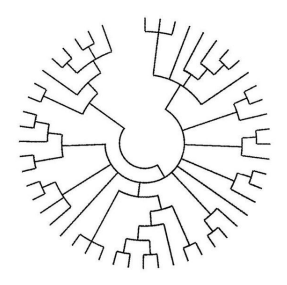
# DOCUMENTATION FOR THE PYTHON LIBRARY PEDIGRAD.PY

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### Contents

Chapter	1. Introduction	1
$\S 1.1.$	About pedigrads and Pedigrad.py	1
$\S 1.2.$	About this documentation	1
$\S 1.3.$	Acknowledgments	2
Chapter	2. Tutorial	3
$\S 2.1.$	Installation and preparation	3
$\S 2.2.$	Pre-ordered sets	4
$\S 2.3.$	Segments and morphisms of segments	6
$\S 2.4.$	Categories of segments	10
$\S 2.5.$	Environment functors	12
$\S 2.6.$	Aligned functors	13
$\S 2.7.$	Sequence alignments and the dynamic programming	17
Chapter	3. Presentation of the module Useful.py	21
$\S 3.1.$	Description of CategoryItem (class)	21
$\S 3.2.$	Description of usf (class item)	23
Chapter	4. Presentation of the module DProgramming.py	31
$\S 4.1.$	Description of Tree (class)	31
$\S 4.2.$	Description of Sequence (class)	34
$\S 4.3.$	Description of Table (class)	35
Chapter	5. Presentation of the module SegmentCategory.py	39
§5.1.	Description of PreOrder (class)	39
§5.2.	Description of SegmentObject (subclass)	45
§5.3.	Description of MorphismOfSegments (subclass)	50
$\S 5.4.$	Description of CategoryOfSegments (class)	53
Chapter	6. Presentation of the module AlignedFunctor.py	57
§6.1.	Description of PointedSet (class)	57
§6.2.	Description of SequenceAlignment (class)	58
§6.3.		66

<u>iv</u> Contents

Chapter	7. Presentation of the module PartitionCategory.py	73
$\S 7.1.$	Description of _image_of_partition	73
$\S 7.2.$	Description of _epi_factorize_partition	73
$\S 7.3.$	Description of _preimage_of_partition	74
$\S 7.4.$	Description of print_partition	75
$\S 7.5.$	Description of _join_preimages_of_partitions	75
§ <b>7.6</b> .	Description of EquivalenceRelation (class)	77
§7.7.	Description of coproduct_of_partitions	79
§7.8.	Description of product_of_partitions	30
$\S 7.9.$	Description of MorphismOfPartitions (class) 8	80
Chapter	8. Presentation of the module AsciiTree.py 8	33
$\S 8.1.$	Description of tree_of_partitions 8	33
$\S 8.2.$	Description of convert_tree_to_atpf	34
$\S 8.3.$	Description of convert_atpf_to_atf	36
$\S 8.4.$	Description of print_atf	37
$\S 8.5.$	Description of print_evolutionary_tree	88
Chapter	9. Presentation of the module Phylogeny.py 8	39
$\S 9.1.$	Description of Phylogenesis (class)	39
$\S 9.2.$	Description of Phylogeny (class)	91
Bibliogra	aphy	)1

### Introduction

#### 1.1. About pedigrads and Pedigrad.py

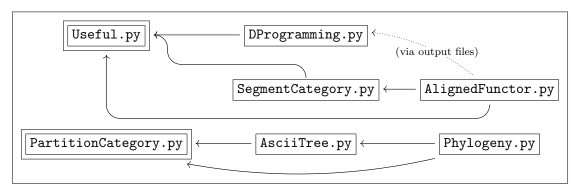
Pedigrads are mathematical tools that were initially created to model genetic mechanisms (see [1, 2, 3]). Mathematically, pedigrads are cone-preserving functors going from a certain class of limit sketches to a given category of values. Different aspects of biology can be encoded depending on the considered category of value. The present python library – Pedigrad.py – provides tools modelling the content described in [2]. Because earlier versions focused on pedigrads in the category of partitions an their use in phylogenetics, the library also comes with modules on partitions and evolutionary trees.

#### 1.2. About this documentation

The present book contains a tutorial (see Chapter 2) explaining how to use the various methods and classes contained in Pedigrad.py as well as a description of the (importable and non-importable) functions and classes of its sub-modules

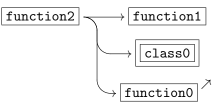
- Useful.py (see Chapter 3)
- DProgramming.py (see Chapter 4)
- SegmentCategory.py (see Chapter 5)
- AlignedFunctor.py (see Chapter 6)
- PartitionCategory.py (see Chapter 7)
- AsciiTree.py (see Chapter 8)
- Phylogeny.py (see Chapter 9)

The dependencies between the different chapters is displayed below.



1. Introduction

Similarly, descriptions of functions given in this documentation will usually start with dependency flow charts showing the modules that they use. Below, we give the example of such a flow chart.



As can be seen above, these flow charts will use three different types of boxes:

- 1) boxes with no specific decoration will usually frame items (such as classes and functions) that belong to the module of the described function;
- 2) double boxes will frame the name of intermediate items that do not have dependencies with other items (final item);
- 3) boxes with arrows in the top-right corner will frame items that are defined in modules external to that of the described function;

We will also give examples and demonstrations within editor mode windows or console mode windows. The editor mode will be used to describe the code of functions and will look as follows.

```
class MyClass:

This is a comment

def __init__(self,arg0,arg1): #This is the constructor of the class

"""

Another comment about the code

"""
```

The console mode will be used for examples and will look as follows.

```
>>> P.obj
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 13]
```

#### 1.3. Acknowledgments

I would like to thank Maxim Wolf and Carles Boix for interesting discussions about DNA and genetics. I would also like to thank Maxim Wolf for answering many of my questions and giving me some of his time.

### **Tutorial**

#### 2.1. Installation and preparation

To install the library, first download the package by clicking on the green button on the right of the screen at https://github.com/remytuyeras/pedigrad-library.



The downloaded package should be a compressed file named pedigrad-library-master.zip. Create a new directory in which you can copy and extract the compressed file using your favorite extraction application.



Enter the extracted directory pedigrad-library-master. Its inside should contain the following files.



In this tutorial, we will create four files:

- b two multiple sequence alignment files (usually with an extension .fa or .fasta);
- ▷ a file in which a pre-ordered set will be specified (preferably with an extension .yml);
- > a python file main.py in which we will write python functions and use the library.

To do this properly, create a new directory in pedigrad-library-master, call it User, and copy the file Pedigrad.py in User.



Now, open the file Pedigrad.py that you copied in User: you will be able to see several instances of the function sys.path.insert in which paths appear in the second argument.

```
import sys
sys.path.insert(0, "Pedigrad_py/Useful/")
from Useful import *
```

Add the text ../ at the beginning of every path passed to the function sys.path.insert, as shown below.

```
import sys
sys.path.insert(0, "../Pedigrad_py/Useful/")
from Useful import *
```

Once the paths are all updated, create four new files with the names align.fa; dprog.fa; main.py; and omega.yml in the directory User. The inside of the directory User should look as follows, where the file Pedigrad.pyc will appear later, after compilation.



Open the file main.py and insert the following piece of code.

```
main.py

1 from Pedigrad import *
```

Pedigrad.py

We are now ready to use the library – proceed to section 2.2.

#### 2.2. Pre-ordered sets

This section demonstrates the use of the class PreOrder (section 5.1), which belongs to the module SegmentCategory.py (section 5). The class PreOrder allows the user to load a preordered structure specified in a text file to the Random-Access Memory (RAM). For the sake of making this tutorial as exhaustive as possible, we will consider the pre-ordered set pictured in (2.1), which will allow us to illustrate various cases. In diagram (2.1), every (dashed or non-dashed) arrow  $x \to y$  represents a relation of the form x > y in the pre-order structure.

For this tutorial, we specify such a pre-ordered structure by inserting the following piece of code in the omega.yml (for the syntax, see the grammar rules shown in section 5.1.3.1).

```
omega.yml
1
     !obj:
2
       - 1;
3
       - 2;
4
       - 3;
5
6
       - 5;
7
     rel:
8
       -2, 3 > 5;
9
       -1, 2 > 3, 4;
10
```

Note that the minimal object! appearing in diagram (2.1) is specified through the symbol! preceding the key word obj:.

2.2. Pre-ordered sets 5

We can load the pre-order structure contained in omega.yml by creating an item of the class PreOrder relative to the file omega.yml (see the example given below). The class PreOrder is equipped with three objects: .relations, .mask and .cartesian. Let us what these do by calling them in the file main.py as follows.

```
main.py

3  Omega = PreOrder("omega.yml")

4  print Omega.relations

5  print Omega.mask

6  print Omega.cartesian
```

The object Omega.relations is a list of lists in which the relations specified in omega.yml are organized in the form of (non-transitive) down-closures. More specifically, every list in Omega.relations is associated with a particular object of the pre-ordered set, located at the beginning of the list, and contains all those objects that are less than or equal to this particular object based on the relation specified in omega.yml.

The object Omega.mask contains a Boolean value specifying whether the file omega.yml creates a formal minimal object through the use of the key word !obj:.

Finally, the object Omega.cartesian is an integer specifying whether the pre-ordered set is the Cartesian product of another pre-ordered set. If this integer is 0, then the Cartesian structure is not relevant to the PeOrder item. Compiling the file main.py should give the output.

```
[['1', '4', '3'], ['2', '5', '3', '4'], ['3', '5'], ['4', '3'], ['5']]
True
0
```

As mentioned earlier, the down-closures contained in Omega.relations are only based on the relations specified in omega.yml and may hence lack certain relations that can only be deduced from transitivity. One can add these relations to the lists of Omega.relations by calling the method .closure, as shown below.

```
7 Omega.closure()
8 print "--after:"
9 print Omega.relations
```

As can now be seen on the standard output, after compilation, the list Omega.relations has been completed with the relations 1>5 and 3>5, which can only be deduced by transitivity from the relations given in omega.yml.

```
--after:
[['1', '4', '3', '5'], ['2', '5', '3', '4'], ['3', '5'], ['4', '3', '5'],
['5']]
```

The class PreOrder is also equipped with a method .geq (short for greater than or equal to – see section 5.1.5). Let us have a look at the behavior of this method on objects that either belong to, or do not belong to the pre-order structure of omega.yml.

```
main.py

10  print "--relation:"
11  print Omega.geq("1","2")
12  print Omega.geq("2","1")
13  print Omega.geq("4","3")
14  print Omega.geq("?","3")
15  print Omega.geq("?","?")
```

As can be seen, the function Omega.geq can only return Truewhen its inputs are elements of the pre-ordered set and are comparable. In any other case, the output is False.

```
--relation:
False
False
True
False
False
False
```

In addition to the method .geq, the class PreOrder possesses an infimum function .inf (section 5.1.6). By definition, for every pair of elements in the pre-ordered set, the method .inf returns the largest element of the pre-ordered set that is less than or equal to the two input elements. Let us see what the following lines of code give us.

```
main.py

16  print "--infimum:"

17  print Omega.inf("2","3")

18  print Omega.inf("1","2")

19  print Omega.inf("?","3")
```

After compilation, we obtain the following text of the standard output. As expected, the infimum of 2 and 3 is 3 because the relation 2 > 3 holds in omega.yml. Similarly, the infimum of 1 and 2 is 4 because 4 is the maximum of the values 4, 3, and 5, which constitute the set of lower bounds of 1 and 2 (see diagram (2.1)). Finally, if the infimum of two values is not well-defined, then the method returns the value stored in Omega.mask, as shown below for the last call of the method .inf.

```
--infimum:
3
4
True
```

Finally, as a set, a pre-ordered set should also equipped with a relation belong to (i.e.  $\in$ ) to check whether a given element belongs to the pre-ordered set. This relation is available for PreOrder item through the method .presence (section 5.1.7). We can, for instance, try the following presence tests.

```
main.py

20 print "--presence test:"
21 print Omega.presence("3")
22 print Omega.presence("1")
23 print Omega.presence("?")
```

As expected (see output below), compiling main.py shows that 3 and 1 are elements of omega.yml and? is not.

```
--presence test:
True
True
False
```

#### 2.3. Segments and morphisms of segments

The present section demonstrates the use of the classes SegmentObject (section 5.2) and MorphismOfSegments (section 5.3), which are both subclasses of the class CategoryItem (section 3.1). The class SegmentObject allows the user to generate segments as defined in [2] and the class MorphismOfSegments allows the user to construct morphisms between

these segments (see [2, section 2]). For this library, the specification of a segment requires a topology and a color map. The former is given by a list of pairs of integers specifying successive intervals in the segments (called *patches* in [2]) and the latter is specified through a list of objects that should (but may not necessarily) belong to a PreOrder item (see section 2.2). In the following picture, the topology is specified above the segment while the color map is shown below the patches.

```
(0,3) \quad (4,5) \quad (6,10) \qquad (16,18) \quad (19,23)
(\circ \circ \circ \circ) (\bullet \bullet) (\bullet \bullet \bullet \bullet \bullet) (\circ \circ \circ \circ) (\bullet \bullet \bullet) (\bullet \bullet \bullet \bullet)
\cdot \text{mask} \quad "2" \quad "1" \quad "3"
```

Note that the patch (11,15) is missing in the topology. This will often happen for 'masked' patches, that is to say patches that we want to associate with the minimal object of the PreOrder item, in order to save space in the RAM.

Let us now show how one can construct a segment with the class SegmentObject. First, insert the following piece of code in the file main.py

```
main.py
24
25
    print "\n--
26
27
     t = list()
28
     c = list()
29
     for i in range(20):
       t = t + [(i,i)]
30
31
       if i%15 in [11]:
32
         c = c + ["?"]
       elif i%2 in [0]:
33
34
         c = c + ["1"]
       elif i%2 in [1]:
35
36
         c = c + ["2"]
```

The list t defines a topology while the list c defines a color map. We can then use these two lists two create a segment as follows.

```
main.py
37  s = SegmentObject(20,t,c)
38  sys.stdout.write("s = ")
39  s.display()
40  print "colors = "+str(s.colors)
```

The integer 20 given in the first argument of the constructor of SegmentObject sets the length of the segment. This length conveniently fits the coverage of the topology t that we defined earlier. Compiling the previous code will display the following text on the standard output.

```
s = (o|o|o|o|o|o|o|o|o|o|o|o|o|o|o|o|o|o|o)

colors = ['1', '2', '1', '2', '1', '2', '1', '2', '1', '2', '1', '?', '1',

'2', '1', '2', '1', '2', '1', '2']
```

If we changed the length of the segment to 25, then we would obtain the following alternative output, where the last five nodes are not colored (or rather masked).

However, changing the length to a high number, say 250, will not display the whole sequence of nodes on the standard output, mostly if these nodes are masked. Instead, the method .display will turn most of the masked nodes into a count, as shown below (the same would happen at the beginning of the segment if the index of the first node was greater than 12).

#### 

Now, the idea is to use the segment s to construct more exotic segments. For instance, we can use the method .merge (section 5.2.6) to merge patches within the topology of s. The color of every new patch is defined as the infimum value of the colors of the merged patches.

In the following piece of code, we merge the patches of s according to 3 different tiling patterns. The triple (0,3,9) indicates that we will merge every set of successive triple of (non-masked) patches from the (non-masked) patch of index 0 to the (non-masked) patch of index 9, the triple (10,2,14) indicates that we will merge every set of successive pairs of (non-masked) patches from the (non-masked) patch of index 10 to the (non-masked) patch of index 14, and the triple (15,3,19) indicates that we will merge every set of successive triples of (non-masked) patches from the (non-masked) patch of index 15 to the (non-masked) patch of index 19.

```
main.py

42    s1 = s.merge([(0,3,9),(10,2,14),(15,3,19)],Omega.inf)
43    sys.stdout.write("s1 = ")
44    s1.display()
45    print "topology = "+str(s1.topology)
46    print "colors = "+str(s1.colors)
```

Compiling outputs the following text, where we can see the newly merged patches. The new colors generated through this process are either 4 = Omega.inf(1,2) or the Boolean value Omega.mask = Omega.inf(1,?), which leads to masking the associated patch.

```
s1 = (ooo|ooo|ooo|oo|oo|oo|oo|oo)

topology = [(0, 2), (3, 5), (6, 8), (9, 9), (12, 13), (14, 14), (15, 17),

(18, 19)]

colors = ['4', '4', '4', '2', '4', '1', '4', '4']
```

In addition of merging patches, we can also 'remove' patches from the topology of a segment, which will lead to consider the patch as masked. We do so by using the method .remove and giving it the indices of the patches to be removed. Note that the method .remove can be quite handy when one needs to display various parts of a very long segments in which the information is distributed sparsely.

To illustrate the use of the method .remove , let us insert the following piece of code in the file main.py.

To make our example slightly more interesting, let us also augment the length of the segment s2 by 4 nodes, as shown below.

Compiling the file main.py gives the following additional text on the standard output.

```
s2 = (ooo|ooo|ooo|o|oo|oo|o|o|o|o|o)

s2 = (ooo|ooo|ooo|o|oo|oo|oo|o|o|o|o|o)

s2 = (ooo|ooo|ooo|ooo|o|oo|oo|o|o|o|o|oooo)

colors = ['4', '4', '4', '2', '1', '1', '2', '1', '2']
```

We can create another segment by appending the following similar piece of code to the file main.py.

```
main.py
61
    s3 = s.merge([(0,3,9),(10,2,14)],Omega.inf)
62
    sys.stdout.write("s3 = ")
63
    s3.display()
64
    s3 = s3.remove([1,5,6,8,10])
    sys.stdout.write("s3 = ")
65
66
    s3.display()
67
    s3.topology = s3.topology+[(24,24)]
    s3.colors = s3.colors+["5"]
68
69
70
    s3.domain = s3.domain+5
71
    sys.stdout.write("s3 = ")
72
    s3.display()
    print "colors = "+str(s3.colors)
73
```

Compiling main.py adds the text to the standard output displayed below; note how the value in s3.domain restricts the display of the patch (24,24) in the second version of s3, but not in its last version.

```
s3 = (\frac{000}{000} \begin{align*} \long \long
```

Let us now move on constructing morphisms between segments (see [2, section 2.5]). In particular, we want to construct a morphism between our previous two segments s2 and s3. To do so, let us add the following piece of code to main.py, where the parameter "id" is treated as the list range(s2.domain), which can be seen as a trivial mapping  $i \mapsto i$ .

```
main.py

75  m = MorphismOfSegments(s2,s3,"id",Omega.geq)

76  print m.defined

77  sys.stdout.write("s = ")

78  m.source.display()

79  sys.stdout.write("t = ")

80  m.target.display()

81  print "f0 = "+str(m.f0)
```

After compiling, we obtain the following additional text on the standard output. The value of m.defined tells us whether the construction of the morphism succeeded or not. The value of m.defined may be False if the morphism does not satisfy the axioms specified

in [2, section 2.5]. In the case of an ill-defined morphism, the list m.f0, which represents the order-preserving function  $f_0: [n_0] \to [n'_0]$  used in [2, section 2.5], should be empty (see section 5.3). Note that failure to construct a morphism of segments is likely to be due to the impossibility of constructing  $f_0$ .

```
True

s = (000|000|000|0|00|0|0|0|0|0|0|0|0000)

t = (000|000|000|0|00|00|0|0|0|00000|0)

f0 = [0, -1, 1, 2, 3, -1, 4, -1, 5]
```

The previous example rather simple, mostly because we used the parameter "id" instead of specifying list f1 that would model an interesting mapping  $i \mapsto f1[i]$ . The reason is that it can sometimes be quite time consuming to specify these types of mappings by hand. Instead, we want to use more powerful tools, such the class CategoryOfSegments, which presented in section 2.4.

#### 2.4. Categories of segments

The present section demonstrates the use of the class SegmentCategory, which models the features of a category of segments [2]. In particular, this class will allow us to be more efficient in the specification of segments (through the method .initial) and morphisms of segments (through the method .homset). As with category of segments, an item of the class SegmentCategory needs to be initialized with a PreOrder item.

In our case, we want to use the PreOrder item Omega defined in section 2.2. We call our category of segments as follows.

```
main.py

82
83     print "\n-----\n"
81
85     Seg = CategoryOfSegments(Omega)
86
```

As with categories, we can know whether there is an identity morphism between two segments. For instance, the following lines of code test whether a segment can be equal to itself. We also test an equality between the two different segments.

```
main.py
87
    s = Seg.initial(18,"1")
88
    s = s.merge([(2,2,8)],Omega.inf)
89
90
    print Seg.identity(s,s)
91
92
    t = Seg.initial(20,"1")
    t = t.merge([(2,3,10),(15,2,18)],Omega.inf)
93
94
95
    print Seg.identity(s,t)
```

As expected, compiling the previous lines of code displays the following output, which tells us that there is an identity between **s** and itself, but not between **s** and **t**.

\_\_\_\_\_

True False We can check that **s** and **t** are indeed different segments by adding the following piece of code to main.py

```
main.py

97  sys.stdout.write("s= ")

98  s.display()

99  sys.stdout.write("t= ")

100  t.display()
```

Compiling gives us the following output.

