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(based on a slides by Stuart Brown, NYU)



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### Modules

- Python functions are divided into three sets
  - A small core set that are always available
  - Some built-in modules such as math and os that can be imported from the basic install (ie. >>> import math)
  - An number of optional modules that must be downloaded and installed before you can import them: code that uses such modules is said to have "dependencies"
  - Most are available in different Linux distributions, or via pypy.org using pip (the Python Package Index)
- Anyone can write new Python modules, and often several different modules are available that can do the same task



## **Object Oriented Code**

- Python implements oject oriented programming
- Classes bundle data and functionality

```
magic method" class MyClass:
    """A simple example class"""

def __init__(self):
    self.data = []
    self._priv = "fuggetaboutit"

def say_hi(self):
    return 'hello world'
    def __str__(self):
    return "yoohoo"+str(self.data[:3]) +"..."
```

```
instantiation
z = MyClass()
z.data = [1,"foo","bar",-5,9]
print(z) #???
```



## The Seq object

- The Seq object class is simple and fundamental for a lot of Biopython work. A Seq object can contain DNA, RNA, or protein.
- It contains a string and a defined alphabet for that string.
- The alphabets are actually defined objects such as

```
IUPACAmbiguousDNA Or IUPACProtein
```

- Which are defined in the Bio.Alphabet module
- A Seq object with a DNA alphabet has some different methods than one with an Amino Acid alphabet

```
>>> from Bio.Seq import Seq
>>> from Bio.Alphabet import IUPAC
>>> my_seq = Seq('AGTACACTGGT', <u>IUPAC.unambiguous_dnapython</u>
```

## SeqRecord Example

http://biopython.org/wiki/SeqRecord

#### Extracting information from a SeqRecord

Lets look in detail at the well annotated SeqRecord objects Biopython creates from a
GenBank file, such as ls\_orchid.gbk, which we'll load using the SeqIO module. This
file contains 94 records:



### SeqIO and FASTA files

- SeqIO is the all purpose file read/write tool for SeqRecords
  - SeqIO can read many file types: http://biopython.org/wiki/SeqIO
- SeqIO has .read() and .write() methods
  - (do not need to "open" file first)
- It can read a text file in FASTA format
- In Biopython, fasta is a type of SeqRecord with specific fields
  - grab the file: NC\_005816.fna, and saved it as a text file in your current directory



## Multiple FASTA Records in one file

- The FASTA format can store many sequences in one text file
- SeqIO.parse() reads the records one by one
- This code creates a list of SeqRecord objects:



# Seq-ing deeper

Seq: an object for containing sequence information. Basic sequence operations.

```
>>> from Bio.Seq import Seq
>>> dir(Seq)
['__add__', '__class__', '__contains__', '__delattr__',
'__dict__', '__dir__', '__doc__', '__eq__',
""" format__', '__ge__', '__getattribute__',
""" getitem__', '__gt__', '__hash__', '__init__',
""" le__', '__len__', '__lt__', '__module__', '__ne__',
""" new__', '__radd__', '__reduce__', '__reduce_ex__',
'__repr__', '__setattr__', '__sizeof__', '__str ',
' subclasshook ', ' weakref ',
'_get_seq_str_and_check_alphabet', 'back_transcribe',
'complement', 'count', 'endswith', 'find', 'lower',
'lstrip', 'reverse complement', 'rfind', 'rsplit',
'rstrip', 'split', 'startswith', 'strip', 'tomutable',
'tostring', 'transcribe', 'translate', 'ungap', 'upper']
```



# Seq Source Code

```
def __init__(self, data, alphabet=Alphabet.generic_alphabet):
        # Enforce string storage
        if not isinstance(data, basestring):
            raise TypeError("The sequence data given to a Seq object should "
                            "be a string (not another Seg object etc)")
        self. data = data
        self.alphabet = alphabet
 def __repr__(self):
        """Returns a (truncated) representation of the sequence for debugging."""
        if len(self) > 60:
            # Shows the last three letters as it is often useful to see if there
            # is a stop codon at the end of a sequence.
            # Note total length is 54+3+3=60
            return "{0}('{1}...{2}', {3!r})".format(self.__class__.__name__,
                                                     str(self)[:54],
                                                     str(self)[-3:],
                                                     self.alphabet)
        else:
            return '{0}({1!r}, {2!r})'.format(self.__class__.__name__,
                                               self._data,
                                               self.alphabet)
```



