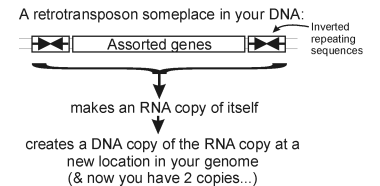
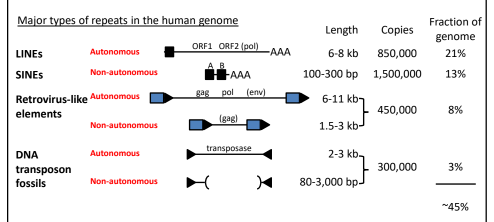
You and your (DNA) parasites



Events like these, happening over and over again, have led to...

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You and your (DNA) parasites



Bottom line: Roughly half of your (and my) genome is the fossil wreckage of genomic parasites.

We know this (in part) from sequence alignments.

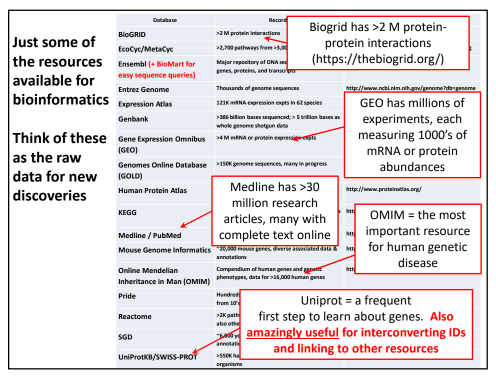
So far, we've talked about

- DNA, RNA and protein sequences
- How to compare sequences to decide if they are related
- Having databases full of sequences and comparing them rapidly (BLAST)

In fact, <u>many</u> such databases exist, so today we'll start with a brief tour of <u>some</u> of the biological data on the web.

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	Database	Records	Address
Just some of the resources available for bioinformatics	BioGRID	>2 M protein interactions	https://thebiogrid.org
	EcoCyc/MetaCyc	>2,700 pathways from >3,000 organisms	http://www.ecocyc.org, http://www.metacyc.org
	Ensembl (+ BioMart for easy sequence queries)	Major repository of DNA sequences, genomes, genes, proteins, and transcripts	http://useast.ensembl.org/index.html
	Entrez Genome	Thousands of genome sequences	http://www.ncbi.nlm.nih.gov/genome?db=genome
	Expression Atlas	121K mRNA expression expts in 62 species	https://ebi.ac.uk/gxa/home/
	Genbank	>386 billion bases sequenced; > 5 trillion bases as whole genome shotgun data	https://www.ncbi.nlm.nih.gov/genbank/
Think of these as the raw data for new discoveries	Gene Expression Omnibus (GEO)	>4 M mRNA or protein expression expts	http://www.ncbi.nlm.nih.gov/geo/
	Genomes Online Database (GOLD)	>150K genome sequences, many in progress	https://gold.jgi.doe.gov/index
	Human Protein Atlas	millions of high-res images of ~17K human proteins across tissues, cancers, & cell lines	http://www.proteinatlas.org/
	KEGG	Most known pathways, in 538 graphical diagrams and >6K organisms (<i>via</i> homology)	http://www.genome.ad.jp/kegg/
	Medline / PubMed	>30 million references	https://www.ncbi.nlm.nih.gov/PubMed/
	Mouse Genome Informatics	~20,000 mouse genes, diverse associated data & annotations	http://www.informatics.jax.org/
	Online Mendelian Inheritance in Man (OMIM)	Compendium of human genes and genetic phenotypes, data for >16,000 human genes	https://www.ncbi.nlm.nih.gov/omim/
	Pride	Hundreds of millions of peptide mass spectra from 10's of thousands of experiments	https://www.ebi.ac.uk/pride/archive/
	Reactome	>2K pathways involving >10K human proteins, also other organisms	https://www.reactome.org/
	SGD	~6,000 yeast genes, diverse associated data & annotations	https://www.yeastgenome.org/
	UniProtKB/SWISS-PROT	>550K hand-curated sequence entries from >9K organisms	https://www.uniprot.org/



Live demo Ensembl->BioMart->filter for [IPR031588], OMIM, Reactome, Human Protein Atlas

It's nice to know that all of this exists, but ideally, you'd like to be able to so something constructive with the data.

That means getting the data inside your own programs.

All of these databases let you download data in big batches, but this isn't always the case, so....

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Let's empower your Python scripts to grab data from the web.

We'll use Python <u>library/module</u> = an optional, specialized set of Python methods

This particular Python module is called *urllib* (Py3) or *urllib2* (Py2)

urllib/urllib2 is:

- A collection of programs/tools to let you to surf the web from inside your programs.
- Much more powerful than the simple tasks we'll do with it.
- More details: <u>https://docs.python.org/3.8/library/urllib.request.html</u> or <u>http://docs.python.org/2/library/urllib2.html</u>

The basic idea:

We first set up a "request" by opening a connection to the URL.

