

HELSINKI UNIVERSITY OF TECHNOLOGY
LABORATORY OF PHYSICS
COMP

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IUPAP Commission 20

Computational Physics

Working Group on Nanoscience

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Courtesy of The Archives, California Institute of Technology.

- How do we write small?
- Information on a small scale
- Better electron microscopes
- The marvellous biological system
- Miniaturizing the computer
- Miniaturization by evaporation
- Problems of lubrication
- A hundred tiny hands
- Rearranging the atoms
- Atoms in a small world

"There Is Plenty of Room
at the Bottom"

Richard P. Feynman
December 1959

10^{-9} m

Three pillars of nanosciences

Manufacture and processing of nanometer-scale structures

- “top-down”: **lithography**
- “bottom-up”: **self-assembly**

Characterisation, imaging and probing

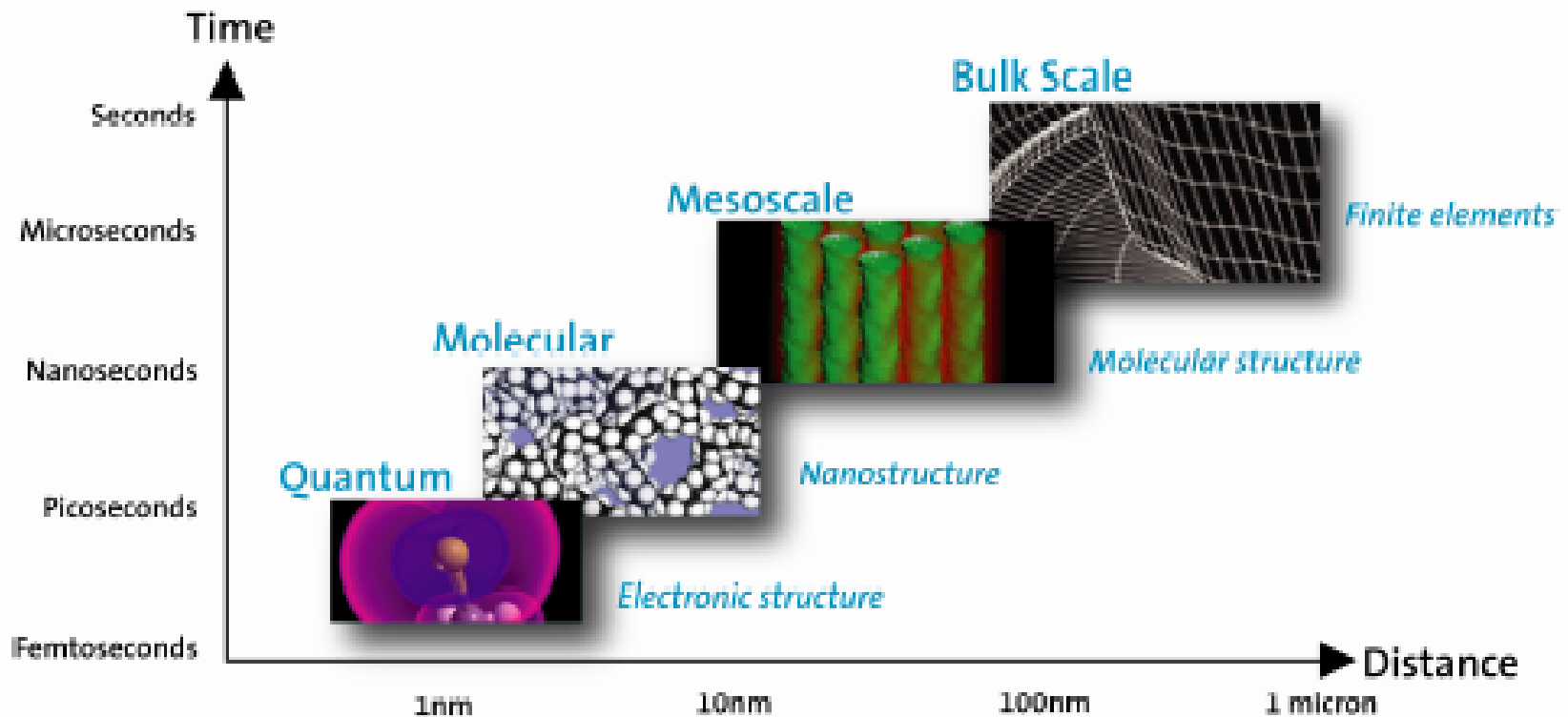
Theory, modelling and simulation

- Predictive computation of physical, chemical and biological functions
- Process design
- Interpretation of characterisation and imaging