- s = (o|o|oo|oo|oo|o|o|o|o|o|o|o|o|o)
- t = (0|0|000|000|000|0|0|0|00|00|00)

If we cannot find an obvious morphism between s and t, we can always use the object .homset of the class PreOrder to enumerate all the existing morphisms going from the segment s to the segment t – this done in the following lines of code.

```
main.py

102 h = Seg.homset(s,t)

103 for i in range(len(h)):

104  print str(i)+") well-defined = "+str(h[i].defined)

105  print "f1 = "+str(h[i].f1)

106  print "f0 = "+str(h[i].f0)
```

```
Compiling the file main.py now gives the list of the 14 morphisms of segments s \rightarrow t.
0) well-defined = True
f1 = [0, 1, 2, 3, 5, 6, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
1) well-defined = True
f1 = [0, 1, 2, 3, 5, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
2) well-defined = True
f1 = [0, 1, 2, 3, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
3) well-defined = True
f1 = [0, 1, 2, 4, 5, 6, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
4) well-defined = True
f1 = [0, 1, 2, 4, 5, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
5) well-defined = True
f1 = [0, 1, 2, 4, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
6) well-defined = True
f1 = [0, 1, 3, 4, 5, 6, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
7) well-defined = True
f1 = [0, 1, 3, 4, 5, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
```

```
8) well-defined = True
f1 = [0, 1, 3, 4, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 1, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
9) well-defined = True
f1 = [0, 2, 3, 4, 5, 6, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 2, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
10) well-defined = True
f1 = [0, 2, 3, 4, 5, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 2, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
11) well-defined = True
f1 = [0, 2, 3, 4, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [0, 2, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
12) well-defined = True
f1 = [1, 2, 3, 4, 5, 6, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [1, 2, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
13) well-defined = True
f1 = [1, 2, 3, 4, 5, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [1, 2, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
14) well-defined = True
f1 = [1, 2, 3, 4, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
f0 = [1, 2, 2, 3, 4, 4, 5, 6, 7, 8, 9, 9, 10, 10, 11]
```

#### 2.5. Environment functors

The present section demonstrates the use of the class Environment (section 6.3), which models the features of environment functors  $E_b^{\varepsilon}$ , as defined in [2, Definition 3.11]. As with environment functors, an Environment item is equipped with a pointed set  $(E,\varepsilon)$  (i.e. a PointedSet item – see section 6.1), a category of segments  $\mathbf{Seg}(\Omega)$  (i.e. a CategoryOfSegments item – see section 5.4) and a truncation element b taken from the pre-ordered set  $\Omega$ . More specifically, initializing an Environment item usually takes the following lines of code.

```
main.py
107
108
109
    E = PointedSet(["-","A","C","G","T"],0)
110
111
    Env = Environment(Seg,E,5,["4"]*5) #[] = white nodes
112
113
    print Env.Seg.preorder.relations
114
    print Env.pset.symbols
115
    print Env.pset.point()
116
    print Env.spec
117
    print Env.b
```

Compiling the previous code adds the following text to the standard output.

\_\_\_\_\_\_

```
[['1', '4', '3', '5'], ['2', '5', '3', '4'], ['3', '5'], ['4', '3', '5'], ['5']]
['-', 'A', 'C', 'G', 'T']
-
5
['4', '4', '4', '4', '4']
```

As a model for a particular type of functors, an **Environment** item is equipped with a pullback operation, which we briefly illustrate before ending this section (see section 6.3 for more details).

$$\{ au\}$$
 -  $\overset{\subseteq}{-}$   $\mathbf{Seg}(\Omega)$ 
 $\downarrow$   $\mathbf{A}E_b^{arepsilon}$ 
 $\mathbf{1}$   $\xrightarrow{\mathsf{input}}$   $\mathbf{Set}$ 

More specifically, we can pullback any sequence of characters, taking its values in the pointed set  $(E, \varepsilon)$ , through the method .segment. Let us consider the following example:

```
main.py
s4 = Env.segment(["A","C","G","T","T","N","C","A","-","C","T"],"1")
s4.display()
```

As can be seen, after compiling, the characters belonging to the pointed sets are associated with non-masked nodes, while characters that do not belong to the pointed set are associated with masked nodes.

(0|0|0|0|0|0|0|0|0|0)

#### 2.6. Aligned functors

The present section demonstrate the use of the class SequenceAlignment (section 6.2), which models the features of a sequence alignment functor, as defined in [2, Definition 3.20]. In particular, we show how the type of construction illustrated in [2, Example 3.22] can be with the functions of this library.

To start with, let us consider the following sequence alignment file.

```
align.fa
1
    >1:A:1
2
    ACTCGATCTCTG?TCGATCGATCG
3
    CCTATCGGATCGATC
4
    >1:B:2
5
    ACTCTAT?TCTATCC-ACTCATCA
6
    CCTACTATCTCGAAA
7
    >1:C:3
8
    ACTCTATC----TCCGACT?ATCA
9
    CCTACTATCTCGAAA
10
    >1:D:1
11
     ACTATTTA?TTTC----TTTTCTA
12
    CCGGGCTGGGGGGG
```

```
align.fa
13
    >2:A:1
14
    ACTCGATCGGATC-AT??CGATCG
15
    CCTATCGGATCGATCG
16
    >2:B:2
17
    ACTCC?CCTA--CTGGGCTGCTCA
18
    CCTACTATCTCGAAAG
19
    >2:C:3
20
    ACTCTCTATC?CTCTATC---TCA
21
    CCTACTATC?CGAAAG
22
    >2:D:1
23
    ACTATT-??T--AC--CTTTTCTA
24
    CCGGGCTTTTCGAAAA
25
    >1:K:1
26
    ACTATCGATCTTTAGCTAGTTCTA
27
    CC-??????----GA
```

We can create a SequenceAlignment item from the pairwise sequence alignments stored in the file align.fa by using the method .seqali, as shown below.

```
main.py

121
122 print "\n----\n"
123
124 Seqali = Env.seqali("align.fa")
```

The SequenceAlignment item Seqali is equipped with two objects .base and .database (see section 6.2). The object .base contains the SegmentObject items at which the sequences given in align.fa can be found in the aligned environment functor. In other words, the object .base contains the objects of the category B designed in [2, Example 3.22]. The object .database contains the lists of sequence alignments associated with each SegmentObject item in .base. In other words, it encodes a preliminary version of the mapping  $T: B \to \mathbf{Set}$  constructed in [2, Example 3.22].

The following code shows how to use the objects .base and .database in order to retrieve the information contained in align.fa.

```
main.py
126
    print "\nDatabase\n"
127
128
    print Seqali.indiv
129
    for i in range(len(Seqali.base)):
      print str(i)+") color: "+str(Seqali.base[i].colors[
130
    Seqali.base[i].parse ])
131
      Seqali.base[i].display()
      for j in range(len(Seqali.database[i])):
132
133
        for k in range(len(Seqali.database[i][j])):
134
          print Seqali.database[i][j][k]
135
        print ""
```

Compiling gives the following output:

#### Database

```
['A', 'B', 'C', 'D', 'K']
0) color: ['1', '2', True, '1', '1']
|o|o|o
ACTCGATCTCTG?TCGATCGATCGCCTATCGGATCGATC
ACTCTAT?TCTATCC-ACTCATCACCTACTATCTCGAAA
masked
ACTATTTA?TTTC----TTTTCTACCGGGCTGGGGGGGG
ACTATCGATCTTTAGCTAGTTCTACC-??????----GA
1) color: ['1', '2', True, '1', True]
({\color{red}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{blue}0}{\color{b
|o|o|o|o
ACTCGATCGGATC-AT??CGATCGCCTATCGGATCGATCG
ACTCC?CCTA--CTGGGCTGCTCACCTACTATCTCGAAAG
masked
ACTATT-??T--AC--CTTTTCTACCGGGCTTTTCGAAAA
masked
```

Alternatively, we can call the images of the underlying aligned environment functor by using the method .image. Below, we use the method .image to retrieve the image of the functor at the SegmentObject item stored in Seqali.base[0] and Seqali.base[1].

```
main.py
137
    print "\nImage\n"
138
    for i in range(len(Seqali.base)):
139
      print "base["+str(i)+"]"
140
      Seqali.base[0].display()
141
      sal = Seqali.eval(Seqali.base[0])
142
      for j in range(len(sal)):
143
         for k in range(len(sal[j])):
144
           print sal[j][k]
145
         print ""
```

Compiling the previous code gives the following text on the standard output:

#### Image

#### base[0]

ACTCGATCTCTG?TCGATCGATCGCCTATCGGATCGATC
ACTCTAT?TCTATCC-ACTCATCACCTACTATCTCGAAA
masked

ACTATTTA?TTTC----TTTTCTACCGGGCTGGGGGGGGACTATCGATCTTTAGCTAGTTCTACC-???????----GA

#### base[1]

```
ACTCGATCGGATC-AT??CGATCGCCTATCGGATCG
ACTCC?CCTA--CTGGGCTGCTCACCTATCTCGAAAG
masked
ACTATT-??T--AC--CTTTTCTACCGGGCTTTTCGAAAA
masked
```

In addition to being able to display the images of the sequence alignment functor, we can also construct the set of objects of its underlying *extending category*, which is defined in [2, Definition 3.25].

```
main.py

147 print "\nExtending_category\n"

148 l= Seqali.extending_category(Seqali.base[0])

149 for i,m in 1:
    print i

151    print m.f1

152    print m.f0
```

Compiling the previous lies of codes gives the list of all the objects of the extending category associated with the SegmentObject item stored in Seqali.base[0] (for conciseness, only the beginning and the end of the output is given)

#### Extending category

```
0
id
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20,
21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38]
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20,
21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38]
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20,
21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38]
. . .
1
[0, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21,
22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39]
[0, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20,
22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21,
22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39]
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21,
22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39]
```

Of course, we can do the same for the SegmentObject item stored in Seqali.base[1].

```
main.py

154 print "\nExtending diagram\n"

155 l= Seqali.extending_category(Seqali.base[1])

156 for i,m in 1:

157 print i

158 print m.f1

159 print m.f0
```

However, this time, the extending category possesses only one object.

# Extending diagram 1 id [0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39]

#### 2.7. Sequence alignments and the dynamic programming

The present section demonstrate the use of the class Table (section 4.3), models the features of a dynamic programming score table for the purpose of aligning two pair of sequences of characters. In particular, it encodes the dynamic programming algorithm that allows the type of analysis discussed in [2, Section 4] for the recognition of biological mechanisms.

The class Table should be used as shown in the following example. First, use the class Sequence (section 4.2) to initialize the table and then use the method .incidence to compute all the possible matches between the two sequences given to the Table item.

```
main.py
160
    print "\n----\n"
161
162
163
    a = list("AGCTAGCTGA")
164
    b = list("GTGGATCGATGA")
165
    A = Sequence("a",a,"1")
166
167
    B = Sequence("b",b,"1")
168
    table = Table(A,B)
169
170
    print "\nincidence"
171
    table.incidence()
    table.stdout()
```

Compiling the file main.py displays the following text on the standard output.

\_\_\_\_\_

Then, the method .fillout should be called to fill out the table with the scores that will be used by the dynamic programming algorithm.

```
main.py
173
    print "\nfillout"
    table.fillout()
174
175 table.stdout()
```

Compiling the file main.py now adds the following text on the standard output.

```
fillout
. | . | G | T | G | G | A | T | C | G | A | T | G | A |
G | 0 | 1 | 1 | 1 |
                1 | 1 | 1 | 2 | 2 | 2 | 2 |
             1 |
C | 0 | 1 | 1 | 1 |
                1 | 1 | 2 | 2 | 2 | 2 | 2 |
             1 |
T | 0 | 1 | 2 | 2 | 2 | 2 | 2 |
                     2 | 2 | 2 | 3 | 3 |
A | 0 | 1 | 2 | 2 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 4 |
T | 0 | 1 | 2 | 3 | 3 | 3 | 4 | 4 | 4 | 4 | 5 | 5 | 5 |
G | 0 | 1 | 2 | 3 | 4 | 4 | 4 | 4 | 5 | 5 | 5 | 6 | 6 |
A | 0 | 1 | 2 | 3 | 4 | 5 | 5 | 5 | 5 | 6 | 6 | 6 | 7 |
```

The method .dynamic\_programming can now be applied to compute all the valid sequence alignments resulting from the previous scores and store them in the file "dprog.fa". As with the function open, the second argument of .dynamic\_programming takes either "w", to rewrite on the file, or "a", to append the outputs to an existing file. To do so, add the following line of code to main.py.

```
main.py
176 table.dynamic_programming("dprog.fa",option = "w",debug = False,display
```

Compiling gives the following data on the standard output and inside the file prog.fa.

```
>0:a:1
AGCTA-G--C--TGA
>0:b:1
-G-T-GGATCGATGA
>1:a:1
AGCT-AG--C--TGA
>1:b:1
-G-TG-GATCGATGA
>2:a:1
AGCTAG---C-TGA
>2:b:1
-G-T-GGATCGATGA
>3:a:1
AGCT--AG-C--TGA
>3:b:1
-G-TGGA-TCGATGA
>4:a:1
AGCT--A-GC--TGA
>4:b:1
-G-TGGAT-CGATGA
```

```
>5:a:1
AGCT--A--GC-TGA
>5:b:1
-G-TGGATCG-ATGA
>6:a:1
AGCT--A--G-CTGA
>6:b:1
-G-TGGATCGA-TGA
```

Finally, it is possible to use the file dprog.fa in the same fashion as we used the file align.fa, in section 2.6. For instance, add the following line of codes and compile.

```
main.py
177
    print "\n----\n"
178
179
    E = PointedSet(["-","A","C","G","T"],0)
180
181
182
    Env = Environment(Seg,E,2,["1"]*2) #[] = white nodes
    Seqali = Env.seqali("dprog.fa")
183
184
185
    print "\nImage\n"
186
187
    print Seqali.indiv
    Seqali.base[0].display()
188
    sal= Seqali.eval(Seqali.base[0])
189
    for j in range(len(sal)):
190
191
      for k in range(len(sal[j])):
192
        print sal[j][k]
193
      print ""
```

The following result should be obtained:

```
Image
```

```
['a', 'b']
(O|O|O|O|O|O|O|O|O|O|O|O|O|O
AGCTA-G--C--TGA
-G-T-GGATCGATGA

AGCT-AG--C--TGA
-G-TG-GATCGATGA

AGCTAG--C--TGA
-G-T-GGATCGATGA

AGCTAG--C--TGA
-G-T-GGATCGATGA

AGCT--AG-C--TGA
-G-TGGA-TCGATGA

AGCT--A-GC--TGA
-G-TGGATCGATGA
```

 ${\tt AGCT--A--GC-TGA}$ 

 $-\mathsf{G}-\mathsf{T}\mathsf{G}\mathsf{G}\mathsf{A}\mathsf{T}\mathsf{C}\mathsf{G}-\mathsf{A}\mathsf{T}\mathsf{G}\mathsf{A}$ 

AGCT--A--G-CTGA

 $-\mathsf{G}-\mathsf{T}\mathsf{G}\mathsf{G}\mathsf{A}\mathsf{T}\mathsf{C}\mathsf{G}\mathsf{A}-\mathsf{T}\mathsf{G}\mathsf{A}$ 

# Presentation of the module Useful.py

#### 3.1. Description of CategoryItem (class)

**3.1.1.** Introduction. The class CategoryItem models the diagrammatic features of the elements of a category. This class is meant to be used as a super class. In particular, the class CategoryItem is equipped with an object .level that can be used to model the polymorphic nature functors, namely both arrows and objects (of the considered category) can be given to a function as CategoryItem items and can be handled differently depending on their value stored in the object .level (see the example of section 3.1.3).

**3.1.2. Structure.** The following tables give a preview of the class CategoryItem. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry	Module section	
object	N/A	
Statistics		
⊳ Importable objects: 3		
⊳ Non-importable objects: 0		
▷ Importable methods: 1		
▷ Non-importable methods: 0		

The following table gives a description of the 3 importable objects of the class:

Objects		
Name	Type	Related sections
.level	int	⊳ section 3.1.3
.source	CategoryItem	⊳ section 3.1.3
.target	CategoryItem	⊳ section 3.1.3

Finally, the following table gives a description of the only importable methods of the class:

Methods			
Name	Input types	Output types	Related sections
init	- int	- self	⊳ section 3.1.3
11116	- list		

**3.1.3.** Description of .\_\_init\_\_ (method). The code of the function .\_\_init\_\_ is equipped with the following inputs, among which one can be optional depending on the value of the first argument.

init		
Inputs	Types	Specifications
level	int	necessary
*args	<pre>list(CategoryItem)</pre>	optional

The method possesses two actions, which we describe below through examples.

	Action 1
Case	If the list args possesses exactly 2 items and args[0].level ==
	level-1 and args[1].level == level-1
Description	In this case, the method allocates the values stored in level,
	args[0] and args[1] to the objects self.level, self.source and
	self.target, respectively.

	Action 2		
Case	If the list args does not possess exactly 2 items or args[0].level !=		
	level-1 or args[1].level != level-1		
Description	In this case, the method allocates the values stored in level to the		
	object self.level. The resulting class item is not equipped with the		
	two objects self.source and self.target.		

The following example shows how the class can be used as a super class to create functions that behave like functors. First, we create two subclasses of CategoryItem modelling objects and arrows in a certain category.

```
>>> class Obj(CategoryItem):
...    def __init__(self,content):
...         super(Obj, self).__init__(0)
...         self.content = content
...
>>> class Arr(CategoryItem):
...         def __init__(self,A,B,content):
...         super(Arr, self).__init__(1,A,B)
...         self.content = content
...
```

Then, we code a functor taking the previous two classes as the same inputs thanks to their shared ancestry.

```
>>> def functor(cat_item):
...     if cat_item.level == 0:
...     print "object:"
...     print cat_item.content
...     print cat_item.level
...     if cat_item.level == 1:
...     print "arrow:"
...     print cat_item.content
...     print cat_item.level
...     print cat_item.source.content
...     print cat_item.source.level
...     print cat_item.target.content
...     print cat_item.target.level
...     print cat_item.target.level
...     print cat_item.target.level
...
```

Finally, the following lines of code shows that the function functor can handle both Obj and Arr items.

```
>>> a = Obj("a")
>>> b = Obj("b")
>>> f1 = Arr(a,b,"a-->b")
>>> functor(a)
object:
a
>>> functor(b)
object:
b
>>> functor(f1)
arrow:
a-->b
1
а
0
b
0
```

#### 3.2. Description of usf (class item)

**3.2.1.** Introduction. The class item usf comes from a non-importable class and is equipped with a collection of functions that turned out to be useful during the construction of the present library.

**3.2.2.** Structure. The following tables give a preview of the class item usf. The table given below describes the various dependencies of the class item.

Dependencies		
Class type	Module section	
_Useful N/A		
Sı	tatistics	
⊳ Importable objects: 0		
⊳ Non-importable objects: 0		
▷ Importable functions: 5		
ightarrow Non-importable functions: $0$		

The following table gives a description of the 5 importable functions of the class item. The type  $\alpha$  shown in the table is a polymorphic type

Functions			
Name	Input types	Output types	Related sections
usf.list_to_string	- list(string)	- string	⊳ section 3.2.3
	-file	- list(string)	⊳ section 3.2.4
usf.read until	- list(string)		
usi.leau_until	- list(string)		
	- bool		
usf.fasta	- string	- list(list(string))	⊳ section 3.2.5
usi.lasta		- list(string)	
usf.add_to	- α	- bool	⊳ section 3.2.6
usi.adu_to	- $list(lpha)$		
	- int	- list(list(int))	⊳ section 3.2.7
usf.inclusions	- int		
usi.inclusions	- int		

**3.2.3.** Description of usf.list\_to\_string (function). The code of the function usf.list\_to\_string is equipped with the following input.

${\tt usf.list\_to\_string}$		
Inputs Types Specifications		
a_list	list(string)	necessary

The method possesses one action, which we describe below through an example.

Action		
Case	Always	
Description	The function coverts the list of strings a_list into the concatenation of	
	its internal strings. If the list is empty, then the empty string is returned.	

The following example illustrates the previous description.

```
>>> usf.list_to_string([])
```

>>> usf.list\_to\_string(["AGGATCGTATG","AGCTAG","AATT","CG","ATTTTATCG"])
AGGATCGTATGAGCTAGAATTCGATTTTATCG

**3.2.4.** Description of usf.read\_until (function). The code of the function usf.read\_until is equipped with the following inputs, among which one is optional.

usf.read_until				
Inputs	Specifications			
a_file	file	necessary		
separators	<pre>list(string)</pre>	necessary		
EOL_symbols	list(string)	necessary		
inclusive = False	bool	optional		

The method possesses one action, which we describe below through an example.

Action		
Case	Always	
Description	This function reads a file until it reads a character belonging to the	
	input list EOL_symbols. It returns the list of pieces of text that were	
	separated by characters belonging to the list separators. If the argu-	
	ment inclusive is set to True, then the last character read in the file	
	is included in the output list. As a result, the output of the function	
	usf.read_until is never empty when the argument inclusive is set to	
	True. On the other hand, if the argument inclusive is set to False,	
	then the last character read in the file is excluded from the output list.	

