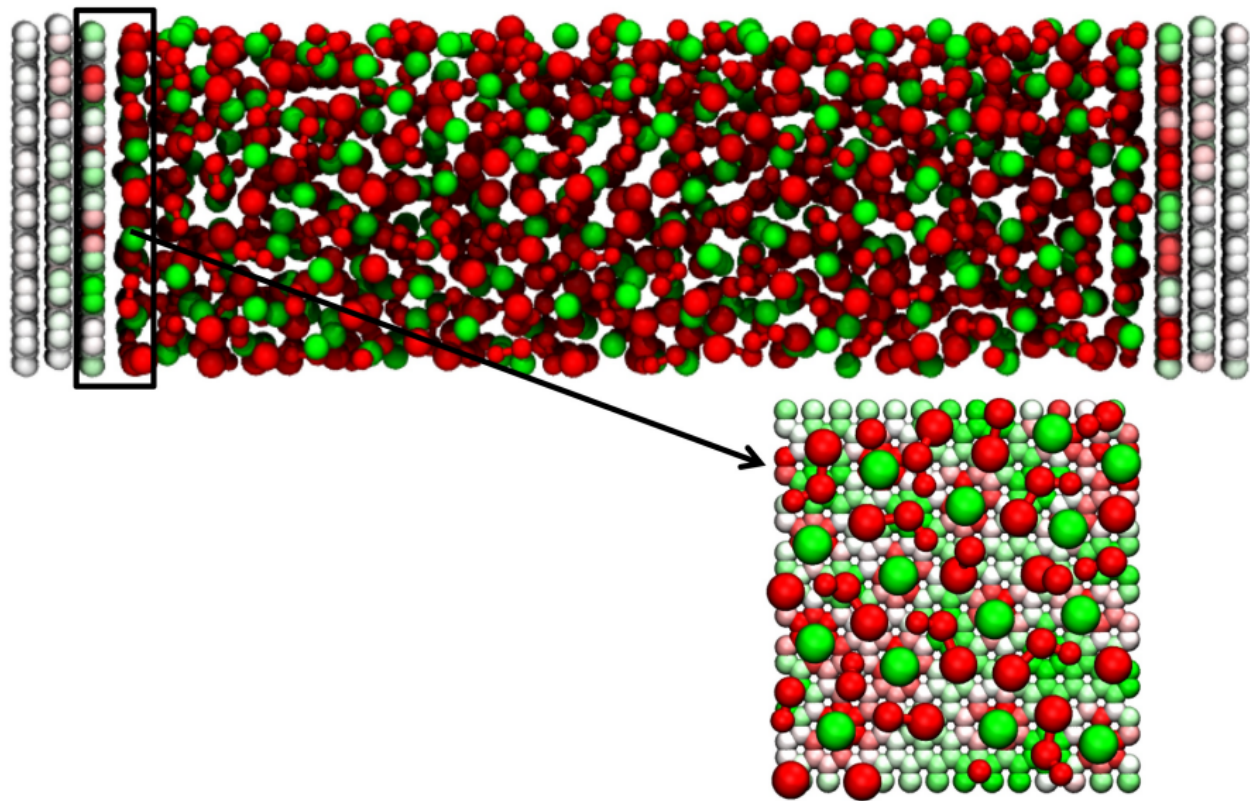
Simulating constant voltage electrodes



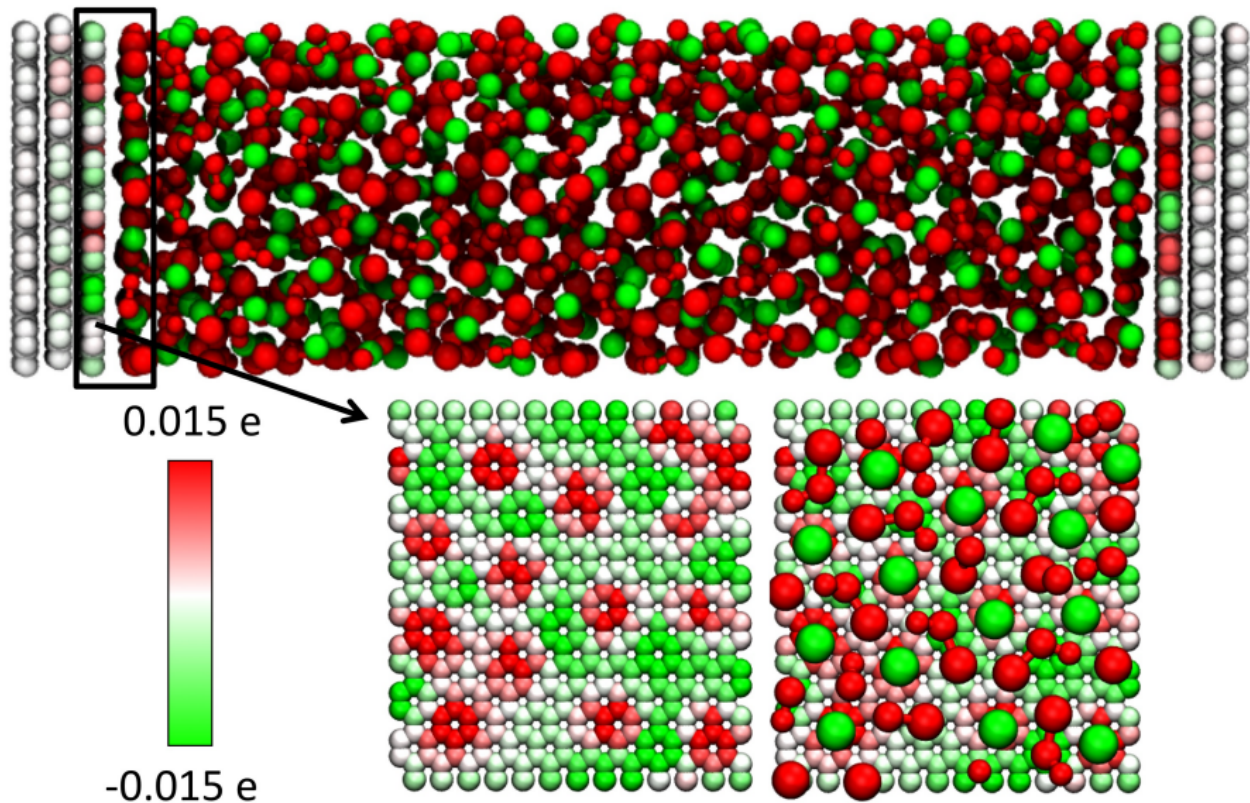
Simulating constant voltage electrodes



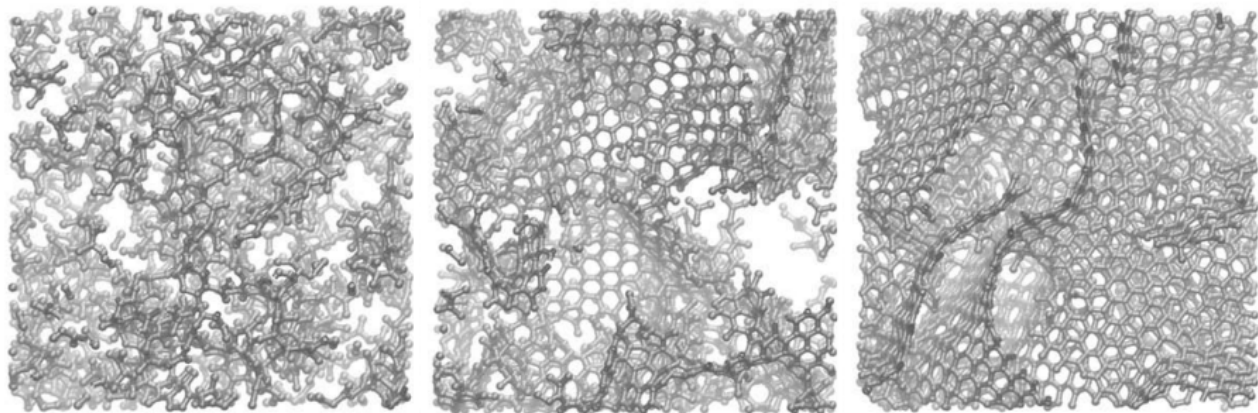
Simulating constant voltage electrodes



Simulating constant voltage electrodes



