

MedeA HT: Automating Calculations with High Throughput Simulations

1 Introduction

Increasing compute power and the robustness of codes such as VASP, LAMMPS, and Gaussian have opened exciting possibilities for large-scale materials screening, materials discovery, and optimization. With its comprehensive databases and flexible software architecture, *MedeA* is perfectly suited for this type of high throughput computations. *MedeA HT* provides a convenient and powerful capability to take full advantage of high-throughput computations. *MedeA HT* consists of three components, namely the *HT-Launchpad*, *HT-Descriptor* and *HT-Correlations*, as illustrated below.

Common to all three components is the use of Structure Lists and the powerful Flowchart stage Foreach Structure Loop . The following describes the Structure Lists and then the two components: *HT-Launchpad* and *HT-Descriptor*.

2 Structure Lists

Purpose and Overview

A Structure List is a single file designed to contain a list of structures and associated properties that a user can create, modify, or use with full control. Being a single file, it can be managed and shared more easily than a collection of distinct structure files. The same list can contain at the same time structures in periodic boundary conditions (PBC, i.e. with cell parameters and eventually some symmetry) often referred to as periodic structure, or without PBC also often referred to as aperiodic structure. Each structure can contain several configurations, that can be for example the output of molecular dynamics simulations.

Structures Lists serve as input for high-throughput calculations and can be extended by computed properties. Together, this information contained in Structure Lists serves as input for the definition and calculation of descriptors. Structures Lists can also be used to compute properties by applying correlations. Furthermore, Structure Lists define the training and validation sets for the *Forcefield Optimizer*.

Properties can be defined for each structure independently, values being defined for each configuration if there are more than one.

A Structure List differs from the structural database used in *InfoMaticA* in the sense that it can contain structures of different classes (periodic or not) with eventually several configurations, and with a highly customizable set of properties. The properties definition and interpretation is therefore controlled by the user according to specific needs. Thus, Structure Lists tend to have the character of dynamic workspaces in contrast to the databases in *InfoMaticA*, which have more of a static character with permanent information such an experimentally determined crystal structure from a scientific publication.

To summarize, Structure List characteristics are:

- A single file for many structures
- Possibly several configurations per structure
- Complete control over the properties associated with each structure.

Such lists can be created automatically, and they can be defined, extended, and modified by the user in a Flowchart or a Structure List Editor.

The default file extension is *.sli*.

Note: Any extension can be used in the end, but it is recommended to use *.sli*.

3 Structure List Editor

This Section describes the features of the structure list editor that are mostly relevant in the context of highthroughput calculations. A comprehensive description of the structure list editor provides the Section MedeA Structure List Editor: Manage Your Structure Repository With Properties.

3.1 Opening

The Structure List Editor allows you to browse and modify the content of a new or an existing Structure List. In the main window's File menu, the structure list editor window is invoked from Structure List Editor .

The following three actions from the File menu of Structure List Editor can be invoked:

- New structure list : will ask for the name and location to store the new empty Structure List file. The file will be initialized, allowing you to start adding structures.
- Open structure list from disk : will ask to select an existing Structure List file on the local file system to display its content.
- Open structure list from job : will ask to select one or more Structure List files from jobs on the current JobServer. Each will be downloaded to the disk (name can be edited to be different from the file in the job) and the last one will be shown in the editor. The editor can be closed with the Close button or the File >> Close menu command.

3.2 Editor Overview

As shown below, the main structure list editor component is a structures table, with descriptive information displayed above the tabular view:

The descriptive part shows the file format, its size, and the full path. One can also control the number of displayed structures in the table, specifying the start and end index (clicking on the Apply button will update the view).

The Properties tab will show all properties in a table, as each structure can have its set defined, and as shown below the property:

DOCUMENTATION DOCUMENTATION

3.3 File Formats

A Structure List has one of the following two formats:

- Text format : data is written in plain ASCII format. One can edit the content with a simple text file editor but with a high risk of erroneous modification. Many molecular dynamics jobs will produce a trajectory file in that format (*Trajectory.dat* or files with *.traj* extension). One cannot apply all available operations with that format.
- SQLite format : information is organized with an SQLite database format, which offers a much higher level of performance when processing and updating lists containing a large number of structures.