Let us illustrate the action of the function usf.read\_until on the following FASTA file.

```
alignment.fa
1
    >1:A:color_1
2
    ACTCGATCTCTG?TCGATCGATCG
3
    CCTATCGGATCGATC
4
    >1:B:color_1
5
    ACTCTAT?TCTATCC-ACTCATCA
6
    CCTACTATCTCGAAA
7
    >1:C:color_1
    ACTCTATC----TCCGACT?ATCA
8
    CCTACTATCTCGAAA
9
10
    >1:D:color_1
    ACTATTTA?TTTC----TTTTCTA
11
    CCGGGCTGGGGGGG
12
```

The following examples illustrate the description that we gave earlier. We start with an example in which the parameter inclusive is not specified.

```
>>> with open("alignment.fa","r") as f:
...    keepgoing = True
...    text = list()
...    while keepgoing:
...    line = usf.read_until(f,[" ",":"],["\n"])
...    if line != []:
...    text.append(line)
...    else:
...    keepgoing = False
...    keepgoing = False
...    for i in range(len(text)):
...    print text[i]
```

Below, we see that setting the parameter separators to [" ",":"] leads to eliminating these characters from the output.

```
['>1', 'A', 'color_1']

['ACTCGATCTCTG?TCGATCGATCG']

['CCTATCGGATCGATC']

['>1', 'B', 'color_1']

['ACTCTAT?TCTATCC-ACTCATCA']

['CCTACTATCTCGAAA']

['>1', 'C', 'color_1']

['ACTCTATC----TCCGACT?ATCA']

['CCTACTATCTCGAAA']

['>1', 'D', 'color_1']

['ACTATTTA?TTTC----TTTTCTA']

['CCGGGCTGGGGGGGGG']
```

We now look at a similar example in which the parameter inclusive is set to True. Notice that the condition if line != [] used in the previous piece of code is now if line != [""].

```
>>> with open("alignment.fa", "r") as f:
       keepgoing = True
       text = list()
       while keepgoing:
         line = usf.read_until(f,[" ",":"],["\n"], inclusive = True)
         if line != [""]:
           text.append(line)
         else:
           keepgoing = False
       for i in range(len(text)):
         print text[i]
['>1', 'A', 'color_1', '\n']
['ACTCGATCTCTG?TCGATCGATCG', '\n']
['CCTATCGGATCGATC', '\n']
['>1', 'B', 'color_1', '\n']
['ACTCTAT?TCTATCC-ACTCATCA', '\n']
['CCTACTATCTCGAAA', '\n']
['>1', 'C', 'color_1', '\n']
['ACTCTATC---TCCGACT?ATCA', '\n']
['CCTACTATCTCGAAA', '\n']
['>1', 'D', 'color_1', '\n']
['ACTATTTA?TTTC----TTTTCTA', '\n']
['CCGGGCTGGGGGGG', '\n']
```

3.2.5. Description of usf.fasta (function). The code of the function usf.fasta is equipped with the following input.

usf.fasta			
Inputs Types Specifications			
name_of_file	string	necessary	

The method possesses one action, which we describe below through examples.

	Action		
Case	Always		
Description	The function takes a FASTA file, whose syntax is described in section		
	3.2.5.1, and returns two lists:		
	▷ names, containing the lists of key words making every sequence		
	labels specified in the file (see section 3.2.5.1);		
	> sequences, containing the sequences associated with every se-		
	quence label.		
	Furthermore, every list associated with a sequence label in names is made		
	of the words that are separated by a colon symbol in the specification		
	of the sequence label.		

3.2.5.1. name\_of\_file (variable). The code contained in name\_of\_file should follow a specific syntax that can be described through the following grammar.

Because this syntax is compatible with the FASTA format, the file should ideally be specified with the extension .fa or .fasta. For example, the file alignment.fa given in section 3.2.4 constitutes a perfect example of the type of file with which the function usf.fasta should be used.

The following example shows what the output of usf.fasta looks like when applied to the file alignment.fa of section 3.2.4.

```
>>> fa = usf.fasta("alignment.fa")
>>> for i in range(len(fa)):
...    print fa[i]
...
[['1', 'A', 'color_1'], ['1', 'B', 'color_1'], ['1', 'C', 'color_1'], ['1',
'D', 'color_1']]
['ACTCGATCTCTG?TCGATCGCCTATCGGATCGATC', 'ACTCTAT?TCTATCC-ACTCATCACCTACTA
TCTCGAAA', 'ACTCTATC----TCCGACT?ATCACCTACTATCTCGAAA', 'ACTATTTA?TTTC----TTT
TCTACCGGGCTGGGGGGGGG']
```

**3.2.6.** Description of usf.add\_to (function). The code of the function usf.add\_to is equipped with the following inputs.

${\tt usf.add\_to}$			
Inputs Types Specifications			
element $\alpha$		necessary	
a_list $list(\alpha)$		necessary	

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	The function appends the element element to the list a_list if this ele-	
	ment is not present in the list. Otherwise, the element is not appended.	

The following examples illustrate the previous description.

```
>>> 1 = [1, 2, 3, 4, 5]

>>> b = usf.add_to(5,1)

>>> print b, 1

False [1, 2, 3, 4, 5]

>>> b = usf.add_to(6,1)

>>> print b, 1

True [1, 2, 3, 4, 5, 6]
```

**3.2.7.** Description of usf.inclusions (function). The code of the function usf.inclusions is equipped with the following inputs.

usf.inclusions			
Inputs	Types Specifications		
start	int	necessary	
domain	int	necessary	
holes	int	necessary	

The method possesses one action, which we describe below through examples.

Action			
Case	Always		
Description	The function computes the list of lists f whose implicit map-		
	pings $i \mapsto f[i]$ represent increasing inclusions from the ordered set		
	$\{0,1,,\mathtt{domain}-\mathtt{holes}-1\}$ to the ordered set $\{\mathtt{start},\mathtt{start}+1\}$		
	$1,,\mathtt{start} + \mathtt{domain} - 1\}$		

The following examples illustrate the previous description. As the reader can notice, the outputs of usf.inclusions enumerate the combinations of (domain - holes) elements among domain elements.

In the following example, we shift the range in which the combinations are computed.

```
\rightarrow >> 1 = usf.inclusions(2,7,3)
>>> for i in range(len(1)):
       print l[i]
. . .
. . .
[2, 3, 4, 5]
[2, 3, 4, 6]
[2, 3, 4, 7]
[2, 3, 4, 8]
[2, 3, 5, 6]
[2, 3, 5, 7]
[2, 3, 5, 8]
[2, 3, 6, 7]
[2, 3, 6, 8]
[2, 3, 7, 8]
[2, 4, 5, 6]
[2, 4, 5, 7]
[2, 4, 5, 8]
[2, 4, 6, 7]
[2, 4, 6, 8]
[2, 4, 7, 8]
[2, 5, 6, 7]
[2, 5, 6, 8]
[2, 5, 7, 8]
[2, 6, 7, 8]
[3, 4, 5, 6]
[3, 4, 5, 7]
[3, 4, 5, 8]
[3, 4, 6, 7]
[3, 4, 6, 8]
[3, 4, 7, 8]
[3, 5, 6, 7]
[3, 5, 6, 8]
[3, 5, 7, 8]
[3, 6, 7, 8]
[4, 5, 6, 7]
[4, 5, 6, 8]
[4, 5, 7, 8]
[4, 6, 7, 8]
[5, 6, 7, 8]
```

# Presentation of the module DProgramming.py

#### 4.1. Description of Tree (class)

**4.1.1.** Introduction. The class Tree models the features of a tree structure. A Tree item is equipped with an object .parent in which it is possible to store information and an object .children through which one can specify descendants. A Tree item can be constructed recursively from the constructor. Tree items whose object .parent is equal to the string "leaf" are distinguished from the rest of the structure and considered as terminal states. For instance, these terminal states are useful if one wants to enumerate all the paths in the tree. For instance, the class Tree is equipped with a method .paths that returns a list of all the paths going from the root to a leaf.

**4.1.2. Structure.** The following tables give a preview of the class **Tree**. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry	Module section	
object	N/A	
Statistics		
⊳ Importable objects: 4		
Non-importable objects: 0		
⊳ Importable methods: 4		
▷ Non-importable methods: 0		

The following table gives a description of the 4 importable objects of the class:

Objects			
Name	Type	Related sections	
.depth	int	$\triangleright$ section 4.1.3	
.level	int	$\triangleright$ section 4.1.3	
.parent	$\alpha$	$\triangleright$ section 4.1.3	
.children	$\mathtt{list}(\mathtt{Tree}(lpha))$	$\triangleright$ section 4.1.3	

Finally, the following table gives a description of the 4 importable methods of the class:

Methods				
Name	Input types	Output types	Related sections	
init	<i>-</i> α	- self	$\triangleright$ section 4.1.3	
	- list $(Tree(lpha))$			
.stdout	- self	- self	$\triangleright$ section 4.1.4	
.levelup	- self	- self	$\triangleright$ section 4.1.5	
.paths	- self	- list $(lpha)$	$\triangleright$ section 4.1.6	

**4.1.3.** Description of .\_\_init\_\_ (method). The code of the function .\_\_init\_\_ is equipped with the following inputs, among which one can be optional depending on the value of the first argument.

init		
Inputs	Types	Specifications
parent	$\alpha$	necessary
*args	$ extstyle{list(Tree}(lpha))$	optional

The method possesses two actions, which we describe below through examples.

Action 1		
Case	If self.parent != "leaf"	
Description	In this case, the method allocates the values stored in the input variables	
	parent and args to the objects self.parent and self.children, re-	
	spectively. The value of the object self.level is set to 0 and the value	
	of the object self.depth is computed as a sum m+1, where m is the	
	maximum value taken among the objects .depth of every Tree item in	
	the list args.	

Action 2		
Case	<pre>If self.parent == "leaf"</pre>	
Description	In this case, the method allocates the value stored in the input vari-	
	able parent to the object self.parent (the object .children is not	
	constructed). The value of the object self.level is set to 0 and the	
	value of the object self.depth is computed as a sum m+1, where m is	
	the maximum value taken among the objects .depth of every Tree item	
	in the list args.	

The following example how the constructor can be used to construct a Tree item for which the value contained in the object .depth is 4.

```
>>> leaf = Tree("leaf")
>>> leaf.depth
>>> a = Tree("a",[leaf])
>>> a.depth
>>> b = Tree("b",[leaf])
>>> c = Tree("c",[leaf])
>>> d = Tree("d",[leaf])
>>> e = Tree("e",[leaf])
>>> ab = Tree("ab",[a,b,leaf])
>>> de = Tree("de",[d,e])
>>> cde = Tree("c(de)",[c,de])
>>> t = Tree("ab(c(de))", [ab,cde])
>>> t.depth
>>> t.level
>>> t.parent
ab(c(de))
```

**4.1.4. Description of .stdout (method).** The code of the function .stdout has no input. The method possesses one action, which we describe below through an example.

Action		
Case	Always	
Description	This method displays the tree structure on the standard output where,	
	for every Tree item within the recursive structure of the Tree item, the	
	line "["+str(self.level)+"] -> "+str(self.parent) is displayed	
	through a recursive search.	

The following example displays the tree constructed in the example of section 4.1.3.

```
>>> t.stdout()
[0] -> ab(c(de))
.[1] -> ab
..[2] -> a
..[2] -> b
.[1] -> c(de)
..[2] -> c
..[2] -> de
...[3] -> d
...[3] -> e
```

**4.1.5.** Description of .levelup (method). The code of the function .levelup has no input. The method possesses one action, which we describe below through examples.

Action			
Case	Always		
Description	This method recursively increments by 1 all the objects .level in the		
	recursive structure of the Tree item.		

The following example shows the effect of the method .levelup on the structure of the tree used in the example of section 4.1.4.

```
>>> t.levelup()
>>> t.stdout()
.[1] -> ab(c(de))
..[2] -> ab
...[3] \rightarrow a
...[3] -> b
..[2] -> c(de)
...[3] -> c
...[3] -> de
....[4] -> d
....[4] -> e
>>> t.levelup()
>>> t.stdout()
..[2] -> ab(c(de))
...[3] -> ab
....[4] -> a
....[4] -> b
...[3] -> c(de)
....[4] -> c
....[4] -> de
.....[5] -> d
.....[5] -> e
```

**4.1.6.** Description of .paths (method). The code of the function .paths has no input. The method possesses one action, which we describe below through an example.

Action			
Case	Always		
Description	This method returns a list of all the paths going from the root		
	(self.parent) to a Tree item, within the recursive structure of the		
	Tree item, whose object .parent is equal to the string "leaf".		

The following example computes a list containing all the paths of the Tree item t constructed in section 4.1.3.

```
>>> t.paths()
[['ab(c(de))', 'ab', 'a'], ['ab(c(de))', 'ab', 'b'], ['ab(c(de))',
'ab'], ['ab(c(de))', 'c(de)', 'c'], ['ab(c(de))', 'c(de)', 'de', 'd'],
['ab(c(de))', 'c(de)', 'de', 'e']]
```

## 4.2. Description of Sequence (class)

- **4.2.1.** Introduction. The class Sequence models the features of a sequence that would belong to a multiple sequence alignment specified in a FASTA file (more specifically, see section 3.2.5.1). Its objects .name and .color contain information that would be specified in the label given before the sequence (according to the FASTA format). The sequence itself is stored in the object .seq.
- **4.2.2. Structure.** The following tables give a preview of the class **PreOrder**. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
object	N/A	
Statistics		
⊳ Importable objects: 3		
▷ Non-importable objects: 0		
▷ Importable methods: 1		
▷ Non-importable methods: 0		

The following table gives a description of the 3 importable objects of the class:

Objects			
Name Type Related section		Related sections	
.name	string	$\triangleright$ section 4.2.3	
.seq	<pre>list(string)</pre>	$\triangleright$ section 4.2.3	
.color	string	$\triangleright$ section 4.2.3	

Finally, the following table gives a description of the only importable method of the class:

Methods			
Name	Input types	Output types	Related sections
	- string	- self	$\triangleright$ section 4.2.3
init	- list(string)		
	- string		

**4.2.3. Description of** .\_\_init\_\_ (**method**). The code of the function .\_\_init\_\_ is equipped with the following inputs.

init			
Inputs	Types	Specifications	
name	string	necessary	
sequence	<pre>list(string)</pre>	necessary	
color	string	necessary	

The method possesses one action, which we describe below through an example.

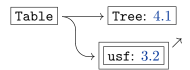
Action		
Case	Always	
Description	The method allocates the values of the input variables name, sequence	
	and color to the objects self.name, self.seq, and self.color, re-	
	spectively.	

The following example shows a canonical use of the constructor of the class **Sequence**.

```
>>> sam = Sequence("Samuel",["A","C","T","T","C","g","g","a","T"],"blue")
>>> sam.name
Samuel
>>> sam. seq
['A','C','T','T','C','g','g','a','T']
>>> sam.color
blue
```

## 4.3. Description of Table (class)

**4.3.1.** Introduction. The code of the class Table uses other functions of the module DProgramming.py as well as external modules.



The class Table models the features of a dynamic programming score table for the purpose of aligning two pair of sequences of characters. In particular, it encodes the dynamic programming algorithm that allows the type of analysis discussed in [2, Section 4] for the recognition of biological mechanisms.

The class Table should be used as shown in the following example. First, use the class Sequence (section 4.2) to initialize the table.

```
>>> a = list("abc[defg]hij")
>>> b = list("abc[gfed]hij")
>>> A = Sequence("a",a,"1")
>>> B = Sequence("b",b,"1")
>>> table = Table(A,B)
```

Then, the method .incidence should be called to compute matches between the two Sequence items given to the constructor of the class Table.

```
>>> table.incidence()
>>> table.stdout()
. | . | a | b | c | [ | g | f | e | d | ] | h | i | j |
0 | 1 | 0 | 0 |
         0 | 0 |
            0 | 0 |
               0 |
0 | 0 | 0 | 0 |
d | 0 | 0 | 0 | 0 |
       0 |
         0 |
          0 |
            0
             | 1 | 0 | 0 |
e | 0 | 0 | 0 | 0 | 0 |
         0 | 0 |
            1 | 0 | 0 | 0 |
   0 | 0 | 0 | 0 |
         0 | 1 | 0 | 0 | 0 | 0 |
```

Then, the method .fillout should be called to fill out the table with the scores used by the dynamic programming algorithm.

```
>>> table.fillout()
>>> table.stdout()
. | . | a | b | c | [ | g | f | e | d | ] | h | i | j |
a | 0 | 1 | 1 | 1 | 1 | 1 |
                    1 |
                      1 | 1 | 1
     1 | 2 |
           2 | 2 |
                 2 |
                    2 |
                      2 |
                         2 |
                            2 |
                               2
c | 0 | 1 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
d | 0 | 1 | 2 | 3 | 4 | 4 | 4 | 4 | 5 | 5
                             I 5
                                  5
e | 0 | 1 | 2 | 3 | 4 | 4 | 4 | 5 | 5 | 5 | 5
f | 0 | 1 | 2 | 3 | 4 | 4 | 5 | 5 | 5 | 5 | 5 | 5
```

Finally, the method .dynamic\_programming can be applied to store all the valid sequence alignments resulting from the previous scores in a file (here taken to be "output.fa"), as shown below. As with the function open, the second argument of .dynamic\_programming takes either "w", to rewrite on the file, or "a", to append the outputs to an existing file.

```
>>> table.dynamic_programming("output.fa",option = "w",debug = False,display
= True)
>0:a:1
abc[defg---]hij
>0:b:1
abc[---gfed]hij
>1:a:1
abc[de-fg--]hij
>1:b:1
abc[--gf-ed]hij
>2:a:1
abc[d-efg--]hij
>2:b:1
abc[-g-f-ed]hij
>3:a:1
abc[-defg--]hij
>3:b:1
abc[g--f-ed]hij
>4:a:1
abc[de-f-g-]hij
>4:b:1
abc[--gfe-d]hij
>5:a:1
abc[d-ef-g-]hij
>5:b:1
abc[-g-fe-d]hij
>6:a:1
abc[-def-g-]hij
>6:b:1
abc[g--fe-d]hij
>7:a:1
abc[d--efg-]hij
>7:b:1
abc[-gfe--d]hij
>8:a:1
abc[-d-efg-]hij
>8:b:1
abc[g-fe--d]hij
>9:a:1
abc[--defg-]hij
>9:b:1
abc[gf-e--d]hij
```

```
>10:a:1
abc[de-f--g]hij
>10:b:1
abc[--gfed-]hij
>11:a:1
abc[d-ef--g]hij
>11:b:1
abc[-g-fed-]hij
>12:a:1
abc[-def--g]hij
>12:b:1
abc[g--fed-]hij
>13:a:1
abc[d--ef-g]hij
>13:b:1
abc[-gfe-d-]hij
>14:a:1
abc[-d-ef-g]hij
>14:b:1
abc[g-fe-d-]hij
>15:a:1
abc[--def-g]hij
>15:b:1
abc[gf-e-d-]hij
>16:a:1
abc[d--e-fg]hij
>16:b:1
abc[-gfed--]hij
>17:a:1
abc[-d-e-fg]hij
>17:b:1
abc[g-fed--]hij
>18:a:1
abc[--de-fg]hij
>18:b:1
abc[gf-ed--]hij
>19:a:1
abc[---defg]hij
>19:b:1
abc[gfed---]hij
```

# Presentation of the module SegmentCategory.py

### 5.1. Description of PreOrder (class)

**5.1.1.** Introduction. The code of the class PreOrder uses external modules.



The class PreOrder models the features of a pre-ordered set. The pre-order relations are specified through either a file or another PreOrder item passed to the constructor. The method .closure makes sure that the pre-order axioms are satisfied and will (re-)compute the transitive closure of the pre-order relations stored in the object .relations if needed; the method .geq returns a Boolean value specifying whether there is a pre-order relation between two given elements of the pre-ordered set; the method .inf returns the infimum of two elements of the pre-ordered set; and the method .presence returns a Boolean value specifying whether an element belongs to the pre-ordered set.

**5.1.2. Structure.** The following tables give a preview of the class PreOrder. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
object	N/A	
Statistics		
⊳ Importable objects: 4		
Non-importable objects: 0		
▷ Importable methods: 5		
▷ Non-importable methods: 2		

The following table gives a description of the 4 importable objects of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Objects			
Name	Type	Related sections	
.relations	$ ext{list}( ext{list}(lpha))$	$\triangleright$ section 5.1.3	
		$\triangleright$ section 5.1.4	
.transitive	bool	$\triangleright$ section 5.1.3	
		$\triangleright$ section 5.1.4	
.mask	bool	$\triangleright$ section 5.1.3	
.cartesian	int	$\triangleright$ section 5.1.3	
		$\triangleright$ section 5.1.5	
		$\triangleright$ section 5.1.6	

Finally, the following table gives a description of the 5 importable methods of the class:

Methods			
Name	Input types	Output types	Related sections
	- string	- self	$\triangleright$ section 5.1.3
init	- int		
	- list(PreOrder)		
.closure	- self	- self	⊳ section 5.1.4
	<b>-</b> α	- bool	$\triangleright$ section 5.1.5
ged	$-\alpha$		
.geq	- list		
	- α	- α	$\triangleright$ section 5.1.6
.inf	$-\alpha$		
. 1111	- list		
.presence	<i>-</i> α	- bool	⊳ section 5.1.7

**5.1.3. Description of** .\_\_init\_\_ (**method**). The code of the function .\_\_init\_\_ is equipped with the following inputs, among which two are optional.

$.\_$ init $\_$			
Inputs	Types	Specifications	
name_of_file	string	necessary	
cartesian = 0	int	optional	
*args	list(PreOrder)	optional	

The method possesses two actions, which we describe below through examples.

Action 1			
Case	If cartesian > 0 and the list args possesses exactly 1 item		
Description	In this case, the list args is assumed to contain a PreOrder item		
	that the method uses to initialize the three objects self.relations,		
	self.transitive, self.mask through componentwise attributions.		
	The object self.cartesian is initialized with the value stored in		
	cartesian.		

Action 2		
Case	If cartesian <= 0 or the list args does not possess exactly 1 item	
Description	In this case, the method uses the file referenced through the variable	
	name_of_file to fill out the objects of the class. In section 5.1.3.1 (see	
	below), we describe the syntax of the file and how it is parsed.	

5.1.3.1. name\_of\_file (variable). The code contained in name\_of\_file should follow a specific syntax that can be described through the following grammar.