Porous electrodes: CDC model structures



- Obtained from quenching a liquid carbon (ReaxFF)
- Different quenching rates: changes in the pore shape and size distribution
- Mimics CDC structures obtained at different temperatures

Example of trajectory

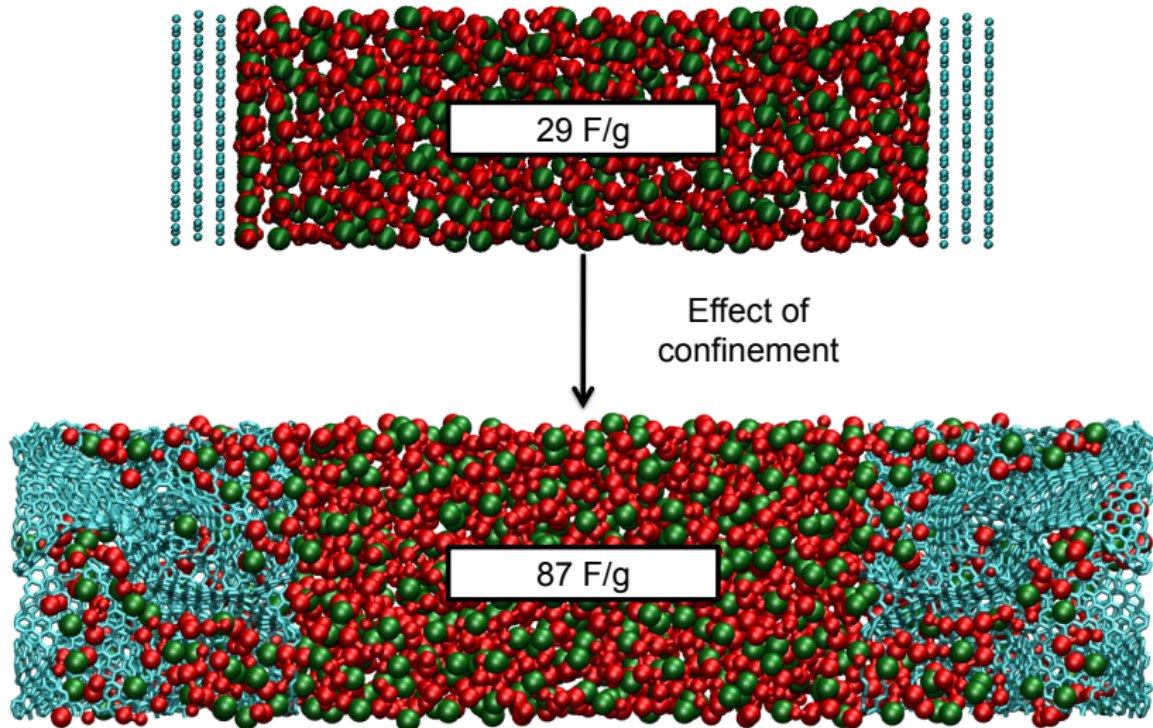
Loading movie

Simulation cell with 2-Dimensional periodic boundary conditions

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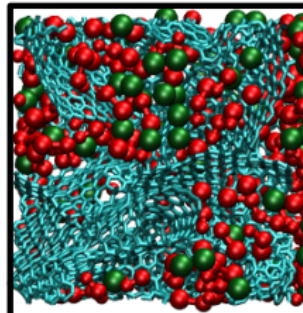
Increase of the capacitance in nanoporous carbons



