

# GEMS, Grid Enabled Molecular Simulations

EGEE 1<sup>st</sup> EU Review – 9<sup>th</sup> to 11<sup>th</sup> February 2005 CERN Antonio Laganà<sup>1</sup>, Osvaldo Gervasi<sup>2</sup> <sup>1</sup> Dept. of Chemistry, University of Perugia <sup>2</sup> Dept. of Math. & Computer Science, University of Perugia

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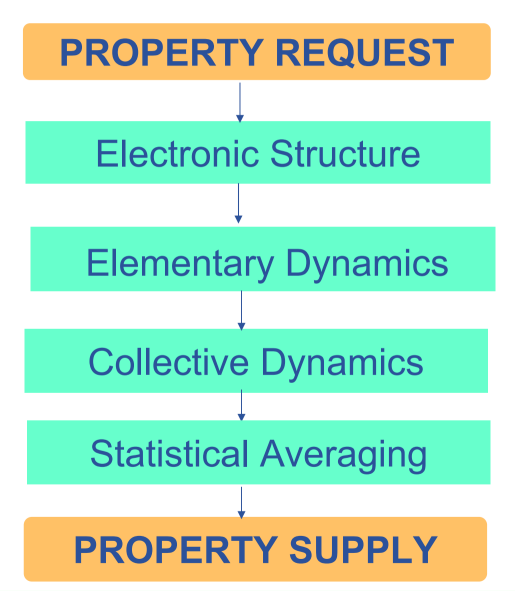
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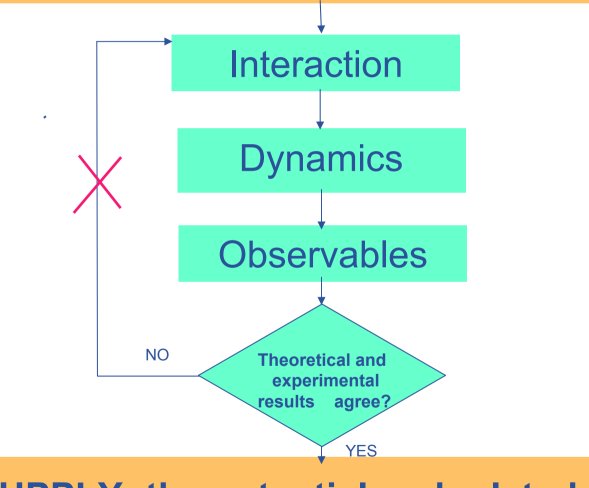
- Overview of GEMS and the prototype application GEMS.0
- Deployment layout, scale of usage, LCG-2 middleware services
- Difficulties, workarounds and alternatives
- Challenges for full production
- The Chemistry and Molecular Science Community
- The grid added value and future work & direction





# **CALCED THE GEMS.0 demo application**

#### **REQUEST:** a potential fitted to beam experiments



### **SUPPLY: the potential and related monitors**

# GEMS.0 implementation on GILDA

- SPECIFIC ASSUMPTIONS OF THE GEMS.0 DEMO
  - Focus on atom diatom systems
  - Take potential energy routine from a library
  - Adopt classical dynamics (trajectories)
- GEMS.0 EXECUTION ON GILDA
  - Set the demo environment
  - Input the atoms involved
  - Input energy, initial vibrational and rotational state
  - Loop on trajectories by generating randomly the other initial conditions
  - Collect results as virtual monitors (in the real application to be compared with the experiment)



$$i\hbar\frac{\partial}{\partial t}\Psi(\{W\},\{w\},t) = \hat{H}\Psi(\{W\},\{w\},t)$$

Separation of electronic and nuclear motions

Electronic Schrödinger equation:

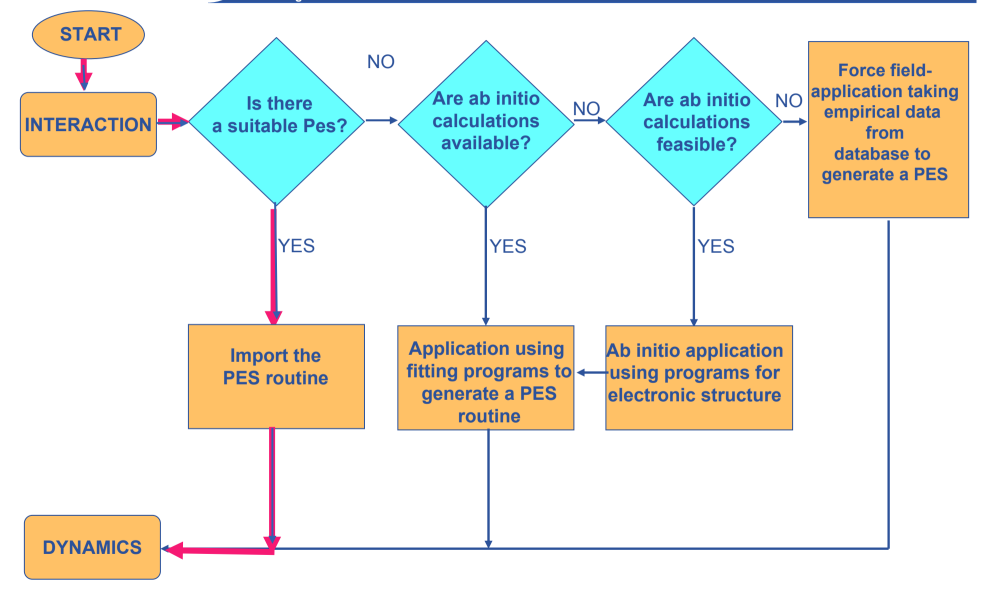
Nuclear Schrödinger equation:

