PyDesc tutorial

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1 PyDesc - general information

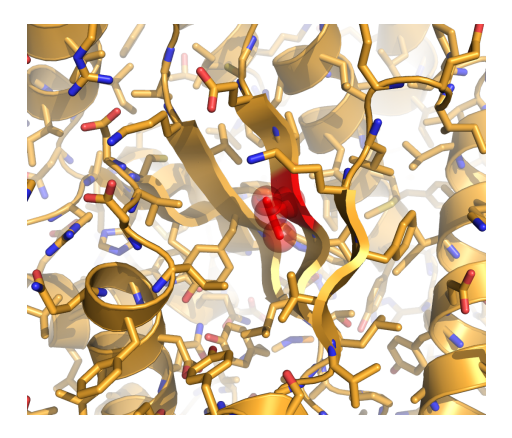
PyDesc is a Python 2.7.x library that provides classes which adapts Python environment to deal with biopolymer structures. It is designed for easy analysis of structures of both, proteins and nucleic acids, mostly using descriptors of local structure (DLS) approach (see chapter 3). Starting from very basic functionality as access to common databases of structures such as PDB, SCOP and CATH, and calculations of different geometrical features of structures or their parts, through combinations of contact criteria and calculation of contact maps, to the structural alignment of multiple structures in descriptor approach or analysis of MD trajectories – everything makes PyDesc useful for bioinformatic tools developers and for fast data analysis for scientists. Basic features of library are written in Python and available to developer for expansion or modification. More advanced and computationally demanding parts are written in C and CUDA C for performance boost. PyDesc requires BioPython for pdb parsing and numpy for most calculations. Additional features are enabled with DSSP, blastp, blastn and prody installed.

2 Basic features

PyDesc is able to load pdb files from common databases over Internet (PDB (including BioUnits), CATH, SCOP) or their local copies as well as single files from local drive. To deal with mmCIF files it loads bundle of pdb files in format provided by PDB. Loaded structures are stored in hierarchical objects, in which chains are stored in structures, mers are stored in chains and mers are simply containers for atoms and pseudoatoms useful in further analysis. It is also possible to create substructures of loaded structures such as ranges of sequential neighbours (called segments), sets of any mers or DLS (see chapter 3) for descriptor approach. Library provides friendly way to work with multiple structures. In case of homologous structures one can easily apply selections from one structure to another. As selection in PyDesc implements set theory approach different operations as addition or intersection of selections are easy to conduct. Dealing with non-homologous structures on the other hand is enabled thanks to modules providing alignment objects, which works on multiple alignments as well and allows to save and load files in common formats such as FASTA, CSV or XML. PyDesc provides calculation of contact maps and set of contact criteria to use for protein-protein and nucleic acid-nucleic acid interactions. It is also possible to combine provided contact criteria and to add user-defined criteria. That enables creation of DLS, which are substructures dependent on contact maps. Module for structural analysis, mostly written in C, provides structure comparison, (multiple) structure alignment and sequential/structural pattern fitting. Structure comparison allows for calculation of RMSD and rotation matrix using Kabsch method. It allows users to visualize superpositions (of both, pairs and multiple structures) using provided PyMOL plug-in. Calculation of structural alignment, by default, uses DLS approach and requires calculation of contact maps. Each of steps of structure analysis can be applied to frames of MD trajectories or states from NMR experiments.

3 Descriptors of Local Structure – DLS

Descriptors of Local Structure (DLS) are sets of mers – residues or nucleotides – representing local physicochemical environment of given mer. Such substructures are useful in structure comparison and computation of alignments, but can be useful in other structural analysis. In approaches using DLS structures are split into smaller overlapping fragments containing central mer (residue or nucleotide) from analyzed biopolymer and all other mers that stay in contact with it. That implies the size and shape of DLS depends on used criteria of contact between mers. The better those criteria describe real interactions between chemical compounds, the more useful obtained DLSs are as approximation of local environment. During calculation of DLS there are two main types of contacts: short and long range. Short range contacts are assumed between mer and their sequential neighbours, so preceding and following mers are included into DLS. Long range contacts depend on applied contact criteria. While defining criteria it is necessary to decide their type (quality, e.g. combination of distance and chemical properties etc.) and their thresholds (quantity, e.g. maximal distance, maximal difference in polarity etc.; see section XXX(ccrits)). Application of such criteria to all mers gives mers that stays in contact with central mer. All short range contacts of such mers are also included as they carry a information about physicochemical context of those mers.