Merlet et al., *Nature Materials*, 11, 306 (2012)

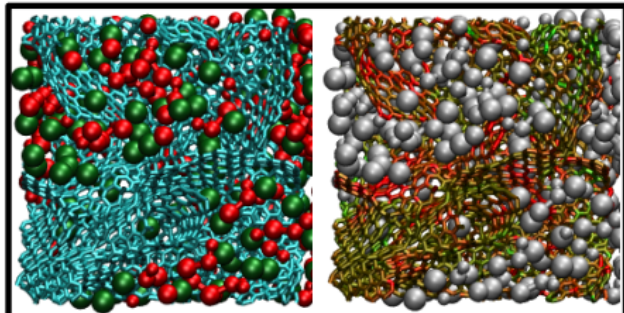
Superionic state

$$\Psi = -0.5 \text{ V}$$



$$N_+ = 104$$
$$N_- = 55$$

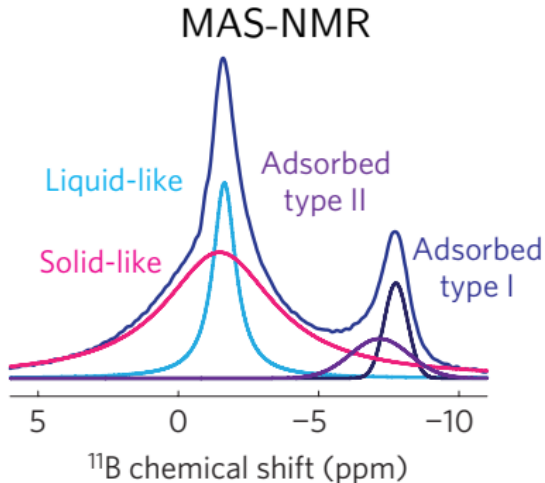
$$\Psi = +0.5 \text{ V}$$



$$N_+ = 66$$
$$N_- = 115$$

Like-like ions interactions screened by the metallic walls
→ superionic state (Kornyshev)

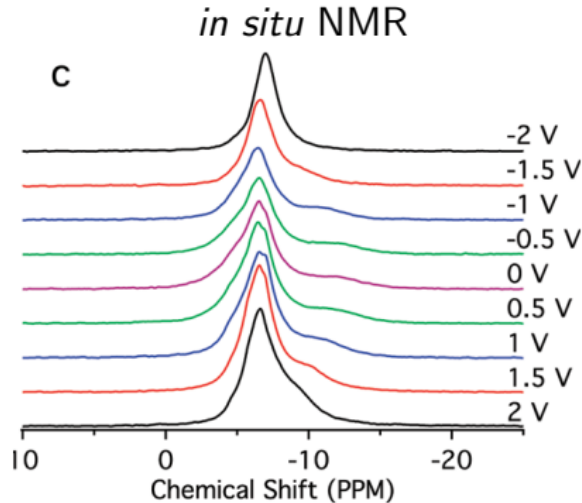
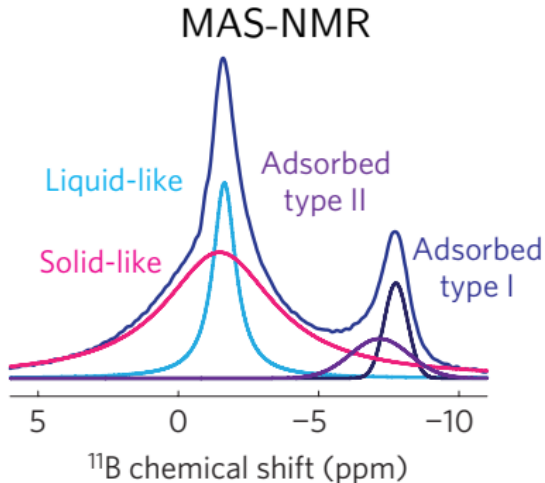
NMR experiments on supercapacitors