 $\hat{H}_{elec}\Psi_{n}(\lbrace w \rbrace; \lbrace W \rbrace) = E_{n}(\lbrace W \rbrace)\Psi_{n}(\lbrace w \rbrace; \lbrace W \rbrace) \quad \hat{H}_{n}\chi_{n}(\lbrace W \rbrace, t) = i\hbar \frac{\partial}{\partial t}\chi_{n}(\lbrace W \rbrace, t)$ 

Statistical averaging for beam conditions

## The INTERACTION module

Enabling Grids for E-sciencE



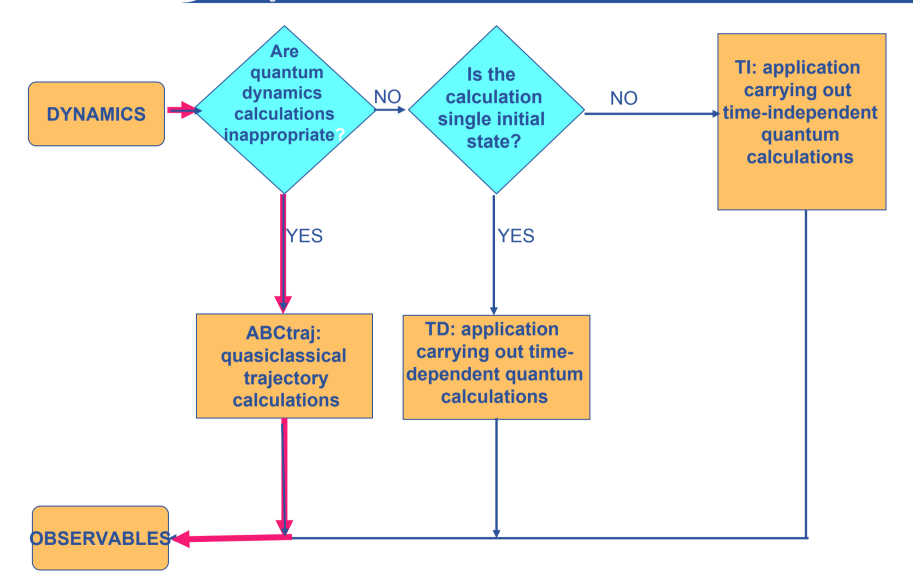
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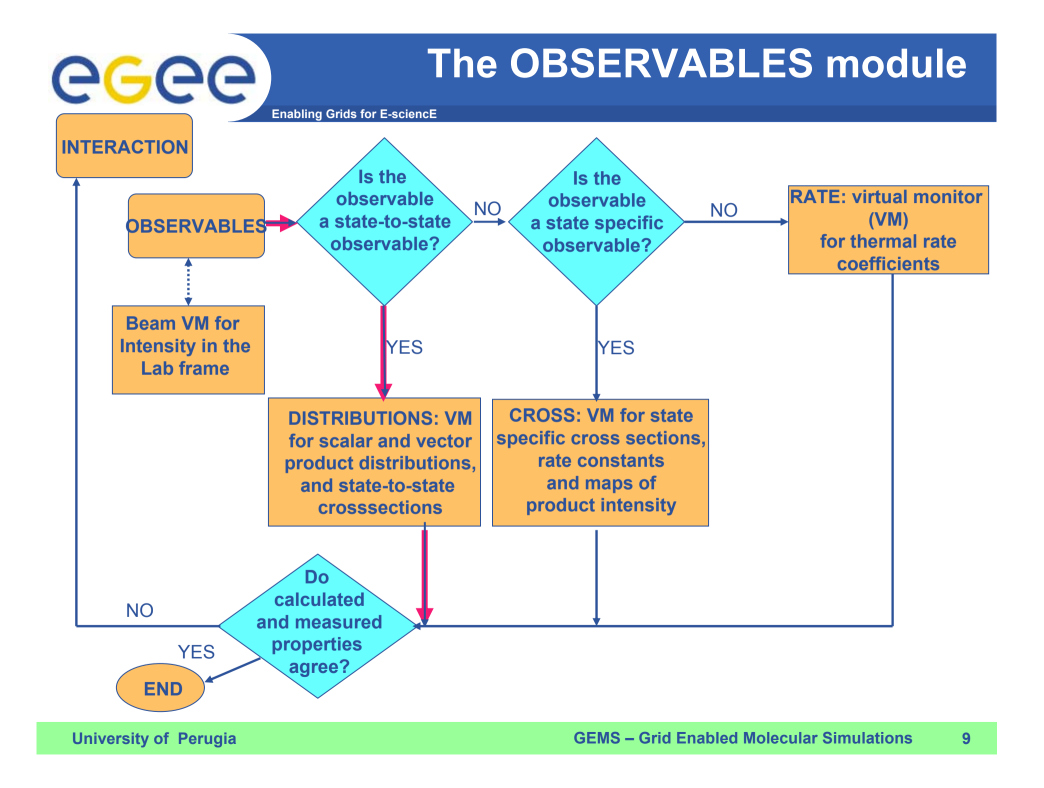
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### The DYNAMICS module