4 Getting PyDesc

PyDesc is a library of classes written in Python 2.7. To install it, one needs Python 2.7 and dependent packages: BioPython, numpy and scipy. There are several way to download and install PyDesc:

ullet Use git

git clone XXX

- \bullet Download ${\tt zip}$ package from git repository home page: [trac.dw]trac.dw
- Use pip to install PyDesc from egg file included in repository.
- Use easy_install:

XXX

5 Getting started

PyDesc provides number of classes to deal with structural analysis. It can be used in development of Python scripts and programs, or as environment for fast analysis in Python interpreter. All the examples are ready to use in Python interpreter.

To check if PyDesc library is available simply import its components in Python:

```
import pydesc.sample
```

Loading structures in PyDesc requires class StructureLoader that is a component of structure module:

```
from pydesc.structure import StructureLoader structure_loader = StructureLoader()
```

That class provides method <code>load_structure</code> to which user can pass reference to database entry or path to a file on local drive and get a Python list of structures from appropriate file:

```
from pydesc.structure import StructureLoader structure_loader = StructureLoader() structures1 = structure_loader.load_structure("2dlc") structures2 = structure_loader.load_structure(code="2dlc") structures3 = structure_loader.load_structure("pdb://2dlc") structures4 = structure_loader.load_structure("2dlc", path="my/directory/2dlc.pdb")
```

In most cases file contains only one structure, but structures from NMR experiments or trajectories stored in pdb files are also able to be loaded using PyDesc. Structure 1 and 2 are downloaded automatically from first database to match the entry type format (PDB in that case). Structure 3 is loaded with forced usage of PDB. Structure 4 loads local file from my/directory and sets structure name to 2dlc.

That is all the basic knowledge needed to load structures. Readers interested in quick start are welcome to skip to part 7. Library features behind structure loading are described below.

During loader initialization User can choose what kind of database handler is to be used while loading structures. Database handler is an object responsible for connection to database. Available handlers are parts of pydesc.dbhandler package and are listed below:

- SCOPHandler provides access to SCOP db.
- PDBHandler provides access to pdb files from PDB.
- PDBBundleHandler provides access to compatibility version of big mmCIF files available as bundle of pdb files with additional text file mapping chains.

- MMCIFHandler provides access to mmCIF files from PDB.
- CATHHandler provides access to CATH db.
- BioUnitHandler provides access to BioUnit part of PDB.
- MetaHandler meta handler containing all necessary handlers.

Objects of that classes can be passed to StructureLoader with following syntax:

```
from pydesc.structure import StructureLoader
from pydesc.dbhandler import PDBHandler
structure_loader = StructureLoader(PDBHandler())
structure_loader = StructureLoader(handler=PDBHandler())
# both lines are equivalents
```

By default MetaHandler is used.

When creating a dbhandler, bear in mind that they can work in three modes:

- 1. Always downloading files via Internet to local cache.
- 2. Always getting file from local copy of DB to local cache.
- 3. Always opening files from local cache.

Modes can be passed as a list to all handlers during their initialization. Subsequent modes will be executed one by one until first success or till the end of mode list, e.g.:

```
from pydesc.structure import StructureLoader
from pydesc.dbhandler import PDBHandler
structure_loader = StructureLoader(PDBHandler(mode=[1,2,3]))
structures = structure_loader.load_structure("2DLC")
```

which is equivalent to default setting, will 1) check local cache for 2dlc.pdb file and load it if possible; otherwise it will 2) try to copy it from local DB copy to cache and load it again, but if that will end in failure, it will 3) download file from online DB to cache. Paths to local copy od DB and cache are stored in configuration manager (see chapter 6).