- Several adsorption modes

Deschamps *et al.*, *Nature Materials* **11**, 306 (2013)

NMR experiments on supercapacitors

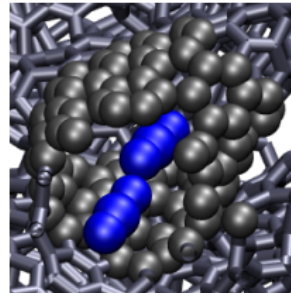
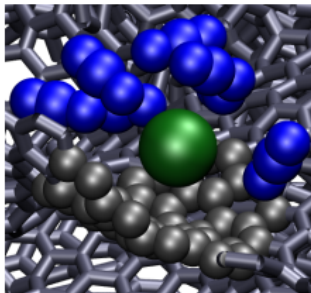
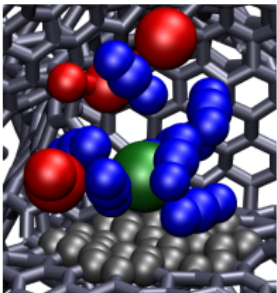
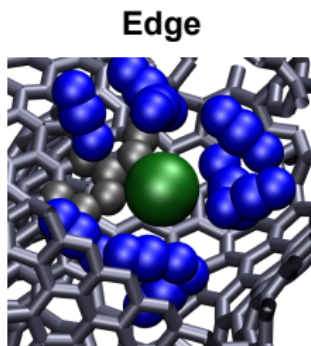
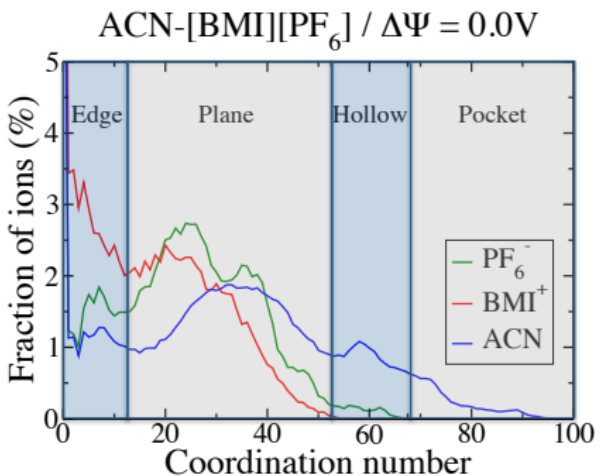
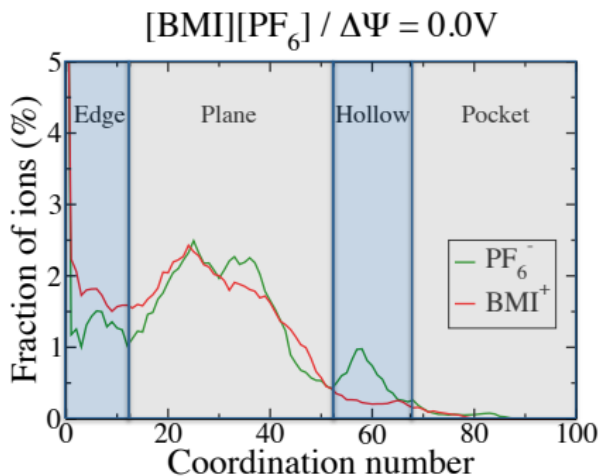


