Modelling high energy density supercapacitors by molecular dynamics simulations

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Forum Teratec - July 2nd, 2014

# Outline

#### 1 Supercapacitors

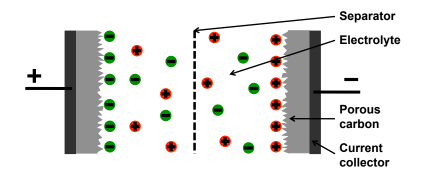
- 2 Molecular simulations
- 3 Molecular origin of supercapacitance

#### 4 Conclusion

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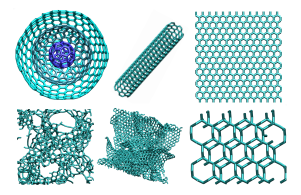
# Supercapacitors: electricity storage devices



- Also called Electrochemical Double Layer Capacitors
- Charge stored through adsorption of ions at the surface of electrode
- No redox reaction in the bulk material  $\rightarrow$  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)

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Modelling of supercapacitors

## Electrolytes

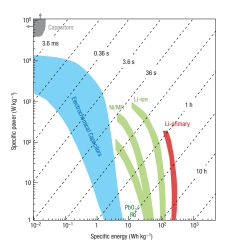
Need to find a good compromise between:

- 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





Collaboration Alstom / Batscap



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

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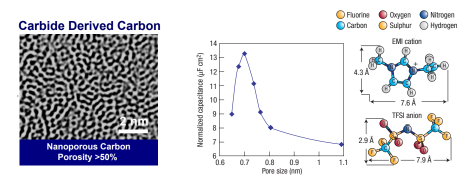
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# Applications



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

# Experimental discovery (Gogotsi & Simon)



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

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

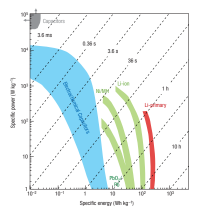
Largeot et al., J. Am. Chem. Soc., 130, 2730 (2008)

# 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 $^{\circ}$ C)



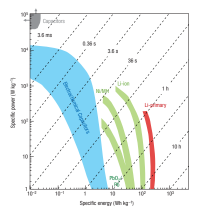
Much research work to be done in materials and electrochemistry, but...

# Challenges

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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
- $\rightarrow$  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

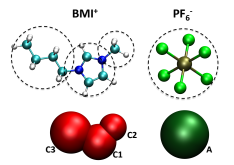
$$m^{i}\frac{\partial^{2}\vec{r}^{i}}{\partial t^{2}} = \sum_{j\neq i}\vec{F}^{j\rightarrow i} = -\frac{\partial V}{\partial \vec{r}^{i}}$$

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

- $\begin{array}{l} \mathsf{PF}_6^-: \text{ hexafluorophosphate} \\ & \to 1 \text{ site} \end{array}$



Interaction potential:

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

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

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

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Modelling of supercapacitors

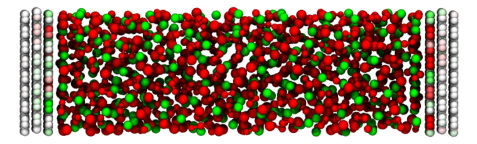


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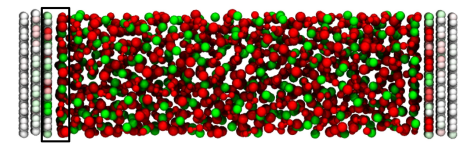
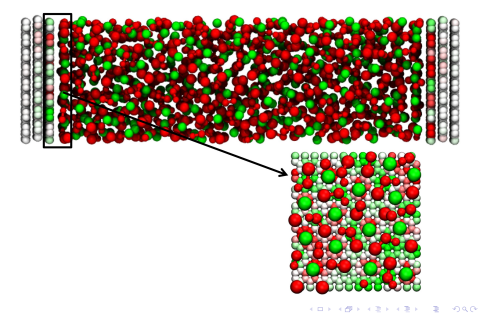
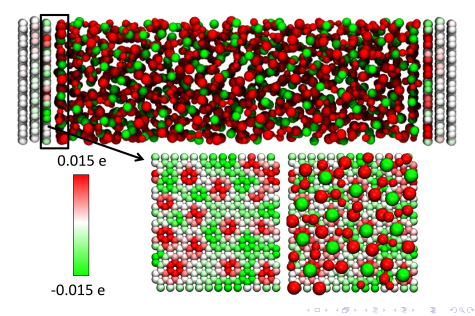
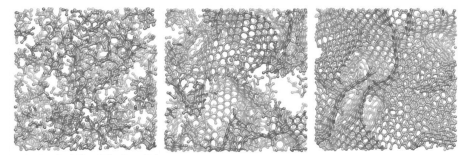