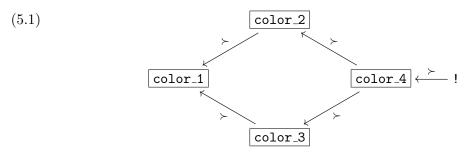
```
\begin{array}{lll} \text{file} & \rightarrow & \text{file} \mid !\text{obj: Obj rel: Rel} \mid obj : \text{Obj rel: Rel} \mid \varepsilon \\ \text{Obj} & \rightarrow & \text{Separator Word Comment Obj} \mid \varepsilon \\ \text{Rel} & \rightarrow & \text{Text} > \text{Text} ; \text{Comment Rel} \mid \text{Comment} \mid \varepsilon \\ \text{Separator} & \rightarrow & \text{not}([0-9]+[@]+[A-Z]+[\_]+[a-z]) \text{ Separator} \mid \varepsilon \\ \text{Word} & \rightarrow & [0-9] \text{ Word} \mid @ \text{Word} \mid [A-Z] \text{ Word} \mid [a-z] \text{ Word} \mid \varepsilon \\ \text{Comment} & \rightarrow & \text{\#Text} \mid \varepsilon \\ \text{Text} & \rightarrow & \text{Separator Word Text} \mid \varepsilon \end{array}
```

Key words		
!obj:/obj:	Indicates that the objects of the pre-ordered set are about to be listed.	
	Each object is defined by a Word variable in the previous grammar.	
!obj:	Adds a formal minimal object to the pre-order structure	
	Specifies that the relations of the pre-ordered set are about to be listed.	
rel:	The relations should only contain variables that were listed after !obj:	
	or obj:.	
#	Indicates that the following piece of text is a comment.	

Because this syntax can be compatible with the YAML grammar, the file can ideally be specified with the extension .yml. Here is an example of such a file.

```
preorder.yml
1
    #This is an example of a pre-ordered set.
2
3
    !obj:#a formal minimal object is added to the pre-order
4
    - color_1 #1st object
5
    - color_2 #2st object
6
    - color_3 #3rd object
7
    - color_4 #4th object
8
9
    rel: #these are generating relations for the preorder.
10
    - color_1 > color_2, color_3;
    - color_1 > color_3, color_1;
11
    - color_3, color_2 > color_4;
12
```

The previous pre-ordered set can be described by atomic relations shown in diagram (5.1), where the additional object! is the formal minimal object added to the pre-order structure.



We now give examples showing how the pre-ordered set preorder.yml is used by the constructor of the class PreOrder.

After calling the constructor, the object .relations will contain the pre-order relations specified in preorder.yml as a list of lists (see below). Every internal list starts with a predecessor of all the elements in that list. There is an internal list for every element specified as an object of the pre-ordered set in preorder.yml.

```
>>> Omega = PreOrder("preorder.yml")
>>> Omega.relations
[['color_1', 'color_2', 'color_3'], ['color_2', 'color_4'], ['color_3',
'color_4'], ['color_4']]
```

The object .mask is set to True if a formal minimal object has been added to the structure via the use of !obj: and is set to False otherwise.

```
>>> Omega.mask
```

#### True

Since we have not specified the value of the parameter cartesian, its value is still 0.

```
>>> Omega.cartesian
0
```

Also, since the method .closure() has not been called yet, the object .transitive is still equal to False.

```
>>> Omega.transitive
```

#### False

Calling the method .closure() will complete the lists inside .relations into down-closures for the first element of every list.

```
>>> Omega.closure()
>>> Omega.relations
[['color_1', 'color_2', 'color_3', 'color_4'], ['color_2', 'color_4'],
['color_3', 'color_4'], ['color_4']]
```

We can now verify that the call of the method .closure() set the value of the object .transitive to True.

```
>>> Omega.transitive
```

True

**5.1.4.** Description of .closure (method). The code of the function .closure has no input. The method possesses two actions, which we describe below through examples.

Action 1		
Case	If self.transitive == False	
Description	In this case, the value of self.transitive is set to True and the method	
	completes every lists contained in self.relations into down-closures	
	for the first element of each list.	

Action 2		
Case	If self.transitive == True	
Description	No action is taken.	

Here is an example that uses the file preorder.yml of section 5.1.3.1 as an input.

**5.1.5.** Description of .geq (method). The code of the function .geq is equipped with the following inputs.

.geq		
Inputs	Types	Specifications
element1	$\alpha$	necessary
element2	$\alpha$	necessary

The method possesses two actions, which we describe below through examples. Before starting each of these actions, the method calls the method .closure() to make sure that the relations contained in the object .relations define transitive down-closures.

Action 1		
Case	If self.cartesian == 0	
Description	The function considers element1 and element2 to be elements belonging	
	to the lists contained in self.relations. The method first looks for	
	element1 at the beginning of every list in self.relations and if a list	
	is found, the method looks for element2 in that list.	

The following example shows how the method compares element that belong or not to the pre-ordered set.

```
>>> Omega = PreOrder("preorder.yml")
>>> Omega.geq("?","color_1")
False
>>> Omega.geq("color_1","?")
False
>>> Omega.geq("color_4","color_4")
True
>>> Omega.geq("color_4","color_1")
False
>>> Omega.geq("color_2","color_3")
False
```

The second action allows us to simulate any Cartesian product structure on the preordered set.

Action 2		
Case	If self.cartesian > 0	
Description	The function considers element1 and element2 to be lists of elements	
	belonging to the lists contained in self.relations. For every element	
	i in range(self.cartesian), the method looks for element1[i] at	
	the beginning of every list in self.relations and if a list is found,	
	it looks for element2[i] in that list. Exceptionally for the present	
	case, any component element2[i] is allowed to take Boolean values. If	
	element2[i] happens to be equal to True, then the method skip the	
	search for element1[i].	

This second example shows how the method .geq extends to the Cartesian product of the pre-order used in the previous example.

```
>>> Cartesian = PreOrder("",2,0mega)
>>> Cartesian.geq(["color_1","color_2"],[Cartesian.mask]*2)
True
>>> Cartesian.geq([Cartesian.mask]*2,["color_1","color_2"])
False
>>> Cartesian.geq(["color_1","color_2"],["color_4","color_4"])
True
>>> Cartesian.geq(["color_1","color_2"],["color_4","color_3"])
False
```

**5.1.6.** Description of .inf (method). The code of the function .inf is equipped with the following inputs.

.inf		
Inputs	Types	Specifications
element1	$\alpha$	necessary
element2	$\alpha$	necessary

The method possesses two actions, which we describe below through examples. Before starting each of these actions, the method calls the method .closure() to make sure that the relations contained in the object .relations define transitive down-closures.

Action 1		
Case	If self.cartesian == 0	
Description	The function considers element1 and element2 to be elements belonging	
	to the lists contained in self.relations. The method first looks for	
	element1 and element2. If one of them is not found, then self.mask	
	is returned. If both elements are present, then the intersection of their	
	down-closures is computed and the first local maxima of the intersection	
	is returned. If the intersection is empty, then self.mask is returned	
	instead.	

The following example shows how the method compares element that belong or not to the pre-ordered set.

```
>>> Omega = PreOrder("preorder.yml")
>>> Omega.inf("?","color_1")
True
>>> Omega.inf("color_2","color_3")
color_4
```

The second action allows us to simulate any Cartesian product structure on the preordered set.

Action 2		
Case	If self.cartesian > 0	
Description	The function considers element1 and element2 to be lists of elements	
	belonging to the lists contained in self.relations. For every element	
	i in range(self.cartesian), the method applies Action 1 to the pair	
	(element1[i],element2[i]). The output is then appended to a list,	
	which is eventually returned.	

This second example shows how the method .geq extends to the Cartesian product of the pre-order used in the previous example.

```
>>> Cartesian = PreOrder("",2,0mega)
>>> Cartesian.inf(["color_1","color_2"],["?"]*2)
[True, True]
>>> Cartesian.inf(["color_1","color_2"],["color_4","color_3"])
['color_4', 'color_4']
```

**5.1.7. Description of .presence (method).** The code of the function .presence is equipped with the following inputs.

.inf		
Inputs	Types	Specifications
element	$\alpha$	necessary

The method possesses one action, which we describe below through an example.

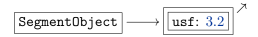
	Action
Case	Always
Description	The function looks for element at the beginning of every list contained in
	self.relations. If element is found, then True is returned, otherwise
	the method returns False.

The following examples shows how the method .presence reacts when given various inputs.

```
>>> Omega = PreOrder("preorder.yml")
>>> Omega.presence("?")
False
>>> Omega.presence("color_2")
True
```

## 5.2. Description of SegmentObject (subclass)

**5.2.1.** Introduction. The code of the class SegmentObject uses external modules.



The (sub)class SegmentObject models the features of segments (as defined in [2, section 2]). A SegmentObject item can be seen as a tape equipped with a read head, whose position is stored in the object .parse and is displayed through the method .display as a red patch of nodes.

#### (0000|0000000|000|0|000000)

The method .patch returns the index of a patch (an area in brackets) containing a node whose position is given as an input. Note that the method .patch starts searching the index associated with the node from where the read head is and only goes in a direction (left or right) specified in the second argument; the method .merge takes a tiling of the segment and merges groups of patches that share the same tiles. The tiling patterns are specified in a list of triples, where each triple contains a start index, a tile length, and an end index; the method .remove removes either a node or a patch (from the segment) at the index given in the first argument: whether a node or a patch is removed is decided upon whether the second argument is equal to "nodes-given" or is not specified.

**5.2.2. Structure.** The following tables give a preview of the class SegmentObject. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
1 0		
CategoryItem   section 3		
Statistics		
⊳ Importable objects: 4		
▷ Non-importable objects: 0		
▷ Importable methods: 5		
Non-importable methods: 1		

The following table gives a description of the 4 importable objects of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Objects		
Name Type		Related sections
.domain int		$\triangleright$ section 5.2.3
.topology	<pre>list(int * int)</pre>	$\triangleright$ section 5.2.3
.colors	$ exttt{list}( exttt{list}(lpha))$	$\triangleright$ section 5.2.3
.parse	int	$\triangleright$ section 5.2.3

Finally, the following table gives a description of the 5 importable methods of the class:

Methods			
Name	Input types	Output types	Related sections
	- int	- self	$\triangleright$ section 5.2.3
init	- list(int * int)		
	- $list(list(lpha))$		
.display	- self	- self	$\triangleright$ section 5.2.4
.display		- stdout	
.patch	- int	- int	$\triangleright$ section 5.2.5
.patch	- string		
merge	- list(int * int * int)	- SegmentObject( $lpha$ )	$\triangleright$ section 5.2.6
.merge	-fun: $\alpha$ * $\alpha$ -> $\alpha$		
.remove	- list	- SegmentObject	$\triangleright$ section 5.2.7
	- string		

**5.2.3. Description of** .\_\_init\_\_ (**method**). The code of the function .\_\_init\_\_ is equipped with the following inputs.

init		
Inputs	Types Specifications	
domain	int	necessary
topology	<pre>list(int * int)</pre>	necessary
colors	$ extstyle{list}(lpha)$	necessary

The method possesses one action, which we describe below through examples.

	Action
Case	Always
Description	The method uses the inputs domain, topology and colors to initial-
	ize the objects self.domain, self.topology and self.colors. The
	object self.parse is given the value 0.

The following examples show how the constructor can be used.

```
>>> t = list()
>>> c = list()
>>> for i in range(5):
...    t = t + [(3*i,3*i+2)]
...    c = c + ["color_1"]
...
>>> s = SegmentObject(20,t,c)
>>> s.domain
20
>>> s.topology
[(0, 2), (3, 5), (6, 8), (9, 11), (12, 14)]
>>> s.colors
['color_1', 'color_1', 'color_1', 'color_1']
>>> s.parse
0
```

**5.2.4.** Description of .display (method). The code of the function .display has no input. The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	Displays on the standard output a segment equipped with a 'read head'	
	displayed as a <b>red</b> patch of nodes whose index matches the index con-	
	tained in self.parse. The intervals defined by the pairs contained in	
	self.topology are displayed as <b>bold</b> patches of nodes while the nodes	
	that are not comprised within pairs in self.topology are displayed	
	using a normal font.	

The following examples show the action of the method .display on the segment s defined at the end of section 5.2.3.

```
>>> s.display()
(ooo|ooo|ooo|ooo|ooooooo)
>>> s.parse = 3
>>> s.display()
(ooo|ooo|ooo|ooo|ooo|ooooooooo)
```

**5.2.5.** Description of .patch (method). The code of the function .patch is equipped with the following inputs, among which one is optional.

.patch		
Inputs	Types	Specifications
position	int	necessary
search = ">1"	string	optional

The method possesses two actions, which we describe below through examples.

	Action 1		
Case	If search == ">"+str(n) where n is of the type int		
Description	In this case, the list self.topology is parsed from the pair at index		
	self.parse to the pair at index len(self.topology)-1 with an incre-		
	mentation equal to n. If the value position is within the range of one		
	of the parsed pairs in self.topology, then the index of the pair (within		
	self.topology) is returned. In any other case, the value -1 is returned.		

Action 2		
Case	If search == "<"+str(n) where n is of the type int	
Description	In this case, the list self.topology is parsed from the pair at index	
	self.parse to the pair at index 0 with an incrementation equal to n.	
	If the value position is within the range of one of the parsed pairs in	
	self.topology, then the index of the pair (within self.topology) is	
	returned. In any other case, the value -1 is returned.	

The following examples show the action of the method .patch on the segment s considered at the end of section 5.2.4.

```
>>> s.display()
(000|000|000|000|000|00000)
>>> s.patch(5,"<1")
>>> s.display()
(000|000|000|000|0000)
>>> s.patch(7,">1")
>>> s.display()
(000|000|000|000|000|00000)
>>> s.patch(11,">2")
-1
>>> s.display()
(000|000|000|000|000|00000)
>>> s.patch(4,"<2")
-1
>>> s.display()
(000|000|000|000|000|0000)
```

5.2.6. Description of .merge (method). The code of the function .merge is equipped with the following inputs.

.merge		
Inputs	Types	Specifications
folding_format	<pre>list(int * int * int)</pre>	necessary
infimum	fun: $\alpha * \alpha \rightarrow \alpha$	necessary

The method possesses one action.

	Action	
Case	Always	
Description	The method returns a new segment that is a merged version of	
	self. Specifically, the patches of self are merged according to	
	the tiling patterns specified in folding_format, in which a tiling	
	pattern is given by a pair (x,1,y). For every pair (x,1,y) in	
	folding_format, the method will transitively merge any subset of	
	patches whose indices in self.topology belong to a same interval of	
	the form $(x+k*1,min(x+(k+1)*1,y))$ . For each merging, the resulting	
	pair that is included in the new topology consists of the first compo-	
	nent of the first merged pair and the last component of the last merged	
	pair. The color associated with that pair is the output of the recur-	
	sive two-by-two application $f(_{-},(_{-},f(_{-},f(_{-},_{-}))))$ of the function $f$	
	:= infimum.	

The following examples will probably make the previous description more explicit. We use the preOrder constructed from the file preorder.yml shown in section 5.1.3.1.

```
>>> Omega = PreOrder("preorder.yml")
>>> t = list()
>>> c = list()
>>> for i in range(5):
      t = t + [(3*i, 3*i+2)]
      if i % 2 == 0:
        c = c + ["color_2"]
      if i % 2 == 1:
         c = c + ["color_3"]
>>> s = SegmentObject(20,t,c)
>>> s.display()
(000|000|000|000|0000)
>>> s1 = s.merge([(0,1,3)], Omega.inf)
>>> s1.display()
(000|000|000|000|000|00000)
>>> s1.colors
['color_2', 'color_3', 'color_2', 'color_3', 'color_2']
>>> s1 = s.merge([(0,2,3)],Omega.inf)
>>> s1.display()
(000000|000000|000|00000)
>>> s1.colors
['color_4', 'color_4', 'color_2']
>>> s1 = s.merge([(0,3,3)],Omega.inf)
>>> s1.display()
(000000000|000|000|00000)
>>> s1.colors
['color_4', 'color_3', 'color_2']
>>> s1 = s.merge([(0,4,3)],Omega.inf)
>>> s1.display()
(00000000000|000|00000)
>>> s1.colors
['color_4', 'color_2']
>>> s1 = s.merge([(0,5,3)],Omega.inf)
>>> s1.display()
(00000000000|000|00000)
>>> s1.colors
['color_4', 'color_2']
```

**5.2.7.** Description of .remove (method). The code of the function .remove is equipped with the following inputs, among which one is optional.

init			
Inputs Types Specification			
a_list	list	necessary	
option = "patches-given"	string	optional	

The method possesses two actions, which we describe below through examples.

Action 1		
Case	If option != "nodes-given"	
Description	In this case, the method removes every patch whose index is an integer	
	i in a_list.	

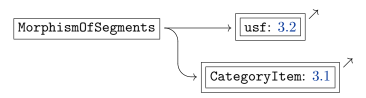
Action 2			
Case	If option == "nodes-given"		
Description	In this case, the method removes every patch whose index is an in-		
	teger i for which there exists an index j in a_list such that i =		
	<pre>self.patch(j).</pre>		

The following examples shows the effects of choosing either two options. We use the segment s defined at the end of section 5.2.6.

```
>>> s.display()
(000|000|000|000|000|00000)
>>> s1 = s.remove([0],"patches-given")
>>> s1.display()
(000|000|000|000|0000)
>>> s1 = s.remove([0], "nodes-given")
>>> s1.display()
(000|000|000|000|0000)
>>> s1 = s.remove([1],"patches-given")
>>> s1.display()
(000|000|000|000|000|00000)
>>> s1 = s.remove([1], "nodes-given")
>>> s1.display()
(000|000|000|000|0000)
>>> s1 = s.remove([2], "patches-given")
>>> s1.display()
(000|000|000|000|0000)
>>> s1 = s.remove([2], "nodes-given")
>>> s1.display()
(000|000|000|000|0000)
>>> s1 = s.remove([0,1,2,6],"patches-given")
>>> s1.display()
(000000000|000|000|00000)
>>> s1 = s.remove([0,1,2,6],"nodes-given")
>>> s1.display()
(000|000|000|000|0000)
```

## 5.3. Description of MorphismOfSegments (subclass)

**5.3.1.** Introduction. The code of the class MorphismOfSegments uses external modules.



The (sub)class MorphismOfSegments models the features of a morphism of segments (as defined in [2]). As such, its constructor takes a source segment, a target segment, a list f1 of increasing integers describing the image of an injection  $i \mapsto f1[i]$  relating the domain of

the source segment to the domain of the target segment, and a pre-order relation compatible with the pre-ordered sets of two the segments.

**5.3.2. Structure.** The following tables give a preview of the class MorphismOfSegments. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
CategoryItem	section 3	
Statistics		
⊳ Importable objects: 3		
▷ Non-importable objects: 0		
▷ Importable methods: 1		
▷ Non-importable methods: 1		

The following table gives a description of the 3 importable objects of the class:

Objects			
Name	Type	Related sections	
.defined	bool	$\triangleright$ section 5.3.3	
.f0 list(int)		$\triangleright$ section 5.3.3	
.f1	<pre>list(int) or string</pre>	$\triangleright$ section 5.3.3	

Finally, the following table gives a description of the only importable method of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Methods			
Name Input types Output types Related s			
init	- SegmentObject(α) - SegmentObject(α) - list(int) - fun: α * α -> bool	- self	⊳ section 5.3.3

**5.3.3. Description of** .\_\_init\_\_ (**method**). The code of the function .\_\_init\_\_ is equipped with the following inputs.

$.\_$ init $\_$			
Inputs	Types	Specifications	
source	${\tt SegmentObject}(\alpha)$	necessary	
target	${\tt SegmentObject}(\alpha)$	necessary	
f1	<pre>list(int) or string</pre>	necessary	
geq	fun: $\alpha * \alpha \rightarrow bool$	necessary	

The method possesses two actions, which we describe below through examples.

Action 1		
Case	If f1 == "id"	
Description	In this case, the super objects self.source and self.target are given	
	the value of the variables source and target, respectively. The object	
	self.f1 is given the value "id" and the object self.f0 will given the	
	list containing the integers self.target.patch(i), for every index i in	
	range(self.source.topology) if self.f0 defines a function. In this	
	case, the object self.defined is set to True.	

Action 2			
Case	If f1 != "id" and f1 is of the type list(int)		
Description	In this case, the super objects self.source and self.target are given		
	the value of the variables source and target, respectively. The object		
	self.f1 is given the list stored in f1 and the object self.f0 will be		
	given the list containing the integers self.target.patch(f1[i]), for		
	every index i in range(self.source.topology) if self.f0 defines a		
	function. In this case, the object self.defined is set to True.		

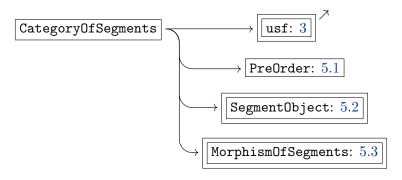
The following examples illustrate the previous actions. We consider the lists t and c defined in section 5.2.6.

```
>>> s = SegmentObject(20,t,c)
>>> t = SegmentObject(25,t,c)
>>> s.display()
(000|000|000|000|0000)
>>> t.display()
(000|000|000|000|000|00000000)
>>> s1 = s.remove([2])
>>> s1.display()
(000|000|000|000|0000)
>>> t1 = t.remove([0,2,3])
>>> t1.display()
(000|000|00000|000|00000)
>>> f1 = range(20)
>>> m = MorphismOfSegments(s1,t1,f1,Omega.geq)
>>> m.defined
True
>>> m.f0
[-1, 0, -1, 1]
>>> m.f1
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
```

Not all morphisms are well-defined. For instance, the following examples contained non-well-defined morphisms of segments. We obtained these examples by changing the topology or the mapping f1. As can be seen, the resulting object .f0 is an empty list.

## 5.4. Description of CategoryOfSegments (class)

**5.4.1.** Introduction. The code of the class CategoryOfSegments uses other functions of the module SegmentCategory.py as well as external modules.



The class CategoryOfSegments models the features of a category of segments. The class is initialized by passing a PreOrder item to its constructor and allows one to know or compute information related to a category structure. The method .identity returns a Boolean value that specificies whether there may exist an identity morphism between two SegmentObject items (these may be saved in different places in the memory); the method .initial returns an local initial object in the category, where the local aspect is determined by the colors of the segment; the method .homset computes the hom-set of a pair of SegmentObject items.