- Several adsorption modes
- Evolution with applied potential

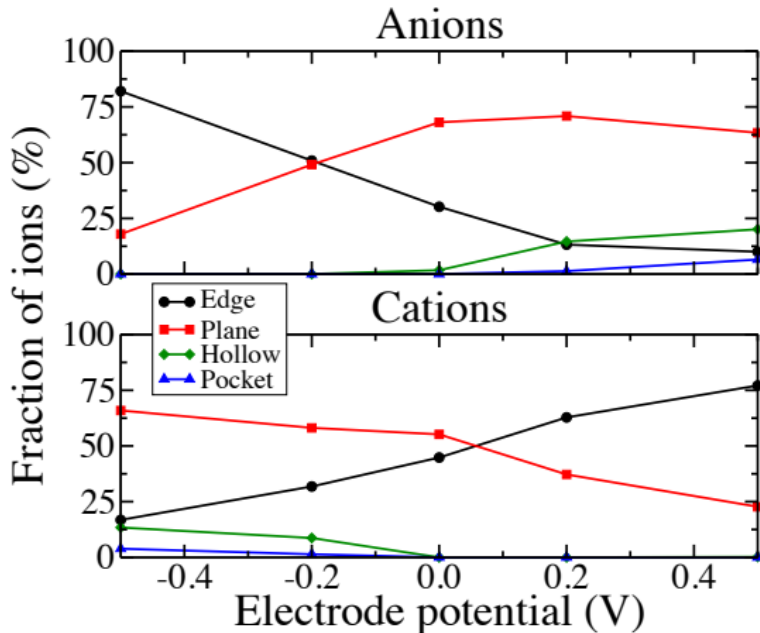
Deschamps *et al.*, *Nature Materials* **11**, 306 (2013)

Wang *et al.*, *J. Am. Chem. Soc.* **133**, 19270 (2011)

A variety of adsorption sites

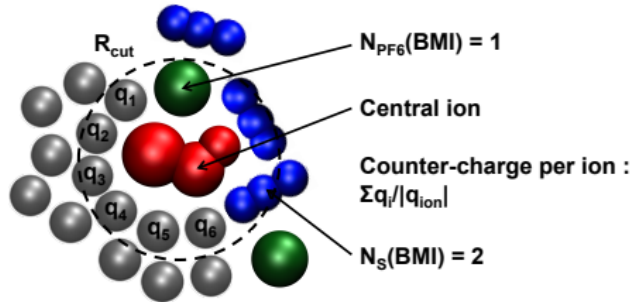


Evolution with applied potential

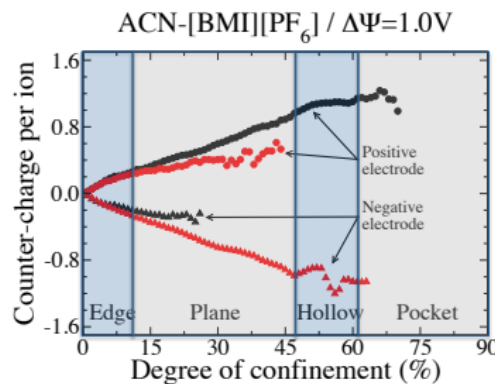
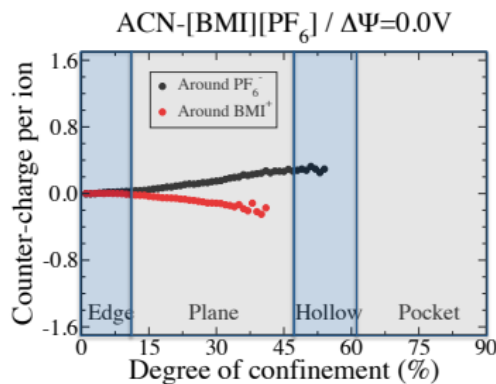


- Counter-ions migrate in confined sites...
- ... While other ions (and the solvent) leave them