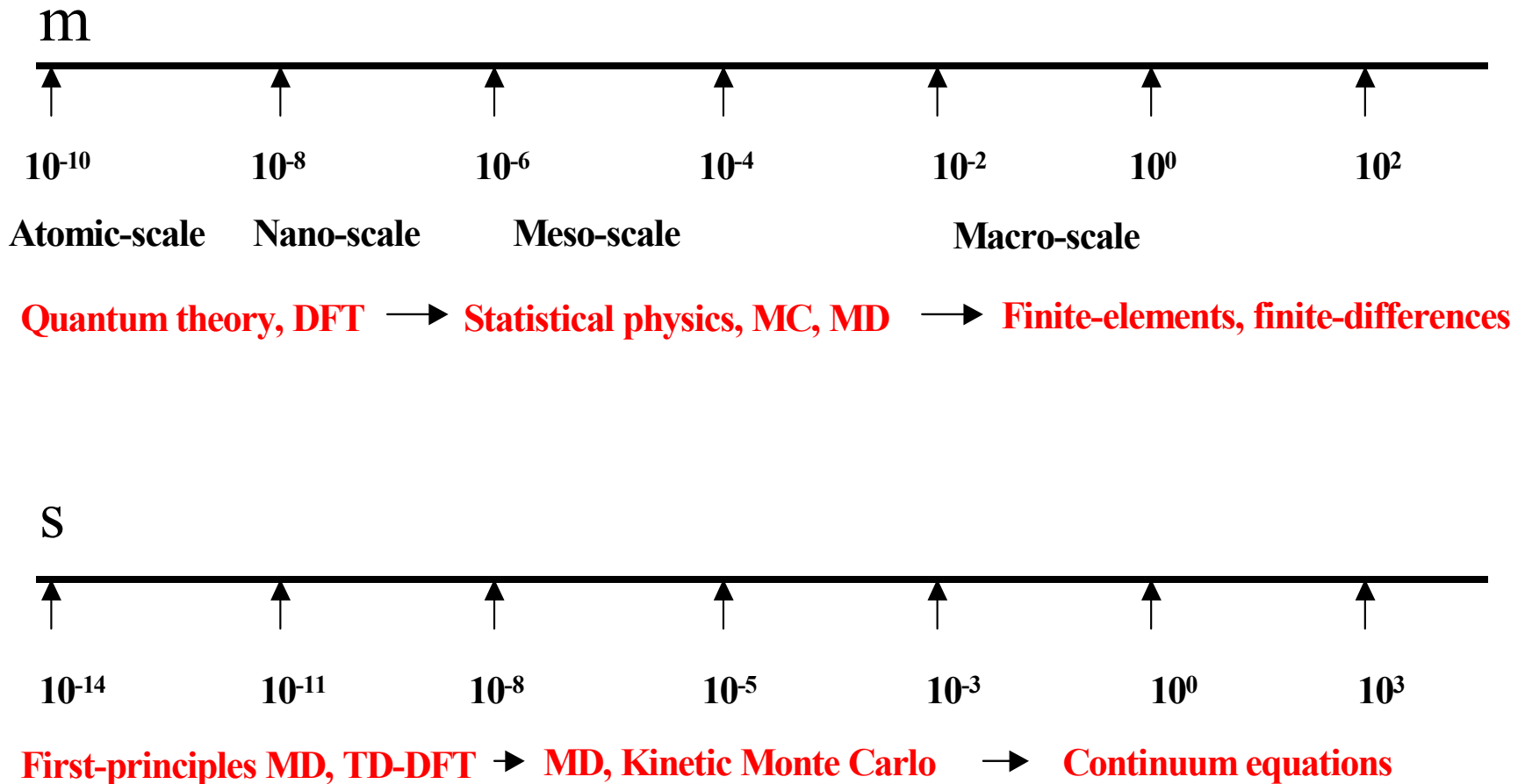
“Mimicking Nature”