By default, the SQLite format is used when creating a new Structure List. It is recommended to convert to SQLite whenever possible, which can be done with the File >> Convert to SQLite/text format. It will create a file of the other type than the one displayed.

3.4 Add Structures

There are several ways to add structures to the current Structure List and they are accessible from the Add structures menu:

The different new structures sources are:

From MedeA: : invokes a dialog to select opened structures in the *MedeA* main window.

From files : opens a window to select individual structure files.

From a SMILES list file : selects an ASCII file (default *.smi* extension) formatted with two columns: the first for a molecule name and the second with the SMILES code.

Structure list from disk : select another Structure List from disk and add all its structures (with their properties) to the end of this one.

Structure list from job: select another Structure List from job and add all its structures (with their properties) to the end of this one.

Split structure(s) into molecule(s) : selects a displayed structure in the *MedeA* main window and analyzes it different molecules (defined by bond connectivity, bond orders, atom elements). An aperiodic structure will be added for each molecule type, and every image will be added as a configuration.

Split structure(s) into molecule(s) with PBC : will do the same but with PBC.

3.5 Display

The information appearing in the structure table can be controlled by the Display menu selectors. The columns Order and Name will always be displayed.

The information displayed can be the structural information (Structural Formula , #atoms : number of atoms, #configurations : number of configurations, Symmetry and Cell parameters when in PBC) or user-defined properties. If a large number of properties is defined, only the first three will be displayed by default, others have to be selected from the Display menu to appear.

In case there are several configurations for a structure, the value shown in the table will correspond to the first configuration. Other configurations' property values can be seen by choosing to show the structure properties in the Properties table.

3.6 Ordering and Numbering Structures

Structures are stored in a given order in the list, which appears in the Order column in the structures table. It is possible to change the order in the table by sorting rows (structures) according to a column from the right-click popup menu commands Sort Ascending or Sort Descending , on a column title. Also, a given structure can be moved up or down by a right-click on its row with the corresponding command in the popup menu.

If the initial order of the rows in the table is changed by one of these methods, the actual structure order in the Structure List file remains unchanged. It is possible to apply the new displayed order to reorder the list internal ordering with the right-click popup menu command Save the order of the rows . This can be useful, for example, to change the processing order in a job.

4 Delete Existing Properties

The Properties >> Delete menu command brings up a small dialog window summarizing the list of all existing properties for all structures. They can be selected for being deleted. Once the OK button is used the data is deleted and the file is updated, *i.e.* the deleted data is no more recoverable.

5 Create a New Property

The Properties >> New Property opens a small dialog window allowing to add a property to all current structures in the list. One has to provide the following:

Name : the property name

Units (optional): units that can apply to this property

Type : has to be one real (default), integer, or string

Size : the number of values for this property, default: 1

Dimension : the values dimension, e.g. '3 3' would mean a 3 by 3 matrix

Default (optional): default value that will be added to all structures initially.

6 Edit Property Values

Values in user-defined properties can be interactively modified in several ways:

- Double click on the displayed value turns it in editing mode. Pressing the ESC key will dismiss the change, OK will confirm it. If there are several configurations for a given structure, this action will update the value of the first configuration.
- Copy, for example, values from a column in a spreadsheet and paste them in the target property column, using the right-click menu command Paste in \langle property $>$. Values will be added down starting from the row where the click was performed. This will update values for the first configuration in the updated structures.
- Call the value editor from the right-click popup menu Edit <property> value . Each configuration value can be edited (changing configuration value can be done with a selector). Changes are applied only once the OK button is used. Of course, this is mostly useful if the structure has more than one configuration.

6.1 Copy property values

When default copy shortcut (usually Ctrl $+/$ C) is pressed, and some rows are selected in the structure table, the content of the selected rows is copied to the clipboard and can later be pasted in e.g. an external spreadsheet. If some structures have more than one configuration, additional values corresponding to these additional configurations will not be included in the clipboard. Columns can be copied individually, using the right-click popup menu on a column title, as shown:

Either the full column can be copied to clipboard or the only values in selected rows.

