# Entrez Direct: E-utilities on the UNIX Command Line

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# **Getting Started**

#### Introduction

Entrez Direct (EDirect) provides access to the NCBI's suite of interconnected databases (publication, sequence, structure, gene, variation, expression, etc.) from a UNIX terminal window. Functions take search terms from command-line arguments. Individual operations are combined to build multi-step queries. Record retrieval and formatting normally complete the process.

EDirect also includes an argument-driven function that simplifies the extraction of data from document summaries or other results that are returned in structured XML format. This can eliminate the need for writing custom software to answer ad hoc questions. Queries can move seamlessly between EDirect commands and UNIX utilities or scripts to perform actions that cannot be accomplished entirely within Entrez.

#### Installation

EDirect will run on UNIX and Macintosh computers that have the Perl language installed, and under the Cygwin UNIX-emulation environment on Windows PCs. To install the EDirect software, copy the following commands and paste them into a terminal window:

```
cd ~
perl -MNet::FTP -e \
    '$ftp = new Net::FTP("ftp.ncbi.nlm.nih.gov", Passive => 1);
    $ftp->login; $ftp->binary;
    $ftp->get("/entrez/entrezdirect/edirect.tar.gz");'
gunzip -c edirect.tar.gz | tar xf -
rm edirect.tar.gz
export PATH=$PATH:$HOME/edirect
```

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```
./edirect/setup.sh
```

This downloads several scripts into an "edirect" folder in the user's home directory. The setup.sh script then downloads any missing Perl modules, and may print an additional command for updating the PATH environment variable in the user's configuration file. Copy that command, if present, and paste it into the terminal window to complete the installation process. The editing instructions will look something like:

```
echo "export PATH=\$PATH:\$HOME/edirect" >> $HOME/.bash_profile
```

The configuration file can instead be modified manually using a text editor, if desired.

#### **Entrez Direct Functions**

Navigation functions support exploration within the Entrez databases:

- **esearch** performs a new Entrez search using terms in indexed fields.
- **elink** looks up neighbors (within a database) or links (between databases).
- **efilter** filters or restricts the results of a previous query.

Records can be retrieved in specified formats or as document summaries:

• **efetch** downloads records or reports in a designated format.

Desired fields from XML results can be extracted without writing a program:

• **xtract** converts XML into a table of data values.

Several additional functions are also provided:

- einfo obtains information on indexed fields in an Entrez database.
- epost uploads unique identifiers (UIDs) or sequence accession numbers.
- **nquire** sends a URL request to a web page or CGI service.

# **Entering Query Commands**

UNIX programs are run by typing the name of the program and then supplying any required or optional arguments on the command line. Argument names are letters or words that start with a dash ("-") character.

In order to begin an Entrez search, the user types "esearch" and then enters the required - db (database) and -query arguments. A query on unqualified search terms:

```
esearch -db pubmed -query "opsin gene conversion"
```

constructs the appropriate Entrez Utilities (E-utilities) URL from the query terms and executes the search. EDirect handles many technical details behind the scenes (avoiding the learning curve normally required for E-utilities programming), and saves the results on the Entrez history server.

# Constructing Multi-Step Queries

EDirect allows individual operations to be described separately, combining them into a multi-step query by using the vertical bar ("|") UNIX pipe symbol. Piping esearch to elink:

```
esearch -db pubmed -query "opsin gene conversion" | elink -related
```

will look up related articles (precomputed PubMed neighbors) of the initial results.

# Writing Commands on Multiple Lines

A query can be continued on the next line by typing the backslash ("\") UNIX escape character immediately before pressing the Return key. Continuing the query links to all protein sequences published in the neighbor articles:

```
esearch -db pubmed -query "opsin gene conversion" | \
elink -related | \
elink -target protein
```

The vertical bar pipe symbol also allows the query to continue on the next line.

# Retrieving PubMed Reports

Piping PubMed query results to efetch and specifying the "abstract" format:

```
esearch -db pubmed -query "lycopene cyclase" |
efetch -format abstract
```

returns a set of reports that can be read by a person:

```
10. PLoS One. 2013;8(3):e58144. doi: 10.1371/journal.pone.0058144. Epub ... Levels of lycopene \beta-cyclase 1 modulate carotenoid gene expression and accumulation in Daucus carota.  
Moreno JC(1), Pizarro L, Fuentes P, Handford M, Cifuentes V, Stange C.  
Author information:  
(1)Departamento de Biología, Facultad de Ciencias, Universidad de Chile, Santiago, Chile.  
Plant carotenoids are synthesized and accumulated in plastids through a highly regulated pathway. Lycopene \beta-cyclase (LCYB) is a key enzyme involved directly in the synthesis of \alpha-carotene and \beta-carotene through the cyclization of lycopene. Carotenoids are produced in both carrot
```

Using "efetch -format medline" instead produces a report that can be entered into common bibliographic management software packages:

```
...
PMID- 23555569
```

```
OWN - NLM
STAT- MEDLINE
DA - 20130404
DCOM- 20130930
LR - 20131121
IS - 1932-6203 (Electronic)
IS - 1932-6203 (Linking)
VI - 8
   - 3
ΤP
DP - 2013
TI - Levels of lycopene beta-cyclase 1 modulate carotenoid gene expression
     and accumulation in Daucus carota.
PG - e58144
LID - 10.1371/journal.pone.0058144 [doi]
AB - Plant carotenoids are synthesized and accumulated in plastids
      through a highly regulated pathway. Lycopene beta-cyclase (LCYB) is a
      key enzyme involved directly in the synthesis of alpha-carotene and
      beta-carotene through the cyclization of lycopene. Carotenoids are
```

### Retrieving Sequence Reports

Nucleotide and protein records can be downloaded in FASTA format:

```
esearch -db protein -query "lycopene cyclase" | efetch -format fasta
```

which consists of a definition line followed by the sequence:

. . .

>gi|735882|gb|AAA81880.1| lycopene cyclase [Arabidopsis thaliana]
MDTLLKTPNKLDFFIPQFHGFERLCSNNPYPSRVRLGVKKRAIKIVSSVVSGSAALLDLVPETKKENLDF
ELPLYDTSKSQVVDLAIVGGGPAGLAVAQQVSEAGLSVCSIDPSPKLIWPNNYGVWVDEFEAMDLLDCLD
TTWSGAVVYVDEGVKKDLSRPYGRVNRKQLKSKMLQKCITNGVKFHQSKVTNVVHEEANSTVVCSDGVKI
QASVVLDATGFSRCLVQYDKPYNPGYQVAYGIIAEVDGHPFDVDKMVFMDWRDKHLDSYPELKERNSKIP
TFLYAMPFSSNRIFLEETSLVARPGLRMEDIQERMAARLKHLGINVKRIEEDERCVIPMGGPLPVLPQRV
VGIGGTAGMVHPSTGYMVARTLAAAPIVANAIVRYLGSPSSNSLRGDQLSAEVWRDLWPIERRRQREFFC
FGMDILLKLDLDATRRFFDAFFDLQPHYWHGFLSSRLFLPELLVFGLSLFSHASNTSRLEIMTKGTVPLA
KMINNLVQDRD

. . .

Additional FASTA -format variants are fasta\_cds\_na, fasta\_cds\_aa, and gene\_fasta.

Sequence records can also be obtained as GenBank (-format gb) or GenPept (-format gp) flatfiles, which have features annotating particular regions of the sequence:

```
LOCUS AAA81880 501 aa linear PLN ...