COMPUTATIONAL PHYSICS

Different modeling and simulation methods address a range of time and size scales

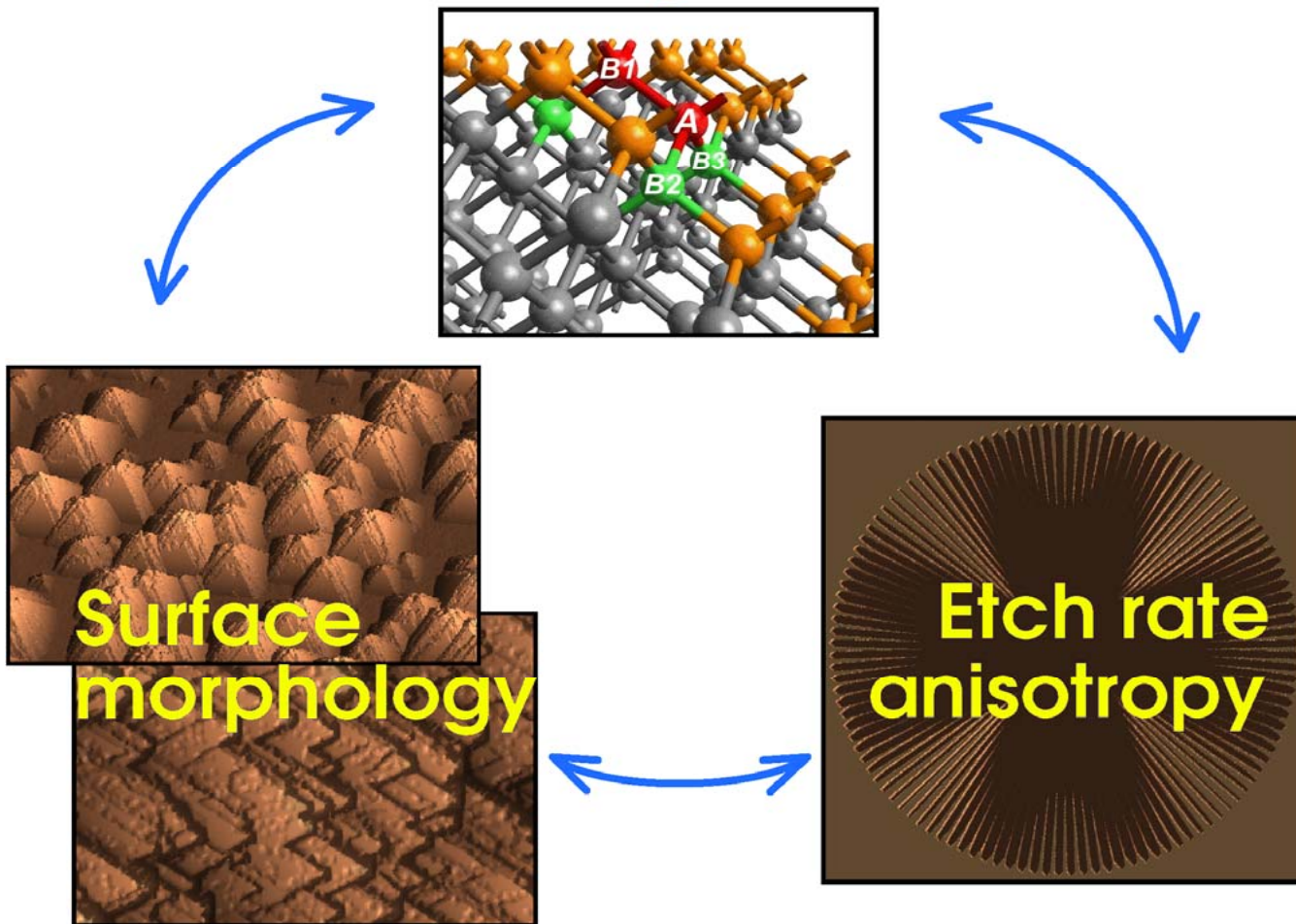


Multiple length and time scales



Bridging the different length scales

Atomistic processes



Interrelation between microscopic, mesoscopic and macroscopic features
of the etching process