6.2 Export Structures

Structures can be exported to individual structure files from the right-click popup menu:

- Individually to an individual structure file (*.sci* extension) with the Export structure <*number*> command, the *<number*> corresponding to the structure number in the row the click is made.
- All structures at once to individual structure files (*.sci* extension) with the Export all structures command. A filename prefix is used and the structure number and configuration (if more than one) are used for the final file name.
- All structures at once to the MaterialsDesign database with the Save All structure(s) to MD database command.

7 HT-Launchpad

The *HT-Launchpad* enables the automatic computation of materials properties for large sets of systems given in a Structure List. The two central items for this capability are (i) the Flowchart stage "ForEach Structure" and (ii) the Structure Lists described above.

The Flowchart stage *ForEach Structure* is conceptually simple yet extremely powerful. A file containing a Structure List is selected and by opening the Flowchart tab the various computational steps are specified in the form of a Flowchart which may contain stages running VASP, Gaussian, MOPAC, LAMMPS, and other steps. The results are added to the existing or new Structure Lists.

A checkbox enables parallel execution of the loop iterations, i.e. performing calculations on simultaneously on a sub-set of structures. The level of parallel execution is controlled by specifying the maximum number of jobs running simultaneously.

By checking the box "Advanced settings", it is possible to skip structures in the loop over structures and to include loops over multiple configurations of a single structure, as shown below.

To add computed properties to a Structure List, the desired item is specified in a dialog box as shown below.

The name in the column "Source" has a specific format such as *\$Eelectronic calc*. A list of these properties is given in this User's Guide in the chapter of MedeA Flowcharts.

The result of such a high-throughput calculation is stored in a Structure List as shown in the example below.

8 HT-Descriptors

The *HT-Descriptor* component of *MedeA HT* enables the definition and evaluation of descriptors. For example, if one is interested in the VASP energy per atom for the compounds given in the previous Structure List, one can define a descriptor which is simply the total energy computed with VASP divided by the total number of atoms.

To this end, open the Structure List with the computed VASP energies and select "Descriptors" from the pull-down menu "Properties".

A dialog box will appear which allows the definition and the evaluation of descriptor. In the present example, the descriptor is called "eatom" and is defined as follows.

Application of this descriptor to the Structure List with the VASP energies results in a new column with the desired values. The display of this and all other columns can be controlled by selecting the corresponding item in the pull-down menu under "Display".

E VASP (eV)

-53.586151

eatom

 -89310252

Symmetry

P42/MNM

atoms

 ϵ

MedeA: structure list edito

In this example, the list is sorted by increasing energies per atom, which is accomplished by a right-click on the header of the column "eatom" and selecting sorting by ascending values.

After this simple example, the following text describes the functionality of the *HT-Descriptor* in detail.

HT-Descriptors enable the user to assign properties and descriptors to each structure in a Structure List, thus allowing classification and ranking of structures. A descriptor can be defined, for example, as a function of computed properties like elastic coefficients and topological properties such as the existence of channels in a structure. Sets of descriptors can be stored in catalogs, reloaded, and applied to newly created structure lists. Thus, Descriptors together with Structure Lists are a powerful tool in the context of high-throughput calculations.

Descriptors computed with this tool are scalar properties that can be defined as functional, using a set of mathematical operators and variables. They can be computed on the fly or submitted to a Flowchart. Values will be computed for every configuration of every structure when applicable.

A catalog is a collection of properties that are jointly defined. This notion is important since properties can be interdependent. They can be saved and loaded from files for later reuse, using buttons Load Catalog and

Save Catalog . The Clear button will delete all existing descriptor definition data from this dialog.

The computation can be performed either interactively within *MedeA* by editing the Structure List or within a Compute descriptors List stage in a flowchart, which can be succeeding a stage used to create a Structure List, or simply to facilitate processing a large Structure List in a time-efficient manner.