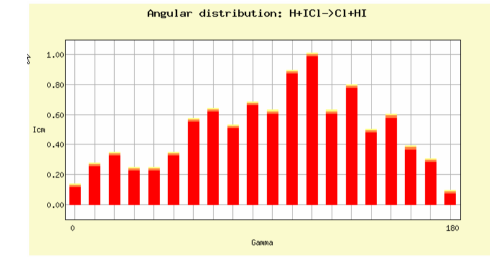
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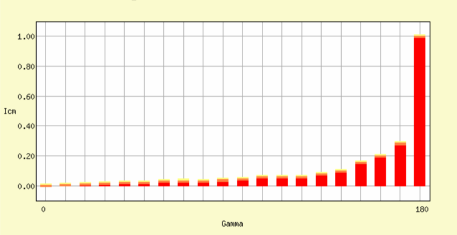


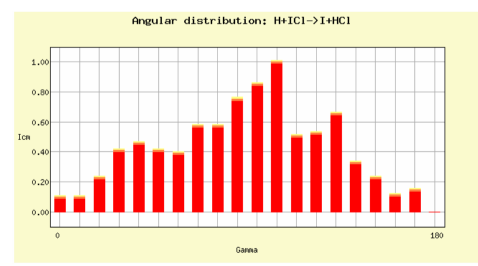




## The Virtual monitors







Angular distribution:  $H + ICl \rightarrow H + ICl$ 



## **Demo deployment layout**

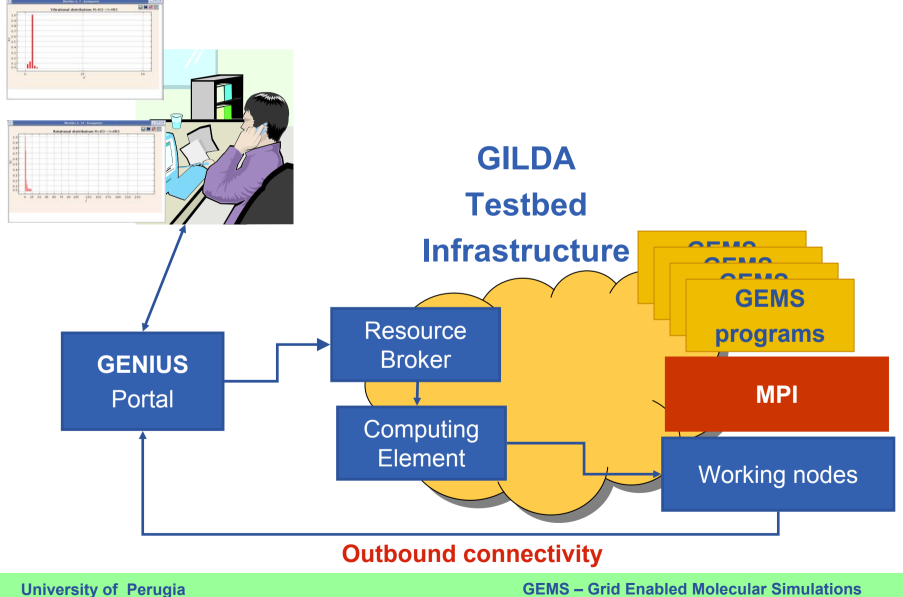
Enabling Grids for E-science

- Sites
  - GILDA testbed sites.
- Key services
  - Resource Broker: grid004.ct.infn.it
  - Computing Element: ce.grid.unipg.it
  - User Interface: grid-tutor.ct.infn.it
  - GENIUS Portal: https://genius.ct.infn.it
  - CompChem VOMS



