

Implementation of the Simulation Data Model

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Abstract

This document aims at providing examples of implementation of SimDM. Two VO-Theory databases are explained in details. First, the STARFORMAT database devoted to MHD simulations. Second, the PDR database devoted to 1D models of interstellar clouds. We present how the results of these simulations have been mapped on the Simulation Data Model. We also present in each case, the pipeline we used to simplify the updates of the databases each time the codes behind the simulation evolves or that new simulations have to be added in the databases. An overview of Millennium is also presented as well as a discussion presenting the relationships between S3 and a service based on SimDM.

Status of This Document

This is an IVOA Note expressing suggestions from and opinions of the authors. It is intended to share best practices, possible approaches, or other perspectives on interoperability with the Virtual Observatory. It should not be referenced or otherwise interpreted as a standard specification.

A list of [current IVOA Recommendations and other technical documents](http://www.ivoa.net/Documents/) can be found at <http://www.ivoa.net/Documents/>.

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1 Introduction

The simulation data model (SimDM) has been developed by the VO-Theory Interest Group at IVOA. Initially, one of the major objectives was to publish 3D + time simulations. Quickly the Data Model, SimDM, evolved to cover larger sets of simulations. In this note, we present two detailed examples of implementation of SimDM: one about 3D+time MHD simulations, and one about 1D stationary models of the atomic and molecular structure of interstellar clouds. These two examples cover two extreme

cases for the implementation of SimDM. We also present the overview of an implementation on Millennium simulations as well as some comments on the relationships between SimDM and S3.

Among the major steps one has to do to develop such VO-Theory services, we recommend to the developers to pay attention to two points at the beginning of the project:

1) The first question any publisher should ask to himself, is what do I want to publish and for who ? Simulations can be quite complex and may provide many heterogeneous quantities. Is it really useful to publish all the computed quantities? Do not forget that the purpose of SimDM is to describe simulations for discovery, not to publish all the results of simulations in a SimDB. Quantities not published in SimDB can still be accessible by a SimDAL access or specific services.

For example, in PDRDB, we publish information's about quantities as line intensities and column densities because observers want to discover interesting models thanks to these quantities. But we do not publish in SimDB local emissivities or local cooling and heating rates for the various physical processes. Nevertheless, these quantities, not published in the database, can still be obtained by a download of the full results of the simulations.

2) The other aspect we would like to highlight is that the developer has to think at the beginning of the project to the future evolutions of the service.

Codes can be modified frequently to add new physics. As a consequence, the quantities published in a VO-Theory service may have to be modified regularly. Obviously, we wish this be as transparent as possible for the scientist publishing his data. To solve this issue, we developed a pipeline to ingest simulations / models in the data bases. For example, in the case of the PDR code, the code produces directly an XML file describing the content of the outputs (metadata). The ingestion pipeline, that makes use of VO-URP, analyzes this XML file, identifies if new quantities are present in it, and eventually modify automatically the database. This way, scientists can publish their results without the help of a software engineer who would have to modify the database each time the code evolves.

Preliminary note : In this document, we use the notion of **SimDB**. In this context, SimDB means an implementation of SimDM in a relational database with a TAP interface. This TAP interface is not used in the implementations Starformat and PDRDB. In the S3 part (section 4), we present relationships between this TAP interface and S3 services.

2 Starformat – MHD simulations

2.1 Description and objectives

StarFormat stands for Star Formation. It is a grouped database assembling multiple simulation models from the results of theoretical projects designed to study the formation of molecular clouds and prestellar dense cores in order to aid in the analysis and interpretation of current and future observations. It contains so far 4 different sets of large MHD simulations using the three-dimensional adaptive mesh refinement (AMR) magneto-hydrodynamical code RAMSES-MHD. Tryouts have been performed and StarFormat will soon host also very different kinds of simulation like turbulence boxes or dense cores from projects using the other popular codes FLASH, GADGET or PHOENIX.

The service proposes a synthesized access to all simulations descriptions, parameters and snapshots.

Post-processings applied to these snapshots in order to identify and characterize specific objects like dense clumps are described extensively and can be queried along the values of a selection of properties. Data cubes extraction from the raw data is proposed as an extra service. TAP access to the content of the whole database will also be provided.

2.2 SimDM implementation on STARFORMAT

The SimDM data model provides a rich and exhaustive representation of all the elements needed to describe extensively the StarFormat simulations.

2.2.1 - Project part

uType: SimDB:simdb/Project

All the simulations are stored and described within different instances of the Resource class: different versions of the RAMSES codes and their runs are respectively stored as instances of the classes Protocol and Experiment. These are grouped as ProjectResources within a “Formation of Molecular Cloud” Project, third kind of possible Resource. Any participant to the work can be associated with his complete coordinates to these different levels of Resource as a Contact assuming different possible roles such as owner, creator, publisher or contributor.

uTypes:

- *Resource: SimDB:simdb/Resource*
- *Project: SimDB:simdb/Project*
- *ProjectResource: SimDB:simdb/ProjectResource*
- *Contact: SimDB:simdb/Contact*

Example of StarFormat instances for the Project part

2.2.2 Protocol part

uType: SimDB:simdb/protocol/Protocol

The RAMSES code used is described within the Simulator class, an instance of the Protocol class. Using The Simulator's various heritage classes, the code is fully described with its different components: all the possible physical processes which can be part of a run (MHD, hydrodynamics, gravitation, heating and cooling, ...) are described within the Physics class; the parameters (AMR levels, boundary and initial conditions, incoming flow velocity, ...) are stored as InputParameters and grouped within a ParameterGroup as ParameterGroupMembers whenever they come as a series of values like for example the jeans refinement lengths. Finally the use of adaptive mesh refinement is specified within the RepresentationObjectType class in which we describe the basic AMR cells that compose the data cube for a RAMSES simulation.

uTypes:

- *Protocol: SimDB:simdb/protocol/Protocol*
- *Simulator: SimDB:simdb/protocol/Simulator*
- *Physics: SimDB:simdb/protocol/Physics*
- *InputParameter: SimDB:simdb/protocol/InputParameter*
- *ParameterGroup: SimDB:simdb/protocol/ParameterGroup*

- *ParameterGroupMember: SimDB:simdb/protocol/ParameterGroupMember*
- *RepresentationObjectType: SimDB:simdb/protocol/RepresentationObjectType*

2.2.3 Experiment part

uType: SimDB:simdb/experiment/Experiment

Any run of the code is then described as a Simulation, which is an instance of Experiment class, and refers to the corresponding Protocol of class Simulator. The specificities of each simulation are described though the uses of the corresponding Experiment's heritage classes referring to the Protocol's corresponding ones: the physical processes applied are notified through instances of AppliedPhysics as references to the Physics elements defined earlier; all the parameters of the run, whether they are numerical values or textual, are given as instances of string or quantity ParameterSetting referring to the previous InputParameter class. The molecular cloud simulated by a run is described as a TargetObjectType that can be characterized by properties further on. An instance of ExperimentRepresentationObject pointing to the previous RepresentationObjectType indicates the use of AMR representation for the simulation and ExperimentProperty instances are used to point to the properties characterizing these representation like the number of cells.

uTypes:

- *Experiment: SimDB:simdb/experiment/Experiment*
- *Simulation: SimDB:simdb/experiment/Simulation*
- *AppliedPhysics: SimDB:simdb/experiment/AppliedPhysics*
- *ParameterSetting: SimDB:simdb/experiment/ParameterSetting*
- *TargetObjectType: SimDB:simdb/experiment/TargetObjectType*
- *ExperimentRepresentationObject: SimDB:simdb/experiment/ExperimentRepresentationObject*
- *RepresentationObjectType: SimDB:simdb/experiment/RepresentationObjectType*

Example of StarFormat instances for the Protocol and Experiment parts

2.2.4 Snapshot part

uType: SimDB:simdb/experiment/Snapshot

Each timestep of the simulation is stored as a Snapshot. For each snapshot, ObjectCollection instances referring to the cloud and the AMR (i.e the run's representation and target objects) are associated and characterized though the use of Statistics instances providing values for the Object's Property classes.

uTypes:

- *Snapshot: SimDB:simdb/experiment/Experiment*
- *ObjectCollection: SimDB:simdb/experiment/ObjectCollection*
- *Statistics: SimDB:simdb/experiment/Statistics*
- *Property: SimDB:simdb/object/Property*

Example of StarFormat instances for the Snapshot and PostProcessing parts

2.2.5 PostProcessor & PostProcessing part

uTypes: SimDB:simdb/protocol/PostProcessor & SimDB:simdb/experiment/PostProcessing

Finally the postprocessing applied to each snapshot in order to extract dense cores out of the molecular cloud is saved as a PostProcessor, another instance of the Protocol class referring to its parent Simulator and which can be described extensively in the same ways as the parent Protocol. Each Snapshot of the previous Simulations has multiple applications of PostProcessings referring to this PostProcessor child protocol and are referring to their parent Experiment as well as to the corresponding Snapshot through the use of the InputDataSet class.

These PostProcessings, or child Experiments, have as many Snapshot as extracted dense cores of matter, qualified as “clumps”. The clumps are described as TargetObjects of the PostProcessing and each individual one is saved as a different Snapshot of time 0 for the PostProcessing Experiment. They can therefore be characterized in identical ways as the cloud for the parent Experiment.

uTypes:

- *PostProcessor: SimDB:simdb/protocol/PostProcessor*
- *PostProcessing: SimDB:simdb/experiment/PostProcessing*
- *InputDataSet: SimDB:simdb/experiment/InputDataSet*

2.3 Implementation SimDB / VO-URP

2.3.1 VO-URP environment

To implement SimDB for StarFormat, we used VO-URP. This is a Java/JPA/PostgreSQL environment deployment tool provided by the Theory Working Group members Gerard Lemson and Laurent Bourges in order to help deploy a SimDB instance or any hosting project based on any UML data model.

One just needs to provide his data model drawn with the tool MagicDraw respecting some specifications including full descriptions, and eventually provide an XSD schema of the data. Then VO-URP generates from these info a full environment with SQL scripts and JAVA/JPA classes matching the model in order to create, handle and manipulate data with the database, either in PostgreSQL or MSSQLServer.

VO-URP also provides a basic web implementation to browse through the data, import or validate data from an XML matching the model.

For StarFormat, we just had to complete the SimDM UML model within MagicDraw with a couple complementary classes in order to provide extra services functionalities for the web interface. After configuring the options for the model, compiling VO-URP provided all these tools thanks to which the full web-service could be implemented. The simple interface to browse the database (SimDB-Browser) was integrated into the administrative part of the website.

2.3.2 Ingestion pipeline

An IDL pipeline was developed in order to compute statistics, images and histograms of properties from the resulting raw data of each snapshot. This tool generates all the specified metadata and resulting files for a snapshot or post-processing and stores them within a VOTable and a tar.gz archive of extra files the theorists wants to provide along with the data. After copying these files on the server, one can import all data into the database using a simple one-click interface within the administrative

part of the website. While the files are stored into a file system on the server, organized as a Project/Simulation/Snapshot/PostProcessing/Object directory tree, the VOTable, organizes the metadata within the SimDM classes. Using VO-URP's java classes and a VOTable parser based on CDS's SAVOT Java libraries this VOTable is then read and stored as classes of SimDM into the database.

A complete JSP servlet interface was also developed thanks to VO-URP's Java classes in order to provide snapshot browsing services on the website: one can easily navigate through the different simulations and their snapshots with a synthesized summary of all recorded data. After this browsing, the user is invited to choose a selection of snapshots and query the post-processings snapshots to identify desired clumps by their properties. Users can then extract as VOTable or ASCII files all the parameters of an experiment or the properties of a clump or a set of extracted clumps from a post-processing.

Connecting the service with RAMSES post-processing command-line routines on some snapshots raw data, extraction of small subsets of resulting data is allowed to the users after identification of a desired clump in order to perform their own analysis of the results.

2.4 Comments

The only real issue that arose through SimDM's implementation was the need within StarFormat to provide histogram statistics of the molecular cloud: the simple use of the Statistics class was not sufficient to provide such feature, except by organizing the values within a property group which did not seem sufficient for a complete representation of such tables of values.

So we derived a natural set of complementary classes as StatisticsTables referring not only to one property but two. StatisticsTableValues providing values for both properties are linked tot this StatisticTable. This way, classes of statistics can be stored as well. Thanks to the VO-URP's functionalities, such change to the model is easy to handle by simply updating the UML model and running again the Java classes generation on the new model.

Apart from this specific need, the genericity and the richness of the Simulation data model allowed a good and simple description of various kinds of simulations considered within the StarFormat project. Even though we are manipulating big sets of data with a lot of characterizing properties needed to identify specific subset results of a simulation, everything could be mapped properly within a database with which a browsing and querying website could easily communicate thanks to the VO-URP project.

3 PDRDB – Micro-physics simulations

PDRDB is a SimDB service aiming at publishing in the Virtual Observatory models of interstellar clouds computed by the Meudon PDR code. The Meudon PDR code, is a 1D stationary code that computes the atomic and molecular structure of interstellar clouds. At each position of the cloud, the code computes several physical quantities as the abundance of hundreds chemical species, level excitations, gas temperature, grains charge and temperature, ... It also computes properties of the whole cloud as column densities, line intensities and spectra that can be used to interpret observations.

Compared to the STARFORMAT project presented in section 2, PDRDB deals with “micro-physics” simulations.

We present here how we used SimDB to publish in the Virtual Observatory PDR models. In section 1, we present the objectives of the service, in section 2, we present the instantiation of the data model. Section 3, provides some technical information on the way we did the development of the service to facilitate its maintenance and further development. Finally, section 4 presents the advantages and the limits of the service.

3.1 Description and objectives

One of our objectives is to publish sets of models corresponding to different interstellar clouds (diffuse clouds, PDRs, extragalactic media) to facilitate the interpretation of observations in the interstellar medium. These developments were motivated by the new generation of instruments HERSCHEL and ALMA. We wish the service provides “observables” as line intensities and column densities but also the chemical and physical structure computed in the models. This is mandatory to make sure the users will have all the information to understand the physics behind the model.

The discovery of simulations is possible in two ways:

- 1) Search using input parameters. For example, a user can search a model of interstellar clouds with a proton density about 1000 cm^{-3} and a UV radiation field about 100 times the interstellar standard radiation field.
- 2) Search using outputs. For example, a user can search all simulations with the column density (or a line intensity) of species X between a minimum and a maximum value. This is useful to quickly obtain some guess to interpret observations.