# Seq Source Code

```
def transcribe(self):
    base = Alphabet._get_base_alphabet(self.alphabet)
    if isinstance(base, Alphabet.ProteinAlphabet):
        raise ValueError("Proteins cannot be transcribed!")
    if isinstance(base, Alphabet.RNAAlphabet):
        raise ValueError("RNA cannot be transcribed!")

if self.alphabet == IUPAC.unambiguous_dna:
        alphabet = IUPAC.unambiguous_rna
elif self.alphabet == IUPAC.ambiguous_dna:
        alphabet = IUPAC.ambiguous_rna
else:
        alphabet = Alphabet.generic_rna
return Seq(str(self).replace('T', 'U').replace('t', 'u'), alphabet))
```



## Create your own sequence class

Base class

```
from Bio. Seg import Seg
class MySeq(Seq):
     def __repr__(self):
         if len(self) > 60:
             # Shows the last three letters as it is often useful to see if there
             # is a stop codon at the end of a sequence.
             # Note total length is 54+3+3=60
             return "{0}('{1}---{2}', {3!r})".format(self.__class__.__name__,
                                                       str(self)[:54],
                                                       str(self)[-3:],
Derived class
                                                       self.alphabet)
         else:
                                 Was " "
{2!!} ':format(self.__class__.__name__,
             return '{0}({1!r},
                                                 self._data,
                                                 self.alphabet)
```

- The new \_\_repr\_\_ overrides the old \_\_repr\_\_
- Everything else remains the same!
- MySeq is a derived class of Seq
- Also, MySeq is inherited from Seq



### Direct Access to GenBank

- BioPython has modules that can directly access databases over the Internet
- The Entrez module uses the NCBI Efetch service
- Efetch works on many NCBI databases including protein

### >>> print(record)

ID: EU490707.1

Name: EU490707

Description: Selenipedium aequinoctiale maturase K (matK) gene, partial cds; chloroplast.

Number of features: 3

/sequence version=1

/source=chloroplast Selenipedium aequinoctiale

/taxonomy=['Eukaryota', 'Viridiplantae',

'Streptophyta', 'Embryophyta', 'Tracheophyta',

'Spermatophyta', 'Magnoliophyta', 'Liliopsida', 'Asparagales', 'Orchidaceae', 'Cypripedioideae

### **BLAST**

- BioPython has several methods to work with the popular NCBI BLAST software
- NCBIWWW.qblast() sends queries directly to the NCBI BLAST server. The query can be a Sequence of this now qbject, FASTA file, or a GenBank ID.

## **XML**

Extensible Markup Language
Used to encode documents in a form that is human readable and machine readable

### Note

To: Tove

From: Jani

#### Reminder

Don't forget me this weekend!



## XML is Extensible

#### Note

To: Tove

From: Jani

#### Reminder

Don't forget me this weekend!

#### Note

To: Tove

From: Jani

Date: 2015-09-01 08:30

Don't forget me this weekend!