DEFINITION lycopene cyclase [Arabidopsis thaliana].

ACCESSION AAA81880

VERSION AAA81880.1 GI:735882

DBSOURCE locus ATHLYC accession L40176.1

KEYWORDS .

SOURCE Arabidopsis thaliana (thale cress)
```

```
ORGANISM Arabidopsis thaliana
           Eukaryota; Viridiplantae; Streptophyta; Embryophyta;
           Tracheophyta; Spermatophyta; Magnoliophyta; eudicotyledons;
           Brassicales; Brassicaceae; Camelineae; Arabidopsis.
REFERENCE
           1 (residues 1 to 501)
 AUTHORS Scolnik, P.A. and Bartley, G.E.
 TITLE Nucleotide sequence of lycopene cyclase (GenBank L40176) from
           Arabidopsis (PGR95-019)
 JOURNAL Plant Physiol. 108 (3), 1343 (1995)
FEATURES
                    Location/Qualifiers
                     1..501
                     /organism="Arabidopsis thaliana"
                     /db xref="taxon:3702"
                     1..501
    Protein
                     /product="lycopene cyclase"
     transit_peptide 1..80
    mat_peptide
                   81..501
                     /product="lycopene cyclase"
     CDS
                     1..501
                     /gene="LYC"
                     /coded_by="L40176.1:2..1507"
ORIGIN
        1 mdtllktpnk ldffipqfhg ferlcsnnpy psrvrlgvkk raikivssvv sgsaalldlv
      61 petkkenldf elplydtsks gyvdlaivgg gpaglavagg vseaglsvcs idpspkliwp
      121 nnygvwydef eamdlldcld ttwsgavvyv degykkdlsr pygrvnrkql kskmlqkcit
      181 ngvkfhqskv tnvvheeans tvvcsdgvki qasvvldatg fsrclvqydk pynpgyqvay
      241 giiaevdghp fdvdkmvfmd wrdkhldsyp elkernskip tflyampfss nrifleetsl
      301 varpglrmed igermaarlk hlqinvkrie edercvipmg gplpvlpqrv vqiqqtaqmv
      361 hpstgymvar tlaaapivan aivrylgsps snslrgdqls aevwrdlwpi errrqreffc
      421 fgmdillkld ldatrrffda ffdlqphywh gflssrlflp ellvfglslf shasntsrle
     481 imtkgtvpla kminnlvqdr d
//
```

# Searching and Filtering

# Restricting Query Results

The current results can be refined by further term searching in Entrez (useful in the protein database for limiting BLAST neighbors to a taxonomic subset):

```
esearch -db pubmed -query "opsin gene conversion" |
elink -related |
efilter -query "tetrachromacy"
```

Results can also be filtered by time. For example, the following statements:

```
efilter -days 60 -datetype PDAT efilter -mindate 1990 -maxdate 1999 -datetype PDAT
```

restrict results to articles published in the previous two months or in the 1990s, respectively.

# Qualifying Queries by Indexed Field

Query terms in esearch or efilter can be qualified by entering an indexed field abbreviation in brackets. Boolean operators and parentheses can also be used in the query expression for more complex searches.

Commonly-used fields for PubMed queries include:

```
[AFFL] Affiliation
[ALL] All Fields
[AUTH] Author
[FAUT] Author - First
[LAUT] Author - Last
[PDAT] Date - Publication
[FILT] Filter
[JOUR] Journal
[LANG] Language
[MAJR] MeSH Major Topic
[SUBH] MeSH Subheading
[MESH] MeSH Terms
[PTYP] Publication Type
[WORD] Text Word
[TITL] Title
[TIAB] Title/Abstract
[UID] UID
```

#### and a qualified query looks like:

```
"Tager HS [AUTH] AND glucagon [TIAB]"
```

Filters that limit search results to subsets of PubMed include:

```
humans [MESH]
pharmacokinetics [MESH]
chemically induced [SUBH]
all child [FILT]
english [FILT]
freetext [FILT]
has abstract [FILT]
historical article [FILT]
randomized controlled trial [FILT]
clinical trial, phase ii [PTYP]
review [PTYP]
```

Sequence databases are indexed with a different set of search fields, including:

```
[ACCN] Accession
[ALL] All Fields
[AUTH] Author
[GPRJ] BioProject
```

```
[ECNO] EC/RN Number
[FKEY] Feature key
[FILT] Filter
[GENE] Gene Name
[JOUR] Journal
[KYWD] Keyword
[MLWT] Molecular Weight
[ORGN] Organism
[PACC] Primary Accession
[PROP] Properties
[PROT] Protein Name
[SQID] SeqID String
[SLEN] Sequence Length
[SUBS] Substance Name
[WORD] Text Word
[TITL] Title
        UID
[UID]
```

and a sample query in the protein database is:

```
"alcohol dehydrogenase [PROT] NOT (bacteria [ORGN] OR fungi [ORGN])"
```

Additional examples of subset filters in sequence databases are:

```
mammalia [ORGN]
mammalia [ORGN:noexp]
cds [FKEY]
lacz [GENE]
beta galactosidase [PROT]
protein snp [FILT]
reviewed [FILT]
biomol genomic [PROP]
dbxref flybase [PROP]
gbdiv phg [PROP]
phylogenetic study [PROP]
sequence from mitochondrion [PROP]
src cultivar [PROP]
srcdb refseq validated [PROP]
150:200 [SLEN]
2000:4000 [MLWT]
```

(The calculated molecular weight (MLWT) field is only indexed for proteins (and structures), not nucleotides.)

# **Examining Intermediate Results**

EDirect stores intermediate results on the Entrez history server. EDirect navigation functions produce a custom XML message with the relevant fields (database, web environment, query key, and record count) that can be read the next command in the pipeline.

The results of each step in a query can be examined to confirm expected behavior before adding the next step. The Count field in the ENTREZ\_DIRECT object contains the

number of records returned by the previous step. A good measure of query success is a reasonable (non-zero) count value. For example:

with 39 protein structures being within the specified molecular weight range and having the desired (X-ray crystallographic) atomic position resolution.

(The QueryKey value is 7 instead of 5 because each elink command obtains the record count by running a separate ESearch query immediately after the ELink operation.)

# Combining Independent Queries

Independent esearch, elink, and efilter operations can be performed and then combined at the end by using the history server's "#" convention to indicate query key numbers. (The steps to be combined must be in the same database.) Subsequent esearch commands can take a -db argument to override the database piped in from the previous step. (Piping the queries together is necessary for sharing the same history thread.) For example, the query:

```
esearch -db protein -query "amyloid* [PROT]" |
elink -target pubmed |
esearch -db gene -query "apo* [GENE]" |
elink -target pubmed |
esearch -query "(#3) AND (#6)" |
efetch -format docsum |
xtract -pattern DocumentSummary -element Id Title
```

uses truncation searching (entering the beginning of a word followed by an asterisk) to return titles of papers with links to amyloid protein sequence and apolipoprotein gene records:

```
23962925 Genome analysis reveals insights into physiology and ... 23959870 Low levels of copper disrupt brain amyloid-\beta homeostasis ... 23371554 Genomic diversity and evolution of the head crest in the ... Novel genetic loci identified for the pathophysiology of ...
```

The use of (#3) AND (#6) instead of (#2) AND (#4) above reflects the need for each elink command to execute a separate ESearch query, which increments the QueryKey, in order to obtain the record count. The -label argument can be used to get around this artifact. The label value is prefixed by a "#" symbol and placed in parentheses in the final search. Thus:

```
esearch -db structure -query "insulin [TITL]" |
elink -target pubmed -label struc_cit |
esearch -db protein -query "insulin [PROT]" |
elink -target pubmed -label prot_cit |
esearch -query "(#struc_cit) AND (#prot_cit)" |
efetch -format uid

will return:

15299880
9235985
9141131
8421693
```

without the need to keep track of the internal QueryKey values.

#### Structured Data

# Advantages of XML Format

The ability to obtain Entrez records in structured XML format, and to easily extract the underlying data, allows the user to ask novel questions that are not addressed by existing analysis software.

The advantage of XML is that many pieces of information are in specific locations in a well-defined data hierarchy. Accessing individual units of data that are fielded by name, such as:

```
<PubDate>2013</PubDate>
<Source>PLoS One</Source>
<Volume>8</Volume>
<Issue>3</Issue>
<Pages>e58144</Pages>
```

requires matching the same general pattern, differing only by the element name. This is much simpler than parsing the units from a long, complex string:

```
1. PLoS One. 2013;8(3):e58144 ...
```

The disadvantage of XML is that data extraction usually requires programming. But EDirect relies on the common pattern of XML value representation to provide a simplified approach to interpreting XML data.

#### Conversion of XML Data into Tabular Form

The xtract function uses command-line arguments to direct the selective conversion of XML data into a tab-delimited table. The -pattern argument divides the results into rows, while placement of data into columns is controlled by -element. A trivial example:

```
xtract -pattern ENTREZ DIRECT -element Count
```

will print the number of records in the current query.

Xtract provides control over data conversion with a divide-and-conquer strategy using separate arguments for element selection, path exploration, conditional processing, and report formatting.

Element selection finds every occurrence of each indicated item, printing values as they are encountered. Exploration control limits selection by context, presenting specified objects one at a time. Conditional processing filters by content, requiring presence (or absence) of a particular data value in order to continue. Finally, custom formatting can override the normal tabular layout of the default output.

The details and ramifications of this flexible approach are discussed in the remainder of this section.

# **Extraction Arguments**

Selection arguments (-element, -first, and -last) extract and print data values from the indicated element names:

```
-element Id -first Name Title
```

Exploration arguments (-pattern, -group, -block, and -subset) limit data extraction to specified regions of the XML, visiting all relevant objects one at a time. This sets a context for data collection, eliminates the need to provide the full path to a data element, and uncouples the concept of "what to look for" from "where to find it":

```
-pattern DocumentSummary
-block Author
```

Each pattern can have multiple groups, each group can have multiple blocks, and each block can have multiple subsets. This design allows nested exploration of complex, hierarchical data to be controlled by a linear chain of command-line argument statements.

Conditional processing arguments restrict exploration statements by object name and value (-if and -unless) or item location (-position):

```
-if Source -equals "J Bacteriol"
-position first
```

These commands are issued immediately after an exploration argument.

Formatting arguments (-ret, -tab, -sep, -pfx, and -sfx) allow extensive customization of the default row/column table presentation:

```
-pfx "\n[" -sfx "]\t" -sep " " -tab "" -ret "\n\n"
```

and apply to subsequent selection statements.

(The "\n" escape sequence indicates a line break, while "\t" specifies a tab character.)

#### **XML Document Summaries**

Entrez provides a document summary in structured XML format for every record. Piping a query to "efetch -format docsum":

```
esearch -db pubmed -query "Garber ED [AUTH] AND PNAS [JOUR]" |
elink -related |
efilter -query "mouse" |
efetch -format docsum
```

will generate an XML document summary set:

```
<DocumentSummarySet status="OK">
  <DbBuild>Build150407-2207m.3/DbBuild>
  <DocumentSummary>
    <Id>19650888</Id>
   <PubDate>2009 Aug 3</PubDate>
   <EPubDate>2009 Aug 3</EPubDate>
   <Source>BMC Microbiol</Source>
    <Authors>
      <Author>
       <Name>Cano V</Name>
        <AuthType>Author</AuthType>
        <ClusterID></ClusterID>
      </Author>
      <Author>
        <Name>Moranta D</Name>
        <AuthType>Author</AuthType>
```

Piping the document summary output to:

```
xtract -outline
```

will give an indented overview of the XML structure hierarchy:

```
DbBuild
DocumentSummary
Id
PubDate
EPubDate
Source
Authors
Author
Name
```

```
AuthType
ClusterID
Author
Name
```

The outline view presents a clear, uncluttered picture of the XML hierarchy that is useful in designing the appropriate command for actual data extraction. Copy and paste from the -outline output to xtract arguments can help avoid typographical errors. Thus:

```
esearch -db pubmed -query "Garber ED [AUTH] AND PNAS [JOUR]" |
elink -related |
efilter -query "mouse" |
efetch -format docsum |
xtract -pattern DocumentSummary -element Id SortFirstAuthor Title
```

returns the PubMed identifier (PMID), first author name, and article title:

```
19650888 Cano V Klebsiella pneumoniae triggers a cytotoxic ...

19262028 Suto J Metabolic consequence of congenital asplenia ...

19248821 Fukumoto N Hypoalgesic behaviors of P/Q-type voltage- ...

18822497 Trishin AV [Protective activity of secreted proteins of ...

18582214 Singh A Generation and characterization of monoclonal ...
```

Using "xtract -synopsis" instead of -outline show the full path to each element. Piping those results to "sort-uniq-count" (see below) produces a table of unique path counts.

# Processing Results with UNIX Utilities

A tab-delimited table can be processed by many UNIX utilities. For example:

```
esearch -db pubmed -query "Garber ED [AUTH] AND PNAS [JOUR]" |
elink -related |
efilter -query "mouse" |
efetch -format docsum |
xtract -pattern DocumentSummary -element Id SortFirstAuthor Title |
sort -t $'\t' -k 2,2f -k 3,3f
```

sorts the results of the previous example by author name and then (if there are multiple publications by the same author) alphabetically by title:

```
17474906 Benghezal M Inhibitors of bacterial virulence ...
19650888 Cano V Klebsiella pneumoniae triggers a cytotoxic ...
17102561 Chatterjee S How reliable are models for malaria vaccine ...
17371870 Clements A Secondary acylation of Klebsiella ...
17142396 Fresno S A second galacturonic acid transferase is ...
16735743 Fresno S The ionic interaction of Klebsiella ...
```

Rather than always having to retype a series of common post-processing instructions, frequently used combinations of UNIX commands can be placed in a function, stored in

an alias file (e.g., the user's .bash\_profile), and executed by name. (The following two functions are now included as scripts with the EDirect software.) For example:

```
WordAtATime() {
  sed 's/[^a-zA-Z0-9]/ /g; s/^ *//' |
  tr 'A-Z' 'a-z' |
  fmt -w 1
}
alias word-at-a-time='WordAtATime'