Once a model has been found, through the web portal, it is possible to extract some quantities from the simulation through services in the web page. When relevant, these data can be sent from the web page to VO-Tools using SAMP. Since the number of computed physical quantities is large and heterogeneous, only the most important ones for the interpretation of observations are provided through the web portal. For the other ones, the user can download the full result of the simulation.

Some steps of the development of PDRDB with SimDB have been a challenge. The major difficulties came from the following specificities:

- a) The code computes a large number of heterogeneous physical quantities:

Together the number of species, populations in levels and lines intensities represent more than several ten thousands quantities. As a consequence, we only store in the database those useful to find a model.

Then, services on the web page can access to the other data stored in filesystem.

b) The code is frequently modified:

We often have to add new species in the chemistry, new physical processes in the code and we have to update frequently the atomic and molecular data used by the PDR code. Some of these modifications produce new physical quantities in the outputs that we want to publish in PDRDB. As scientists, we do not wish to modify the database each time the code is modified. The database has to update itself depending on the new ingested simulations. This has been solved structuring and documenting the outputs of the codes and developing a sophisticated ingestion pipeline using VO-URP. This is explained in section 2.4.

3.2 SimDM implementation on PDRDB

The simulation DM is an abstract DM. At some steps of the implementation, we did some choices but other ones could have been possible.

3.2.1 - Protocol part

uType: SimDB:simdb/protocol/Protocol

The protocol part defines the code used to produce the experiments. As many other microphysics codes, the Meudon PDR code can be divided in three parts:

- 1) Fortran source codes
- 2) Data files. They contain chemical reaction rates, atomic and molecular data as Einstein coefficients, collision rates, levels energies of atoms and molecules, etc. In the case of the Meudon PDR code, this represents several tens of thousands values. Some of them are well known, other ones are subject to important uncertainties.
- 3) Input parameters (proton density in the cloud, intensity of the UV radiation field, metallicities, flux of cosmic rays, some flags to control the algorithms or the way some physical processes are computed,)

3.2.1.1 Protocol class

uType: SimDB:simdb/protocol/Protocol

The first question is to determine if the Data have to be considered as input parameters or not.

If they were considered as input parameters, this would allow easy queries on the SimDB service to search for simulations produced with different values for atomic and molecular data. For example, it would be possible to publish sets of results with different chemical rates and to search simulations in the database with queries on some of these rates. This could be useful, but it would then be very difficult for publishers to manage a so large number of input parameters.

In the case of PDRDB, we considered that is a not a mandatory service we wish to provide in a VO context and, to simplify the implementation, we choose to not consider these data as input parameters.

As a consequence, we consider that the protocol in PDRDB is the combination of the Fortran source codes and all the atomic and molecular data and chemical reaction rates.

3.2.1.2 InputParameter

uType: SimDB:simdb/protocol/InputParameter

The InputParameter class is used to list the input parameter of the code. In our case, this corresponds to the parameters that are frequently modified at each run to explore space parameters. As mentioned above, we do not consider atomic and molecular data or chemical reaction rates as InputParameter.

InputParameters are for example the density of the cloud, the intensity of the incident UV radiation field on the cloud, the properties of the grains, the metallicities, etc.

3.2.1.3 ParameterGroup

uType: SimDB:simdb/protocol/ParameterGroup

Because the number of input parameters is important in PDRDB, several ParameterGroups are defined. They correspond for example to «metallicities» that contains the InputParameters He/H, C/H, O/H, ... or Grains properties that contains the parameters of grains (minimum size, maximum size, ...)

3.2.1.4 - Algorithms

uType: SimDB:simdb/protocol/Algorithms

The Algorithms class is used to provide some information on the main algorithms used in the code. PDR codes take into account a large number of physical processes with very different algorithms. This class is not very useful for us because the list of algorithms used in the code is not really a way to define the capabilities of the code. More precise information is required. For example we can say that the code uses an «exact radiative transfer method» but the important information is to know if overlapping of lines is taken into account or if radiation emitted by dust can affect non locally level populations of molecules. This information is provided in the documentation of the code but not in the SimDB service.

3.2.1.5 - Physical processes

uType: SimDB:simdb/protocol/Physics

Even if not precise enough (see 2.1.3), this can be used to provide an overview of the main physical processes in the code.

Example: Radiative transfer.

3.2.2 Experiment

uType: SimDB:simdb/experiment/Experiment

An experiment corresponds to a published run of the code.

3.2.2.1 Snapshot

uType: SimDB:simdb/experiment/Snapshot

The Meudon PDR code is a stationary 1D code. We do not have snapshots with the meaning it can have in the context of cosmological simulations for instance. In the SimDB implementation, we consider that for each experiment we have a single snapshot that corresponds to the output of the code, the stationary state.

3.2.2.2 TargetObject

uType: SimDB:simdb/experiment/TargetObjectType

TargetObjects represents the astrophysical object that has been simulated. For PDRDB, most of the time this is «Interstellar Cloud». That could also be «disks» when a model has been run to simulate a protoplanetary disk.

3.2.3 Object

uType: SimDB:simdb/object/Object

This part is a heritage of the protocol and of the experiment parts. We describe first, the ObjectType and Properties for the protocol, then for the experiment.

3.2.3.1 – RepresentationObjectType and Properties for the protocol

uType: SimDB:simdb/protocol/RepresentationObjectType

At each position of the grid, the code computes several quantities for the gas and the grains. We consider two ObjectTypes : gas and grains with the attribute Mesh Cell.

The properties of the RepresentationObjectType are all the quantities computed locally. In the Meudon PDR code, we have several thousands of these properties. Only the most important ones to search models in the database are implemented.

Examples for the RepresentationObjectType, gas, with in bold face, those that are stored in the database:

- **gas temperature, proton density, and ionization degree**
- **abundances of H, H₂, C⁺, C, CO, ...**
- levels populations of CO in J=0, J=1, ... of H₂ in v=0 J=0, ...
- local line emissivity ($\text{erg cm}^{-3} \text{ s}^{-1} \text{ Ang}^{-1} \text{ std}^{-1}$) of H₂ from v=0 J= 2 to v=0 J=0
- heating rate by photoelectric effect, by cosmic rays, ...
- density of energy per bin of wavelength
- ...

Examples for the RepresentationObjectType : grains

- **size of grains**
- **temperature of grains**
- **charge of grains**

Associated to an experiment, the statistics provide the mean, max, nominal values of these properties. These numbers in the database are used to provide some information on the run.

3.2.3.2 – TargetObjectType and Properties for the Experiment.

uType: SimDB:simdb/experiment/TargetObjectType

The properties of the TargetObjectType correspond to the quantities computed for the whole simulated astrophysical object. In PDRDB they correspond to integrated quantities as column densities or line intensities. They are very important in this service because they are the quantities observers are looking for.

Example of TargetObjectType properties :

- column density of H, H₂, CO, ...
- column density of H₂ in level v=0, J=0, in level v=0, J=1, ...
- emergent line intensity of CO from J=2 to J=1, ...

The statistics provide the mean, max, nominal values of these properties. These numbers in the database are used to search simulations on outputs.

3.2.3.3 PropertyGroup

uType: SimDB:simdb/object/PropertyGroup

Because of the large number of properties, we group them using the PropertyGroups. So we have for example a group “abundances” that contains a list of properties, the abundances per element. In the same way, we have a group “column densities”.

3.2.4 - Projects

uType: SimDB:simdb/Project

Projects are used to group together groups of experiments to facilitate searches in the PDRDB and its management. The way they are used is subjective.