### **BLAST XML**

```
MULTISPECIES: bacteriocin sakacin-P [Lactobacillus]
 Sequence ID: WP 004271264.1 Length: 61 Number of Matches: 1
 ▶ See 11 more title(s)
 Range 1: 1 to 61 GenPept Graphics
                                                                                     ▼ Next Match ▲ Previous Match
                   Expect Method
                                                                Identities
                                                                                 Positives
 122 bits(307) 1e-35 Compositional matrix adjust.
                                                               61/61(100%)
                                                                                61/61(100%) 0/61(0%)
                MEKFIELSLKEVTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWN
MEKFIELSLKEVTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWN
MEKFIELSLKEVTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWN
 Sbjct 1
 Query 61
 Sbjct 61 K 61
Bownload ∨ GenPept Graphics
 MULTISPECIES: bacteriocin [Lactobacillus]
 Sequence ID: WP 056946897.1 Length: 61 Number of Matches: 1
 ▶ See 3 more title(s)
 Range 1: 1 to 61 GenPept Graphics
                                                                                     ▼ Next Match ▲ Previous Match
                                                                Identities
                                                                                 Positives
                                                                60/61(98%) 61/61(100%) 0/61(0%)
 120 bits(302) 8e-35 Compositional matrix adjust.
                MEKFIELSLKEYTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWN
MEKFIELSLKEYTAITGGKYYGNGVHCGKYSCTVDWGTAIGNIGNNAAANWATGGNAGWN
MEKFIELSLKEYTAITGGKYYGNGVHCGKYSCTVDWGTAIGNIGNIGNNAAANWATGGNAGWN
 Query 61
Sbjct 61 K 61
Bownload → GenPept Graphics
 bacteriocin mundticin [Enterococcus pallens]
 Sequence ID: WP_010758561.1 Length: 62 Number of Matches: 1
 ▶ See 3 more title(s)
                                                                                     ▼ Next Match ▲ Previous Match
 Range 1: 1 to 61 GenPept Graphics
                                                                                  Positives
                    Expect Method
                                                                  Identities
 91.3 bits(225) 4e-23 Compositional matrix adjust.
                                                                 45/61(74%) 50/61(81%) 0/61(0%)
                MEKFIELSLKEVTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWN
M+ LS KE+ ITGGKYYGNG+ CGK+SC+VDWG AIG IGNNAAANWATGG AGWN
MQNIKALSAKELIEITGGKYYGNGLSCGKYSCSVDWGKAIGIIGNNAAANWATGGAAGWN
 Query 61
                    61
 Sbict 61 K
```



### **BLAST XML**

Sbict 61 K 61

```
<Hit num>3</Hit num>
  <Hit id>gi|498453073|ref|WP 010758561.1|</Hit id>
  <Hit def>bacteriocin mundticin [Enterococcus pallens] &gt;qi|486858092|qb|E0H91048.1| bacterioc
in mundticin [Enterococcus pallens ATCC BAA-351] &qt;qi|508242815|qb|E0U16245.1| bacteriocin mund
ticin [Enterococcus pallens ATCC BAA-351] &qt;qi|1105345489|qb|0JG79015.1| bacteriocin mundticin
[Enterococcus pallens]</Hit def>
  <Hit accession>WP 010758561</Hit accession>
  <Hit len>62</Hit len>
  <Hit hsps>
    <Hsp>
       <Hsp num>1</Hsp num>
       <Hsp bit-score>91.2781/Hsp bit-score>
       <Hsp score>225</Hsp score>
       <Hsp evalue>4.38515e-23/Hsp evalue>
       <Hsp query-from>1</Hsp query-from>
       <Hsp query-to>61</Hsp query-to>
       <Hsp hit-from>1</Hsp hit-from>
       <Hsp hit-to>61</Hsp hit-to>
       <Hsp query-frame>0</Hsp query-frame>
       <Hsp hit-frame>0</Hsp hit-frame>
       <Hsp identity>45</Hsp identity>
       <Hsp positive>50</Hsp positive>
       <Hsp gaps>0</Hsp gaps>
       <Hsp align-len>61</Hsp align-len>
       <Hsp qseq>MEKFIELSLKEVTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWNK</Hsp qseq>
       <Hsp hseq>MQNIKALSAKELIEITGGKYYGNGLSCGKYSCSVDWGKAIGIIGNNAAANWATGGAAGWNK</Hsp hseq>
       <Hsp midline>M+
                              LS KE+ ITGGKYYGNG+ CGK+SC+VDWG AIG IGNNAAANWATGG AGWNK</Hsp midline>
     </Hsp>
  </Hit hsps>
 bacteriocin mundticin [Enterococcus pallens]
 Sequence ID: WP 010758561.1 Length: 62 Number of Matches: 1
 ▶ See 3 more title(s)

▼ Next Match  

▲ Previous Match
 Range 1: 1 to 61 GenPept Graphics
            Expect Method
                                     Identities
                                               Positives
 91.3 bits(225) 4e-23 Compositional matrix adjust. 45/61(74%) 50/61(81%) 0/61(0%)
         MEKFIELSLKEVTAITGGKYYGNGVHCGKHSCTVDWGTAIGNIGNNAAANWATGGNAGWN
M+ LS KE+ ITGGKYYGNG+ CGK+SC+VDWG AIG IGNNAAANWATIG AGWN
MQNIKALSAKELIEITGGKYYGNGLSCGKYSSCSVDWGKAIGIIGNNAAANWATIGAAGWN
 Query 61 K 61
```