8.1 Defining a descriptor

When the side button New is pressed, the dialog is configured to define a new descriptor. The following elements need to be provided:

- Name : It must obey the following rules: contain only lower case letters, numbers and the '_' character. The first character has to be a lower-case letter
- Expression : The actual descriptor as a mathematical expression. One can directly type the expression or use the drop-down menu buttons to insert functions or structural data as a variable. The syntax rules are explained later
- Description : A describing text that will be saved in the catalog file, if the catalog is saved. The purpose is to allow the descriptor's author to provide explanations to any other user that could use the catalog

file.

Pressing the OK button below the Description will register the descriptor in the catalog and, it will appear in the catalog box. As property names in Structure Lists are not restricted to lower cases and allow other characters, one may wish to give a more detailed name for the property using the Assigned property column. One can edit it by double-clicking on it and type *Enter* to validate the new name, or *Escape* to dismiss it.

Syntax Rules

A descriptor's expression obeys arithmetical edition rules: balanced parentheses, operands, and operators properly defined and ordered. The expression has to comply with the following syntax rules:

- *Functions* have to contain only upper-case letters and the '_' character. All accessible functions can be inserted at the cursor position with the drop-down button menus related to functions. Arguments are enclosed within parentheses and separated by commas if more than one argument is needed. The opening parentheses must follow immediately after the last function character, e.g. POWER(a,3.0)
- *Mathematical Operators*: the + / * signs are recognized. The power operator is accessible by the *POWER(,)* function. The *AND* or *&&* (Boolean AND) and *OR* or // (Boolean OR) are also recognized in Boolean conditions when applicable.
- *Operands* can be an integer or real numbers or property names. Property names can contain only lower-case letters, numbers, and the '.' character. Different types of properties can be used, and the drop-down menu button allows you to access by category. Some properties have a unit, and in that case, the default unit is mentioned in the list. If one wishes to use different compatible units, it has to be enclosed in parentheses right after the property name, e.g. *volume(mˆ3)*. The property value will be automatically converted to the provided units.
- Expression elements have to be *separated by spaces*.

Clicking on a descriptor in the display list will display its name, its expression, and comments. A descriptor can be deleted by clicking the Delete button, or edited (change its name, expression or comments) with the Edit button.

In interactive mode, pressing the Compute button will start the computation of all descriptors in the catalog. When applicable the values will be stored in the list under the assigned properties. If the latter is not alcreated defined for a structure, it will be automatically created. If it alcreated exists, the content will be updated with newly computed values. If for some structures in the list, the descriptor cannot be computed, no value will be evaluated and the corresponding property will be left empty. For example, if a descriptor relies on the cell volume, it will not be computed for non-periodic structures in the present list.

8.2 Functions

Functions are of different categories, each corresponding to a drop-down menu button. They are recognized as upper case character strings.

Generic Mathematical Functions

Name Definition

ABS() Returns the absolute value *ACOS()* Returns the arc cosine *ASIN()* Returns the arc sine *ATAN()* Returns the arc tangent *ATAN2()* Returns the arc tangent of the two arguments *CEILING()* Returns the smallest integer value not less than the argument *COS()* Returns the cosine *COT()* Returns the cotangent *DEGREES()* Convert radians to degrees *EXP()* Exponential of *FLOOR()* Returns the largest integer value not greater than the argument *LOG()* Returns the natural logarithm of the first argument *LOG10()* Returns the base-10 logarithm of the argument *PI()* Returns the value of pi *POWER()* Returns the argument raised to the specified power *RADIANS()* Returns argument converted to radians *ROUND()* Round the argument *SIGN()* Returns the sign of the argument *SIN()* Returns the sine of the argument *SQRT()* Returns the square root of the argument

TAN() Returns the tangent of the argument

Atom-Related Functions

Some aggregation functions are available to operate on atomic properties, listed in the below. Atomic properties can only be used within such aggregation functions.