**5.4.2. Structure.** The following tables give a preview of the class CategoryOfSegments. The table given below describe the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
object	N/A	
Statistics		
⊳ Importable objects: 1		
▷ Non-importable objects: 0		
▷ Importable methods: 4		
▷ Non-importable methods: 0		

The following table gives a description of the only importable object of the class:

Objects		
Name Type Related sections		
.preorder PreOrder		$\triangleright$ section 5.4.3

Finally, the following table gives a description of the 4 importable methods of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Methods			
Name	Input types	Output types	Related sections
init	- PreOrder	- self	⊳ section 5.4.3
.identity	- SegmentObject	- bool	⊳ section 5.4.4
.identity	- SegmentObject		
.initial	- int	- SegmentObject	$\triangleright$ section 5.4.5
. IIII CIAI	- $list(list(lpha))$		
.homset	- SegmentObject	- list(MorphismOfSegments)	$\triangleright$ section 5.4.6
.Homset	- SegmentObject		

**5.4.3. Description of** .\_\_init\_\_ (method). The code of the function .\_\_init\_\_ is equipped with the following inputs.

init		
Inputs	Specifications	
preorder	PreOrder	necessary

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	The method allocates the value of the variable preorder to the object	
	self.preorder	

The following example shows how the constructor of the class CategoryOfSegments can be used with the class PreOrder (see section 5.1). We use the file preorder.yml displayed in section 5.1.3.1.

```
>>> Omega = PreOrder("preorder.yml")
>>> Seg = CategoryOfSegments(Omega)
>>> Seg.preorder.relations
[['color_1', 'color_2', 'color_3'], ['color_2', 'color_4'], ['color_3',
'color_4'], ['color_4']]
>>> Seg.preorder.mask
True
```

**5.4.4. Description of .identity (method).** The code of the function .identity is equipped with the following inputs.

.identity		
Inputs Types Specification		
segment1	SegmentObject	necessary
segment2	SegmentObject	necessary

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	The method tests whether each type of objects .domain, topology and	
	colors are equal for both inputs segment1 and segment2.	

In the following example, we use the file preorder.yml displayed in section 5.1.3.1 and the method .initial detailed in section 5.4.5.

If we now use the method .remove, detailed in section 5.2.7, to change the colors of t, then the identity does not hold anymore, as sown in the following example.

```
>>> t = t.remove([15])
>>> t.display()
(o|o|o|o|o|o|o|o|o|o|o|o|o|o|o|o|o)
>>> Seg.identity(s,t)
False
```

**5.4.5. Description of .initial (method).** The code of the function .initial is equipped with the following inputs.

.initial			
Inputs   Types   Specifications			
domain	int	necessary	
color $\alpha$		necessary	

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	The method returns the SegmentObject item whose object .topology	
	consists of the pairs (i,i) for every i in range(domain) and whose	
	object .colors is equal to the list [color]*domain.	

The following example shows what a SegmentObject item generated by the method .initial looks like. We use the file preorder.yml displayed in section 5.1.3.1 and the constructor of the lass PreOrder (see section 5.1).

**5.4.6.** Description of .homset (method). The code of the function .homset is equipped with the following inputs

.homset		
Inputs	Types	Specifications
source	SegmentObject	necessary
target	SegmentObject	necessary

The method possesses one action, which we describe below through examples. We use the file preorder.yml displayed in section 5.1.3.1 and the method .initial detailed in section 5.4.5.

Action		
Case	Always	
Description	The method returns a list containing all MorphismOfSegments items for	
	which the object .source is equal to source and the object .target	
	is equal to target relative to the preorder structure contained in	
	self.preorder.	

The following example shows what the output of the method .homset looks like. We use the file preorder.yml displayed in section 5.1.3.1, the method .initial detailed in section

5.4.5, the method .merge detailed in section 5.2.6 and the constructor of the class PreOrder (see section 5.1).

```
>>> Omega = PreOrder("preorder.yml")
>>> Seg = CategoryOfSegments(Omega)
>>> s = Seg.initial(8, "color_1")
>>> s = s.merge([(0,2,7)],Omega.inf)
>>> t = Seg.initial(12, "color_1")
>>> t = t.merge([(0,3,10)], Omega.inf)
>>> s.display()
(00|00|00|00)
>>> t.display()
(000|000|000|00|0)
>>> for m in Seg.homset(s,t):
       print m.f1
. . .
[0, 1, 3, 4, 6, 7, 9, 10]
[0, 1, 3, 4, 6, 8, 9, 10]
[0, 1, 3, 4, 7, 8, 9, 10]
[0, 1, 3, 5, 6, 7, 9, 10]
[0, 1, 3, 5, 6, 8, 9, 10]
[0, 1, 3, 5, 7, 8, 9, 10]
[0, 1, 4, 5, 6, 7, 9, 10]
[0, 1, 4, 5, 6, 8, 9, 10]
[0, 1, 4, 5, 7, 8, 9, 10]
[0, 2, 3, 4, 6, 7, 9, 10]
[0, 2, 3, 4, 6, 8, 9, 10]
[0, 2, 3, 4, 7, 8, 9, 10]
[0, 2, 3, 5, 6, 7, 9, 10]
[0, 2, 3, 5, 6, 8, 9, 10]
[0, 2, 3, 5, 7, 8, 9, 10]
[0, 2, 4, 5, 6, 7, 9, 10]
[0, 2, 4, 5, 6, 8, 9, 10]
[0, 2, 4, 5, 7, 8, 9, 10]
[1, 2, 3, 4, 6, 7, 9, 10]
[1, 2, 3, 4, 6, 8, 9, 10]
[1, 2, 3, 4, 7, 8, 9, 10]
[1, 2, 3, 5, 6, 7, 9, 10]
[1, 2, 3, 5, 6, 8, 9, 10]
[1, 2, 3, 5, 7, 8, 9, 10]
[1, 2, 4, 5, 6, 7, 9, 10]
[1, 2, 4, 5, 6, 8, 9, 10]
[1, 2, 4, 5, 7, 8, 9, 10]
```

# Presentation of the module AlignedFunctor.py

## 6.1. Description of PointedSet (class)

**6.1.1.** Introduction. The class PointedSet models the features of a pointed set. It is made of a list .symbols, representing the underlying set, and an integer .index indicating where the point is locted whithin .symbols.

**6.1.2.** Structure. The following tables give a preview of the class PreOrder. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
object N/A		
Statistics		
⊳ Importable objects: 2		
⊳ Non-importable objects: 0		
⊳ Importable methods: 2		
⊳ Non-importable methods: 0		

The following table gives a description of the 2 importable objects of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Objects		
Name Type Related sections		
.symbols	$ extstyle{list}(lpha)$	$\triangleright$ section 6.1.3
.index	int	⊳ section 6.1.3

Finally, the following table gives a description of the 2 importable methods of the class:

Methods			
Name	Input types	Output types	Related sections
init	- $list(lpha)$	- self	$\triangleright$ section 6.1.3
•	- int		
.point	- self	<i>-</i> α	$\triangleright$ section 6.1.3

**6.1.3.** Description of .\_\_init\_\_ (method). The code of the function .\_\_init\_\_ is equipped with the following inputs.

init			
Inputs Types Specifications			
symbols	string	necessary	
index	int	necessary	

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	The method allocates the value stored in the input variable symbols	
	to the object self.symbols. If the index is in the range of the list	
	self.symbols, then the value of the input variable index is passed to	
	the object self.index. Otherwise, the object self.index is set to 0.	

In the following example, we see how the object .index can be used with the object .symbols to return the 'point' of the PointedSet item.

```
>>> E = PointedSet(["-","A","C","G","T"],0)
>>> E.symbols
['-','A','C','G','T']
>>> E.index
0
>>> E.symbols[E.index]
```

**6.1.4.** Description of .point (method). The code of the function .point has no input. The method possesses one action, which we describe below through an example.

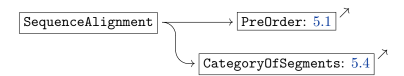
Action		
Case	Always	
Description	iption The method returns the point self.symbols[self.index] of the struc-	
	ture.	

The following example ilustrates the use of the method .point.

```
>>> E = PointedSet(["A","C","G","T","epsilon"],4)
>>> E.point()
epsilon
```

## 6.2. Description of SequenceAlignment (class)

**6.2.1.** Introduction. The code of the class SequenceAlignment uses external modules.



The class SequenceAlignment models the features of a sequence alignment functor, as defined in [2]. The images of the sequence alignment functor are stored in the object .database and can be queried throught the method .eval. The method also computes the images of the right Kan extension of this functor through the method .ran (TO BE CODED). The objects of the extending category (see [2, Definition 3.25]), which is used to compute this right Kan extension, can be obtained through the method .extending\_category.

**6.2.2. Structure.** The following tables give a preview of the class **SequenceAlignment**. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
object	N/A	
Statistics		
⊳ Importable objects: 5		
▷ Non-importable objects: 0		
▷ Importable methods: 4		
▷ Non-importable methods: 0		

The following table gives a description of the 5 importable objects of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Objects			
Name	Type	Related sections	
.env	Environment	$\triangleright$ section 6.2.3	
.Seg	list	$\triangleright$ section 6.2.3	
.indiv	list	$\triangleright$ section 6.2.3	
.base	<pre>list(SegmentObject)</pre>	$\triangleright$ section 6.2.3	
.database	$ extsf{list}(lpha)$	$\triangleright$ section 6.2.3	

Finally, the following table gives a description of the 4 importable methods of the class:

Methods			
Name	Input types	Output types	Related sections
init	- Environment - CategoryOfSegments - list(SegmentObject) - list(\alpha)	- self	⊳ section 6.2.3
.eval	- SegmentObject	- $lpha$ or list	$\triangleright$ section 6.2.4
.extending_category	- SegmentObject	- list(int* MorphismOfSegments)	⊳ section 6.2.5
.ran	- CategoryItem	- $lpha$ or list	$\triangleright$ section 6.2.6

**6.2.3.** Description of .\_\_init\_\_ (method). The code of the function .\_\_init\_\_ is equipped with the following inputs.

init			
Inputs	Types	Specifications	
env	Environment	necessary	
indiv	list	necessary	
base	<pre>list(SegmentObject)</pre>	necessary	
database	$ extstyle{list}(lpha)$	necessary	

The method possesses one action, which we describe below through an example.

Action		
Case	Always	
Description	The method allocates the values stored in the input variables env, indiv,	
	base and database to the objects self.env, self.indiv, self.base	
	and self.database. It also uses the object self.env.spec to compute	
	the (self.env.spec)-fold Cartesian product of the pre-ordered set en-	
	coded by self.env.Seg.preorder and allocates to the object self.Seg	
	the CategoryOfSegments item that can be constructed from it.	

The following examples show how to use the constructor of the class in combination with the classes PreOrder (see section 5.1), CategoryOfSegments (see section 5.4), PointedSet (see section 6.1) and Environment (see section 6.3). We use the file preorder.yml displayed in section 5.1.3.1.

```
>>> Omega = PreOrder("preorder.yml")
>>> Seg = CategoryOfSegments(Omega)
>>> E = PointedSet(["-","A","C","G","T"],0)
>>> Env = Environment(Seg,E,2,["color_2"]*2)
>>> base = [Env.Seg.initial(8,["color_1"]*2), Env.Seg.initial(8,["color_2"]*2),
Env.Seg.initial(5,["color_2"]*2), Env.Seg.initial(7,["color_3"]*2)]
```

In practice, the following list should contain actual sequence alignments, namely each string item should be replaced with a succession of string items representing the sequences of a sequence alignment.

```
>>> database = [[["Alignment:1:1"], ["Alignment:1:2"]], [["Alignment:2:1"],
["Alignment:2:2"]], [["Alignment:3:1"],["Alignment:3:2"],
["Alignment:3:3"]], [["Alignment:4:1"],["Alignment:4:2"]]]
```

The following code shows how to recover the data passed to the constructor using the various objects of the class.

```
>>> Seqali = SequenceAlignment(Env,["A","B","C","D"],base,database)
>>> Segali.indiv
['A', 'B', 'C', 'D']
>>> for i in range(len(Seqali.base)):
       print str(i)+") color: "+ str(Seqali.base[i].colors[
Segali.base[i].parse ])
       Seqali.base[i].display()
       for j in range(len(Seqali.database[i])):
         for k in range(len(Seqali.database[i][j])):
           print Seqali.database[i][j][k]
       print ""
0) color: ['color_1', 'color_1']
(0|0|0|0|0|0|0|0)
Alignment:1:1
Alignment:1:2
1) color: ['color_2', 'color_2']
(0|0|0|0|0|0|0)
Alignment:2:1
Alignment:2:2
           ['color_2', 'color_2']
2) color:
```

```
(o|o|o|o|o)
Alignment:3:1
Alignment:3:2
Alignment:1:3

3) color: ['color_3', 'color_3']
(o|o|o|o|o|o|o)
Alignment:4:1
Alignment:4:2
```

**6.2.4.** Description of .eval (method). The code of the function .eval is equipped with the following input.

.eval			
Inputs Types Specifications			
segment	SegmentObject	necessary	

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	The method returns the image of the sequence alignment functor for the	
	given input SegmentObject item. If the input does not belong to the	
	list stored in self.base, then the empty list is returned.	

The following example shows that the images of the SegmentObject items stored in the object .base recovers the elements stored in the object .database. We use the Environment item Env and the SequenceAlignment item Sequli constructed in section 6.2.3.

```
>>> for i in range(len(Seqali.base)):
       print "image for base[" + str(i) +"]"
       Seqali.base[i].display()
       sal = Seqali.eval(Seqali.base[i])
       for j in range(len(sal)):
         for k in range(len(sal[j])):
           print sal[j][k]
         print ""
image for base[0]
(0|0|0|0|0|0|0|0)
Alignment:1:1
Alignment:1:2
image for base[1]
(0|0|0|0|0|0|0|0)
Alignment:2:1
Alignment:2:2
```

```
image for base[2]
(o|o|o|o|o)
Alignment:3:1
Alignment:3:2
Alignment:3:3

image for base[3]
(o|o|o|o|o|o|o
Alignment:4:1
Alignment:4:2
```

For every SegmentObject item that is not in the object .base, the method .eval will return the empty list, as shown below.

**6.2.5.** Description of .extending\_category (method). The code of the function .extending\_category is equipped with the following input.

.extending_category			
Inputs Types Specifications			
segment	SegmentObject	necessary	

The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	This method computes the set of objects of the extending category de-	
	fined in [2, Definition 3.25]. More specifically, the method returns the	
	list of pairs (i,m) such that m is a MorphismOfSegments item whose	
	source object is segment and whose target object is self.base[i].	

The following example shows an example of calculation of an extending category.

```
>>> diag = Seqali.extending_category(Env.Seg.initial(7,["color_1"]*2))
>>> for i,m in diag:
...    print i
...    print m.f1
...    print m.f0
...    m.source.display()
...    m.target.display()
...    for j in range(len(Seqali.database[i])):
...    for k in range(len(Seqali.database[i][j])):
...    print Seqali.database[i][j][k]
...    print ""
```

```
0
[0, 1, 2, 3, 4, 5, 6]
[0, 1, 2, 3, 4, 5, 6]
(0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:1:1
Alignment:1:2
[0, 1, 2, 3, 4, 5, 7]
[0, 1, 2, 3, 4, 5, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(0 \mid 0 \mid 0 \mid 0 \mid 0 \mid 0 \mid 0)
Alignment:1:1
Alignment:1:2
0
[0, 1, 2, 3, 4, 6, 7]
[0, 1, 2, 3, 4, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:1:1
Alignment:1:2
0
[0, 1, 2, 3, 5, 6, 7]
[0, 1, 2, 3, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:1:1
Alignment:1:2
[0, 1, 2, 4, 5, 6, 7]
[0, 1, 2, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(0|0|0|0|0|0|0|0)
Alignment:1:1
Alignment:1:2
[0, 1, 3, 4, 5, 6, 7]
[0, 1, 3, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:1:1
Alignment:1:2
```

```
0
[0, 2, 3, 4, 5, 6, 7]
[0, 2, 3, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:1:1
Alignment:1:2
[1, 2, 3, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0)
(0|0|0|0|0|0|0)
Alignment:1:1
Alignment:1:2
[0, 1, 2, 3, 4, 5, 6]
[0, 1, 2, 3, 4, 5, 6]
(0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:2:1
Alignment:2:2
[0, 1, 2, 3, 4, 5, 7]
[0, 1, 2, 3, 4, 5, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(o|o|o|o|o|o|o|o)
Alignment:2:1
Alignment:2:2
[0, 1, 2, 3, 4, 6, 7]
[0, 1, 2, 3, 4, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:2:1
Alignment:2:2
[0, 1, 2, 3, 5, 6, 7]
[0, 1, 2, 3, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(o|o|o|o|o|o|o|o)
Alignment:2:1
Alignment:2:2
```

```
1
[0, 1, 2, 4, 5, 6, 7]
[0, 1, 2, 4, 5, 6, 7]
(0 \mid 0 \mid 0 \mid 0 \mid 0 \mid 0 \mid 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:2:1
Alignment:2:2
[0, 1, 3, 4, 5, 6, 7]
[0, 1, 3, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0 | 0)
Alignment:2:1
Alignment:2:2
[0, 2, 3, 4, 5, 6, 7]
[0, 2, 3, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(o|o|o|o|o|o|o|o)
Alignment:2:1
Alignment:2:2
[1, 2, 3, 4, 5, 6, 7]
[1, 2, 3, 4, 5, 6, 7]
(0 | 0 | 0 | 0 | 0 | 0)
(o|o|o|o|o|o|o|o)
Alignment:2:1
Alignment:2:2
3
id
[0, 1, 2, 3, 4, 5, 6]
(0 | 0 | 0 | 0 | 0 | 0 | 0)
(0 | 0 | 0 | 0 | 0 | 0)
Alignment:4:1
Alignment:4:2
```

**6.2.6.** Description of .ran (method). The code of the function .ran is equipped with the following input.

.ran		
Inputs	Specifications	
cat_item		necessary
B = ""	string	optional

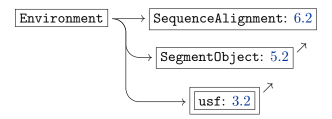
The method possesses one action, which we describe below through examples.

Action		
Case	Always	
Description	This method computes the images of the right Kan extension of the	
	functor encoded by the method self.eval.	

TO BE CODED

### 6.3. Description of Environment (class)

**6.3.1.** Introduction. The code of the class Environment uses other functions of the module AlignedFunctor.py as well as external modules.



The class Environment models the features of an aligned environment functor, as defined in [2, section 3]. As with aligned environment functors, this structure is associated with a PointedSet item .pset. The class is also equipped with a fiber operation (pullback along a point in the image of the aligned environment functor)

$$\{ au\}$$
  $\stackrel{\subseteq}{-}$   $\mathbf{Seg}(\Omega)$ 
 $\downarrow$   $\mathbf{A}E_b^{arepsilon}$ 
 $\mathbf{1}$   $\stackrel{\mathsf{input}}{\longrightarrow}$   $\mathbf{Set}$ 

and a sequence alignment functor constructor call. Specifically, the method .segment returns a segment that is the pullback of the aligned environment functor above any input list that represents an element in one of its images. If the input list contains a character that is not in the object .pset.symbols, then the node associated with that character is masked in the returned segment. Finally, the class Environment is equipped with a method .seqali that constructs a sequence alignment functor from a file of sequence alignments, as shown in the discussion of [2, Example 3.22].

**6.3.2. Structure.** The following tables give a preview of the class **Environment**. The table given below describes the various dependencies of the class.

Dependencies		
Superclass ancestry   Module section		
object	N/A	
Statistics		
▷ Importable objects: 4		
▷ Non-importable objects: 0		
⊳ Importable methods: 3		
Non-importable methods: 0		

The following table gives a description of the 4 importable objects of the class. The type  $\alpha$  shown in the table is a polymorphic type.

Objects		
Name	Type	Related sections
.Seg	${\tt CategoryOfSegments}(lpha)$	$\triangleright$ section 6.3.3
.pset	$\texttt{PointedSet}(\alpha)$	$\triangleright$ section 6.3.3
.spec	int	$\triangleright$ section 6.3.3
.b	$ extsf{list}(lpha)$	$\triangleright$ section 6.3.3

Finally, the following table gives a description of the 3 importable methods of the class. The type  $\beta$  shown in the table is a polymorphic type.

Methods			
Name	Input types	Output types	Related sections
init	- CategoryOfSegments( $\alpha$ ) - PointedSet( $\alpha$ ) - int - list( $\alpha$ )	- self	⊳ section 6.3.3
.segment	- list( $eta$ ) - $lpha$	- self	⊳ section 6.3.4
.seqali	- string	- SequenceAlignment	⊳ section 6.3.5

**6.3.3. Description of** .\_\_int\_\_ (**method**). The code of the function .\_\_init\_\_ is equipped with the following inputs.

$.\_$ init $\_$		
Inputs	Types	Specifications
Seg	CategoryOfSegments(lpha)	necessary
pset	$\texttt{PointedSet}(\alpha)$	necessary
exponent	int	necessary
threshold	list(lpha)	necessary

The method possesses two actions, which we describe below through examples.

Action 1	
Case	If ∀ i such that self.Seg.preorder.presence(self.b[i]) == True
Description	In this case, the method allocates the values stored in the input vari-
	ables Seg, pset, exponent, and threshold to the objects self.Seg,
	self.pset, self.exponent, and self.threshold, respectively. In ad-
	dition, the method checks whether the length of the list self.b is equal
	to self.spec. If not, then self.b is completed into such a list by
	appending copies of the value self.Seg.preorder.mask to it.