- Number of jobs: 10 per day
- Storage: 100KB 50GB depending on the type of computational engine used and the chemical system studied:
  - Trajectory calculations: <100KB</li>
  - Time Dependent Quantum: 10GB
  - Time Independent Quantum: 50GB
- RAM: 100KB 2GB
  - Trajectory calculations: <100K</li>
  - Time Dependent Quantum: 1GB
  - Time Independent Quantum: 2GB
- Success rate: 98%







- We have encountered the following problems:
  - Not all sites did publish the environment variables related to the outbound connectivity of the Working Nodes.
  - The job activation time is high, expecially if an high number of working nodes is requested.
  - Not all sites have enough resources (disk + RAM) for Quantum Chemistry codes.
  - In some sites MPI is not available.
  - The Firewall sometimes is misconfigured and blocks the outbound connectivity in the EGEE standard port range: 20000-25000.
- The workarounds were
  - Alert people involved to properly configure the environment variables.
  - Alert people to open the EGEE standard ports on the Firewall.



 Running at weekends computers of teaching laboratories (trajectories).

 Making a combined use of SP4 at CINECA and CRAY T3E at Edinburgh (3D quantum).



- What's missing to extend the application to production
  - Define the policy for the usage of commercial suites of programs. At present each site buys its own license and opens the access to the VO users. The commercial licence requirement for a job need to be specified in a JDL statement.
  - Credits for contributions to development and service.
  - Extend the suites of programs to other components of the molecular simulator.
  - Full deployment of the VO and consortium.
- As Production Service, the expected usage profile is:
  - Users: 100.
  - Sites: 30.
  - CPU consumption (per job): 100 1000 hours.
  - Disk Storage (per job): 10 MB 800 GB.
  - RAM Storage (per job): 100 KB 200 GB.



## The Chemistry community

Enabling Grids for E-sciencE

Simbex

Murqm

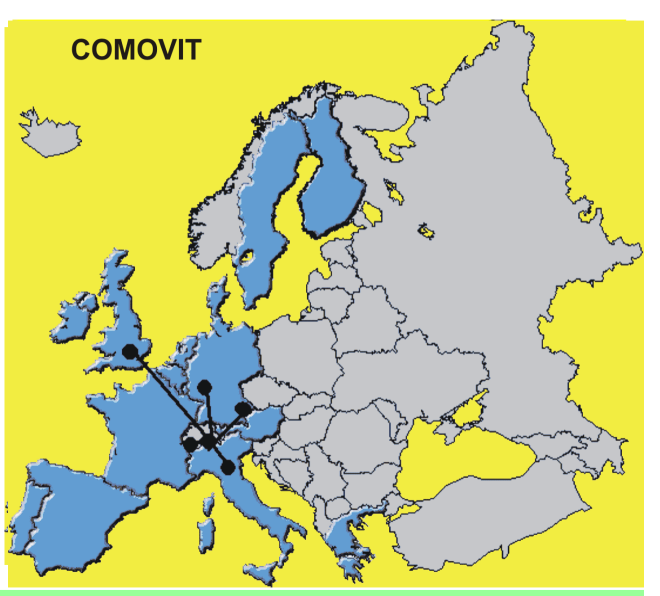
Dirac

Elchem

Icab

Dysts

Comovit





# LABS per NATIONALITY (51)

Enabling Grids for E-sciencE

- 1 Isr,PI,Sk,NI
- 2 Cz,Ch, Fr, Dk, A, Sw, No
- 3 Hu
- **4** Gr
- 5 E
- 6 D, Uk,
- 9 I



- 3 Computer Centers
- New electronic structure programs (MOLCAS, DIRAC, DALTON, COLUMBUS, MR-CCSD).
- New Dynamics programs (AMD, TPS, KMC, condensed phase).
- Chemical knowledge semantic web (molecular structures, apparatuses, processes).



- Software integration into distributed workflows
  - to assemble applications out of various (different or complementary) distributed competences coordinated via the grid (in the prototype electronic structure, elementary dynamics, statistical averaging, interfacing the experiment)

#### Computational Campaigns

 to evaluate properties depending on the fate of few out of millions, billions or more events by distributing the execution of the computations on the grid (in the prototype particular reactive events having specific energetic or steric properties)

#### Collaborative Engineering of knowledge

 to handle (when is the case also in a privacy protecting fashion) chemical information and knowledge including training and production of new knowledge (for the prototype the design of virtual chemical processes or laboratory experiments)



- A consortium agreement will be signed by the new participants of CompChem VO.
- A dedicated Portal for GEMS will be implemented to run on the production environment.
- Various computational engines will be implemented in order to offer grid services based on the a priori simulation of molecular systems.
- Molecular Virtual Reality components will be combined with Human-size Virtual Reality to offer multiscale realistic representations of material structures and processes.
- Design appropriate cost models for supporting research, project and service activities.