Name Definition

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SUM ATOM(<*expr*>*)* sum over atoms of expression <*expr*> *PROD ATOM(*<*expr*>*)* product over atoms of expression <*expr*> *AVG ATOM(*<*expr*>*)* average over atoms of expression <*expr*> *MIN ATOM(*<*expr*>*)* minimum over atoms of expression <*expr*> *MAX ATOM(*<*expr*>*)* maximum over atoms of expression <*expr*> *MATCH ATOM SUM(*<*condition*>*,*<*expr*>*)* sum over atoms matching <*condition*> of expression <*expr*>

MATCH ATOM PRODUCT(<*condition*>*,*<*expr*>*)* product over atoms matching <*condition*> of expression <*expr*>

MATCH ATOM AVG(<*condition*>*,*<*expr*>*)* average over atoms matching <*condition*> of expression <*expr*>

MATCH ATOM MIN(<*condition*>*,*<*expr*>*)* minimum over atoms matching <*condition*> of expression <*expr*>

MATCH ATOM MAX(<*condition*>*,*<*expr*>*)* maximum over atoms matching <*condition*> of expression <*expr*>

Ligand-Related Functions

These functions operate on the ligands (or nearest neighbors). Ligand atomic properties can only be used within such aggregation function. Furthermore, these functions can only be defined within an atom aggregation function.

Name Definition

SUM LIGAND(<*expr*>*)* sum over ligands of expression <*expr*>

PROD LIGAND(<*expr*>*)* product over ligands of expression <*expr*>

AVG LIGAND(<*expr*>*)* average over ligands of expression <*expr*>

MIN LIGAND(<*expr*>*)* minimum over ligands of expression <*expr*>

MAX LIGAND(<*expr*>*)* maximum over ligands of expression <*expr*>

MATCH LIGAND SUM(<*condition*>*,*<*expr*>*)* sum over ligands matching <condition> of expression $<$ expr $>$

MATCH LIGAND PRODUCT(<*condition*>*,*<*expr*>*)* product over ligands matching <*condition*> of expression <*expr*>

MATCH LIGAND AVG(<*condition*>*,*<*expr*>*)* average over ligands matching <*condition*> of expression <*expr*>

MATCH LIGAND MIN(<*condition*>*,*<*expr*>*)* minimum over ligands matching <*condition*> of expression <*expr*>

MATCH LIGAND MAX(<*condition*>*,*<*expr*>*)* maximum over ligands matching <*condition*> of expression <*expr*>

Void-Related Functions

A quick void space analysis can be performed on periodic structures. The following aggregation functions allow accessing the void properties. Void related properties can only be defined within these functions.

Name Definition

SUM VOID(<*expr*>*)* sum over voids of expression <*expr*> *AVG VOID(*<*expr*>*)* average over voids of expression <*expr*> *MIN VOID(*<*expr*>*)* minimum over voids of expression <*expr*> *MAX VOID(*<*expr*>*)* maximum over voids of expression <*expr*> *MATCH VOID SUM(*<*condition*>*,*<*expr*>*)* sum over voids matching <*condition*> of expression <*expr*> *MATCH VOID AVG(*<*condition*>*,*<*expr*>*)* average over voids matching <*condition*> of expression <*expr*> *MATCH VOID MIN(*<*condition*>*,*<*expr*>*)* minimum over voids matching <*condition*> of expression <*expr*> *MATCH VOID MAX(*<*condition*>*,*<*expr*>*)* maximum over voids matching <*condition*> of expression <*expr*>

8.3 Variables

Function operands can be either actual numbers or variables referring to structure properties. Atomic properties are defined for most per-element tabulated properties. Atomic properties related to nearest neighbors or ligands are specific, and if they appear in the descriptor definition they will trigger the computation of nearest neighbors for that structure. By definition, the ligands and voids properties and function will also trigger that computation.

Recursive Definition

Within the same catalog, one can refer to another descriptor directly by its name. The recursive dependency will be detected, and the computation order will follow it. Of course, cycling dependencies will cause an error.

Existing Descriptors

Properties alcreated defined in the list can be used as variables to define new descriptors. Only scalar numerical properties can be understood as existing descriptors. To refer to existing descriptors, the *@* is used, and apart from that, the syntax should still be lower case letters and numbers. For this reason, an automatic mapping from existing property names is done by lowering letter case and removing other characters, e.g. "Exp. Density" should be referred to as *@expdensity*.