SortUniqCountRank() {
  sort -f |
  uniq -i -c |
  perl -pe 's/\s*(\d+)\s(.+)/$1\t$2/' |
  sort -t $'\t' -k 1,1nr -k 2f
}
alias sort-uniq-count-rank='SortUniqCountRank'
```

Titles can be passed to a pair of these UNIX alias commands:

```
esearch -db pubmed -query "Casadaban transposition immunity" |
elink -related |
efetch -format docsum |
xtract -pattern DocumentSummary -element Title |
word-at-a-time |
sort-uniq-count-rank
```

to generate a table of word occurrence counts, sorted by frequency:

```
296
     of
175
     the
114
   transposition
102 and
   in
94
    mu
93
83
    а
61
    dna
61
     tn3
55
    transposon
```

# **Output Format Customization**

The line break between -pattern objects can be overridden with -ret, and the tab character between fields can be replaced by -tab.

The -sep argument is used to distinguish multiple elements of the same type and control their separation independently of the -tab argument. For example:

```
esearch -db gene -query "deuteranopia" |
efetch -format xml |
xtract -pattern Entrezgene \
```

```
-element Gene-track_geneid Gene-ref_locus \
-sep "|" -element Gene-ref syn E
```

combines all synonyms for a gene into a single column, separated by vertical bars:

```
2652 OPN1MW CBD GCP GOP CBBM COD5 OPN1MW1
5956 OPN1LW CBP RCP ROP CBBM COD5
```

The -sep value also applies to unrelated -element items that are grouped with commas. Otherwise the -tab value delineates individual fields.

Groups or fields are preceded by the -pfx value and followed by the -sfx value, both of which are initially empty.

#### Pubmed Article XML Records

The PubmedArticle object (for -db pubmed) has a more detailed structure than the DocumentSummary:

```
esearch -db pubmed -query "tetrachromacy" |
efetch -format xml |
xtract -outline
```

More information is fielded, including author names, dates, and the abstract:

```
PubmedArticle
 MedlineCitation
    PMID
    DateCreated
      Year
      Month
      Day
    DateCompleted
      Year
      Month
      Day
    DateRevised
      Year
      Month
      Day
    Article
      Journal
        ISSN
        JournalIssue
          Volume
          Issue
          PubDate
            Year
            Month
            Day
        Title
        ISOAbbreviation
      ArticleTitle
```

```
Pagination
MedlinePgn
Abstract
AbstractText
CopyrightInformation
AuthorList
Author
LastName
ForeName
Initials
AffiliationInfo
Affiliation
Author
LastName
LastName
```

Using this information to craft a new xtract statement:

```
esearch -db pubmed -query "tetrachromacy" |
efetch -format xml |
xtract -pattern PubmedArticle -element MedlineCitation/PMID LastName
```

results in a table of all authors for each record:

23393278	Sabbah	Troje	Gray	Hawryshyn
20884587	Jordan	Deeb	Bosten	Mollon
18230593	Koshitaka	Kinoshita	Vorobyev	Arikawa
17685813	Wachtler	Doi	Lee	Sejnowski
16086150	Goldsmith	Butler		

(Note that "-element MedlineCitation/PMID" uses the "Parent/Child" construct to prevent the display of additional PMID items that may occur later in CommentsCorrections objects.)

The -first or -last arguments can be used instead of -element, if appropriate.

# **Exploration of XML Sets**

Individual PubmedArticle objects can be retrieved directly by efetch:

```
efetch -db pubmed -id 20643751 -format xml
```

The resulting XML has authors with separate fields for last name and initials:

Without being given any guidance about context, an -element statement with "Initials" and "LastName" arguments:

```
efetch -db pubmed -id 1413997,6301692,781293 -format xml | xtract -pattern PubmedArticle -element MedlineCitation/PMID \ -element Initials LastName
```

will explore the current record for each argument separately, and thus print all author initials followed by all author last names:

1413997	RK	CR	JS	Mortimer	Contopoulou	King
6301692	MA	NR	Krasnow	Cozzarelli		
781293	MJ	Casadaban				

Inserting a -block statement redirects data exploration to present each author one at a time. Subsequent -element statements only see the current author's values:

```
efetch -db pubmed -id 1413997,6301692,781293 -format xml |
xtract -pattern PubmedArticle -element MedlineCitation/PMID \
-block Author -element Initials LastName
```

which restores the correct association of initials and last name:

1413997	RK	Mortimer	CR	Contopoulou	JS	King
6301692	MA	Krasnow	NR	Cozzarelli		
781293	MJ	Casadaban				

Adding a -sep statement to replace the normal tab between group members, and using a comma to combine the two arguments ("Initials,LastName") into a group:

```
efetch -db pubmed -id 1413997,6301692,781293 -format xml |
xtract -pattern PubmedArticle -element MedlineCitation/PMID \
   -block Author -sep " " -element Initials,LastName
```

results in more desirable formatting of author names:

```
1413997 RK Mortimer CR Contopoulou JS King
6301692 MA Krasnow NR Cozzarelli
781293 MJ Casadaban
```

# **Exploring Separate XML Regions**

Multiple -block statements can be used in a single xtract to explore different areas of the XML. This limits element extraction to the desired subregions, and allows disambiguation of fields with identical names.

Combining independent fields with commas allows them to be treated as sets. The tab that normally separates these can be replaced with a -sep argument:

```
efetch -db pubmed -id 6092233,4640931,4296474 -format xml |
xtract -pattern PubmedArticle -element MedlineCitation/PMID \
   -block AuthorList -sep "/" -element LastName "#Author" \
   -block PubDate -sep " " -element Year,Month MedlineDate \
   -block DateCreated -sep "-" -element Year,Month,Day |
sort -t $'\t' -k 3,3n -k 2,2f
```

This generates a table that allows easy parsing of author last names, counts the number of authors present, and prints the date each record was published and the date it was entered into PubMed, sorting the results by author count:

```
      4296474
      Friedmann
      1
      1968 Apr
      1968-06-05

      4640931
      Tager/Steiner
      2
      1972 Dec
      1973-02-15

      6092233
      Calderon/Contopoulou/Mortimer
      3
      1984 Jul-Aug
      1984-12-13
```

(Note that the PubDate object can exist either in a structured form:

```
<PubDate>
  <Year>1968</Year>
  <Month>Apr</Month>
  <Day>25</Day>
</PubDate>
```

(with the Day field frequently absent), or in a string form:

```
<PubDate>
<MedlineDate>1984 Jul-Aug</MedlineDate>
</PubDate>
```

but would not contain a mixture of both types, so the directive:

```
-element Year, Month MedlineDate
```

will only contribute a single column to the output.)

# Nested Exploration of Subsets Within XML Sets

Medical Subject Headings (MeSH terms) in a record may be assigned subheadings:

Visiting each MeSH term with a -block statement, and adding a -subset statement within the -block, allows nested exploration of the subheadings for the current MeSH term:

```
efetch -db pubmed -id 6162838 -format xml |
xtract -pattern PubmedArticle -tab "" -element MedlineCitation/PMID \
    -block MeshHeading -pfx "\n" -tab "" -element DescriptorName \
    -subset QualifierName -pfx " / " -tab "" -element QualifierName
```

and creates a list of MeSH terms with associated subheadings:

```
6162838

Base Sequence

DNA, Recombinant

Escherichia coli / genetics
...