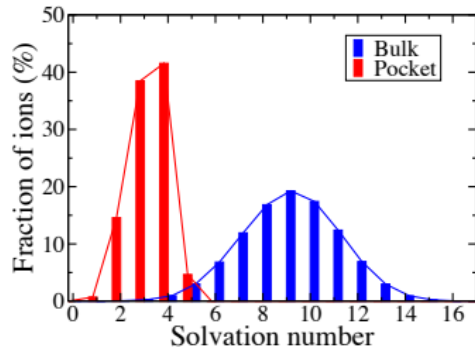
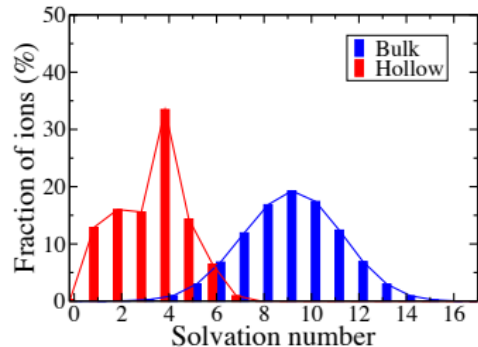
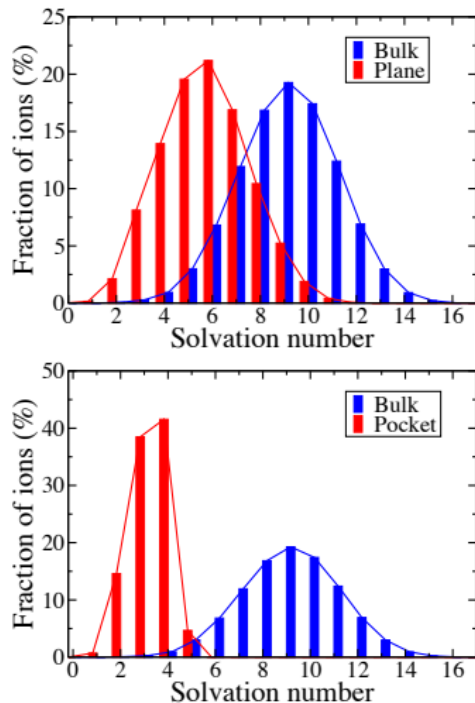
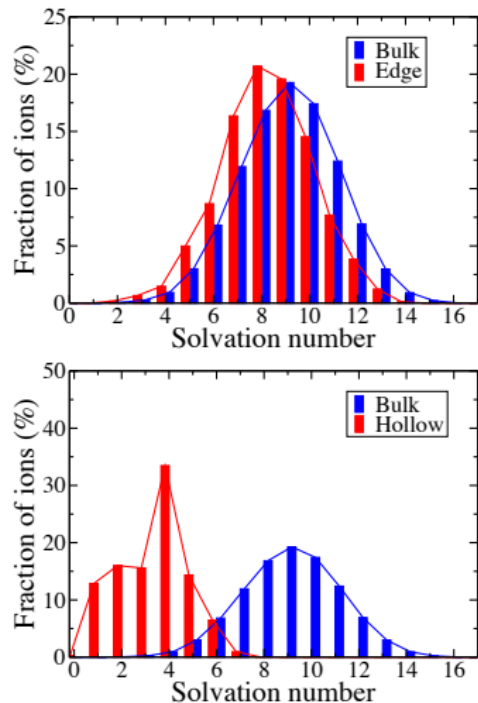
Impact of local morphology on supercapacitor properties



The local charge on the electrode is greater in highly confined sites



Desolvation of ions in nanoporous carbons



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Conclusion & Perspectives

- Simulations performed at constant applied potential
 - Realistic structures of CDCs
 - Nanoporous carbons:
 - screening by the metallic walls → superionic state
 - absence of overscreening due to confinement → better efficiency
 - role of the local morphology
 - Simulations of charge/discharge processes
-
- Influence of the ionic sizes (development of new models)
 - Porous carbon structure (evaluation of new models)
 - Ionic liquids dissolved in other solvents

Acknowledgements

- Céline Merlet, Clarisse Péan, Benjamin Rotenberg (PHENIX)
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