	Action 2
Case	If $\exists$ i such that self.Seg.preorder.presence(self.b[i]) == False
Description	In this case, the method allocates the values stored in the input vari-
	ables Seg, pset, exponent, and threshold to the objects self.Seg,
	self.pset, self.exponent, and self.threshold, respectively, and re-
	places every element self.b[i] for which the case stated above holds
	with the Boolean value self.Seg.preorder.mask. If the length of the
	list self.b is not equal to self.spec, then self.b is completed into
	such a list by appending copies of the value self.Seg.preorder.mask
	to it.

The following examples show how to use the constructor of the class in combination with the classes PreOrder (see section 5.1), CategoryOfSegments (see section 5.4), and PointedSet (see section 6.1). We use the file preorder.yml displayed in section 5.1.3.1.

```
>>> Omega = PreOrder("preorder.yml")
>>> Seg = CategoryOfSegments(Omega)
>>> E = PointedSet(["-","A","C","G","T"],0)

We start by illustrating a case in which Action 1 is applied by the method.
>>> Env = Environment(Seg,E,4,["color_2"]*4)
>>> Env.Seg.preorder.relations
[['color_1', 'color_2', 'color_3'], ['color_2', 'color_4'], ['color_3', 'color_4'], ['color_4']]
>>> Env.pset.symbols
['-', 'A', 'C', 'G', 'T']
>>> Env.pset.point()
-
>>> Env.spec
4
>>> Env.b
['color_2', 'color_2', 'color_2', 'color_2']
```

The following example illustrates a case in which the object .b does not contain valid elements and whose length is too short.

**6.3.4.** Description of .segment (method). The code of the function .segment is equipped with the following inputs.

init		
Inputs	Types	Specifications
a_list	list(eta)	necessary
color	$\alpha$	necessary

The method possesses one action, which we describe below through an example.

Action	
Case	Always
Description	The method takes a list of symbols a_list whose elements are assumed
	to belong to the object self.Env.pset.symbols and returns the pull-
	back (a SegmentObject item) of the underlying environment functor
	(defined by self) above the element represented by a_list. Every ele-
	ment of the list a_list that is not a symbol in self.Env.pset.symbols
	is removed from the topology of the SegmentObject item.

The following example shows what the effect of the method .segment on a list of strings looks like. We use the Environment item defined at the end of section 6.3.3.

```
>>> r = Env.segment(["A","T","G","ERROR","-","-","G","?","?"],"color_1")
>>> r.display()
(o|o|o|o|o|o|o|o)
>>> r.colors
['color_1', 'color_1', 'color_1', 'color_1', 'color_1']
```

**6.3.5.** Description of .seqali (method). The code of the function .seqali is equipped with the following input.

.seqali					
Inputs	Types	Specifications			
name_of_file	string	necessary			

The method possesses one action, which we describe below through examples.

Action						
Case	Always					
Description	The method copies the construction given [2, Example 3.22] and pro-					
	duces a sequence alignment functor (i.e. a SequenceAlignment item -					
	see section 6.2) from a FASTA file of sequence alignments. The input					
	file, referenced through the input variable name_of_file, should fol-					
	low the format specified in section 3.2.5.1. More specifically, every se-					
	quence label succeeding the usual key symbol > should take the form					
	group_label:individual:color, where group_label indicates a ref-					
	erence number gathering sequences under a same sequence alignment,					
	where individual is the name of the individual from which the sequence					
	comes, and color is the color that will be used to color the segment as-					
	sociated with the sequence.					

Let us consider the following FASTA file of sequence alignments whose sequence labels takes the form group\_label:individual:color.

```
alignment.fa
1
    >1:A:color_1
2
    ACTCGATCTCTG?TCGATCGATCG
3
    CCTATCGGATCGATC
4
    >1:B:color_1
5
    ACTCTAT?TCTATCC-ACTCATCA
6
    CCTACTATCTCGAAA
7
    >1:C:color_1
8
    ACTCTATC---TCCGACT?ATCA
9
    CCTACTATCTCGAAA
10
    >1:D:color_1
11
    ACTATTTA?TTTC----TTTTCTA
12
    CCGGGCTGGGGGGG
13
    >2:A:color_1
14
    ACTCGATCGGATC-AT??CGATCG
15
    CCTATCGGATCGATCG
16
    >2:B:color_1
17
    ACTCC?CCTA--CTGGGCTGCTCA
    CCTACTATCTCGAAAG
19
    >2:C:color_1
20
    ACTCTCTATC?CTCTATC---TCA
21
    CCTACTATC?CGAAAG
22
    >2:D:color_1
    ACTATT-??T--AC--CTTTTCTA
23
24
    CCGGGCTTTTCGAAAA
```

The following example shows what the effect of the applying the method .seqali on the previous file is. In particular, we display the outputs of the method .eval applied on the SegmentObject items contained in the object .base.

```
>>> Seqali = Env.seqali("alignment.fa")
>>> for i in range(len(Seqali.base)):
      print "image for base[" + str(i) + "]"
      Seqali.base[i].display()
      print "color: "+str(Seqali.base[i].colors[Seqali.base[i].parse])
      sal = Seqali.eval(Seqali.base[i])
      for j in range(len(sal)):
        for k in range(len(sal[j])):
         print sal[j][k]
       print ""
image for base[0]
|o|o|o
       ['color_1', 'color_1', 'color_1', 'color_1']
ACTCGATCTCTG?TCGATCGATCGCCTATCGGATCGATC
ACTCTAT?TCTATCC-ACTCATCACCTACTATCTCGAAA
ACTCTATC---TCCGACT?ATCACCTACTATCTCGAAA
ACTATTTA?TTTC----TTTTCTACCGGGCTGGGGGGGG
```

image for base[1]

color: ['color\_1', 'color\_1', 'color\_1']

ACTCGATCGGATC-AT??CGATCGCCTATCGGATCGATCG

ACTCC?CCTA--CTGGGCTGCTCACCTACTATCTCGAAAG

ACTCTCTATC?CTCTATC---TCACCTACTATC?CGAAAG

ACTATT-??T--AC--CTTTTCTACCGGGCTTTTCGAAAA

### Presentation of the module PartitionCategory.py

#### 7.1. Description of \_image\_of\_partition

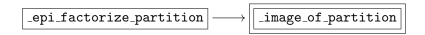
The function <u>\_image\_of\_partition</u> takes a list of elements and returns the list of its elements without repetition in the order in which they can be accessed from the left to the right.

```
1 def _image_of_partition(partition):
2    """ the source code of this function can be found in iop.py """
3    return the_image

    This corresponds to returning the image object of the underlying partition of the list.

>>> print(_image_of_partition([3,3,2,1,1,2,4,5,6,5,2,6]))
[3, 2, 1, 4, 5, 6]
>>> print(_image_of_partition(['A',4,'C','C','G',4,0,0,1,'a','A']))
['A', 4, 'C', 'G', 0, 1, 'a']
```

#### 7.2. Description of \_epi\_factorize\_partition



The function <code>\_epi\_factorize\_partition</code> relabels the elements of a list with non-negative integers. It starts with the integer 0 and allocates a new label by increasing the previously allocated label by 1. The first element of the list always receives the label 0 and the highest integer used in the relabeling equals the length of the image (section 7.1) of the list decreased by 1.

```
1 def _epi_factorize_partition(partition):
2 """ the source code of this function can be found in efp.py """
3 return epimorphism
```

Even though a list already encodes an epimorphism, the goal of the function

is to return a canonical *choice* of epimorphism.

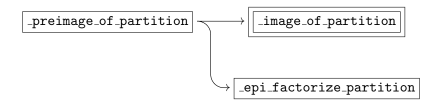
$$S \xrightarrow[e(f)]{f} \operatorname{Im}(f) \xrightarrow{\cong} K$$

This choice ensures that two partitions characterized by the same set of universal properties are *equal* as python lists. In [4, Appendix A], this type of construction is formulated in terms of a factorization system in the category of sets (see the diagram above).

```
>>> p = [3,3,2,1,1,2,4,5,6,5,2,6]
>>> print(_epi_factorize_partition(p))
[0, 0, 1, 2, 2, 1, 3, 4, 5, 4, 1, 5]
>>> im = _image_of_partition(p)
>>> print(_epi_factorize_partition(im))
[0, 1, 2, 3, 4, 5]
>>> print(_epi_factorize_partition(['A',4,'C','C','G',4,0,0,1,'a','A']))
[0, 1, 2, 2, 3, 1, 4, 4, 5, 6, 0]
```

Note that the function does not literally relabel the input list, but allocates a new space in the memory to store the relabeled list.

#### 7.3. Description of \_preimage\_of\_partition



The function \_preimage\_of\_partition takes a list and returns the list of the lists of indices that index the same element.

```
1 def _preimage_of_partition(partition):
2    """ the source code of this function can be found in piop.py """
3    return the_preimage
```

From the point of view of partitions [4, Appendix A], the returned list the\_preimage (see the code above) is the preimage of the underlying epimorphism  $f: S \to K$  of the input partition, where the preimage of f is defined as the K-indexed set of the fibers of the epimorphism.

$$\mathsf{PreIm}(f) = \{f^{-1}(k)\}_{k \in K}$$

Note that, from an implementation viewpoint, the set K might not be equipped with an obvious order relation, which makes it difficult to define the preimage of  $f: S \to K$  as a python list. To rectify this flaw, the preimage is computed with respect to the canonical epimorphism  $e(f): S \to \mathsf{Im}(f)$  whose codomain is equipped with the natural order on integers (see section 7.2).

$$PreIm(f) := \{e(f)^{-1}(k)\}_{k \in Im(f)}$$

```
>>> p = ['a', 'a',2,2,3,3, 'a']
>>> print(_preimage_of_partition(p))
[[0, 1, 6], [2, 3], [4, 5]]
>>> print(_epi_factorize_partition(p))
[0, 0, 1, 1, 2, 2, 0]
```

```
>>> p = [2,1,0,6,5,4,2,1,0]
>>> print(_preimage_of_partition(p))
[[0, 6], [1, 7], [2, 8], [3], [4], [5]]
>>> print(_epi_factorize_partition(p))
[0, 1, 2, 3, 4, 5, 0, 1, 2]
```

In the first example given above:

- the list [0,1,6] is the fiber of the element 'a' and its index in the preimage is 0;
- the list [2,3] is the fiber of the element 2 and its index in the preimage is 1;
- the list [4,5] is the fiber of the element 3 and its index in the preimage is 2.

The preimage will always order its fibers with respect to the order in which the elements of the input list appear.

#### 7.4. Description of print\_partition

```
oxed{ print\_partition } \longrightarrow oxed{ \_preimage\_of\_partition }
```

The function print\_partition is a debug function that takes a list of elements and prints its preimage on the standard output.

```
1 def print_partition(partition):
2  print(_preimage_of_partition(partition))
```

See section 7.3 for examples.

#### 7.5. Description of \_join\_preimages\_of\_partitions

```
\boxed{\_\texttt{join\_preimages\_of\_partitions}} \longrightarrow \boxed{\texttt{\_image\_of\_partition}}
```

The function \_join\_preimages\_of\_partitions takes two lists of lists of indices (the indices can be repeated and should only be non-negative integers) as well as a Boolean value and returns the list of the maximal unions of internal lists that intersect within the concatenation of the two input lists (see the examples below).

```
1 def _join_preimages_of_partitions(preimage1,preimage2,speed_mode):
2    """ the source code of this function can be found in jpop.py """
3    return the_join
```

While the two input lists preimage1 and preimage2 could be two outputs of the procedure

for two input lists of the same length, the Boolean value speed\_mode would indicate whether one of the two input lists may contain at least two different sublists with the same index, as shown below.

Note that the global variable FAST is reserved to this use.

```
>>> print(FAST)
True
```

From the point of view of partitions, the composition of <code>\_preimage\_of\_partition</code> with <code>\_join\_preimages\_of\_partitions</code> would amount to computing the coproduct of two partitions. Since a category of partitions is also a partially ordered set, this coproduct is also the <code>join</code> of the two partitions, which explains the name of the procedure.

For illustration, if we consider the following two lists of lists of indices

```
>>> p1 = [[0, 3], [1, 4], [2]]
>>> p2 = [[0, 1], [2], [3], [4]]
```

we can notice that

- the internal list [0,3] of p1 intersects with the internal lists [0,1] and [3] in p2;
- the internal list [0,1] of p1 intersects with the internal lists [1,4] and [1] in p2;
- the internal list [1,4] of p1 intersects with the internal list [4] in p2;

and

- the internal list [2] of p1 only intersects with the internal list [2] in p2,

so that we have

```
>>> print(_join_preimages_of_partitions(p1,p2,FAST))
[[1, 4, 0, 3], [2]]
```

In terms of implementation, the program

considers each internal list of p1 and searches for the lists of p2 that intersect it. If an intersection is found between two internal lists, it merges the two internal lists in p1 and empties that of p2 (the list is emptied and *not* removed in order to preserve a coherent indexing of the elements of p2). The function continues until all the possible intersections have been checked.

Here is a detail of what program (7.1) does with respect to the earlier example:

The element 0 of [0,3] is searched in the list [0,1] of p2;

The element 0 is found:

The lists [0,3] and [0,1] are merged in p1 and [0,1] is emptied from p2 as follows:

```
p1 = [[0, 3, 1], [1, 4], [2]]
p2 = [[], [2], [3], [4]]
```

Because the element 0 has now been found in p2 and the third input was set to FAST, no other sublist of p2 is supposed to contain the element 0 and the search of the element 0 stops here. Note that if not(FAST) were given in the third argument, then the earlier union operation would also be operated on the remaining sublists of p2.

The element 3 of [0, 3] is searched in the list [] of p2;

The element 3 is not found (continues);

The element 3 of [0, 3] is searched in the list [2] of p2;

The element 3 is not found (continues);

The element 3 of [0, 3] is searched in the list [3] of p2;

The element 3 is found;

The lists [0,3] and [3] are merged in p1 and [3] is emptied from p2 as follows:

```
p1 = [[0, 3, 1], [1, 4], [2]]
p2 = [[], [2], [], [4]]
```

The element 3 has now been found in p2 and does not need to be searched again.

All elements of the initial list [0, 3] have been searched.

The first lists of p1 is appended to p2 in order to ensure the transitive computation of the maximal unions through the next interations.

The list [0, 3, 1] of p1 is emptied as follows:

```
p1 = [[], [1, 4], [2]]
p2 = [[], [2], [], [4], [0, 3, 1]]
```

Repeat the previous procedure with respect to the list [1, 4] of p1. We obtain the following pair:

```
p1 = [[], [], [2]]
p2 = [[], [2], [], [], [1, 4, 0, 3]]
```

Repeat the previous procedure with respect to the remaining list [2] of p1. We obtain the following pair:

```
p1 = [[], [], []]
p2 = [[], [], [], [], [1, 4, 0, 3], [2]]
```

The function stops because there is no more list to process in p1. The output is all the non-empty lists of p2; i.e. [[1, 4, 0, 3], [2]]

Note that, because of the iterative nature of the previous algorithm, the procedure

```
_{\rm join\_preimages\_of\_partitions}(-,-,-)
```

does not necessarily presents its output in the same way as the procedure

```
_preimage_of_partition(-)
```

does. For instance, while the index 0 will always be contained in the first list of the output of \_preimage\_of\_partition, it might not be contained in the first list of the output of \_join\_preimages\_of\_partitions as illustrated below.

```
>>> 1 = _preimage_of_partition([1,2,4,5,1,2,3,2,2,1,3])
>>> print(1)
[[0, 4, 9], [1, 5, 7, 8], [2], [3], [6, 10]]
>>> m = _preimage_of_partition([1,2,5,4,2,2,5,6,5,7,8])
>>> print(m)
[[0], [1, 4, 5], [2, 6, 8], [3], [7], [9], [10]]
>>> print(_join_preimages_of_partitions(1,m))
[[3], [6, 10, 2, 1, 5, 7, 8, 0, 4, 9]]
```

Interestingly, the procedure \_join\_preimages\_of\_partitions can also be used to compute the intersection-free closure of a set of sets, as shown below.

```
>>> a = [[1,0,2,1,3,2,5,4,6],[15,15,18,0,13],[7,11,12,22],[23,12]]
>>> closure_of_a = _join_preimages_of_partitions(a,a,not(FAST))
>>> print(closure_of_a)
[[15, 18, 0, 13, 1, 2, 3, 5, 4, 6], [23, 12, 7, 11, 22]]
```

#### 7.6. Description of EquivalenceRelation (class)

```
 \boxed{ \texttt{EquivalenceRelation} } \longrightarrow \boxed{ \texttt{\_join\_preimages\_of\_partitions} }
```

The class EquivalenceRelation possesses two objects, namely

- .classes (list of lists);
- .range (integer);

and three methods, namely

- .\_\_init\_\_ (constructor);
- .closure;
- .quotient.

The constructor .\_\_init\_\_ takes between 1 and 2 arguments: the first argument should either be an empty list or a list of lists of indices (i.e. non-negative integers) and the second argument, which is optional when the first argument is not an empty list, should be an integer that is greater than or equal to the maximum index contained in the first input, if it exists.

```
1 class EquivalenceRelation:
 2
     #The objects of the class are:
 3
     #.classes (list of lists);
 4
     #.range (integer);
 5
     #The following constructor takes between 1 and 2 arguments,
     #the first one being a list and the second being an integer.
 6
 7
     def __init__(self,*args):
     """ the source code of this constructor can be found in cl_er.py """
 8
     def closure(self):
10
       self.classes = _join_preimages_of_partitions(self.classes,
   self.classes,not(FAST))
11
     def quotient(self):
12
     """ the source code of this function can be found in cl_er.py """
13
       return the_quotient
```

If the first input is not empty, then it is stored in the object .classes while the object .range receives:

- either the second input, when this second input is given;
- or the maximum index contained in the first input when no second input is given.

```
>>> eq1 = EquivalenceRelation([[0,1,2,9],[7,3,8,6],[4,9,5]])
>>> print(eq1.classes)
[[0, 1, 2, 9], [7, 3, 8, 6], [4, 9, 5]]
>>> print(eq1.range)
9
>>> eq2 = EquivalenceRelation([[0,1,2,9],[7,3,8,7],[9,15]],18)
>>> print(eq2.classes)
[[0, 1, 2, 9], [7, 3, 8, 7], [9, 15]]
>>> print(eq2.range)
18
```

If the first input is empty, then the second argument is required. In this case, the object .classes receive the lists containing all the singleton lists containing the integers from 0 to the integer given in the second argument, which is stored in the object .range.

```
>>> eq3 = EquivalenceRelation([],5)
>>> print(eq3.classes)
[[0], [1], [2], [3], [4], [5]]
```

The method .closure() replaces the content of the object .classes with the transitive closure of its classes. After this procedure, the object .classes describes an actual equivalence relation (modulo the singleton equivalence classes, which do not need to be specified for obvious reasons).

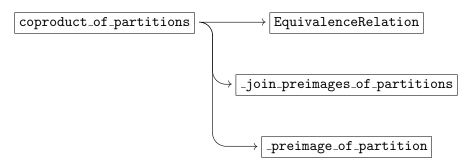
```
>>> eq1.closure()
>>> print(eq1.classes)
[[7, 3, 8, 6], [4, 9, 5, 0, 1, 2]
```

```
>>> eq2.closure()
>>> print(eq2.classes)
[[7, 3, 8], [9, 15, 0, 1, 2]]
```

The method .quotient() returns a list of integers whose length is equal to the integer contained in the object .range decreased by 1 and whose non-trivial fibers are those contained in the object .classes after a call of the method .closure().

```
>>> print(eq1.quotient())
[1, 1, 1, 0, 1, 1, 0, 0, 0, 1]
>>> print(eq2.quotient())
[1, 1, 1, 0, 2, 3, 4, 0, 0, 1, 5, 6, 7, 8, 9, 1, 10, 11, 12]
```

#### 7.7. Description of coproduct\_of\_partitions



The function coproduct\_of\_partitions takes two lists of the same legnth and returns their coproduct (or join) as partitions. Specifically, the procedure outputs the quotient of the join of their preimages (see the piece of code given below). If the two input lists do not have the same length, then an error message is outputted and the program is aborted.

```
1 def coproduct_of_partitions(partition1,partition2):
     if len(partition1) == len(partition2):
 2
 3
       #Returns the coproduct of two partitions as the quotient of the
       #equivalence relation induced by the join of the preimages
 4
 5
       #of the two partitions.
       the_join = EquivalenceRelation(_join_preimages_of_partitions(
   _preimage_of_partition(partition1),_preimage_of_partition(partition2),
  FAST))
 7
       return the_join.quotient()
8
9
       print("Error: in coproduct_of_partitions: lengths do not match.")
10
       exit()
```

Note that the outputs of the procedure coproduct\_of\_partitions do not necessarily belong to the set of outputs of the procedure \_epi\_factorize\_partition. The reason comes from the way in which the procedure \_join\_preimages\_of\_partitions is implemented (see the end of section 7.5).

```
>>> 1 = [1,2,4,5,1,2,3,2,2,1,3]

>>> m = [1,2,5,4,2,2,5,6,5,7,8]

>>> c = coproduct_of_partitions(1,m)

>>> print(c)

[1, 1, 1, 0, 1, 1, 1, 1, 1, 1]

>>> print(_epi_factorize_partition(c))

[0, 0, 0, 1, 0, 0, 0, 0, 0, 0]
```

#### 7.8. Description of product\_of\_partitions

```
	ext{product\_of\_partitions} \longrightarrow 	ext{$-$epi\_factorize\_partition}
```

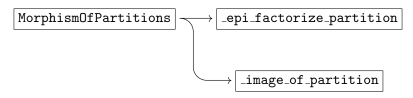
The function product\_of\_partitions takes two lists and returns a list that is the relabeling of the zipping of the two lists (i.e. the list of pairs of elements with corresponding indices in each of the input lists) via the procedure \_epi\_factorize\_partition.

```
1 def product_of_partitions(partition1,partition2):
     #The following line checks if the product of the two lists is possible.
     if len(partition1) == len(partition2):
3
4
       #Constructs the list of pairs of element with the
5
       #same index in the two lists, and then relabels
6
       #the pairs using _epi_factorize_partition.
7
       return _epi_factorize_partition(zip(partition1,partition2))
8
9
       print("Error: in product_of_partitions: lengths do not match.")
10
       exit()
```

The function outputs an error if the two input lists do not have the same length.

```
>>> product_of_partitions([1,1,1,1,2,3],['a','b','c','c','c','c','c'])
[0, 1, 2, 2, 3, 4]
>>> product_of_partitions([1,1,1,1,2],['a','b','c','c','c','c','c'])
Error: in product_of_partitions: lengths do not match.
```

#### 7.9. Description of MorphismOfPartitions (class)



The class MorphismOfPartitions possesses three objects, namely

```
- .arrow (list)- .source (list)- .target (list)
```

and a constructor .\_\_init\_\_ The consructor .\_\_init\_\_ takes two lists as well as an optional argument and stores, in the object .arrow, the list that describes, if it exists, the (unique) morphism of partitions from the first input list (seen as a partition) to the second input list (seen as a partition). If the morphism does not exist, then the method returns an error message unless the value False was given as a third input.