RNA, Messenger / genetics

Transcription, Genetic

beta-Galactosidase / genetics / metabolism
```

#### Selection of Attributes

The MeSH term and subheading fields actually have major topic attributes:

```
<MeshHeading>
    <DescriptorName MajorTopicYN="N">beta-Galactosidase</DescriptorName>
    <QualifierName MajorTopicYN="Y">genetics</QualifierName>
    <QualifierName MajorTopicYN="N">metabolism</QualifierName>
</MeshHeading>
...
```

that can be selected as "DescriptorName@MajorTopicYN" or "@MajorTopicYN":

```
efetch -db pubmed -id 6162838 -format xml |
xtract -pattern PubmedArticle -tab "" -element MedlineCitation/PMID \
    -block MeshHeading -pfx "\n" -sep "|" -tab "" \
        -element DescriptorName@MajorTopicYN,DescriptorName \
        -subset QualifierName -pfx " / " -sep "|" -tab "" \
        -element "@MajorTopicYN,QualifierName"
```

The major topic value is placed before each MeSH term or subheading:

```
6162838
N|Base Sequence
```

```
Y|DNA, Recombinant
N|Escherichia coli / N|genetics
...
N|RNA, Messenger / Y|genetics
N|Transcription, Genetic
N|beta-Galactosidase / Y|genetics / N|metabolism
```

The results can be processed by the UNIX stream editor "sed":

```
sed -e 's/N|//g' -e 's/Y|/*/g'
```

to display an asterisk for major ("starred" MeSH term) concepts:

```
6162838
Base Sequence
*DNA, Recombinant
Escherichia coli / genetics
...
RNA, Messenger / *genetics
Transcription, Genetic
beta-Galactosidase / *genetics / metabolism
```

# Recording Values in Variables

A value can be recorded in a variable and then displayed multiple times as needed. Variables are indicated by a hyphen followed by a string of capital letters or digits. The variable "-PMID" is referred to as "&PMID" in an -element argument. For example:

```
efetch -db pubmed -id 1413997,6301692,781293 -format xml |
xtract -pattern PubmedArticle -PMID MedlineCitation/PMID \
   -block Author -element "&PMID" \
   -sep " " -tab "\n" -element Initials,LastName
```

produces a list of authors, with the PMID in the first column of each row:

```
1413997 RK Mortimer
1413997 CR Contopoulou
1413997 JS King
6301692 MA Krasnow
6301692 NR Cozzarelli
781293 MJ Casadaban
```

#### Variable Initialization

Variables can be initialized with a literal value in parentheses:

```
efetch -db pubmed -id 1413997,6301692,781293 -format xml |
xtract -pattern PubmedArticle -element MedlineCitation/PMID \
   -block Author -sep " " -tab "" \
   -element "&COM" Initials,LastName -COM "(, )"
```

This can be used as a placeholder to prevent missing data from shifting columns in a table, or to have additional control over output formatting:

```
1413997 RK Mortimer, CR Contopoulou, JS King
6301692 MA Krasnow, NR Cozzarelli
781293 MJ Casadaban
```

All variables are reset when the next record is processed.

# **Conditional Processing**

Xtract provides -if and -unless arguments that filter by element name or name plus data value. For example:

```
esearch -db pubmed -query "Cozzarelli NR [AUTH]" |
efetch -format xml |
xtract -pattern PubmedArticle -if "#Author" -eq 3 \
   -block Author -if LastName -is-not Cozzarelli \
    -sep ", " -tab "\n" -element LastName, Initials |
sort | uniq
```

will select papers with exactly 3 authors and print the coauthor names:

```
Ackerman, RS
Adams, DE
Alexandrov, AI
Arimondo, PB
Bauer, WR
```

Multiple conditions are specified with -and and -or commands:

```
-if @score -equals 1 -or @score -starts-with 0.9
```

The -else command can supply alternative -element or -lbl instructions to be run if the condition is not satisfied:

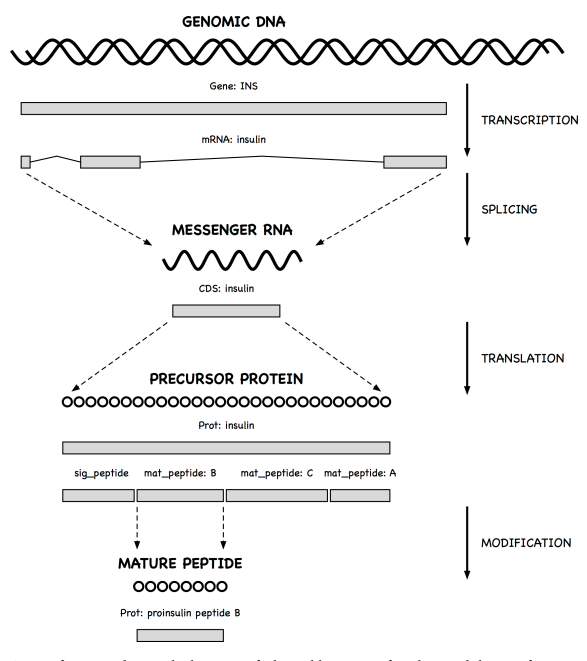
```
-if MapLocation -element MapLocation -else -lbl "\-"
```

Parallel -if and -unless statements can be used to provide a more complex response to alternative conditions that includes nested exploration.

# Sequence Records

# NCBI Data Model for Sequence Records

The NCBI represents sequence records in a data model that is based on the central dogma of molecular biology. Sequences, including genomic DNA, messenger RNAs, and protein products, are "instantiated" with the actual sequence letters, and are assigned identifiers (e.g., accession numbers) for reference. Features carry information about the biology of a given region, with a location that refers to specific intervals on a particular sequence. Some features may also point to the product sequence of a particular transformation.



A gene feature indicates the location of a heritable region of nucleic acid that confers a measurable phenotype. An mRNA feature on genomic DNA represents the exonic and untranslated regions of the message that remain after transcription and splicing. A coding region (CDS) feature has a product reference to the translated protein.

Since messenger RNA sequences are not always submitted with a genomic region, CDS features (which model the travel of ribosomes on transcript molecules) are traditionally annotated on the genomic sequence, with locations that encode the exonic intervals.

Features display specific biological annotation in qualifiers. For example, the name of a gene is shown in the /gene qualifier. A qualifier can be dynamically generated from

underlying data for the convenience of the user. Thus, the sequence of a mature peptide may be extracted from the mat\_peptide feature's location on the precursor protein and displayed in a /peptide qualifier, even if a mature peptide is not instantiated.

# Sequence Records in INSDSeq XML

Sequence records can be retrieved in an XML version of the GenBank or GenPept flatfile. The query:

efetch -db protein -id 26418308,26418074 -format gpc