Examples :

- Diffuse clouds : could be a project grouping a set of models with different input parameters corresponding to parameters of diffuse interstellar clouds.
- Damped Lyman Alpha Systems : could be a group of simulations with extragalactic metallicities to study the chemistry of diffuse intergalactic medium.
- Horsehead nebula : could be a project grouping all the models (from different protocols) of the horsehead nebula.

SimDB classes used for PDR's implementation:

3.3 Implementation SimDB / VO-URP

In this section we present how we implemented SimDB on PDRDB. This is not the only way this could have been done, nevertheless, this may give some ideas on how to avoid some difficulties in the development of VO services on microphysics simulations.

PDRDB is composed of :

- a database
- an ingestion pipeline
- Services

3.3.1 - Database

The PDR DataBase was built as a full implementation of the Simulation Data Model as a PostgreSQL database communicating with a Java/JPA interface thanks to the use of the VO-URP environment tool provided by the Theory Working Group members Gerard Lemson and Laurent Bourges.

3.3.2 - Ingestion of data in PDRDB

The ingestion of data in PDRDB has been one of the major difficulty from the beginning of the project: The Meudon PDR code is frequently modified to compute new physical or chemical processes. As a consequence, the SimDB service had to be flexible enough so that we avoid, each time we wish to introduce in PDRDB a new protocol or new experiments (with eventually new physical quantities in the outputs, so new properties) to have to modify manually the database.

Indeed, we want the scientist register new simulations in a very transparent way.

This difficulty should be common to many microphysics codes. We partially avoided it thanks to :

- 1) a modification of the outputs of the code to describe in details the list of physical quantities computed in the code.
- 2) the use of templates in the ingestion pipeline. These templates contain the list of the quantities that have to be extracted from the outputs of the code and imported in the database. They are in XML format and can be easily modified by any scientist.

3.3.2.1 Metadata describing the outputs of the PDR code

To develop PDRDB, we had to modify the output format of the PDR code. Previously the code produced a binary file containing all the computed quantities. This binary file could be difficult to read on different architectures and the physical quantities were not properly described in it.

We modified the PDR code so that it produces 2 output files :

1. a FITS file containing the data. FITS has been chosen because :
 - i. it is a format easy to read on any architecture
 - ii. data are structured in tables and so it is simple to extract specific quantities or to write new ones in a structured form.

Note that FITS may not be the best choice because of its limitations. HDF5 or some other formats could have been more appropriate.

2. a XML file describing the content of the FITS file. This XML file follows the syntax of VO-TABLE because many routines have been developed in the VO to deal with them (savot for example).

An ID that is present in the FITS file as keyword and in the XML file identifies each quantity. Associated to this ID, the XML file contains a human readable name of the quantity, a full description, units, UCDS, ...

These two files contain all the quantities computed by the code, the input parameters of the model, and some of the atomic and molecular data used to produce the simulation.

The XML file also provides the organization of properties as groups. Each group has also an ID. For example, there are hundreds properties “ABUNDANCE_X” where X is the name of the chemical species and a group ABUNDANCE.

3.3.2.2 – Templates and ingestion pipeline

To document the simulations within the database all results from the code are not necessary, so we first made choices on which parameters and properties to give access to and wrote a protocol and a experiment XML templates giving a short description of all basic groups of elements to include within

the database for a version of the code and a run.

Thanks to the JSP servlet interface developed with VO-URP's JPA classes to communicate with SimDB, we created an easy administrative interface which allows adding of new code versions and new models in a few clicks : after adding textual descriptions and placed the XML+FITS output within a specific directory, the ingestion pipeline matches the model fields along the templates definition, reads the values and stores them within the correct classes of the data model.

Thanks to this interface, whenever we want to introduce new quantities in PDRDB, one only needs to add the corresponding ID in the template file.

3.3.3 Services

In order to help users browsing data, we provide them some tools for visualization and interoperability. Concerning the spectra :

3.3.3.1 – Visualization

The PDRDB website allows the user to display, compare and save spectra available for a given experiment. This functionality is a java applet implemented thanks to the JFreechart library. Getting and plotting data is a 2-step process.

First of all we query the server to know which types of spectra are available for a given PDR experiment (incident or emergent with different angles for example). The response is an xml document. Then the applet parses this document to create its own GUI according to the type of spectra available.

The user can now query the service to get a spectrum. It is returned by the server as a simple csv file containing 2 columns (wavelength/intensity). The plotting is managed by the JFreechart API (we use it to draw histograms on the Starformat database too).

The data can be saved as ASCII or VOTable files.

3.3.3.2 - Interoperability

All the displayed spectra can be sent to another VO applications through a SAMP hub. This functionality has been implemented thanks to the WebSampConnector library developed by VO-Paris Data Centre. It consists in an java applet that can connect to a SAMP hub, send and receive messages, and some JavaScript functions that create the interaction between the web page and the applet.

To send data from the visualizer to the SAMP applet, we have to establish a communication between the 2 applets. This is achieved thanks to the JSObject (netscape.javascript.JSObject).

This object can call JavaScript functions from the web page containing the applet :

```
//array containing the parameters of the JavaScript function
String[] params = {"v", id, id, votableUrl};
//call the function directly by its name
jsInterface.call("sendSampMsg", params);
```

All the selected spectra in the interface are then sent to all the VO applications connected to the hub.

4 - Millenium run¹: simulation, post-processing and publication

In 2004, the *VIRGO consortium*² ran a cosmological **simulation** modeling a large part of the universe. For this simulation they used a particular version of the **simulation code** *Gadget*³ written by Volker Springel. This simulation code needs a certain (large) number of **input parameters** to be set. Some of these refer to physical concepts, such as parameters defining the cosmology (Hubble constant, density parameter Ω in dark matter, cosmological constant, and baryons etc) some numerical such as the gravitational smoothing length. The simulation started from initial conditions generated from a predefined power spectrum in the particular cosmology. The simulation code approximates various **physical processes** including gravity, hydrodynamics, star formation and black hole formation using appropriate **numerical algorithms** (TreePM, SPH). The simulation code allows one to choose which physical processes one wants to include. For the Millennium Run only gravity was included. The processes act on **simulation objects** (here point particles) that on their own, or in aggregate represent **real world objects** (here clusters, or Large Scale Structure). The simulation objects have a number of properties, some constant (identifier, mass) some variable that are evolved forward in time by the simulation code. The code allows me to choose which properties one may want to calculate (here [x, y, z, vx, vy, vz,]).

The simulation produces **results** that consist of collections of the simulation objects, one for each type of simulation object that is included. In this simulation each output data set is a **snapshot** taken at a particular time. The results are stored in multiple **files** (512 per snapshot). These files have a complex **format**, the particles are ordered using an index based on a space-filling curve, and the files contain hash tables that act as indexes to optimize retrieval of spatial sub volumes.

The Millennium simulation contained about 10 billion particles in a box of size 500Mpc/h. It had 64 snapshots, including the initial conditions that were calculated using a separate code. Each snapshot was about 400GB.

The snapshots produced by the simulation were used as input for various types of **post-processing**. For example a **cluster finder** using the *friends-of-friends* (FOF) algorithm was run on each of the snapshots. This produces *FOF groups*, which are dark matter halos with more complex properties (radius, various types of mass, velocity dispersion etc). The results are stored again in 512 files per snapshot.