The canonical epimorphisms associated with the partitions of the first and second input lists (see section 7.2) are stored in the objects .source and .target, respectively.

```
1 class MorphismOfPartitions:
2  #The objects of the class are:
3  #.arrow (list);
4  #.source (list);
5  #.target (list).
6  def __init__(self,source,target,*args):
7  """ the source code of this constructor can be found in cl_mop.py """
```

The list that is contained in the object .arrow is the image of the function  $\mathtt{source} \mapsto \mathtt{target}(\mathtt{source})$  that can be constructed from the parametrization  $t \mapsto \mathtt{source}[t]$  and  $t \mapsto \mathtt{target}[t]$ , which are given by the list structure of the two input lists  $\mathtt{target}$  and  $\mathtt{source}$ . This is illustrated below in more detail.

For illustration, let us consider the following pair of lists.

```
>>> p1 = [0,1,2,3,3,4,5]
>>> p2 = [0,1,2,3,3,3,1]
```

To construct the function  $source \mapsto target(source)$ , we can first try to construct the graph (source[t], target[t]), for which we use the procedure zip.

```
>>> p3 = zip(p1,p2)
>>> print(p3)
[(0, 0), (1, 1), (2, 2), (3, 3), (3, 3), (4, 3), (5, 1)]
```

The image of the zipping is then as follows:

```
>>> p4 = _image_of_partition(p3)
>>> print(p4)
[(0, 0), (1, 1), (2, 2), (3, 3), (4, 3), (5, 1)]
```

We can see that for each pair (x,y) in p4, every component x is mapped to a unique image y so that p4 defines the *graph* of the morphism of partitions between p1 and p2. The constructor .\_\_init\_\_ then records, in the object .arrow, the second projections of the pairs contained in p4, which also corresponds to the image of the underlying graph encoded by p4.

```
>>> m = MorphismOfPartitions(p1,p2)
>>> print(m.arrow)
[0, 1, 2, 3, 3, 1]
```

If there exists no morphism from the first input list to the second input list, then the function outputs an error message.

For example, if we modify p2 as follows

```
\Rightarrow p2 = [0,1,2,3,6,3,1]
```

then the image of the zipping of p1 and p2 is as follows:

```
>>> p4 = _image_of_partition(zip(p1,p2))
>>> print(p4)
[(0, 0), (1, 1), (2, 2), (3, 3), (3, 6), (4, 3), (5, 1)]
```

As can be seen, the argument 3 is 'mapped' to two different images, namely 3 and 6. In this case, the constructor .\_\_init\_\_ exits the program with an error message.

Here is a complete example summarizing the previous explanation.

```
>>> m = MorphismOfPartitions([1,6,5,3,3,4,2],[1,2,5,4,4,4,2])
>>> print(m.source)
[0, 1, 2, 3, 3, 4, 5]
>>> print(m.target)
[0, 1, 2, 3, 3, 3, 1]
>>> print(m.arrow)
[0, 1, 2, 3, 3, 1]
>>> m = MorphismOfPartitions([1,6,5,3,3,4,2],[1,2,5,4,6,4,2])
Error: in MorphismOfPartitions.__init__: source and target are not compatible.
>>> m = MorphismOfPartitions([1,6,5,3,3,4,2],[1,2,5,4,6,4,2],False)
```

Finally, note that the constructor of MorphismOfPartitions can be used to test whether there is a morphism between two partitions by using the key words try and except and setting the thrid argument to False, as illustrated below.

```
>>> try:
>>>
      m = MorphismOfPartitions([1,6,5,3,3,4,2],[1,2,5,4,4,4,2],False)
      print("True")
>>>
>>> except:
      print("False")
>>>
True
>>> try:
>>>
      m = MorphismOfPartitions([1,6,5,3,3,4,2],[1,2,5,4,6,4,2],False)
      print("True")
>>>
>>> except:
      print("False")
>>>
False
```

# Presentation of the module AsciiTree.py

#### 8.1. Description of tree\_of\_partitions

$$\boxed{\texttt{tree\_of\_partitions}} \longrightarrow \boxed{\texttt{MorphismOfPartitions}}^{\nearrow}$$

The function tree\_of\_partitions takes a list of lists whose pairs of successive lists can be related via MorphismOfPartitions items (see section 7.9)

pair of succ. lists 
$$\underbrace{[l_1, l_2, l_3, \dots, l_n]}_{\text{pair of succ. lists}}$$

and returns the actual lists of MorphismOfPartitions items between these.

```
1 def tree_of_partitions(partitions):
2    """ the source code of this function can be found in top.py """
3    return the_tree
```

The input list should always start with the target of the first arrow, then present its source, which should also be the target of the next arrow, etc.

```
[0, 0, 1, 2, 2, 3]
| arrow num 3: [0, 1, 1, 2]
| V

[0, 0, 1, 1, 1, 2]
| arrow num 2: [0, 0, 1]
| V

[0, 0, 0, 0, 0, 0, 1]
| arrow num 1: [0, 0]
| V

[0, 0, 0, 0, 0, 0, 0]
```

#### 8.2. Description of convert\_tree\_to\_atpf

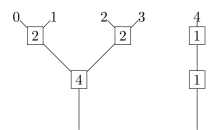


The function convert\_tree\_to\_atpf takes a list of MorphismOfPartitions item (as returned by the procedure tree\_of\_partitions) and converts it into its associated ascii tree pre-format (abbrev. atpf), which is a regular expression of type ATPF given by the following double grammar rules.

```
 \begin{array}{lll} & & \text{The grammar terms} & \text{and their associated weights} \\ & & \text{ATPF} := [\mathsf{Tree}_1, \mathsf{Tree}_2, \dots, \mathsf{Tree}_k]; \\ & & \text{Tree} := \big( \mathsf{weight}(\mathsf{Tree}), [\mathsf{Tree}_1, \mathsf{Tree}_2, \dots, \mathsf{Tree}_k] \big), \\ & & \text{Tree} := \big( \mathsf{weight}(\mathsf{Tree}), [\mathsf{Leaf}_1, \mathsf{Leaf}_2, \dots, \mathsf{Leaf}_k] \big), \\ & \text{Leaf} := \big( \mathsf{weight}(\mathsf{Leaf}), l \big), \text{ where } l \text{ is a list} \\ & & \text{weight}(\mathsf{Leaf}) := \mathsf{len}(l); \\ \end{array}
```

The idea of such a construction is to give access to the number of leaves contained in each fork of a tree. This type of information will later be used to display trees with ascii characters on the console. For illustration, the following line gives the example of an atpf that describes the forest displayed below it.

```
atpf = [(4, [(2, [0, 1]), (2, [2, 3])]), (1, [(1, [4])])]
```



Note that the number of levels in the trees can be linked to what could be called the *depth* of the bracketing structure of the atpf. Specifically, we define the *depth* of an afpt according

to the following recursive equations, relative to the definition of the terms given above.

```
\begin{split} \operatorname{depth}(\operatorname{ATPF}) &:= \max\{\operatorname{depth}(\operatorname{Tree}_i) \mid i=1,\ldots,k\}; \\ \operatorname{depth}(\operatorname{Tree}) &:= \max\{\operatorname{depth}(\operatorname{Tree}_i) \mid i=1,\ldots,k\}+1; \\ \operatorname{depth}(\operatorname{Tree}) &:= \max\{\operatorname{depth}(\operatorname{Leaf}_i) \mid i=1,\ldots,k\}+1; \\ \operatorname{depth}(\operatorname{Leaf}) &:= 1; \end{split}
```

The procedure convert\_tree\_to\_atpf then returns a list and an integer, where the list is the atpf of the input (i.e. of the tree) and the integer is equal to the depth of the atpf, which is equal to len(tree)+1.

```
1 def _convert_tree_to_atpf(tree):
2    """ the source code of this function can be found in ctta.py """
3    return (the_atpf,len(tree)+1)
```

Since the depth is only returned for parsing purposes (see section 8.3 and section 8.4), the main task of the procedure convert\_tree\_to\_atpf is to compute the atpf associated with the input list of composable MorphismOfPartitions items by considering the successive preimages of the objects .arrow of each MorphismOfPartitions item contained in the list.

For illustration, consider the following list of partitions:

```
>>> a = [0,1,0,0,0,0]
>>> b = [0,2,0,0,0,1]
>>> c = [0,4,2,3,3,5]
```

This list can be associated with an obvious sequence of MorphismOfPartitions item that is constructed by the procedure tree\_of\_partitions (see section 8.1). Below, we make the steps of this construction explicit for the list [a, b,c].

First, one wants to consider the preimage of the last partition c. This preimage is outputted by the function \_preimage\_of\_partition (see section 7.3) as shown below.

```
>>> _preimage_of_partition(c)
[[0], [1], [2], [3, 4], [5]]
```

The internal lists appearing in the previous list are important for our next step. More specifically, we want to follow the recursive definition of the grammar of atpfs (given above) and take the lists [0], [1, 2], [3, 4], and [5] to be the initial values of the recursion so that the first level of the atpf is as shown below, where the red numbers are the weight of the Leaf terms (see grammar rules above).

```
the_atpf = [(1, [0]), (1, [1]]), (1, [2]]), (2, [3, 4]), (1, [5])]
```

For the next level, we need to compute the preimage of the object .arrow encoding the morphism  $c \to b$ . Recall that the list contained in this object encodes the mapping rules associated with the morphism  $c \to b$ . Below, we display this mapping up to relabeling of the lists c and b as [0,1,2,3,3,4] and [0,1,0,0,0,2], respectively.

```
>>> f = MorphismsOfPartitions(c,b)
>>> for i in range(len(f.arrow)):
... print("f: "+str(i)+" |--> "+str(f.arrow[i]))
f: 0 |-> 0
f: 1 |-> 1
f: 2 |-> 0
f: 3 |-> 0
f: 4 |-> 2
```

Now, since b has three elements in its image, the MorphismOfPartitions item f possesses three fibers, which are given by the following list of lists.

```
>>> fiber = _preimage_of_partition(f.arrow)
>>> print(fiber)
[[0, 2, 3], [1], [4]]
```

Following the atpf grammar and, more specifically, the syntax for the Tree terms, we now want to use the list fiber computed above to create the next level of the atpf. The idea is to replace the intergers living in fiber with the elements of the list the\_atpf. Precisely, we want to replace:

```
fiber[0][0] = 0 with the_atpf[0]
fiber[0][1] = 2 with the_atpf[2]
fiber[0][2] = 3 with the_atpf[3]
fiber[1][0] = 1 with the_atpf[1]
fiber[2][0] = 4 with the_atpf[4]
```

Doing so, the list fiber is turned into the following list.

```
fiber = [[(1, [0]), (1, [2]]), (2, [3, 4])], [(1, [1]])], [(1, [5])]]
```

To complete the construction of the next level of the atpf, there remains to compute the weight for each internal list. Precisely, we can see that

```
the weight of [(1, [0]), (1, [2]]), (2, [3, 4])] is 1+1+2 = 4;
the weight of [(1, [1]])] is 1;
the weight of [(1, [5])]] is 1.
```

We then equip each list with its weight by using tuples, as shown below.

```
the_atpf = [(4,[(1, [0]), (1, [2]]), (2, [3, 4])]), (1,[(1, [1]]))], (1,[(1, [5])])]
```

We then repeat the previous procedure with, this time, the fiber of the morphism  $b \rightarrow a$  and the earlier list the\_atpf so that the final atpf is of the following form.

```
the_atpf = [(5, [(4, [(1, [0]), (1, [2]), (2, [3, 4])]), (1, [(1, [5])])]), (1, [(1, [1], [1])])]
```

The user that verify that the previous list conresponds to the output of the following command.

```
>>> print(convert_tree_to_atpf(tree_of_partitions([a,b,c]))[0])
```

#### **8.3.** Description of convert\_atpf\_to\_atf

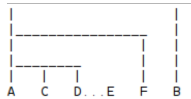
The function <code>convert\_atpf\_to\_atf</code> takes an atpf and its depth (see section 8.2) and returns the associated ascii tree format (abbrev. atf), which is a modified version of an atpf in which one substracts all the weights by the rightmost weight of the next level, as shown by the following grammar rules

```
\begin{split} & \texttt{ATPF} := [\texttt{Tree}_1, \texttt{Tree}_2, \dots, \texttt{Tree}_k]; \\ & \texttt{Tree} := \big( (\texttt{weight}(\texttt{Tree}), \texttt{weight}(\texttt{Tree}) - \texttt{weight}(\texttt{Tree}_k)), [\texttt{Tree}_1, \texttt{Tree}_2, \dots, \texttt{Tree}_k] \big); \\ & \texttt{Tree} := \big( (\texttt{weight}(\texttt{Tree}), \texttt{weight}(\texttt{Tree}) - \texttt{weight}(\texttt{Tree}_k)), [\texttt{Leaf}_1, \texttt{Leaf}_2, \dots, \texttt{Leaf}_k] \big); \\ & \texttt{Leaf} := \big( (\texttt{weight}(\texttt{Leaf}), 0), l \big), \text{ where } l \text{ is a list}; \\ \end{split}
```

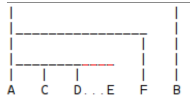
Note that this function uses the depth of the atpf in order to differentiate between the leaves and the intermediate levels of the tree, which require two different types of treatement.

```
1 def convert_atpf_to_atf(atpf,depth):
2    """ the source code of this function can be found in cata.py """
3    return the_atf
```

The reason for this is that the procedure print\_atf (see section 8.4) is to display ascii trees whose trunks are on the left of the screen, as show below.



Substracting the rightmost weights of the atpf from the weight placed below it, in the tree, allows print\_atf to know when it needs to stop printing the horizontal level of the tree. Intuitively, the following pictures shows what atpf would look like without convertion into an atf, where the red underscore symbols sticking out toward the right symbolize the amount of weight subtracted in the atf.



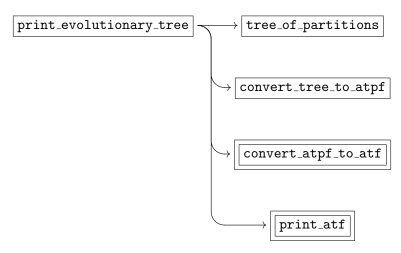
#### 8.4. Description of print\_atf

The function print\_atf takes an atf and its depth and prints the ascii tree associated with the atf on the standard output.

```
1 def print_atf(atf,depth):
2  """ the source code of this function can be found in patf.py """
```

The following example shows how print\_atf can be combined with the procedures convert\_tree\_to\_atpf and convert\_atpf\_to\_atf.

#### 8.5. Description of print\_evolutionary\_tree



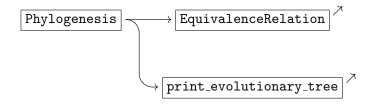
The function print\_evolutionary\_tree takes a list of lists whose pairs of successive lists can be related via MorphismOfPartitions items and returns the tree encoded by this sequence of morphisms.

```
1 def print_evolutionary_tree(atf,depth):
2  #Returns a sequence of morphisms of partitions.
3  tree = tree_of_partitions(partitions)
4  #Returns an ascii tree pre-format and its depth.
5  atpf = convert_tree_to_atpf(tree)
6  #Returns the ascii tree format of the atpf.
7  atf = convert_atpf_to_atf(*atpf)
8  #Prints the atf on the standard output.
9  print_atf(atf,atpf[1])
```

See the example given in section 8.5 to see what this procedure does.

# Presentation of the module Phylogeny.py

#### 9.1. Description of Phylogenesis (class)



The class Phylogenesis possesses two objects, namely

- .taxon (non-negative integer);
- .history (list of lists of indices);

and three methods, namely

- .\_\_init\_\_ (constructor)
- .partitions
- .print\_tree

A Phylogenesis item is meant to be part of another structure called a Phylogeny (see section 9.2). The first object .taxon of the class Phylogenesis stores an integer that allows us to identify the Phylogenesis item with respect to other Phylogenesis items in the Phylogeny structure. On the other hand, the object .history stores a list of lists encoding a historical record of the coalescence events between the Phylogenesis item and the other taxa contained by the Phylogeny structure. The list of lists contained in the object .history should:

- start with a singleton list containing the integer representing the taxon itself;
- be such that every list should contain its predecessor list;

```
1 class Phylogenesis:
 2
     #The objects of the class are:
 3
     #.taxon (non-negative integer);
     #.history (list of lists of indices);
     def __init__(self,history):
 5
     """ the source code of this constructor can be found in cl_pgs.py """
 6
 7
     def partitions(self):
     """ the source code of this function can be found in cl_pgs.py """
 8
 9
       return partitions
10
     def print_tree(self):
     #Returns the evolutionary tree described by the list of lists outputted
11
12
     #by the procedure partitions().
13
       return print_evolutionary_tree(self.partitions())
```

The constructor .\_\_init\_\_ takes a non-empty list of lists whose first list is a singleton and allocates

- the index contained in the first internal list of the input list to the object .taxon,
- the input list to the object .history.

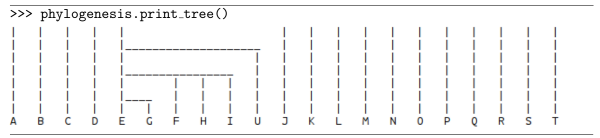
Before terminating, the procedure checks whether the list of lists is made of non-negative integers and whether each list preceding another is contained in the successor list. Below, we give the example of an initialization of a Phylogenesis item.

```
>>> history = [[4],[4,6],[5,4,7,8,6],[6,5,4,7,8,20]]
>>> phylogenesis = Phylogenesis(history)
>>> print(phylogenesis.taxon)
4
>>> print(phylogenesis.history[len(phylogenesis.history)-1])
[6, 5, 4, 7, 8, 20]
```

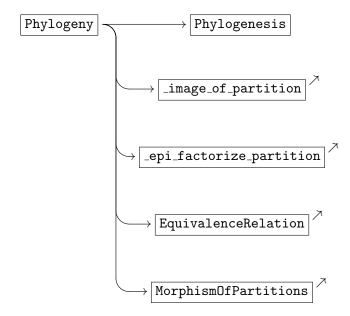
The method .partitions() returns the sequence of partitions induced by the list of lists contained in the object .history over the set of indices ranging from 0 to the maximum index of the last list of the object .history.

```
>>> p = phylogenesis.partitions()
>>> for i in range(len(p)):
>>> print(p[i])
[1, 2, 3, 4, 0, 0, 0, 0, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 0]
[1, 2, 3, 4, 0, 0, 0, 0, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16]
[1, 2, 3, 4, 0, 5, 0, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]
[1, 2, 3, 4, 0, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20]
```

The method .print\_tree() returns the evolutionary tree associated with the sequence of partitions returned by .partition() (for more intuition, see section 8).



#### 9.2. Description of Phylogeny (class)



The class Phylogeny possesses one object, namely

- .phylogeneses (list of Phylogenesis items)

and ten methods, namely

- .\_\_init\_\_ (constructor)
- .coalescent
- .extend
- .count\_uniformity
- .boolean\_partition
- .make\_friends
- .set\_up\_friendship
- .score
- .choose
- .set\_up\_competition

The object .phylogenesis is supposed to contained a list of Phylogenesis items. The taxon associated with the i-th phylogenesis should be indexed by the interger i itself and any label appearing in the Phylogenesis items of the list should have its own Phylogenesis item included in the list.

```
1 class Phylogeny:
2  #The objects of the class are:
3  #.phylogeneses (lists of Phylogenesis items);
4  def __init__(self,phylogeneses):
5  """ the source code of this constructor can be found in cl_pgs.py """
```

**9.2.1.** First generation. Below, we will often use the term *first generation* of the phylogenesis of a certain taxon t to refer to the last list contained in the list

```
self.phylogeneses[t].history
```

relative to the indexing order associated with the list structure.

**9.2.2.** Constructor. The constructor .\_\_init\_\_ takes a list of lists of lists containing nonnegative integers and use every internal list of the input to create a Phylogenesis item, which is stored in the object .phylogeneses.