Other argument passed to StructureLoader initializator is parser, which should be object with method get_structure returning instances of structures as defined in Biopython Bio.PDB module. By default MetaParser from pydesc.dbhandler module is used. That object contains PDBParser and mmCIFParser from Biopyton Bio.PDB module, instances of both classes can be used on their own, if only passed handler returns appropriate files. E.g. one can use PDBHandler, which provides only pdb files to parser, therefore parser could be Bio.PDB.PDBParser.

When using non-default parser make sure that it can handle type of files returned by used dbhandler.

6 Configuration

There is a bunch of global settings influencing PyDesc library behaviour. They are all stored in pydesc.config module in static class ConfigManager with show_config method that shows tree of configuration:

```
import pydesc.structure
from pydesc.config import ConfigManager
ConfigManager.show_config()
```

All settings are stored in separate branches of names more or less corresponding with modules available in PyDesc. Branches containing appropriate configuration are created on module import. Settings important for certain function of library will be mentioned in appropriate chapters. Here we will present how to change and use settings using example of structure loaders mentioned in previous chapter.

```
import pydesc.structure
from pydesc.config import ConfigManager

print ConfigManager.dbhandler.cachedir
```

Setting cachedir in dbhandler branch of configuration manager is a directory in which cached structure files are stored. Lets assume that User conducts an experiment on set of structures. At some point he decides to test some other parameters in new directory, and wants to run modified scripts on the same set of structures. To avoid downloading and caching the same structures in new directory, User adds the following line:

```
ConfigManager.dbhandler.cachedir = '../old/directory/biodb/'
# equivalent:
ConfigManager.dbhandler.cachedir.set('../old/directory/biodb/')
```

7 Dealing with structures

Let us assume that one has already loaded a structure:

```
from pydesc.structure import StructureLoader
structure_loader = StructureLoader()
structures = structure_loader.load_structure("2dlc")
structure = structures[0]
```

As doing so in Python interpreter every time just to test some simple commands is arduous, for the sake of tuition PyDesc provides get function in sample module, which can be used instead.

```
from pydesc.sample import get
structure = get("2dlc")
```

That function returns only one structure from list of loaded structures. By default it is the first one, but User can pass index of structure to be returned as second argument. Note that lists in Python are indexed from 0.

7.1 Getting mers and substructures

PyDesc structures and substructures consist of mers (residue, nucleotide, ligands or ions), which consist of atoms. Each entity is represented by separate iterable object, so one way to get to building blocks of any level is to loop over their elements:

```
for mer in structure:
    print mer
    for atom in mer:
        print atom
```

Often we are not interested in whole structure, but in only one chain of it. Objects representing chains are created automatically and are stored in chains attribute of Structure instance. Class Structure also provides get_chain method, which takes chain name as str:

```
print structure.chains
print structure.get_chain('X')
```

Sample structure 2DLC consists of chains "X" and "Y".

Chains in PyDesc are very similar to whole structures and any other part of structure we will deal with in that tutorial. All of them provide method for getting their mers using Python syntax, which behaves slightly different than its equivalent for python sequential types:

```
chainX = structure.get_chain('X')
print structure[0], structure[0].ind
print structure[1], structure[1].ind
print chainX[0], chainX[0].ind
print chainX[75], chainX[75].ind
```

Mers in PyDesc are indexed from 1. Index 0 always returns first mer of the (sub)structure. PyDesc mer id, stored in ind attribute of mer, is created once at the creation of structure the mer belongs to and identifies mer in that particular structure, thus picking chain from the middle of the structure leads to shifted indexing. Attempts to pick mer of index not occurring in (sub)structure leads to IndexError. This is very useful in some cases but can be confusing for new Users. In above example chain X starts with mer 75. If one wants to index elements of PyDesc structure as in Python, simply convert PyDesc structure to Python sequential type, e.g. list or tuple:

```
print tuple(chainX)[0], list(chainX)[1]
```