We then save the response in a variable and print it.

If it can't connect to the site, it'll print out a helpful error message instead of the page.

You can more or less use the commands in a cookbook fashion....

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For example:

```
import urllib.request # include the urllib.request module

url = "https://www.utexas.edu/"

try: # this 'try' statement tells Python that we might expect an error.

request = urllib.request.urlopen(url) # setup a request
page = request.read().decode('utf-8') # save the response
print(page) # show the result to the user

except urllib2.URLError: # handle a page not found error
print("Could not find page.")

→ Run this...
```

Python 3 version

```
For example:
import urllib2
                                       # include the urllib2 module
url = "https://www.utexas.edu/"
                    # this 'try' statement tells Python that we might expect an error.
   request = urllib2.urlopen(url)
                                       # setup a request
   page = request.read()
                                       # save the response
   print(page)
                                       # show the result to the user
except urllib2.URLError:
                                       # handle a page not found error
   print("Could not find page.")
→ Run this...
                                                                         Python 2 version
```

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That was (more or less) a static web page.

Let's try one that requires some sort of action, for example by entering a document id or an id code for a sequence.

Many web pages pass this information along in the web URL itself...

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Here's a complete Python program to retrieve a single entry from Medline:

Python 3 version

Here's a complete Python program to retrieve a single entry from Medline:

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```
If you run that program, you should get back...
<!DOCTYPE html>
.....lots of metadata.....
OWN - NLM
STAT- MEDLINE
DCOM- 20010322
                                                  the Medline entry for the human
LR - 20210108
IS - 0028-0836 (Print)
                                                        genome sequence paper
IS - 0028-0836 (Linking)
VI - 409
IP - 6822
DP - 2001 Feb 15
TI - Initial sequencing and analysis of the human genome.
PG - 860-921
AB - The human genome holds an extraordinary trove of information about human
   development, physiology, medicine and evolution. Here we report the results of an
   international collaboration to produce and make freely available a draft sequence of
   the human genome. We also present an initial analysis of the data, describing some
   of the insights that can be gleaned from the sequence.
{\sf FAU} - Lander, E {\sf S}
AU - Lander ES
AD - Whitehead Institute for Biomedical Research, Center for Genome Research, Cambridge,
   MA 02142, USA. lander@genome.wi.mit.edu
     [and so on]
```

```
If you run that program, you should get back...
>>>
<!DOCTYPE html>
....lots of metadata.....
OWN - NLM
STAT- MEDLINE
DCOM- 20010322
                                                We just printed it. We could have
LR - 20210108
IS - 0028-0836 (Print)
                                                saved it or extracted data from it.
IS - 0028-0836 (Linking)
VI - 409
                                                                For example...
IP - 6822
DP - 2001 Feb 15
TI - Initial sequencing and analysis of the human genome.
AB - The human genome holds an extraordinary trove of information about human
   development, physiology, medicine and evolution. Here we report the results of an
   international collaboration to produce and make freely available a draft sequence of
   the human genome. We also present an initial analysis of the data, describing some
   of the insights that can be gleaned from the sequence.
FAU - Lander, ES
AU - Lander ES
AD - Whitehead Institute for Biomedical Research, Center for Genome Research, Cambridge,
   MA 02142, USA. lander@genome.wi.mit.edu
```

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Here's our Python program again to retrieve a single entry from Medline. How would we modify this to count the authors?

Python 3 version

Here's our Python program again to retrieve a single entry from Medline. How would we modify this to count the authors? import urllib.request pmid = 11237011 # Insert the pmid where the {} are in the following URL: url = "https://pubmed.ncbi.nlm.nih.gov/?term={0}[uid]&format=pubmed".format(pmid) # there might be an error! **Medline begins** request = urllib.request.urlopen(url) author lines with page = request.read().decode('utf-8') "AU - ", so... print(page.count("AU - ")) <</pre> except urllib.request.URLError: # handle page not found error print("Could not connect to Medline!") So, there were 256 authors on one (of → Run this, & get ... >>> 256 the two) human genome papers

Python 3 version

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 Queries to Medline or any other NCBI database, including GenBank, are described at:

http://www.ncbi.nlm.nih.gov/books/NBK3862/

(& for that matter, <u>all</u> of medline is downloadable)

- You can often figure out the form of the URL just by looking something up in a database, then noting the address of the web page with the data.
- This very simple approach could easily be the basis for:
 - a home-made web browser
 - a program to consult biological databases in real time
 - a program to map the internet, etc.
- Of course, with this kind of power available to you, the imagination reels...