### **BLAST**

- BioPython has several methods to work with the popular NCBI BLAST software
- NCBIWWW.qblast() sends queries directly to the NCBI BLAST server. The query can be a Sequence yet?

  qbject, FASTA file, or a GenBank ID.

```
>>> from Bio.Blast import NCBIWWW
>>> from Bio import SeqIO

>>> query = SeqIO.read("test.fasta", format="fasta")

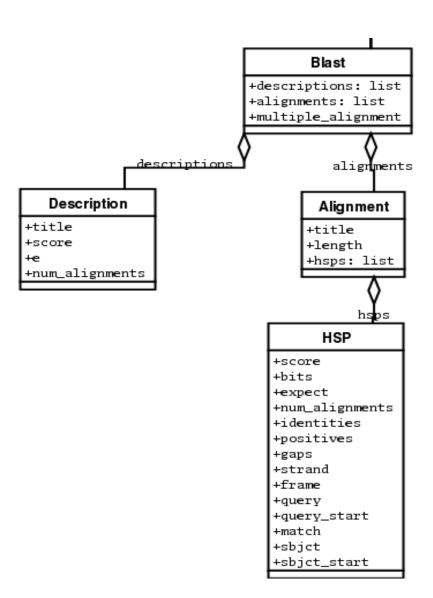
>>> result_handle = NCBIWWW.qblast("blastn", "nt",
query.seq)
>>> blast_result = result_handle.read()
>>> blast_file = open("my_blast.xml", "w") # create an xm output file
>>> blast_file.write(blast_result)
```

### Parse BLAST Results

- It is often useful to obtain a BLAST result directly (local BLAST server or via Web browser) and then parse the result file with Python.
- Save the BLAST result in XML format
  - NCBIXML.read() for a file with a single BLAST result (single query)
  - NCBIXML.parse() for a file with multiple BLAST results (multiple queries)



## **BLAST Record Object**





## View Aligned Sequence

```
>>> from Bio.Blast import NCBIXML
    >>> handle = open("my_blast.xml")
    >>> blast record = NCBIXML.read(handle)
    >>> for hit in blast record.alignments:
             for hsp in hit.hsps:
                 print hit.title
                 print hsp.expect
                 print (hsp.query[0:75] + '...')
                 print(hsp.match[0:75] + '...')
                 print(hsp.sbjct[0:75] + '...')
qi|731383573|ref|XM 002284686.2| PREDICTED: Vitis vinifera cold-regulated 413
plasma membrane protein 2 (LOC100248690), mRNA
2.5739e-53
{	t ATGCTAGTATGCTCGGTCATTACGGGTTTTGGCACT-CATTTCCTCAAATGGCTCGCCTGCCTTTGCGGCTATTTAC\dots}
1111 11 11111 ...
ATGCCATTAAGCTTGGTGGTCTGGGCTTTGGCACTACATTTCTTGAG-TGGTTGGCTTCTTTTTGCTGCCATTTAT...
```



## Many Matches

- Often a BLAST search will return many matches for a single query (save as an XML format file)
- NCBIXML.parse() can return these as BLAST record objects in a list, or deal with them directly in a for loop.

## Illumina Sequences

 Illumina sequence files are usually stored in the FASTQ format. Similar to FASTA, but with an additional pair of lines for the quality annotation of each base.

```
@
SR
R3
50
95
3.
Μ
ΕN
DE
00
47
C6
Μ
N8
AA
XX
:1:
```

16



## Get a file by FTP in Python

```
>>> from ftplib import FTP
>>> host="ftp.sra.ebi.ac.uk"
>>> ftp=FTP(host)
>>> ftp.login()
'230 Login successful.'
ftp.cwd('vol1/fastq/SRR020/SRR020192')
'250 Directory successfully changed.'
>>> ftp.retrlines('LIST')
                 ftp 1777817 Jun 24 20:12
-r--r-- 1 ftp
SRR020192.fastq.gz
'226 Directory send OK.'
```

>>> ftn rotrhinary/'DETD CDD020102 facta az'

n