Users are allowed to get parts or slices of structures:

```
seg = structure [75: 82]
for mer in seg: print mer
```

Note that mer 82 is a part of returned Segment instance! That is another difference between PyDesc and Python sequential types indexing. That means that expressions:

```
seg1 = structure [75: 75]
mer75 = structure [75]
```

return different type of objects. First one returns Segment instance of length 1, containing only 75th mer. Second one returns mer itself, residue in that case. Structures can be easily joined using + operator:

```
mix = structure [75: 82] + structure [84:85] + structure [90:90]
```

This kind of substructures are instances of UserStructure objects and has all the methods already presented here for Chain, Segment, Structure classes. Note that adding single mer to substructure requires adding 1-mer length segment (structure[90:90]). Attempt to add mer (structure[90]) will end up in AttributeError.

7.2 Getting atoms and pseudoatoms

Mers are sets of atoms and pseudoatoms (like geometrical center of side chain). Iteration over mer instance gives subsequent atoms and pseudoatoms:

```
for atom in structure [90]:
print atom
```

To get certain atom, one can use syntax similar to one used in Python dictionaries:

```
structure [90]['CA']
```

It works also for pseudoatoms, which usually are also an attribute of mer:

```
structure [90]['cbx']
structure [90].cbx
```

cbx is a pseudoatom defined for residues used by PyDesc default contact definition. It is vector $\langle C_{\alpha}, C_{\beta} \rangle$ extended by 1 Å.

There are also alternative iterators for mer, available using mer methods:

- __iter__ default, iterating over all atoms and pseudoatoms.
- iter_atoms iterating over atoms only.
- iter_atomsbb iterating over backbone atoms only.
- iter_atomsnbb iterating over non-backbone atoms only.

Distinguish of backbone and non-backbone atoms bases on setting backbone_atoms in branch ConfigManager.monomer.*, where asterisk is to be replaced with name of chainable mer, basically residue or nucleotide. Configuration stores any sequential type containing names of atoms as strings (with no spaces). E.g.:

```
from pydesc.config import monomer
from pydesc.config import ConfigManager

print ConfigManager.monomer.residue.backbone_atoms
```

```
('N', 'CA', 'C')
```

7.3 DLS, Contacts, Elements

There is also special kind of substructures containing parts of structure that are not sequential neighbours: DLSs (see chapter 3). That kind of structures depend on contact maps, which are matter of next chapter (9), here we will use default settings to calculate them.

To create DLS one needs to provide Element subclass instance (ElementChainable, ElementOther) of central mer and elements of all mers that stay in contact with central mer. Element chainable is special kind of Segment of odd number of mers with distinguished central mer. By default elements are segments of length 5 and depends on setting pydesc.config.ConfigManager.element.element_chainable_length. Elements of mers in contact with central mer should delivered as attributes of Contact instances, so second object needed to form a DLS is list of contacts. All classes, Element adn subclasses, Contact, as well as ProteinDescriptor or NucleotideDescriptor can be imported from pydesc.structure module.

Assuming that one knows the list of PyDesc indexes of mers that stays in contact with certain mer (list cnt1 in example below), lets say 90 in 2DLC, creation of descriptor goes as follow:

Contacts could be read from contact map calculated with PyDesc module. Similarly to this, one could create a nucleotide descriptor using NucleotideDescriptor class with appropriate element and contacts list. As that procedure is complicated and could be executed very often, each subclass of AbstractDescriptor provide static method of descriptor building basing on contact map calculated for structure from which mer comes from:

```
from pydesc.structure import StructureLoader
from pydesc.structure import Element
from pydesc.structure import ProteinDescriptor

structure = StructureLoader().load_structure('2dlc')[0]
structure.set_contact_map()

element90 = Element(structure[90])
descriptor90 = ProteinDescriptor.build(element90)
```