These files with FOF groups are used as input for the SUBFIND code that detects substructures in the FOF groups, which correspond to bound “sub-halos”. These sub-halos evolve in time by merging with each other. The resulting merger trees are detected using a further post-processing algorithm that uses the particle data and the sub-halo files.

These merger trees are the input for a simulation code that implements galaxy formation using so called semi-analytical algorithms. These algorithms model physical processes such as gas cooling, star formation and evolution, supernovae feedback etc.

All the post-processing results are also stored in a **relational database**, which is published through a **web application** that allows users to submit SQL queries to the database.

The simulation code Gadget and the Millennium Run Simulation itself are mapped to the data model in full detail in two XML documents on the GoogleCode specification site.

¹ <http://www.mpa-garching.mpg.de/millennium>

² <http://www.virgo.dur.ac.uk>

³ <http://www.mpa-garching.mpg.de/gadget>

5 - S3

S3 is a proposal for a simple protocol aimed at querying and retrieving results of, or even running, “micro-simulations”. S3 is defined by a number of simple HTTP GET requests :

1. **Metadata:** Returns the parameters defining the service.
2. **Data query:** Returns information about files that exist for given (ranges of) parameters.
3. **Retrieve file:** Returns a particular file.

The result of a **Metadata** query is a VOTable describing the service in natural language and a list of PARAM elements, one for each parameter accepted by the Data query request. These parameters may correspond to parameters of the model⁴, but can also contain extra service specific parameters.

The **data query** finds models corresponding to particular (ranges of) values for these parameters, selected ones of which can be downloaded using the **Retrieve file** request.

Most of the S3 protocol can be described using SimDB/DM concepts. In the next section we will give this interpretation. It will then be seen that in theory the discovery, i.e. Data Query part of any S3 service could be completely supported by the SimDB protocol. In the next section we show the role S3 might play as a simplified protocol when the context is constrained.

5.1 Data model for a S3 service

One can argue that a S3 service is a SimDB/WebService giving access to results of SimDB/Simulations (“models” in S3) that are all run using the same SimDB/Simulator.

The **Metadata** request returns an English description and a list of parameters, some of which are SimDB/InputParameters, others may correspond to SimDB/Property-s used to characterize SimDB/TargetObject-s.

The **Data query** part searches for SimDB/Simulations executed with the protocol and having particular SimDB/ParameterSetting.value-s or SimDB/Characterisation.value-s (in case the S3/parameter corresponds to a value obtained after the experiment was run). Each S3 model presumably produces a single result. Partially to accommodate S3 we have introduced the generic SimDB/Result type in the model, since the SimDB/Snapshot class is used to describe 3+1D simulations.

So we see that the SimDM data model contains most concepts required in a potential S3 data model. Some features of S3 could be added to SimDB to complete this support. For example a new “S3” literal in the ServiceType enumeration and the more generic Result class has already been added. SimDB/WebService might be declared (using a reference) to “implement a” (rather than “be a”) SimDB/Protocol in its own right, which allows it to define its own extra parameters.

From the SimDB model we can also derive suggestions as to what might be missing from S3. For example a description of the type and contents of the results is missing, i.e. there is no SimDB/RepresentationObjectType collection, nor a SimDB/TargetObjectType or SimDB/TargetObjectProcess. Curation is not included either, nor a possible aggregating SimDB/Project type. Particularly for “micro-simulations” a parameter study may be a frequent type of experiment, which can be described this way.

These features can be added so that S3 services and their underlying experiments can be found in a SimDB implementation. The current S3 protocol does not address that issue, but it seems clear that without much work, and using the SimDB approach this is easily accommodated.

⁴ A “model” in S3 can correspond both the Simulator and Simulation concepts in SimDB

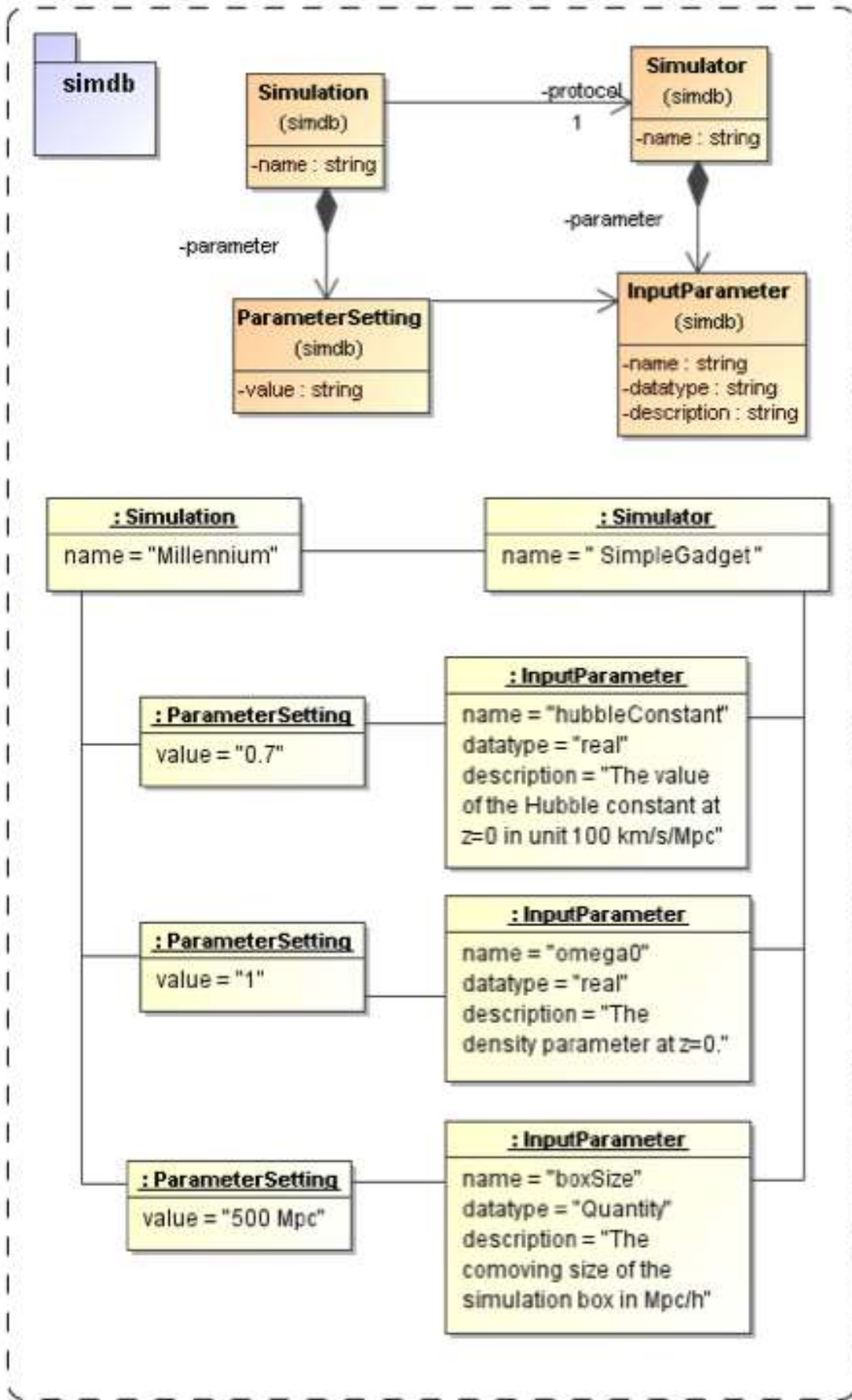
5.2 S3 as a TAP service.