Multiscale computational methods

<u>Method</u>	<u>Type of information</u>	<u>Time scale</u>	<u>Length scale</u>
Quantum Monte Carlo/ED	Microscopic	-	~100 atoms
Density-functional theory	Microscopic	-	~ 1000 atoms
Ab initio molecular dynamics	Microscopic	$t < 10 \text{ psec}$	~ 100 atoms
Semi-empirical molecular dynamics	Microscopic	$t < 1 \text{ nsec}$	~ 100000 atoms
Kinetic Monte Carlo	Microscopic to mesoscopic	$1 \text{ psec} < t < 1 \text{ hour}$	~1 micrometer
Rate equations	Averaged	$0.1 \text{ sec} < t < \infty$	All
Continuum equations	Macroscopic	$1 \text{ sec} < t < \infty$	> 10 nm

Commission conferences (C20)

International Conference in Computational Physics (CCPXXXX)

- CCP2003 Xian, China
- CCP2004 Genoa, Italy
- CCP2005 Los Angeles, USA

CCP2003

*Postponed to 2004 due to the SARS epidemic;
difficulties in scheduling*

CCP2004

1. Nanoscience in plenary talks

- electron transport and dissipation in nanoscale devices (R. Car)
- optical, electrical and mechanical properties of nanostructures (S. Louie)
- resonating valence-bond wave functions: from lattice models to realistic simulations (S. Sorella)

2. Nanoscience in invited talks

- molecular nanostructures
- semiconductor/metal interfaces
- carbon nanotubes
- large molecules at metal interfaces

- partially folded states of proteins
 - from nanodiamonds to nanotube growth
 - wave function optimisation for accurate quantum Monte Carlo
 - doped helium clusters
 - multiscale algorithms
-

Three major themes:

- condensed-matter and materials physics,
especially nano (including soft and biological matter) (50 %)
- statistical physics and complexity (30 %)
- particle physics, astrophysics and cosmology (20 %)

CCP2005

1. Nanoscience in plenary talks

- physics and chemistry in the non-scalable and emergent regimes (Landman)
- soft matter and biomaterials (Klein)
- dielectric response and polarization (Vanderbilt)