It is also possible to pass other contact map to that method as second argument. For structures containing both, protein and nucleic acids, it may be convenient to use method build provided by AbstractDescriptor class, which takes element and decides what type of descriptor to build:

```
from pydesc.structure import StructureLoader
from pydesc.structure import Element
from pydesc.structure import AbstracDescriptor

structure = StructureLoader().load_structure('2dlc')[0]
structure.set_contact_map()

element90 = Element(structure[90]) #residue
descriptor90 = AbstractDescriptor.build(element90)

element20 = Element(structure[20]) #nucleotide
descriptor20 = AbstractDescriptor.build(element20)

print descriptor90
print descriptor20
```

Often it is necessary to create all possible DLSs for given structure, which can be easily done with create_descriptors static method, which returns generator for subsequent DLSs:

```
from pydesc.structure import StructureLoader
from pydesc.structure import AstractDescriptor

structure = StructureLoader().load_structure('2dlc')[0]
structure.set_contact_map()
DLSgen = AbstractDescriptor.create_descriptors(structure)
DLSs = list(DLSgen)
```

For mers, for which DLS cannot be created None is returned. That allows for convenient usage of zip function on result:

```
DLSgen = AbstractDescriptor.create_descriptors(structure)
for mer, its_DLS in zip(structure, DLSgen):
    print mer, its_DLS
```

8 Selections

9 Contact maps

9.1 Basics, default criteria

Contact map contains information about contacts between mers in structure. To create one, one needs to know criteria of contact. First let us look at contact maps with default PyDesc criteria. PyDesc provides methods for checking values, getting contacts and saving all non-zero values in CSV file:

```
from pydesc.structure import StructureLoader

structure = StructureLoader().load_structure('2dlc')[0]

structure.set_contact_map()

print structure.contact_map
with open("2dlc_cmap.csv", "w") as fhobj:
    structure.contact_map.dump(fhobj)

print structure.contact_map.get_monomer_contacts(90)

print structure.contact_map.get_contact_value(90, 50)
print structure.contact_map.get_contact_value(90, 96)
print structure.contact_map.get_contact_value(90, 281)
```

Data stored in contact map can be passed to any file-like object including streams and files. It calls write method on passed object and feeds it with string containing as manly lines, as many contacts there is in structure (times two, as they are assumed to be symmetrical), in each line placing two PDB ids and value that can be 1 or 2.

As contact criteria in PyDesc uses tree-value logic the latter mean reliable contact, and the former – possible contact. Zero denote no contact. Criterion usually consist of some quality requirements (what features of mer should be taken into account) and quality requirements (threshold, below which criterion is satisfied). By default PyDesc uses:

• for residues of indexes i and j:

$$distance(C_{\alpha,i}, C_{\alpha,j}) \leq 6.00 \pm 0.50 \text{Å}$$

OR

$$distance(C_{\beta x,i}, C_{\beta x,j}) \le 6.50 \pm 0.50 \text{Å}$$

$$distance(C_{\beta x,i}, C_{\beta x,j}) - distance(C_{\alpha,i}, C_{\alpha,j}) \leq 0.75 \pm 0.00 \text{Å}$$

where.

 $C_{\alpha,a}$ – coordinates of alpha carbon of a-th residue,

 $C_{\beta x,a}$ – coordinates of extended beta carbon of a-th residue.

• for nucleotides of indexes i and j:

$$distance(R_{c,i}, R_{c,j}) \le 6.25 \pm 0.00 \text{Å}$$

OR

$$distance(R_{c,i}, I) \leq 7.40 \pm 0.00 \text{Å} \land distance(R_{c,j}, I) \leq 7.40 \pm 0.00 \text{Å}$$

where:

 $R_{c,a}$ – a-th nucleotide ring center,

I – ion coordinates.

Standard error notation was used here to indicate range in which contacts get value of 1.

9.2 Simple and combined pre-defined criteria

All pre-defined contact criteria and abstract classes are implemented in contacts module. For example first part of default criterion for residues measures distance of their alpha carbons and checks if it is under the threshold. It could be used alone for contact map calculation:

```
from pydesc.structure import StructureLoader
from pydesc.contacts import CaContact

structure = StructureLoader().load_structure('2dlc')[0]
structure.set_contact_map(CaContact())
print structure.contact_map.get_monomer_contacts(90)
```

Note that with such criterion there is no contacts between any nucleotides (residues 1-74 for 2DLC).