8.4 Coordination properties

Whenever a property or a function related to ligands (nearest neighbors) is included in one of the catalog descriptors expression, a quick search for nearest neighbors will be triggered. This neighbor analysis comprises two steps:

- Firstly, all neighboring atoms that are not Voronoi neighbors are excluded. Voronoi cells are defined around each atom, as the portion of space where each point is closer to that atom than any other. Atomic extents are defined by atomic radii, and this Voronoi analysis provides a superior criterion for neighbor selection when a structure contains species with differing atomic radii. (Because if the segment from a particular atom to a neighboring atom intersects the Voronoi cell of a third atom, that potential neighbor can be discarded).
- Secondly, reduced distances between an atom and its neighbors are computed as the distance divided by the sum of atomic radii. The largest gap in the neighbor's reduced distances distribution is considered to separate nearest neighbors from further neighbors, and this criterion is employed to detect nearest neighbors.

As mentioned, covalent atomic radii are the default. However, it is also possible to use an alternative list of radii that is composed of covalent radii for elements found mostly in neutral condition and ionic radii that are the most frequent for atoms elements like Li. For simplification, this alternative list is labeled as Ionic (the default is labeled Covalent) and can be selected before actually computing descriptors where coordination comes into play.

Once the coordinated neighbors have been computed, the coordinated geometry class can be computed. To that end, both the distance to neighbors and the angles between each vector joining the atoms to its coordinated neighbors are analyzed. As in many cases, there will not be an exact match in distances and angles to a given regular coordination geometry, two additional values are also computed, namely

• *Distance dispersion* the maximum distance deviation from the average neighbor distances.

$$
d_{\rm{disp}} = {\rm{max}}_i \frac{\mid d_i - d_{\rm{avg}} \mid}{d_{\rm{avg}}}
$$

• *Angular dispersion* the maximum angle deviation from all the ideal case angles.

$\alpha_{\text{atom}} = \max_i |\alpha_{\text{actual}} - \alpha_{\text{ideal}}|$

As a result, each atom of each structure is characterized by the following:

- *coordination* (no units): the coordination number.
- *coordination confidence* (no units): the coordination confidence score. In some cases, the systematic discrimination of neighboring atoms as the ligand is not unquestionable: some neighboring atoms might or might not be ligands. This confidence score accounts for this uncertainty: its lower limit is 0 meaning no confidence and 1 means that the coordination number is established.
- *coordination class* (no units): the coordination class geometry code (a string representation can be used instead of the integer code), see the coordination class table.
- *coordination ddispersion* (no units): coordination distance deviation mentioned above.
- *coordination adeviation* (degrees): angular deviation

The coordination class values can be inserted in the expression editor with the Coordination class drop-down menu button.

8.5 Voids Properties

Voids are characteristics of voids topology: empty spaces such as cages, layers, or tunnels. Whenever a property or a function related to voids is included in the catalog descriptors expression, a quick search for voids is triggered. The search algorithm for the voids analysis is based on a multi-scale cell subdivision. The idea is to approximate the void space with void subcells up to a given precision. In the case of a cubic cell, these subcells are cubes. According to this analysis, connected sets of void subcells (subcells that are entirely included in void space) are identified as voids volume. They are characterized by:

- *void volume* (Angˆ3): the sum of the void subcell volume.
- *cage diameter* (Ang): the diameter of the largest sphere that could stand in the void.

This estimation is based on the subcells sizes decomposition and should be understood as a lower limit. A finer analysis would provide an actual higher estimate in most cases.

• *cross diameter* (Ang): the diameter of the largest sphere that could diffuse through the void, using the periodic conditions of the cell. This estimation is based on the subcells sizes decomposition and should be understood as a lower limit. A finer analysis would provide an actual higher estimate in most cases.

As these properties rely on approximation, an inherent uncertainty value is available for each of these diameters, labeled e.g. *delta cage diameter*. These properties can be inserted with the Voids properties drop-down button. Each structure may have any number of voids, each belonging to one of these 3 classes:

- *0*: cage closed void, no diffusion is possible. The cross diameter is 0 Ang.
- *1*: channel there is at least one crossing path in the void. The cross diameter is higher than 0.
- *2*: layer the void divides the cell space in 2 distinct parts (except the void itself).