The following lines show how a phylogeny can be created via the consutructor of the class.

```
>>> p = list()
>>> for i in range(8):
     p.append([[i], [i, (i+10) % 8], [i, (i+10) % 8, (i+5*i+13) % 8]]))
>>> pgy = Phylogeny(p)
>>> for i in range(len(pgy.phylogeneses)):
      print("taxon: "+str(pgy.phylogeneses[i].taxon))
>>>
      print(pgy.phylogeneses[i].history)
taxon:
[[0], [0, 2], [0, 2, 5]]
taxon:
[[1], [1, 3], [1, 3, 3]]
taxon:
        2
[[2], [2, 4], [2, 4, 1]]
taxon:
[[3], [3, 5], [3, 5, 7]]
taxon: 4
[[4], [4, 6], [4, 6, 5]]
taxon:
[[5], [5, 7], [5, 7, 3]]
taxon:
[[6], [6, 0], [6, 0, 1]]
taxon:
[[7], [7, 1], [7, 1, 7]]
```

**9.2.3. Elementary methods.** Phylogeny items can be updated, modified or analyzed through the methods .coalescent; .extend; and .make\_friends, which we present in the following sections.

```
6
     def coalescent(self):
 7
     """ the source code of this constructor can be found in cl_pgs.py """
8
       return coalescent
     def extend(self, extension):
9
10
     """ the source code of this constructor can be found in cl_pgs.py """
         return False
11
     def make_friends(self,taxon):
12
     """ the source code of this constructor can be found in cl_pgs.py """
13
14
       return (friends,coalescence_hypothesis)
```

**9.2.4.** Coalescent. The method .coalescent() returns the list of the last lists of the objects .history of each Phylogenesis item contained in the object .phylogeneses (*i.e.* what one would like to understand as the *first generations* of the current phylogeny). The k-th list of the outputted list is therefore the first generation (or last list) of the history of taxon k (see example below).

```
>>> coalescent = pgy.coalescent()
>>> for i in range(len(coalescent)):
      print("1st generation of " + str(i) + "'s history:
    str(coalescent[i]))
1st generation of 0's history:
                                 [0, 2, 5]
1st generation of 1's history:
                                 [1, 3, 3]
1st generation of 2's history:
                                [2, 4, 1]
1st generation of 3's history:
                                 [3, 5, 7]
                                 [4, 6, 5]
1st generation of 4's history:
                                 [5, 7, 3]
1st generation of 5's history:
1st generation of 6's history:
                                 [6, 0, 1]
1st generation of 7's history:
                                 [7, 1, 7]
```

- **9.2.5.** Extend. The method .extend takes a list of pairs of the form (t,1) where t is the label of a taxon of the Phylogeny item (accessible through pgy.phylogeneses[-].taxon) and 1 is a list of taxa and it updates the object .phylogeneses as follows:
  - for all pairs (t,1) contained in the input passed to .extend:
    - 1) ▷ if every list 1 contains the last list of self.phylogeneses[t].history, ▷ if at least one of the lists 1 strictly contains the last list of

```
self.phylogeneses[t].history,
```

- → then every list 1 is appended to the list self.phylogeneses[t].history and the value True is returned;
- 2) if there is no strict inclusion of the last list of self.phylogeneses[t].history into 1, then the object .phylogeneses is not modified and the value False is returned;
- 3) otherwise, an error message is returned and the procedure exit the program;
- if an update has happened, then for all other taxa t of the phylogeny that do not appear in the input of .extend, the last list of self.phylogeneses[t].history (i.e. the first generation of the history of the phylogenesis of t) is again repeated (i.e. appended again) in the list self.phylogeneses[t].history.

The following code lines illustrate these various cases for the example used in the previous sections (starting from section 9.2.2).

```
>>> extension = [(5,[3,7,7,5]),(7,[7,1])]
>>> flag = pgy.extend(extension)
>>> print(flag)
False
>>> for i in range(len(pgy.phylogeneses)):
     print(pgy.phylogeneses[i].history)
[[0], [0, 2], [0, 2, 5]]
[[1], [1, 3], [1, 3, 3]]
[[2], [2, 4], [2, 4, 1]]
[[3], [3, 5], [3, 5, 7]]
[[4], [4, 6], [4, 6, 5]]
[[5], [5, 7], [5, 7, 3]]
[[6], [6, 0], [6, 0, 1]]
[[7], [7, 1], [7, 1, 7]]
>>> extension = [(5,[3,7,7,5]),(7,[7,1]),(1,[1,3,4,5])]
>>> flag = pgy.extend(extension)
>>> print(flag)
```

```
True
>>> for i in range(len(pgy.phylogeneses)):
      print(pgy.phylogeneses[i].history)
[[0], [0, 2], [0, 2, 5], [0, 2, 5]]
[[1], [1, 3], [1, 3, 3], [1, 3, 4, 5]]
[[2], [2, 4], [2, 4, 1], [2, 4, 1]]
[[3], [3, 5], [3, 5, 7], [3, 5, 7]]
[[4], [4, 6], [4, 6, 5], [4, 6, 5]]
[[5], [5, 7], [5, 7, 3], [3, 7, 5]]
[[6], [6, 0], [6, 0, 1], [6, 0, 1]]
[[7], [7, 1], [7, 1, 7], [7, 1]]
>>> extension = [(-1,["error"])]
>>> pgy.extend(extension)
>>> for i in range(len(pgy.phylogeneses)):
      print(pgy.phylogeneses[i].history)
[[0], [0, 2], [0, 2, 5], [0, 2, 5]]
[[1], [1, 3], [1, 3, 3], [1, 3, 4, 5]]
[[2], [2, 4], [2, 4, 1], [2, 4, 1]]
[[3], [3, 5], [3, 5, 7], [3, 5, 7]]
[[4], [4, 6], [4, 6, 5], [4, 6, 5]]
[[5], [5, 7], [5, 7, 3], [3, 7, 5]]
[[6], [6, 0], [6, 0, 1], [6, 0, 1]]
[[7], [7, 1], [7, 1, 7], [7, 1]]
>>> extension = [(5,[1,4,80]),(7,["error"])]
>>> pgy.extend(extension)
>>> for i in range(len(pgy.phylogeneses)):
      print(pgy.phylogeneses[i].history)
        in Phylogeny.extend: the extension is not compatible with the
phylogenesis of taxon 5
```

- **9.2.6.** Make friends. The method .make\_friends takes the label of a taxon (i.e. a nonnegative integer) and returns a pair of lists (friends, hypothesis) where
  - the list friends contains all those taxa that have not coalesced with the input taxon, which means that there are not in the first generation of the phylogenesis of the taxon;
  - the list hypothesis contains the (sorted) lists obtained by making the union of the first generation of the input taxon with the first generation of one of the taxon in friends.

For illustration, consider the phylogeny constructed in section 9.2.5. Below, we first recall the structure of that phylogeny.

```
>>> for i in range(len(pgy.phylogeneses)):
>>> print(pgy.phylogeneses[i].history)

[[0], [0, 2], [0, 2, 5], [0, 2, 5]]

[[1], [1, 3], [1, 3, 3], [1, 3, 4, 5]]

[[2], [2, 4], [2, 4, 1], [2, 4, 1]]

[[3], [3, 5], [3, 5, 7], [3, 5, 7]]

[[4], [4, 6], [4, 6, 5], [4, 6, 5]]
```

```
[[5], [5, 7], [5, 7, 3], [3, 7, 5]]
[[6], [6, 0], [6, 0, 1], [6, 0, 1]]
[[7], [7, 1], [7, 1, 7], [7, 1]]
   The output of the method .make_friends for taxa 1, 2 and 7 is as follows.
>>> friends_made = pgy.make_friends(1)
>>> friends = friends_made[0]
>>> coalescence_hypothesis = friends_made[1]
>>> for i in range(len(friends)):
      print("taxa 1 and "+ str(friends[i])+" have ancestor
    "+str(coalescence_hypothesis[i]))
taxa 1 and 0 have ancestor [0, 1, 2, 3, 4, 5]
taxa 1 and 2 have ancestor [1, 2, 3, 4, 5]
taxa 1 and 6 have ancestor [0, 1, 3, 4, 5, 6]
taxa 1 and 7 have ancestor [1, 3, 4, 5, 7]
>>> friends_made = pgy.make_friends(2)
>>> friends = friends_made[0]
>>> coalescence_hypothesis = friends_made[1]
>>> for i in range(len(friends)):
      print("taxa 2 and "+ str(friends[i])+" have ancestor
    "+str(coalescence_hypothesis[i]))
taxa 2 and 0 have ancestor [0, 1, 2, 4, 5]
taxa 2 and 3 have ancestor [1, 2, 3, 4, 5, 7]
taxa 2 and 5 have ancestor [1, 2, 3, 4, 5, 7]
taxa 2 and 6 have ancestor [0, 1, 2, 4, 6]
taxa 2 and 7 have ancestor [1, 2, 4, 7]
>>> friends_made = pgy.make_friends(7)
>>> friends = friends_made[0]
>>> coalescence_hypothesis = friends_made[1]
>>> for i in range(len(friends)):
      print("taxa 7 and "+ str(friends[i])+" have ancestor
    "+str(coalescence_hypothesis[i]))
taxa 7 and 0 have ancestor [0, 1, 2, 5, 7]
taxa 7 and 2 have ancestor [1, 2, 4, 7]
taxa 7 and 3 have ancestor [1, 3, 5, 7]
taxa 7 and 4 have ancestor [1, 4, 5, 6, 7]
taxa 7 and 5 have ancestor [1, 3, 5, 7]
taxa 7 and 6 have ancestor [0, 1, 6, 7]
```

**9.2.7.** Algorithm for constructing phylogenies. The next set of methods are procedures meant to be used to implement the algorithm described in [4], which constructs a phylogeny according of the definition given thereof.

```
15
     def set_up_friendships(self):
16
     """ the source code of this constructor can be found in cl_pgs.py """
17
       return (friendships, coalescence_hypotheses)
     def score(self,partitions,friendship_network):
15
16
     """ the source code of this constructor can be found in cl_pgs.py """
17
       return score_cardinality_adjusted
     def choose(self,scores):
18
19
     """ the source code of this constructor can be found in cl_pgs.py """
20
       return result
```

```
21  def set_up_competition(self,best_fit):
22  """ the source code of this constructor can be found in cl_pgs.py """
23  return coalescence_hypothesis
```

9.2.8. Set up friendships. The method .set\_up\_friendships() returns a pair of lists (friendships,hypotheses) containing the lists of the two different outputs of the method .make\_friends for every taxon of the phylogeny. More specifically,

- friendships is the list of lists whose t-th list contains the first output of the procedure self.make\_friends for taxon t;
- hypotheses is the list of lists whose t-th list contains the second output of the procedure self.make\_friends for taxon t;

The following code lines give a description of the output of .set\_up\_friendships() for the phylogeny used in section 9.2.6.

```
>>> friendships_made = pgy.set_up_friendships()
>>> friendships = friendships_made[0]
>>> coalescence_hypotheses = friendships_made[1]
>>> for t in range(len(friendships)):
      for r in range(len(friendships[t])):
        print("taxa " + str(t) + " and " + str(friendships[t][r])+" have
>>>
    ancestor "+str(coalescence_hypotheses[t][r]))
taxa 0 and 1 have ancestor [0, 1, 2, 3, 4, 5]
taxa 0 and 3 have ancestor [0, 2, 3, 5, 7]
taxa 0 and 4 have ancestor [0, 2, 4, 5, 6]
taxa 0 and 6 have ancestor [0, 1, 2, 5, 6]
taxa 0 and 7 have ancestor [0, 1, 2, 5, 7]
taxa 1 and 0 have ancestor [0, 1, 2, 3, 4, 5]
taxa 1 and 2 have ancestor [1, 2, 3, 4, 5]
taxa 1 and 6 have ancestor [0, 1, 3, 4, 5, 6]
taxa 1 and 7 have ancestor [1, 3, 4, 5, 7]
taxa 2 and 0 have ancestor [0, 1, 2, 4, 5]
taxa 2 and 3 have ancestor [1, 2, 3, 4, 5, 7]
taxa 2 and 5 have ancestor [1, 2, 3, 4, 5,
taxa 2 and 6 have ancestor [0, 1, 2, 4, 6]
taxa 2 and 7 have ancestor [1, 2, 4, 7]
taxa 3 and 0 have ancestor [0, 2, 3, 5, 7]
taxa 3 and 1 have ancestor [1, 3, 4, 5, 7]
taxa 3 and 2 have ancestor [1, 2, 3, 4, 5, 7]
taxa 3 and 4 have ancestor [3, 4, 5, 6, 7]
taxa 3 and 6 have ancestor [0, 1, 3, 5, 6, 7]
taxa 4 and 0 have ancestor [0, 2, 4, 5, 6]
taxa 4 and 1 have ancestor [1, 3, 4, 5, 6]
taxa 4 and 2 have ancestor [1, 2, 4, 5, 6]
taxa 4 and 3 have ancestor [3, 4, 5, 6, 7]
taxa 4 and 7 have ancestor [1, 4, 5, 6, 7]
taxa 5 and 0 have ancestor [0, 2, 3, 5, 7]
taxa 5 and 1 have ancestor [1, 3, 4, 5, 7]
taxa 5 and 2 have ancestor [1, 2, 3, 4, 5, 7]
taxa 5 and 4 have ancestor [3, 4, 5, 6, 7]
taxa 5 and 6 have ancestor [0, 1, 3, 5, 6, 7]
```

```
taxa 6 and 2 have ancestor [0, 1, 2, 4, 6]
taxa 6 and 3 have ancestor [0, 1, 3, 5, 6, 7]
taxa 6 and 4 have ancestor [0, 1, 4, 5, 6]
taxa 6 and 5 have ancestor [0, 1, 3, 5, 6, 7]
taxa 6 and 7 have ancestor [0, 1, 6, 7]
taxa 7 and 0 have ancestor [0, 1, 2, 5, 7]
taxa 7 and 2 have ancestor [1, 2, 4, 7]
taxa 7 and 3 have ancestor [1, 3, 5, 7]
taxa 7 and 4 have ancestor [1, 4, 5, 6, 7]
taxa 7 and 5 have ancestor [1, 3, 5, 7]
taxa 7 and 6 have ancestor [0, 1, 6, 7]
```

**9.2.9.** Scoring system. The method .score takes a list of lists of non-negative integers (i.e. partitions), call it partitions, and a pair of lists, say (friendships, hypotheses), where

- friendships is a list of lists;
- hypotheses is a list of length len(friendships) whose t-th element is a list of length len(friendships[t]) whose elements are lists of integers ranging from 0 to

```
len(self.phylogeneses)-1
```

(preferrably sorted from smallest to greatest);

and returns a list of length len(friendships) whose t-th element is a list of triples of the form (r,large,exact) where

- r runs over the elements of friendships[t],
- large is the large score [4] of the hypothetical ancestor hypotheses[t] [r] within the set of ancestors contained in hypotheses[t] for the list of partitions given in the input,
- exact is the exact score [4] of the hypothetical ancestor hypotheses[t] [r] within the set of ancestors contained in hypotheses[t] for the list of partitions given in the input.

This means that large is the number of partitions belonging to the first input list partitions for which there is a morphism of partitions  $x.quotient() \rightarrow partitions[i]$  where we take

```
x = EquivalenceRelation([hypotheses[t][r]],len(self.phylogeneses)-1)
```

and exact is the number of partitions that were counted in the large score of r such that if these partitions belong to the large score of any other element s in friendships[t], then either the equality

```
hypotheses[t][r] = hypotheses[t][s]
```

holds or the intersection of hypotheses[t][r] with hypotheses[t][s] is empty.

The second input of the method .score can, for instance, be taken to be the output of the procedure

```
self.set_up_friendships().
```

For example, consider the following file containing a sequence alignment.

### 2 ACGCTAGCGCGATCGATCGATCGATC 4 ACGACTTAGCGGATCTGATACTCCCTCGATC 6 ACGACCTAGCGGATCTTATAACTCACCGATC

- 8 ACGCTAGCGGCTGATAACGATCGTATCGATC
- 9 >Eric

7 >Doug

1 >Alice

5 >Carles

- 10 ACGCATGCGCGATCGACGGATCGTATCTATC
- 11 >Fred
- 12 ACGACCTAGCAGATTTCTAATCTCAACGATC
- 13 >Garv
- 14 ACGCGAGCATCTGAACACGATTGTAACGATC
- 15 >Haley
- 16 ACGCTACGCGACGATCGGCTTTAGATCGATC

Let us now construct the list local of partitions induced by each non-trivial column of the previous alignment, as shown below.

Align.fa

```
>>> from efp import _epi_factorize_partition
>>> pre_local = usf.fasta("Align1.fa")[1]
>>> local = list()
>>> for j in range(len(pre_local[0])):
       column = list()
         for i in range(len(pre_local)):
           column.append(pre_local[i][j])
         r = _epi_factorize_partition(column)
         if r != [0]*len(column):
           local.append(r)
. . .
```

Then, we can use the method .score, as shown below, with the Phylogeny item pgy considered in section 9.2.6.

```
>>> 1 = pgy.score(local,pgy.set_up_friendships())
>>> for t in range(len(1)):
      for (r,large,exact) in l[t]:
>>>
        print("taxa " + str(t) + " and " + str(r)+" coalesce with large
>>>
    score "+str(large)+" and with exact score "+str(exact))
taxa 0 and 1 coalesce with large score 1 and with exact score 1
taxa 0 and 3 coalesce with large score 2 and with exact score 0
taxa 0 and 4 coalesce with large score 1 and with exact score 0
taxa 0 and 6 coalesce with large score 1 and with exact score 0
taxa 0 and 7 coalesce with large score 1 and with exact score 0
taxa 1 and 0 coalesce with large score 1 and with exact score 0
taxa 1 and 2 coalesce with large score 2 and with exact score 1
taxa 1 and 6 coalesce with large score 0 and with exact score 0
taxa 1 and 7 coalesce with large score 0 and with exact score 0
```

```
taxa 2 and 0 coalesce with large score 2 and with exact score 2
taxa 2 and 3 coalesce with large score 0 and with exact score 0
taxa 2 and 5 coalesce with large score 0 and with exact score 0
taxa 2 and 6 coalesce with large score 0 and with exact score 0
taxa 2 and 7 coalesce with large score 0 and with exact score 0
taxa 3 and 0 coalesce with large score 2 and with exact score 0
taxa 3 and 1 coalesce with large score 0 and with exact score 0
taxa 3 and 2 coalesce with large score 0 and with exact score 0
taxa 3 and 4 coalesce with large score 2 and with exact score 1
taxa 3 and 6 coalesce with large score 1 and with exact score 0
taxa 4 and 0 coalesce with large score 1 and with exact score 0
taxa 4 and 1 coalesce with large score 1 and with exact score 0
taxa 4 and 2 coalesce with large score 1 and with exact score 0
taxa 4 and 3 coalesce with large score 2 and with exact score 1
taxa 4 and 7 coalesce with large score 0 and with exact score 0
taxa 5 and 0 coalesce with large score 2 and with exact score 0
taxa 5 and 1 coalesce with large score 0 and with exact score 0
taxa 5 and 2 coalesce with large score 0 and with exact score 0
taxa 5 and 4 coalesce with large score 2 and with exact score 1
taxa 5 and 6 coalesce with large score 1 and with exact score 0
taxa 6 and 2 coalesce with large score 0 and with exact score 0
taxa 6 and 3 coalesce with large score 1 and with exact score 0
taxa 6 and 4 coalesce with large score 0 and with exact score 0
taxa 6 and 5 coalesce with large score 1 and with exact score 0
taxa 6 and 7 coalesce with large score 2 and with exact score 1
taxa 7 and 0 coalesce with large score 1 and with exact score 0
taxa 7 and 2 coalesce with large score 0 and with exact score 0
taxa 7 and 3 coalesce with large score 1 and with exact score 0
taxa 7 and 4 coalesce with large score 0 and with exact score 0
taxa 7 and 5 coalesce with large score 1 and with exact score 0
taxa 7 and 6 coalesce with large score 2 and with exact score 1
```

**9.2.10.** Choose. The method .choose takes a list of lists of triples (r,1,e) where 1 and e are non-negative integers and returns a list of lists whose *i*-th list is the list of those elements r of the *i*-th internal list of the input list for which the associated pairs (1,e) are equal to the greatest local maxima of the function  $\Gamma: (e,1) \mapsto (1,e)$  ordered by the lexicographical order and relative to the pairs of the *i*-th internal list of the input list (see the example below and [4, Definition 3.14]).

```
>>> k1 = [[("a",1,0),("b",2,0),("c",1,1),("d",3,3)]]
>>> choose = pgy.choose(k1)
>>> print(choose)
[['d']]
>>> k2 = [[("a",1,0),("b",2,0),("c",1,1),("d",3,3),("e",3,3)]]
>>> choose = pgy.choose(k2)
>>> print(choose)
[['d', 'e']]
>>> k3 = [[("a",1,0),("b",2,0),("c",8,1),("d",3,3),("e",3,3)]]
>>> choose = pgy.choose(k3)
>>> print(choose)
[['c']]
```

```
>>> k4 = [[("a",1,0),("b",9,0),("c",8,1),("d",3,3),("e",3,3)]]
>>> choose = pgy.choose(k4)
>>> print(choose)
[['b']]
>>> k = k1 + k2 + k3 + k4
>>> choose = pgy.choose(k)
>>> print(choose)
[['d'], ['d', 'e'], ['c'], ['b']]
```

The following example shows the type of output that one gets if one gives the list 1 of the example of section 9.2.9 to the method .choose.

**9.2.11.** Set up competition. The method <code>.set\_up\_competition</code> takes a list of lists of integers whose length must be equal to the length of <code>self.phylogeneses</code> (i.e. the number of taxa of the phylogeny) and returns a list of lists of integers whose length is also equal to the length of <code>self.phylogeneses</code> and whose t-th internal list is the union of the t-th list of <code>self.coalescent()</code> with the r-th list of <code>self.coalescent()</code> for every element r in the t-th internal list of the input list.

The following example shows the ouput of the method .set\_up\_competition when it is given the list choose constructed in the last example of section 9.2.10.

### **Bibliography**

- [1] R. Tuyéras, (2017), Category theory for genetics, arXiv:1708.05255
- [2] R. Tuyéras, (2018), Category theory for genetics I: mutations and sequence alignments, arXiv:1805.07002
- [3] R. Tuyéras, (2018), Category theory for genetics II: genotype, phenotype and haplotype, arXiv:1805.07004
- [4] R. Tuyéras, (2018), Category theory for genetics III: natural selection, evolution and phylogeny.