By default that criterion uses threshold equals to 6.00 and so called *unde-cidable range* of 0.50, which can be changed simply by passing them to criterion object initializator:

```
structure.set_contact_map(CaContact(8., .75))
print structure.contact_map.get_monomer_contacts(90)
```

If no arguments is passed to initializator, settings from configuration manager:

```
structure.set_contact_map(CaContact())

print structure.contact_map.get_monomer_contacts(90)

ConfigManager.contacts.ca_contact_distance = 8.0

ConfigManager.contacts.ca_contact_undecidable_range = 0.75

structure.set_contact_map(CaContact())

print structure.contact_map.get_monomer_contacts(90)
```

Difference between calling initializator with arguments and changing default settings is in behaviour of already existing contact criterion:

```
ConfigManager.contacts.ca_contact_distance = 8.0
ConfigManager.contacts.ca_contact_undecidable_range = 0.75
ca_crit_SD = CaContact()
ca_crit_8_75 = CaContact(8., .75)

structure.set_contact_map(ca_crit_SD)
print structure.contact_map.get_monomer_contacts(90)
structure.set_contact_map(ca_crit_8_75)
print structure.contact_map.get_monomer_contacts(90)
```

Result is the same, but in case of any change in settings:

```
ConfigManager.contacts.ca_contact_distance = 6.0
ConfigManager.contacts.ca_contact_undecidable_range = 0.50

structure.set_contact_map(ca_crit_SD)
print structure.contact_map.get_monomer_contacts(90)
structure.set_contact_map(ca_crit_8_75)
print structure.contact_map.get_monomer_contacts(90)
```

the ca_crit_SD will change values on the fly.

Second part of default criterion is a combined criterion, first part of which is basically the same as previously described CaContct, just calculating distance between cbx with different threshold. For that purpose there is another class CbxContact in appropriate module. One could be interested in criterion a bit easier than default one: alternative of ca i cbx criteria. With class ContactsAlternative it is as simple as:

```
from pydesc.contacts import ContactsAlternative
from pydesc.contacts import CaContact
from pydesc.contacts import CbxContact
ccrit = ContactsAlternative(CaContact(), CbxContact())
```

One can pass as many contact criteria as needed. PyDesc delivers also ContactsConjunction and ContactExclusiveDisjunction classes, which can also be called with Python operators:

```
from pydesc.contacts import CaContact
from pydesc.contacts import CbxContact

alt = CaContact() | CbxContact() #alternative
con = CaContact() & CbxContact() #conjunction
exd = CaContact() ^ CbxContact() #exclusive disjuntion
```

It is also possible to easily enerate negative criteria with Not class decorator, which for combined criteria regires a bit tricky syntax:

```
from pydesc.contacts import ContactsAlternative
from pydesc.contacts import CaContact
from pydesc.contacts import CbxContact

n_Ca = Not(CaContact)()
n_alt = Not(ContactsAlternative)(CaContact(), CbxContact())
```

9.3 Defining own simple criteria

In most cases it is sufficient to use pre-defined classes of criteria. PyDesc provides following **abstract** (one should not create instance of those classes, only create classes that inherit from them) classes:

- ContactCriterion generic criterion.
- PointsDistanceCriterion criteria basing on distance between (pseudo)atoms of choice.
- SetDistanceCriterion criteria based on set of distances between more than two pairs of atoms from different mers.
- VectorDistanceCriterion criteria based on orientation of vectors distinguished in mers.

• DihedralAngleCriterion – criteria based on dihedral angels between planes spanned on mer atoms.