These class numbers correspond to the *void class* property.

8.6 Examples

Mass Density

It can be defined as:

Mass density=SUM ATOM(atomic mass(Mg))/volume(mˆ3)

Let's say we want to compute the density of materials in the case all hydrogen atoms are removed. It can be achieved by taking into consideration all the atoms of the unit cells that are not hydrogen i.e. which atomic

number is greater than 1. That's what we call Not H density with descriptor name *not h density* and its definition is using a filtering criterion ($atomic_number > 1$):

not h density=MATCH ATOM SUM(atomic number>*1,atomic mass(Mg))/volume(mˆ3)*

Now, to illustrate how computed properties can be recursively defined, let us define the ratio of in-cell hydrogen mass density over total mass density, that we name H density ratio with the following definition:

h density ratio=(mdensity-not h density)/mdensity

Coordination

To illustrate the coordination geometry, we focus on C atoms in tetrahedral coordination. We will also pay attention to the deviation (in angle and distance) from the exact tetrahedral conformation. To this end, we define these 3 properties.

Tetrahedral C: the count of C atoms in tetrahedral conformation: *tetrac=MATCH ATOM SUM(atomic number=6 && coordination class=COORD 4 TETRAHEDRAL,1)*

Tetrahedral C angle deviation: The maximum tetrahedral C angle deviation *tetrac adev=MATCH ATOM MAX(atomic number=6 && coordination class=COORD 4 TETRAHEDRAL, coordination adeviation)*

Tetrahedral C distance dispersion: The maximum tetrahedral C distance dispersion *tetrac ddev=MATCH ATOM MAX(atomic number=6 && coordination class=COORD 4 TETRAHEDRAL,coordination ddispersion)*

Note that the filtering condition are using *&&*, the logical AND operator, and that a coordination constant has been used (the 440 value could have been used instead).

Ligands

Here are some properties related to atom-ligand properties. First the average atom-ligand distance, that defined as a double average, first level on all atoms, second level on all ligands (for each atom):

Avgdistligand=AVG ATOM(AVG LIGAND ligand distance(Ang)))

Then the average number of ligands per atom:

avgnligand=AVG ATOM(SUM LIGAND(1))

Finally, the average *atom ligand atom*, excluding the hydrogens (as atom and as *ligand*), which implies 2 filtering average functions:

avgdistligand not h=MATCH ATOM AVG(atomic number>*1, MATCH LIGAND AVG(ligand atomic number*>*1,ligand distance(Ang)))*

Voids

Here is a list of simple descriptors for voids:

caged (void cage diameter): The maximum (of any void) cage diameter: *MAX VOID(cage diameter(Ang))*

crossd (void cross diameter): The maximum (of any voids) cross diameter: *MAX VOID(cross diameter(Ang))*

ncages (number of cages): The number of cages in the system: *MATCH VOID SUM(void class=0,1)*

nchannels (number of channels): The number of distinct channels in the system: *MATCH VOID SUM(void class=1,1)*

nlayers (number of layers): The number of distinct layers in the system: *MATCH VOID SUM(void class=2,1)*

nvoids nNumber of voids): The number of distinct voids in the structure: *SUM VOID(1)*

voidv (void volume ratio): The void volume over total volume ratio: *SUM VOID(void volume)/volume*

8.7 MPM

The description above refers to operations on a Structure List. *MedeA HT* also offers the ability to create descriptors operating on entire crystallographic structural databases available in *MedeA InfoMaticA*, *i.e*. on hundreds of thousands of structural entries. This is called Materials Property Mining (MPM). While operations on Structure Lists are dynamic and can involve newly computed properties such as DFT total energies or elastic coefficients, MPM is designed to work with pre-computed properties such as coordination numbers and voids. The properties available within MPM are computed in advance and delivered with the databases.

Analogous to the definition of descriptors illustrated above, MPM allows the construction of descriptors which then can be applied to all entries in the *InfoMaticA* databases containing the relevant structural information (i.e. atomic positions in stoichiometric compounds). Thus, MPM is an extremely powerful data mining tool, which can be used in conjunction with the descriptors operating on Structure Lists.