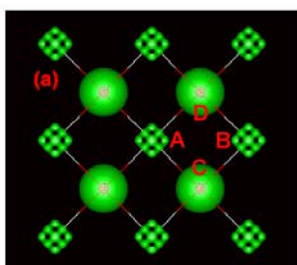
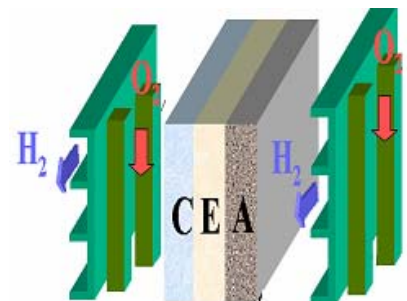
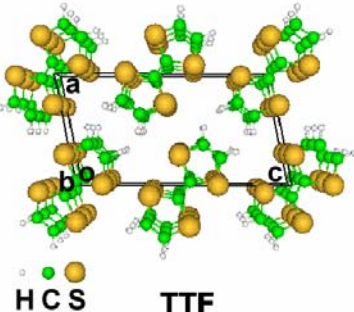
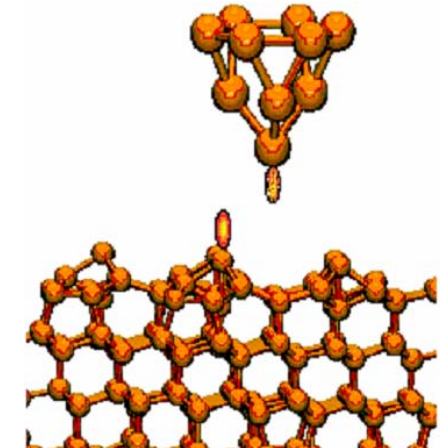
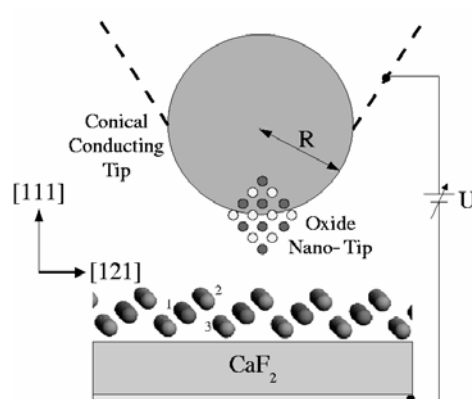
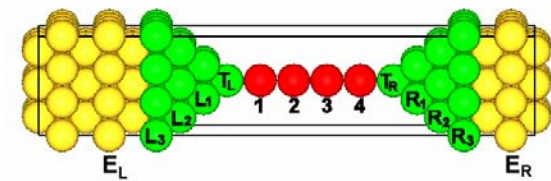
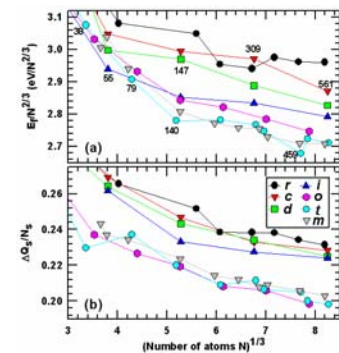
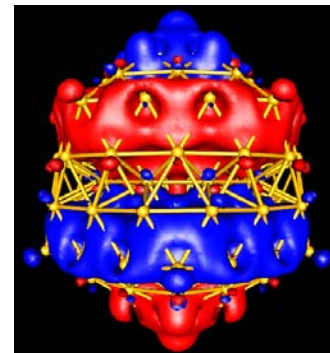
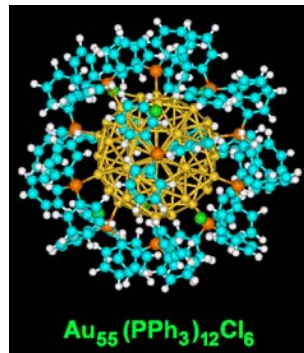
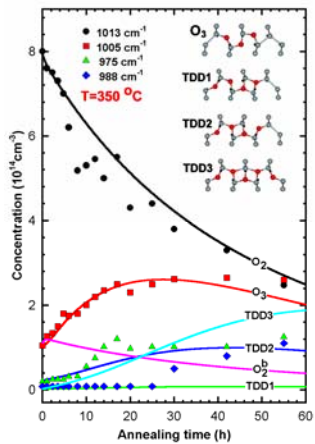
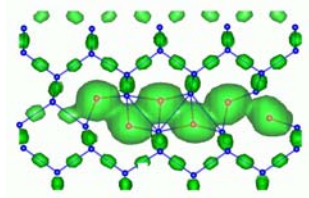
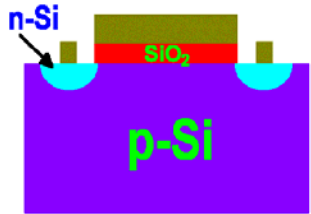
2. Nanoscience in invited talks

- carbon nanotubes (several talks)
- quantum dots
- semiconductor nanoparticles
- nanowires and nanocorras
- metallic nanoparticles
- future directions in the simulation of nanomaterials

Major themes:

- condensed matter and materials, including nano (50 %)
- biological physics (15 %)
- statistical physics and complexity (15 %)
- quantum information and computing (15 %)
- algorithms (5 %)

Some recent research areas



"The general theory of quantum mechanics is now almost complete. The underlying physical laws necessary for the mathematical theory of a large part of physics and the whole of chemistry are thus completely known, and the difficulty is only that the exact application of these laws leads to equations much too complicated to be soluble."