To further analyse the relation between S3 and SimDB it is useful to interpret S3 as a TAP service. In this interpretation the Metadata query returns the metadata of a single table the columns of which correspond to the parameters of the S3 service. The S3 data query can be translated into a particular ADQL query⁵ against that table. Here we may have to assume that the table contains also the access reference to the model file represented by a row in the table.

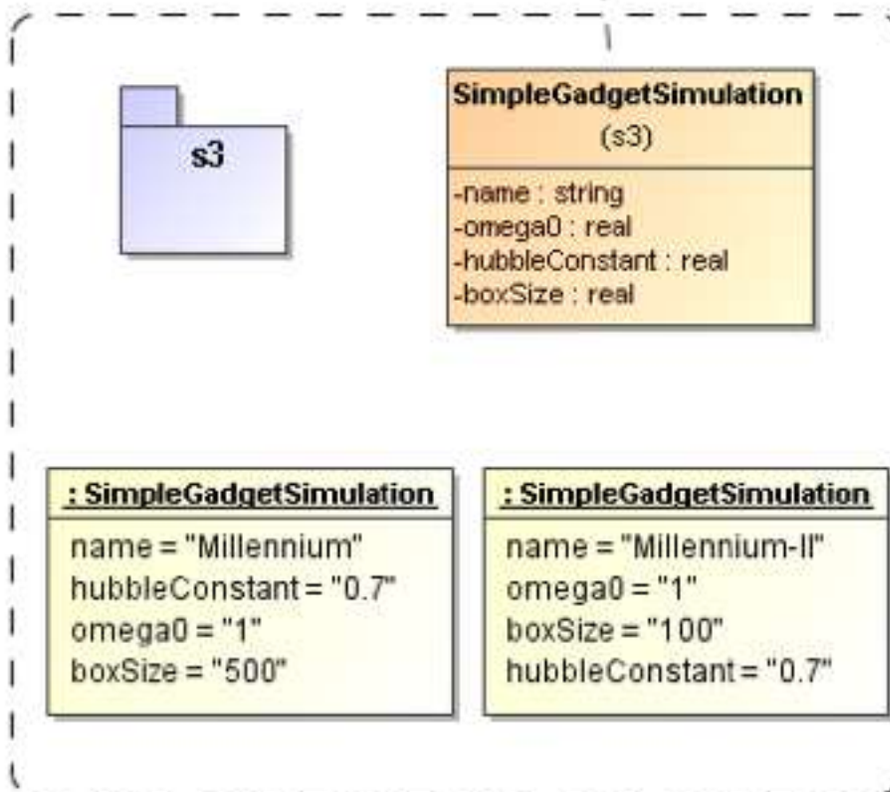
We can compare this “S3-TAP” service to the SimDB-TAP service. The difference is solely in the data model. To illustrate the difference, but also the correspondence between these models consider Figures below. The first one shows a greatly simplified cut from the SimDB model, concentrating on parameters and ignoring some tricky details. In addition to the classes we show an instance diagram defining a Simulator named “SimpleGadget” and a simulation named “Millennium” that was run using the SimpleGadget simulator. We define some InputParameters on the Simulator and add corresponding ParameterSettings to the Simulation.

The second figure shows a data model that we might have created if our task had been to only model “SimpleGadget” simulations. In that case the definition of the model, its “meta-data” if you wish, will contain knowledge about the SimpleGadget simulator that therefore does not have to be represented in the model itself. The model for simulations can now be made much more explicit. We can define a Gadget2Simulation class, which contains named and typed attributes representing the input parameters.

⁵ In fact the ParamQuery “mode” of TAP may be even closer to S3.



This class describes simulations run with the "SimpleGadget" simulation code. It allows the following attributes to be set:
 name: the name of the simulation
 omega0: The density parameter at z=0.
 hubbleConstant: The value of the Hubble constant at z=0 in unit 100 km/s/Mpc
 boxSize: The comoving size of the simulation box in Mpc/h



The information content in the two instance diagrams is basically the same. But obviously the second model is much simpler. In the SimDB version of the model all information is contained in the instances (the data) and can be retrieved from there. One only needs to understand the concept of a Simulator and a Simulation etc. In the S3 model the information is distributed over instances and model definition (metadata).

The SimDB model can capture simulations run according to many different protocols; the S3 version supports only SimpleGadget simulations. If one only is interested in SimpleGadget simulations though, clearly the S3 version of the model is easier to use.

Appendix A: SimDM Classes Instantiation

The table below summarises some key elements defined as instances of the different classes of SimDB within the two different implementations.

Class	PDR	Starformat
Project	Diffuse Clouds PDRs Extragalactic Clouds	Molecular Clouds Dense cores Turbulence
Protocol	PDR 1.4 – Chemistry10 PDR 1.4 – Drcnosd	Ramses-MHD 2 Ramses-MHD 3
RepresentationObject	AMR cells (gas, grains)	AMR cells
TargetObject	Interstellar cloud	Molecular cloud
Snapshot	Single result (stationary)	Snapshot (time iteration)
Property (Represent.)	Abundance H, H2, ... Gas temperature ...	n/a
Property (TargetObj.)	Column density H, H2, ... Line intensities	Statistics on: Temperature, mass, density, ...
PropertyGroup	Abundances Column densities	n/a
PostProcessing	n/a	Clump extractions
TargetObject	n/a	Clump
Property (TargetObj.)	n/a	Temperature Mass Density

References

- [1] SimDM proposed recommendation
<http://www.ivoa.net/internal/IVOA/SimRFC/PR-SimulationDataModel-v.1.00-20110906.pdf>