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# Porous electrodes: CDC model structures



- 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

Palmer et al., Carbon 48, 1116 (2010)

## Example of trajectory

Loading movie

#### Simulation cell with 2-Dimensional periodic boundary conditions

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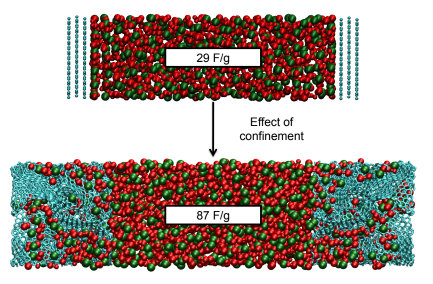
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## Increase of the capacitance in nanoporous carbons



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

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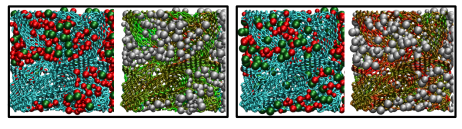
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Superionic state

$$\Psi = -0.5 V$$

#### $\Psi = + 0.5 V$

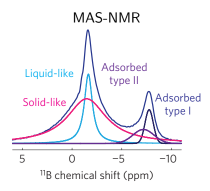


Like-like ions interactions screened by the metallic walls  $\rightarrow$  superionic state (Kornyshev)

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

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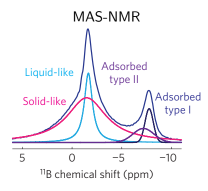
#### NMR experiments on supercapacitors

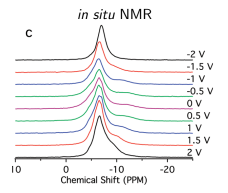


• Several adsorption modes

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

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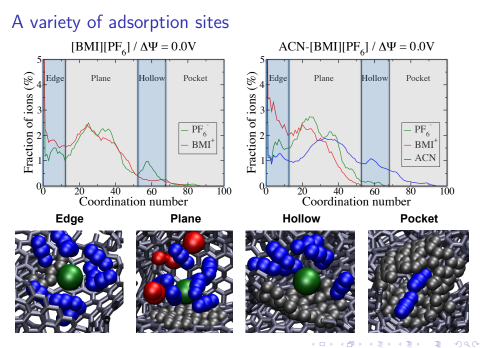




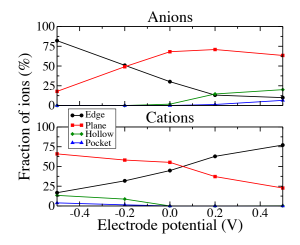
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Evolution with applied potential

Deschamps et al., Nature Materials 11, 306 (2013) Wang et al., J. Am. Chem. Soc. 133, 19270 (2011)

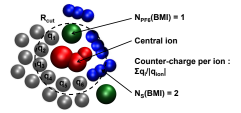


# Evolution with applied potential

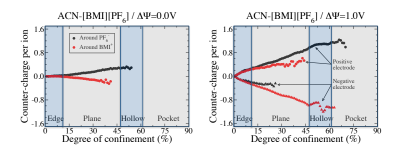


- Counter-ions migrate in confined sites...
- ... While other ions (and the solvent) leave them

# Impact of local morphology on supercapacitor properties

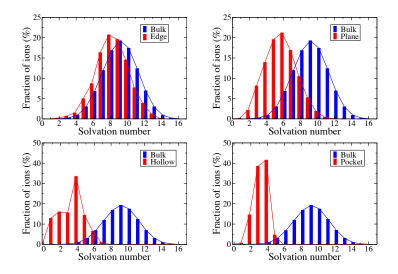


The local charge on the electrode is greater in highly confined sites



Merlet et al., Nature Communications, 4, 2701 (2013)

#### Desolvation of ions in nanoporous carbons



Merlet et al., Nature Communications, 4, 2701 (2013)

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# Conclusion & Perspectives

- Simulations performed at constant applied potential
- Realistic structures of CDCs
- Nanoporous carbons:
  - -screening by the metallic walls  $\rightarrow$  superionic state
  - -absence of overscreening due to confinement  $\rightarrow$  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

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