P.A.M. Dirac, 1929

1902-1984

Physics Nobel Prize 1933 (with E. Schrödinger)

"It therefore becomes desirable that approximate practical methods of applying quantum mechanics should be developed, which can lead to an explanation of the main features of complex atomic systems without too much computation."

P.A.M. Dirac, Proc. Royal Soc. London A 123, 714 (1929)

Density-Functional Theory (DFT)

$$E_{ks}[\{\psi_i(\mathbf{r})\}] = -\frac{1}{2} \sum_i f_i \int \psi_i \nabla^2 \psi_i d\mathbf{r} + \frac{1}{2} \sum_{i \neq j} \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r} d\mathbf{r}' \\ + \int V_{ion}(\mathbf{r})\rho(\mathbf{r})d(\mathbf{r}) + E_{xc}[\rho(\mathbf{r})] + E_{ion}(\{\mathbf{R}_I\})$$

Kohn-Sham Eqs. self-consistently solved

$$\left(-\frac{1}{2} \nabla^2 + V_{ion}(\mathbf{r}) + V_H(\mathbf{r}) + V_{xc}(\mathbf{r}) \right) \psi_i(\mathbf{r}) = \varepsilon_i \psi_i(\mathbf{r}) \\ \rho(\mathbf{r}) = \sum_i |\psi_i|^2$$

First-principles Molecular Dynamics

$$F_I = -\frac{dE_{ks}}{d\mathbf{R}_I} = -\frac{\partial E_{ks}}{\partial \mathbf{R}_I} - \sum_i \frac{\partial E_{ks}}{\partial \psi_i} \cdot \frac{d\psi_i}{d\mathbf{R}_I} - \sum_i \frac{\partial E_{ks}}{\partial \psi_i} \cdot \frac{d\psi_i^*}{d\mathbf{R}_I}$$

Density functional theory provides a very efficient way for the application to both solids and molecules, with higher accuracy than Hartree-Fock.

DFT Methods

all-electron full potential
all-electron muffin-tin
pseudopotential
jellium

beyond LDA

GGA
LDA

GW
time-dependent DFT
LDA+U
self-Interaction corr. (SIC)
sx-LDA

$$\left(-\frac{1}{2} \nabla^2 + V_{ion}(\mathbf{r}) + V_H(\mathbf{r}) + V_{xc}(\mathbf{r})\right) \psi_i(\mathbf{r}) = \varepsilon_i \psi_i(\mathbf{r})$$

fully-relativistic
scalar-relativistic
non-relativistic

spin-polarized
collinear spin
non-collinear spin

spin-nonpolarized

periodic
non-periodic

plane waves
atomic orbitals
GTO, STO, numerical
augmented waves
FLAPW, PAW, LMTO, ASW
fully numerical

Scientific software used at COMP

QUANTUM SCALE

QMC

Quantum Monte Carlo

FINGER VASP
CASTEP WIEN2k
FLEUR MIKA
SIESTA

Density-functional
theory

ATOMISTIC SCALE

TB-MOLDY MOLDY

Molecular dynamics

MESOSCOPIC SCALE

KMC
TAPAS

Kinetic Monte Carlo
Cellular automata

CONTINUUM SCALE

ELMER

Finite-element methods

C20 (Greetings from Risto Nieminen)

CCP2007 : Conference on Computational Physics

5-8 September, 2007
Université Libre de Bruxelles
Vrije Universiteit Brussel

Tentative Sponsors :

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Instituts Internationaux de Physique et de Chimie fondés par Ernest Solvay

C20 (Greetings from Risto Nieminen)

Topics will include

- Materials Science, (electronic structure, ab-initio MD, Quantum MC..)
- Soft Matter (Polymers, membranes, proteins..)
- Biological Physics
- Computational Statistical Physics, equilibrium and non-equilibrium (with a particular emphasis on hydrodynamical flows by particle methods)
- Astrophysics
- Fluids and Plasmas
- Particle Physics
- New Methods in computations
- New technical challenges in large-scale computing: new architectures and development of middleware, grid computing, problems raised by code development and maintenance...