Appendix A: XML serializations

A.1 Protocol for STARFORMAT

```
<?xml version="1.0" encoding="UTF-8" standalone="yes"?>
<ns2:aSimulator xmlns:ns2="http://www.ivoa.net/xml/SfsDB/0.9-20101102">
  <identity publisherDID="Ramses3-_-MHD_02" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/Simulator/2"/>
    <name>Ramses3 - MHD_02</name>
    <description><p>&The simulations have been performed with the &lt;a
href="http://magnet.ens.fr/"&gt;RAMSES-MHD code&lt;/a>; (&lt;a
href="http://adsabs.harvard.edu/abs/2002A&26A...385..337T"&gt;Teyssier 2002, A&A, 385,
337&lt;/a>;, &lt;a href="http://adsabs.harvard.edu/abs/2006A&26A...457..371F"&gt;Fromang
et al., A&A, 457, 371&lt;/a>). This is a mesh refinement code, implying that it can
increase locally the spatial resolution by adding new cells in the computation. It uses the
Godunov method and constraint transport method to maintain the divergence of the magnetic
field equal to zero.&lt;/p></description>
    <referenceURL>http://magnet.ens.fr/code</referenceURL>
    <created>2011-01-10T17:33:25.113+01:00</created>
    <updated>2011-01-10T17:34:14.609+01:00</updated>
    <status>published</status>
    <contact>
      <identity publisherDID="patrick.hennebelle@ens.fr" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/Contact/5"/>
        <role>publisher</role>
        <party publisherDID="patrick.hennebelle@ens.fr" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/Party/8"/>
      </contact>
      ...
    <code>http://magnet.ens.fr/</code>
    <version>MHD_02</version>
    <parameterGroup>
      <identity publisherDID="m_refine" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/ParameterGroup/2"/>
        <description>During the run, the mass within one cell should be smaller than a given
value. This value is  $10 \times (2^{\text{levelmin}} + 1)^{\text{sup}} \times m_{\text{refine}} \times 3^{\text{sup}}$ . If the density becomes
larger than this value, then the resolution is increased.</description>
        <name>Refinement based on density threshold</name>
        <member>
          <identity publisherDID="m_refine-_-0000" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/ParameterGroupMember/39"/>
            <parameter publisherDID="m_refine-_-0000" xmlId="InputParameter_31"/>
          </member>
          <member>
            <identity publisherDID="m_refine-_-0001" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/ParameterGroupMember/30"/>
              <parameter publisherDID="m_refine-_-0001" xmlId="InputParameter_46"/>
            </member>...
        </parameterGroup>
      ...
    <representation>
      <identity publisherDID="amr" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/RepresentationObjectType/1"/>
        <name>AMR</name>
        <description>The code is based on adaptive mesh refinement scheme.</description>
        <property>
          <identity publisherDID="cell_level-_-0000" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/object/Property/14" xmlId="Property_14"/>
            <name>Number of cells per level-_-0000</name>
            <datatype>real</datatype>
            <cardinality>1</cardinality>
            <description>Number of cells per AMR level. Below levelmin, no cell is
active.</description>
            <isEnumerated>>false</isEnumerated>
          </property>
          ...
        </propertyGroup>
    </aSimulator>
```

```

    <identity publisherDID="cell_level" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/object/PropertyGroup/1"/>
    <name>Number of cells per level</name>
    <description>Number of cells per AMR level. Below levelmin, no cell is
active.</description>
    <member>
        <identity publisherDID="cell_level-_0000" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/object/PropertyGroupMember/16"/>
        <property publisherDID="cell_level-_0000" xmlId="Property_14"/>
    </member>
    ...
</propertyGroup>
<type>hierarchical mesh cell</type>
<label>Hierarchical_Mesh_Cell</label>
</representation>
<parameter>
    <identity publisherDID="boxlength" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/InputParameter/53"/>
    <name>Box size</name>
    <datatype>real</datatype>
    <cardinality>1</cardinality>
    <description>The actual size for one dimension of the boxcube.</description>
    <isEnumerated>>false</isEnumerated>
    <label>phys.length</label>
</parameter>
...
<algorithm>
    <identity publisherDID="AMR" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/Algorithm/1"/>
    <name>Adaptive Mesh Refinement</name>
    <description>amr</description>
    <label>Adaptive_Mesh_Refinement</label>
</algorithm>
<physicalProcess>
    <identity publisherDID="mhd" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/Physics/2"/>
    <name>Magneto Hydro Dynamics</name>
    <description>Magneto-hydrodynamics is treated in this simulation. This implies that the
gas is subject to Lorentz forces while the evolution of the magnetic field is dictated by
the induction equation.</description>
    <label>magnetohydrodynamics</label>
</physicalProcess>
...
</ns2:aSimulator>

```

A.2 - Experiment from STARFORMAT

```

<?xml version="1.0" encoding="UTF-8" standalone="yes"?>
<ns2:aPostProcessing xmlns:ns2="http://www.ivoa.net/xml/SfsDB/0.9-20101102">
    <identity publisherDID="DECAY-_DECAY-_00030-_clumps_100000" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/experiment/PostProcessing/5"/>
    <name>00030-_clumps_100000</name>
    <description7>
    <referenceURL>http://roxxor.obspm.fr/StarFormat</referenceURL>
    <created>2011-01-10T17:33:26.296+01:00</created>
    <status>published</status>
    <contact>
        <identity publisherDID="patrick.hennebelle@ens.fr" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/Contact/22"/>
        <role>owner</role>
        <party publisherDID="patrick.hennebelle@ens.fr" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/Party/19"/>
    </contact>
    ...
</parameter>

```



```

</target>
<result xsi:type="ns4:GenericResult" xmlns:ns4="http://www.ivoa.net/xml/SfsDB/0.9-
20101102/simdb/experiment" xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance">
  <identity publisherDID="DECAY-_-DECAY-_-00030-_-clumps_100000-_-0000"
ivoId="ivo://www.g-vo.org/SfsDB#SimDB:simdb/experiment/GenericResult/74"/>
  <product>
    <identity publisherDID="DECAY-_-DECAY-_-00030-_-clumps_100000-_-0000-_-Clump"
ivoId="ivo://www.g-vo.org/SfsDB#SimDB:simdb/experiment/Product/74"/>
    <numberOfObjects>1</numberOfObjects>
    <characterisation>
      <identity publisherDID="dens" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/experiment/StatisticalSummary/317"/>
      <statistic>nominal</statistic>
      <numericValue>
        <value>252.16485596</value>
        <unit>cm<sup>3</sup></unit>
      </numericValue>
      <axis publisherDID="dens" xmlId="Property_125"/>
    </characterisation>
    <characterisation>
      <identity publisherDID="mass" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/experiment/StatisticalSummary/2266"/>
      <statistic>nominal</statistic>
      <numericValue>
        <value>0.12910716013</value>
        <unit>solar mass</unit>
      </numericValue>
      <axis publisherDID="mass" xmlId="Property_142"/>
    </characterisation>
    <characterisation>
      <identity publisherDID="posx" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/experiment/StatisticalSummary/964"/>
      <statistic>nominal</statistic>
      <numericValue>
        <value>0.04296875</value>
        <unit>pc</unit>
      </numericValue>
      <axis publisherDID="posx" xmlId="Property_128"/>
    </characterisation>
    ...
    <objectType publisherDID="Clump" xmlId="TargetObjectType_4"/>
  </product>
</result>
...
<protocol publisherDID="Rameses3 MHD Clump Extraction-_-MHD_02_Clumps" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/protocol/PostProcessor/4"/>
  <primaryExperiment publisherDID="DECAY-_-DECAY" ivoId="ivo://www.g-
vo.org/SfsDB#SimDB:simdb/experiment/Simulation/3"/>
</ns2:aPostProcessing>