PointsDistanceCriterion is the class of most widely used type of geometrical criteria. Extending this class is the best way to get own criteria dependent on distance between (pseudo)atom of choice. For example, lets define criterion for side chain geometrical center, which is stored in rc attribute of any chainable mer (for sake of example, as that criterion is implemented in RcContact class):

```
from pydesc.contacts import PointsDistanceCriterion

class RcCrt(PointsDistanceCriterion):
    monomer_hallmark = 'rc'

crt = RcCrt(4., 1.)
```

It requires only static attribute monomer_hallmark set to string containing name of attribute to be used in distance calculation. Of course, that is more likely to be useful for types of mers defined by Users, with their own pseudoatoms.

If no arguments will be passed to initializator — class will automatically seek for settings in configuration manager in pydesc.config.ConfigManager.contacts branch. One important setting is *attribute*_contact_distance, and second — *attribute*_contact_undecidable_range, where *attribute* will be replaced by class monomer_hallmark attribute.

Similarly for SetDistanceCriterion one needs to define names of mers attributes that are lists or dicts of atoms to be searched for:

```
from pydesc.contacts import PointsDistanceCriterion

class PaACrt(SetDistanceCriterion):
    monomer_hallmark = 'pseudoatoms'
    monomer_hallmark2 = 'atoms'

crt = PaACrt(criterion_distance=4.5, undecidable_range=.5,
    num_of_checked_pairs=2)
```

Initializator of such criteria can be feed with threshold and undecidable range, but also with number of pairs to meet those criteria. By default it is set to 2.

9.4 Defining own mers, pseudoatoms and contact criteria

By deafult PyDeck works on all-atom representation and is meant to be a tool in protein and nucleic acid structures analysis, but can be also used for other representations and for other molecules. In both cases it is more convenient to define new kinds of mers. Mers in PyDesc are stored in pydesc.monomer module. Main class, Monomer is abstract and has two subclasses: MonomerChainable and MonomerOther, first of which is, again, an abstract superclass for Nucleotide and Residue classes, while later is superclass for Ligand and Ion. Lets assume

we have some pdb file containing protein structure, but in representation containing only carbons alpha, beta and some pseudoatoms representing side chain, called SSn, where n is number of that pseudoatom (e.g. MARTINI representation). As that representation lacks important backbone atoms – by default PyDesc would fail to load it, unless new subclass of MonomerChainable is defined. We will be interested in, lets say, contact map of that protein with contact criterion defined as distance between SS1 pseudoatoms, but only for serines. Therefore we need:

As you can notice, this class simply calls Monomer initializator (instead of MonomerChainable to avoid questions about backbone atoms) and defines a new property, ss1, which returns atom called SS1. In PyDesc everything that is read from pdb file is atom and to make it accessible for contact criteria – one needs to define a property returning that atom.

Now movin og to contact criteria. As mentioned before, Users criterion contact should be subclass of classes provided in pydesc.contacts module. Usually contact criteria are classes that use two methods to tell if two mers are in contact or not: _calculate_distance and _is_in_contact. Both should take two mer instances as arguments. First one returns distance or array of distances, dependent on criterion type. Second one should call first one and compare results obtained for given mers with thresholds. For this case we should write:

```
from pydesc.contacts import ContactCriterion
class SerSS1Crt(ContactCriterion):
    def __init__(self, threshold, undecidable_range):
        self.thr = threshold
        self.undr = undecidable_range
    def _calculate_distance(self, m1, m2, **kwargs):
        return (m1.ss1 - m2.ss2).calculate_length()
    def _is_in_contact(self, m1, m2, **kwargs):
        if m1.name != m2.name != 'SER':
            return 0
        dist = self._calculate_distance(m1, m2)
        if dist < self.thr - self.undr:
            return 2
        elif dist < self.thr + self.undr:
            return 1
        return 0
```

With both those classes we are ready to load file with such reduced representation and calculate contact map for criterion concerning SS1 pseudoatoms in serines only. Structure in that representation is provided as <code>IOStream</code> (file-like object) within <code>pydesc.sample</code> module as result of <code>get_MARTINI_structure</code>, so to test it use:

```
from pydesc.sample import get_MARTINI_structure
s = get_MARTINI_structure()
s.set_contact_map(SerSS1Crt(5.0, 0.5))
```

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- 11 Structure comparison
- 11.1 OverFit
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