```

A.3 - Protocol for PDRDB

```
<?xml version="1.0" encoding="UTF-8" standalone="yes"?>
<ns2:aSimulator xmlns:ns2="http://www.ivoa.net/xml/PDRDB/0.5-20090814">
  <identity publisherDID="PDR_1_4_rev_345_Drcnosd_chi" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/Simulator/348"/>
    <name>PDR 1.4 rev 345 Drcnosd.chi</name>
    <description>&lt;b&gt;PDR 1.4&lt;/b&gt;&nbsp;: The 1.4 version corresponds to the PDR
code described in Gonzalez Garcia M. et al. (2008, A&A, 485, 127).&lt;br&gt;It includes
a new radiative transfer method computing in details I.R. pumping due to line and dust
emission.&lt;br&gt;&lt;b&gt;Drcnosd.chi&lt;/b&gt; : This chemistry is a deuterated
chemistry. It is adapted to the study of simple species in diffuse clouds, and in particular
to D and HD.&lt;/description>
    <referenceURL>http://pdr.obspm.fr</referenceURL>
    <created>2010-05-12T19:05:23.779+02:00</created>
    <updated>2010-05-12T19:05:23.779+02:00</updated>
    <status>private</status>
    <contact>
      <identity publisherDID="Franck.LePetit@obspm.fr" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/Contact/348"/>
        <name>Franck Le Petit</name>
        <address>LUTH, Observatoire de Paris-Meudon - 5, place Jules Janssen - F-92 195 Meudon
Cedex</address>
        <email>Franck.LePetit@obspm.fr</email>
        <telephone>01 45 07 75 66</telephone>
        <role>owner</role>
      </contact>
      <code>http://pdr.obspm.fr/PDRcode_Download.html</code>
      <version>1.4</version>
      <parameterGroup>
        <identity publisherDID="CLOUD_PARAMETERS" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/ParameterGroup/15"/>
          <description>TODO : Franck</description>
          <name>Cloud parameters</name>
          <member>
            <identity ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/ParameterGroupMember/172"/>
              <parameter publisherDID="DU_MODPARAM_R.FD_AVMAX" xmlId="InputParameter_56055"/>
            </member>
            <member>
              <identity ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/ParameterGroupMember/143"/>
                <parameter publisherDID="DU_MODPARAM_R.FD_NHINIT" xmlId="InputParameter_56020"/>
              </member>
            ...
          </parameterGroup>
          ...
        <representation>
          <identity ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/RepresentationObjectType/430"/>
            <name>Adaptive Mesh Cell</name>
            <description/>
            <label>Adaptive_Mesh_Cell</label>
          </representation>
          <representation>
            <identity publisherDID="GAS" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/RepresentationObjectType/351"/>
              <name>Gas</name>
              <description>Gas related properties</description>
              <property>
                <identity publisherDID="GAS_DU_OPTDEPTH.FD_OPTDEPTH" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/object/Property/55872"/>
                  <name>Optical depth</name>
                  <datatype>double</datatype>
                  <cardinality>1</cardinality>
                  <description/>
                </property>
              ...
            </propertyGroup>
            <identity publisherDID="DU_HEATING" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/object/PropertyGroup/2650"/>
              <name>Heating rate</name>
```

```

    <description>Heating rates by various processes in erg cm-3 s-1</description>
    <member>
      <identity ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/object/PropertyGroupMember/55690"/>
      <property publisherDID="GAS_DU_HEATING.FD_TOTALHEAT" xmlId="Property_55741"/>
    </member>
    ...
  </propertyGroup>
  ...
  <type>simple mesh cell</type>
  <label>gas</label>
</representation>
<parameter>
  <identity publisherDID="DU_MODPARAM_R.FD_AVMAX" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/InputParameter/56055" xmlId="InputParameter_56055"/>
  <name>Av of the cloud</name>
  <datatype>double</datatype>
  <cardinality>1</cardinality>
  <description>&lt;b>Size of the cloud &lt;br>in visual extinction given
as input parameter of the model &lt;br>Fortran variable : Avmax</description>
</parameter>
<parameter>
  <identity publisherDID="DU_MODPARAM_R.FD_NHINIT" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/InputParameter/56020" xmlId="InputParameter_56020"/>
  <name>Proton density (initial)</name>
  <datatype>double</datatype>
  <cardinality>1</cardinality>
  <description>&lt;b>Initial proton density &lt;br> given as input
parameters. &lt;br> &lt;i>Unit: cm-3 &lt;br> Note: If a density profil has
been used this parameter has no importance.</description>
</parameter>
<parameter>
  <identity publisherDID="DU_MODPARAM_I.FD_ISONESIDE" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/InputParameter/56027" xmlId="InputParameter_56027"/>
  <name>Flag : 1 or 2 side(s) model</name>
  <datatype>int</datatype>
  <cardinality>1</cardinality>
  <description>Flag: &lt;br>0 - two-side model &lt;br>1 - semi-infinite
cloud &lt;br>Determined by the value of radp in input parametersFortran variable :
isoneside</description>
  <isEnumerated>true</isEnumerated>
  <validValue>
    <identity ivoId="ivo://www.g-vo.org/PDRDB#SimDB:simdb/object/ValidValue/54"/>
    <description>Radiation field illuminates both sides of the cloud</description>
    <label>two-side model</label>
  </validValue>
  <validValue>
    <identity ivoId="ivo://www.g-vo.org/PDRDB#SimDB:simdb/object/ValidValue/52"/>
    <description>Radiation field illuminates the observer's side of the
cloud</description>
    <label>semi-infinite cloud</label>
  </validValue>
</parameter>
  ...
<algorithm>
  <identity publisherDID="" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/Algorithm/3"/>
  <name>Exact Radiative Transfer Method</name>
  <description/>
  <label>Exact_Radiative_Transfer_Method</label>
</algorithm>
<physicalProcess>
  <identity publisherDID="radiative_transfer" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/protocol/Physics/3"/>
  <name>radiative transfer</name>
  <description>should be read from a vocabulary file</description>
  <physicslabel>radiative_transfer</physicslabel>
</physicalProcess>
  ...
</ns2:aSimulator>

```

A.4 - Experiment for PDRDB

```
<?xml version="1.0" encoding="UTF-8" standalone="yes"?>
<ns2:aSimulation xmlns:ns2="http://www.ivoa.net/xml/PDRDB/0.5-20090814">
  <identity publisherDID="n1e3r1e2A20_10" ivoId="ivo://www.g-
vo.org/PDRDB#SimDB:simdb/experiment/Simulation/2"/>
  <name>n1e3r1e2A20_10</name>
  <description>Model of PDR done with the code version <b>PDR 1.4</b>;
(2009)<br/>The 1.4 version corresponds to the PDR code described in Gonzalez Garcia M.
et al. (2008, A&A, 485, 127).<br/><b>Geometry</b>; One
side model<br/><b>ISRF</b>; Draine's
units<br/><b>Thermal balance</b>;
solved<br/><b>Chemistry</b>;
chimie08.chi<br/><b>Recommendations</b>; Can be used to
interpret observations in PDRs for small molecules. Recommended for H2 emission. Must not be
used to interpret observations of carbonated chains above 2 carbons.</description>
  <referenceURL>http://pdr.obspm.fr</referenceURL>
  <created>2010-05-12T18:31:12.029+02:00</created>
  <updated>2010-05-12T18:31:12.029+02:00</updated>
  <status>private</status>
  <contact>
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vo.org/PDRDB#SimDB:simdb/Contact/2"/>
    <name>Franck Le Petit</name>
    <address>LUTH, Observatoire de Paris-Meudon - 5, place Jules Janssen - F-92 195 Meudon
Cedex</address>
    <email>Franck.LePetit@obspm.fr</email>
    <telephone>01 45 07 75 66</telephone>
    <role>owner</role>
  </contact>
  <targetObject>
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    <description>Cloud related properties</description>
    <property>
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      <cardinality>1</cardinality>
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      <ucd>phys.columnDensity</ucd>
    </property>
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    <propertyGroup>
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      <description>Column densities of chemical species in cm-2</description>
      <member>
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    <spatialSizePhysical>
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      <unit>mag</unit>
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  </product>
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  <type>min</type>
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  <objectType publisherDID="CLOUD" xmlId="TargetObjectType_3"/>
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      <unit>mag</unit>
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  <property>
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vo.org/PDRDB#SimDB:simdb/experiment/ExperimentProperty/19"/>
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vo.org/PDRDB#SimDB:simdb/object/Property/142"/>
  </property>
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  <type publisherDID="GAS" ivoId="ivo://www.g-
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  <protocol publisherDID="PDR_1_4_chimie08_chi" ivoId="ivo://www.g-
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