```
returns a set of INSDSeq objects:
  <INSDSet>
    <INSDSeq>
      <INSDSeq_locus>AAN78128</INSDSeq_locus>
      <INSDSeq_length>17</INSDSeq_length>
      <INSDSeq_moltype>AA</INSDSeq_moltype>
      <INSDSeq_topology>linear</INSDSeq_topology>
      <INSDSeq_division>INV</INSDSeq_division>
      <INSDSeq_update-date>03-JAN-2003</INSDSeq_update-date>
      <INSDSeq_create-date>10-DEC-2002</INSDSeq_create-date>
      <INSDSeq_definition>alpha-conotoxin ImI precursor, partial [Conus
         imperialis]</INSDSeq_definition>
      <INSDSeq_primary-accession>AAN78128</INSDSeq_primary-accession>
      <INSDSeq_accession-version>AAN78128.1</INSDSeq_accession-version>
      <INSDSeq_other-seqids>
        <INSDSeqid>gb|AAN78128.1|</INSDSeqid>
        <INSDSeqid>gi|26418308</INSDSeqid>
      </INSDSeq_other-seqids>
      <INSDSeq_source>Conus imperialis</INSDSeq_source>
      <INSDSeq_organism>Conus imperialis</INSDSeq_organism>
      <INSDSeq_taxonomy>Eukaryota; Metazoa; Lophotrochozoa; Mollusca;
         Gastropoda; Caenogastropoda; Hypsogastropoda; Neogastropoda;
         Conoidea; Conidae; Conus</INSDSeq_taxonomy>
      <INSDSeq_references>
        <INSDReference>
```

# INSDSeq XML presents biological features and qualifiers (shown here in GenPept format):

```
FEATURES

source

1..17

/organism="Conus imperialis"

/db_xref="taxon:35631"

/country="Philippines"

Protein

<1..17

/product="alpha-conotoxin ImI precursor"

mat_peptide

5..16

/product="alpha-conotoxin ImI"

/note="the C-terminal glycine of the precursor is post translationally removed"
```

```
/calculated_mol_wt=1357
/peptide="GCCSDPRCAWRC"

1..17
/coded_by="AY159318.1:<1..54"
/note="nAChR antagonist"
```

in a structured feature table:

```
<INSDFeature>
 <INSDFeature_key>mat_peptide</INSDFeature_key>
  <INSDFeature_location>5..16</INSDFeature_location>
  <INSDFeature intervals>
    <INSDInterval>
      <INSDInterval_from>5</INSDInterval_from>
      <INSDInterval_to>16</INSDInterval_to>
      <INSDInterval_accession>AAN78128.1</INSDInterval_accession>
    </INSDInterval>
  </INSDFeature_intervals>
  <INSDFeature_quals>
    <INSDQualifier>
      <INSDQualifier_name>product</INSDQualifier_name>
      <INSDQualifier_value>alpha-conotoxin ImI</INSDQualifier_value>
    </INSDOualifier>
    <INSDQualifier>
      <INSDQualifier_name>note</INSDQualifier_name>
      <INSDQualifier_value>the C-terminal glycine of the precursor is
         post translationally removed</INSDQualifier_value>
    </INSDQualifier>
    <INSDQualifier>
      <INSDQualifier_name>calculated_mol_wt</INSDQualifier_name>
      <INSDQualifier_value>1357</INSDQualifier_value>
    </INSDOualifier>
    <INSDQualifier>
      <INSDQualifier_name>peptide</INSDQualifier_name>
      <INSDQualifier_value>GCCSDPRCAWRC</INSDQualifier_value>
    </INSDQualifier>
  </INSDFeature_quals>
</INSDFeature>
```

Feature and qualifier names are indicated in data values, not XML element tags, and require -if and -equals to select the desired object and content. The xtract -insd argument simplifies this process, as shown below.

# Generating Qualifier Extraction Commands

Because obtaining specific qualifier values from INSDSeq XML is somewhat more complex than previous cases, the xtract -insd argument can be used to generate extraction instructions.

Running xtract -insd in an isolated command prints a new xtract statement that can then be copied, edited if necessary, and pasted into other queries. Running the -insd command within a multi-step pipe dynamically executes the constructed query.

Providing an optional (complete/partial) location indication, a feature key, and then one or more qualifier names:

```
xtract -insd complete mat_peptide "%peptide" product peptide
```

creates a new xtract statement that will produce a table of qualifier values from mature peptide features with complete locations. The statement starts with instructions to record the accession and find features of the indicated type:

```
xtract -pattern INSDSeq -ACCN INSDSeq_accession-version \
  -group INSDFeature -if INSDFeature_key -equals mat_peptide \
  -unless INSDFeature_partial5 -or INSDFeature_partial3 \
  -clr -pfx "\n" -element "&ACCN"
```

Each qualifier then generates custom extraction code that is appended to the growing query. For example:

```
-block INSDQualifier \
  -if INSDQualifier_name -equals peptide \
  -element INSDQualifier value
```

Incorporating the xtract -insd command in a query for marine snail venom peptides:

```
esearch -db pubmed -query "conotoxin" |
elink -target protein |
efilter -query "mat_peptide [FKEY]" |
efetch -format gpc |
xtract -insd complete mat_peptide "%peptide" product peptide
```

produces a table with columns for accession number, calculated peptide length, product name, and peptide sequence:

```
AGO59814.1 32 del13b conotoxin DCPTSCPTTCANGWECCKGYPCVRQHCSGCNH
AAO33169.1 16 alpha-conotoxin GIC GCCSHPACAGNNQHIC
ADB65788.1 20 conotoxin Cal 16 LEMQGCVCNANAKFCCGEGR
AAN78128.1 12 alpha-conotoxin ImI GCCSDPRCAWRC
AAF23167.1 31 BeTX toxin CRAEGTYCENDSQCCLNECCWGGCGHPCRHP
ADB65789.1 20 conotoxin Cal 16 LEMQGCVCNANAKFCCGEGR
AAN78279.1 21 conotoxin Vx-II WIDPSHYCCCGGGCTDDCVNC
ABW16858.1 15 marmophin DWEYHAHPKPNSFWT
```

Piping the results to a series of UNIX commands:

```
grep -i conotoxin |
awk -F '\t' -v 'OFS=\t' '{if ( 10 <= $2 && $2 <= 30 ) print}' |
sort -t $'\t' -u -k 3,4 |
sort -t $'\t' -k 2,2n -k 3,3f |
cut -f 1,3- |
column -s $'\t' -t</pre>
```

filters by product name, limits the results to a specified range of peptide lengths, removes redundant accessions, sorts the table by peptide length, deletes the length column, and aligns the columns for cleaner printing:

```
AAN78128.1 alpha-conotoxin ImI
                                              GCCSDPRCAWRC
AAN78127.1 alpha-conotoxin ImII
                                              ACCSDRRCRWRC
ADB43130.1 conotoxin Cal la
                                            KCCKRHHGCHPCGRK
ADB43131.1 conotoxin Cal 1b
                                            LCCKRHHGCHPCGRT
AAO33169.1 alpha-conotoxin GIC
                                            GCCSHPACAGNNQHIC
ADB43128.1 conotoxin Cal 5.1
                                            DPAPCCOHPIETCCRR
                                        PECCSHPACNVDHPEICR
AAD31913.1 alpha A conotoxin Tx2
ADB43129.1 conotoxin Cal 5.2
                                            MIQRSQCCAVKKNCCHVG
ADD97803.1 conotoxin Cal 1.2
                                              AGCCPTIMYKTGACRTNRCR
ADDUDINES.1 CONOTOXIN Cal 16 LEMQGCVCNANAKFCCGEGR
AAD31912.1 alpha A conotoxin Tx1 PECCSDPRCNSSHPELCGGRR
AAN78279.1 CONOTOXIN We-TT
AAN78279.1 conotoxin Vx-II
                                              WIDPSHYCCCGGGCTDDCVNC
ADB43125.1 conotoxin Cal 14.2
                                              GCPADCPNTCDSSNKCSPGFPG
ADD97802.1 conotoxin Cal 6.4
                                              GCWLCLGPNACCRGSVCHDYCPR
CAH64846.1 four-loop conotoxin
                                              CRPSGSPCGVTSICCGRCSRGKCT
AAD31915.1 O-superfamily conotoxin TxO2
                                              CYDSGTSCNTGNQCCSGWCIFVCL
AAD31916.1 O-superfamily conotoxin TxO3
                                              CYDGGTSCDSGIOCCSGWCIFVCF
AAD31920.1 omega conotoxin SVIA mutant 1 CRPSGSPCGVTSICCGRCYRGKCT
AAD31921.1 omega conotoxin SVIA mutant 2 CRPSGSPCGVTSICCGRCSRGKCT
ABE27010.1 conotoxin fe14.1
                                              SPGSTICKMACRTGNGHKYPFCNCR
ABE27011.1 conotoxin fe14.2
                                              SSGSTVCKMMCRLGYGHLYPSCGCR
                                       SSGSTVCKMMCRLGYGHLYPSCGCR
GPGSAICNMACRLGQGHMYPFCNCN
ABE27007.1 conotoxin p114.1
ABE27008.1 conotoxin p114.2
ABE27009.1 conotoxin p114.3
                                            GPGSAICNMACRLEHGHLYPFCHCR
                                              GPGSAICNMACRLEHGHLYPFCNCD
```

#### For records where a particular qualifier is missing:

```
esearch -db protein -query "RAG1 [GENE] AND Mus musculus [ORGN]" |
efetch -format gpc |
xtract -insd source organism strain |
sort -t $'\t' -u -k 2,3
```

#### a dash is inserted as a placeholder:

P15919.2	Mus musculus	_
AA061776.1	Mus musculus	129/Sv
NP_033045.2	Mus musculus	C57BL/6
XP_006499075.1	Mus musculus	C57BL/6J
EDL27655.1	Mus musculus	mixed
BAD69530.1	Mus musculus castaneus	_
BAD69531.1	Mus musculus domesticus	BALB/c
BAD69532.1	Mus musculus molossinus	MOA

# Sequence Coordinates

#### Gene Positions

An understanding of sequence coordinate conventions is necessary in order to use gene positions to retrieve the corresponding chromosome subregion with efetch or with the UCSC browser.

Sequence records displayed in GenBank or GenPept formats use a "one-based" coordinate system, with sequence position numbers starting at "1":

```
1 catgccattc gttgagttgg aaacaaactt gccggctagc cgcatacccg cggggctgga 61 gaaccggctg tgtgcggcca cagccaccat cctggacaaa cccgaagacg tgagtgaggg 121 tcggcgagaa cttgtgggct agggtcggac ctcccaatga cccgttccca tccccaggga 181 ccccactccc ctggtaacct ctgaccttcc gtgtcctatc ctcccttcct agatcccttc
```

Under this convention, positions refer to the sequence letters themselves:

```
C A T G C C A T T C
1 2 3 4 5 6 7 8 9 10
```

and the position of the last base or residue is equal to the length of the sequence. The ATG initiation codon above is at positions 2 through 4, inclusive.

For computer programs, however, using "zero-based" coordinates can simplify the arithmetic used for calculations on sequence positions. The ATG codon in the 0-based representation is at positions 1 through 3. (The UCSC browser uses a hybrid, half-open representation, where the start position is 0-based and the stop position is 1-based.)

Software at NCBI will typically convert positions to 0-based coordinates upon input, perform whatever calculations are desired, and then convert the results to a 1-based representation for display. These transformations are done by simply subtracting 1 from the 1-based value or adding 1 to the 0-based value.

#### Coordinate Conversions

Retrieving the docsum for a particular gene:

```
esearch -db gene -query "BRCA2 [GENE] AND human [ORGN]" | efetch -format docsum
```

returns the chromosomal position of that gene in 0-based coordinates:

```
</GenomicInfoType>
```

Piping the document summary to an xtract command:

xtract -pattern GenomicInfoType -element ChrAccVer ChrStart ChrStop obtains the accession and 0-based coordinate values:

```
NC_000013.11 32315479 32399671
```

EFetch has -seq\_start and -seq\_stop arguments to retrieve a gene segment, but these expect the sequence subrange to be in 1-based coordinates.

To address this problem, two additional efetch arguments, -chr\_start and -chr\_stop, allow direct use of the 0-based coordinates:

```
efetch -db nuccore -format gb -id NC_000013.11 \
-chr_start 32315479 -chr_stop 32399671
```

and eliminate the need for writing a UNIX shell command to increment the two values.

Xtract has numeric extraction commands to assist with coordinate conversion. Selecting fields with an -inc argument:

```
xtract -pattern GenomicInfoType -element ChrAccVer -inc ChrStart ChrStop
```

obtains the accession and 0-based coordinates, then increments the positions to produce 1-based values:

```
NC_000013.11 32315480 32399672
```

EDirect knows the policies for sequence positions in all relevant Entrez databases (e.g., gene, snp, dbvar), and provides additional shortcuts for converting these to other conventions. For example:

```
xtract -pattern GenomicInfoType -element ChrAccVer -1-based ChrStart ChrStop
```

understands that gene ChrStart and ChrStop fields are 0-based, sees that the desired output is 1-based, and translates the command to convert coordinates using the -inc argument. Similarly:

```
-element ChrAccVer -ucsc-based ChrStart ChrStop
```

leaves the 0-based start value unchanged but increments the original stop value to produce the half-open form that can be passed to the UCSC browser:

```
NC 000013.11 32315479 32399672
```

# Complex Objects

# Heterogeneous Data

XML objects can contain a heterogeneous mix of components. For example:

```
efetch -db pubmed -id 21433338,17247418 -format xml
```

returns a mixture of book and journal records:

The "Parent/\*" construct is used to visit the individual components, even though they may have different names. Piping the XML output to:

```
xtract -pattern "PubmedArticleSet/*" -element "*"
```

separately prints the entirety of each XML component:

```
<PubmedBookArticle><BookDocument> ... </PubmedBookData></PubmedBookArticle>
<PubmedArticle><MedlineCitation> ... </PubmedData></PubmedArticle>
```

Use of the "Parent/Child" construct can isolate objects of the same name that differ by their location in the XML hierarchy. For example:

```
efetch -db pubmed -id 21433338,17247418 -format xml |
xtract -pattern "PubmedArticleSet/*" \
  -group "BookDocument/AuthorList" -tab "\n" -element LastName \
  -group "Book/AuthorList" -tab "\n" -element LastName \
  -group "Article/AuthorList" -tab "\n" -element LastName
```

writes separate lines for book/chapter authors, book editors, and article authors:

```
Fauci Desrosiers
Coffin Hughes Varmus
Lederberg Cavalli Lederberg
```

Simply exploring with individual arguments:

```
-group BookDocument -block AuthorList -element LastName
```

would visit the editors (at BookDocument/Book/AuthorList) as well as the authors (at BookDocument/AuthorList), and print names in order of appearance in the XML:

```
Coffin Hughes Varmus Fauci Desrosiers
```

(In this particular example the book author lists could be distinguished by using -if "@Type" -equals authors or -if "@Type" -equals editors, but exploring by "Parent/Child" is a general position-based approach.)

# **Recursive Definitions**

Certain XML objects returned by efetch are recursively defined, including Taxon in TaxaSet (-db taxonomy) and Gene-commentary in Entrezgene (-db gene). Thus, they can have nested objects with the same XML tag.

Retrieving a set of taxonomy records:

```
efetch -db taxonomy -id 9606,7227 -format xml
```

produces XML with nested Taxon objects (marked below with line references) for each rank in the taxonomic lineage:

```
<TaxaSet>
1
      <Taxon>
        <TaxId>9606</TaxId>
        <ScientificName>Homo sapiens</ScientificName>
       <LineageEx>
2
          <Taxon>
            <TaxId>131567</TaxId>
            <ScientificName>cellular organisms</ScientificName>
            <Rank>no rank</Rank>
3
          </Taxon>
          <Taxon>
            <TaxId>2759</TaxId>
            <ScientificName>Eukaryota</ScientificName>
            <Rank>superkingdom</Rank>
5
          </Taxon>
          . . .
        </LineageEx>
6
      </Taxon>
      <Taxon>
        <TaxId>7227</TaxId>
        <ScientificName>Drosophila melanogaster</ScientificName>
      </Taxon>
    </TaxaSet>
```

Xtract tracks XML object nesting to determine that the <Taxon> start tag on line 1 is actually closed by the </Taxon> stop tag on line 6, and not by the first </Taxon> encountered on line 3.

When a recursive object is given to an exploration command, selection of data using the element command:

```
efetch -db taxonomy -id 9606,7227,10090 -format xml |
xtract -pattern Taxon \
  -element TaxId ScientificName GenbankCommonName Division
```

does not examine fields in the internal objects, and returns information only for the main entries:

```
9606 Homo sapiens human Primates
7227 Drosophila melanogaster fruit fly Invertebrates
10090 Mus musculus house mouse Rodents
```

The "\*/Child" construct will skip past the outer start tag:

```
efetch -db taxonomy -id 9606,7227,10090 -format xml |
xtract -pattern Taxon -block "*/Taxon" \
   -tab "\n" -element TaxId,ScientificName
```

to visit the next level of nested objects individually:

```
131567 cellular organisms
2759 Eukaryota
33154 Opisthokonta
```

Recursive objects can be fully explored with a double-star-slash prefix:

```
esearch -db gene -query "DMD [GENE] AND human [ORGN]" |
efetch -format xml |
xtract -pattern Entrezgene -block "**/Gene-commentary" \
   -tab "\n" -element Gene-commentary_type@value,Gene-commentary_accession
```

which visits every child object regardless of nesting depth:

```
genomic NC_000023
mRNA XM_006724469
peptide XP_006724532
mRNA XM_011545467
peptide XP_011543769
```

# **Advanced Topics**

# Storing Common Phrases in Alias Files

Long or complicated search phrases can be saved in a file to avoid having to retype (or copy and paste) the full text for each query. Each line of the file has a shortcut keyword, a tab character, and the expanded search term. Shortcuts are referenced by placing them in parentheses after prefixing with a pound ("#") sign.

For example, given a file named "q\_aliases" containing:

```
jour_filt [MULT] AND ncbijournals [FILT]
  trans_imm (transposition OR target) immunity

the esearch line in:
  esearch -alias q_aliases -db nlmcatalog -query "Science (#jour_filt)" |
  efetch -format docsum |
```

```
xtract -pattern DocumentSummary -element ISOAbbreviation \
  -subset ISSNInfo -sep "|" -element issn,issntype
```

#### will be expanded to:

```
esearch -db nlmcatalog -query "Science [MULT] AND ncbijournals [FILT]" with the query producing:
```

```
J. Zhejiang Univ. Sci. 1009-3095|Print 1009-3095|Linking
Science (80-) 0193-4511|Print 0193-4511|Linking
Science 0036-8075|Print 1095-9203|Electronic ...
```

An alias file can also be read in a separate instruction at the beginning of a pipeline or script:

```
eproxy -alias q aliases
```

For maximum flexibility, separate eproxy commands can be piped together to load multiple shortcut files, as long as the shortcut strings are all unique.

# Additional EDirect Options

ESearch and EFilter can be given a -sort argument to specify the order of results when the records are retrieved:

```
esearch -db pubmed -query "opsin gene conversion" -sort "last author" | efetch -format docsum | xtract -pattern DocumentSummary -element Id LastAuthor PubDate Title
```

ELink can return links to the citation list using "-name pubmed\_pubmed\_citedin", but only for publications with full text deposited in PubMed Central (PMC). For example, the query:

```
esearch -db pubmed -query "Beadle GW [AUTH]" | elink -related -name pubmed_pubmed_citedin | efetch -format docsum | xtract -pattern Author -element Name | sort-uniq-count-rank | head -n 10
```

produces a ranked list of the ten most cited authors:

Beadle GW
Ephrussi B
Glass NL
Hawley RS
Mitchell MB
PERKINS DD
Tatum EL
Mitchell HK
YANOFSKY C
Langley CH

Similarly, "-name pubmed\_pubmed\_refs" returns an article's reference list, again for publications deposited in PMC.

ELink has several command modes, and these can be specified with the -cmd argument. When not using the default "neighbor\_history" command, elink will return an eLinkResult XML object, with the links for each UID presented in separate blocks. For example:

```
esearch -db pubmed -query "Hoffmann PC [AUTH] AND dopamine [MAJR]" |
elink -related -cmd neighbor |
xtract -pattern LinkSetDb -element Id
```

will show the original PMID in the first column and related article PMIDs in subsequent columns:

```
      1504781
      11754494
      3815119
      1684029
      14614914
      12128255
      ...

      1684029
      3815119
      1504781
      8097798
      17161385
      14755628
      ...

      2572612
      2903614
      6152036
      2905789
      9483560
      1352865
      ...
```

When the elink command "prlinks" is used with "ref" mode, it can obtain HTML containing or referencing full text articles directly from the publishers. The UNIX "xargs" command calls elink separately for each identifier:

```
epost -db pubmed -id 22966225,19880848 |
efilter -query "freetext [FILT]" |
efetch -format uid |
xargs -n 1 elink -db pubmed -cmd prlinks -mode ref -http get -id
```

The elink -batch flag will bypass the Entrez history mechanism for large queries.

# **Xtract Special Topics**

Self-closing tags of the standard form:

```
<Na-strand/>
```

or alternative form:

```
<Na-strand></Na-strand>
```

have no text content and thus cannot be selected with an -element command. If the tag contains an attribute:

```
<Seq-interval_strand>
  <Na-strand value="plus"/>
</Seq-interval_strand>
```

it can be selected by matching on the specified value:

```
-group Seq-interval_strand \
  -block Seq-interval_strand -if Na-strand@value -equals plus -lbl "+" \
  -block Seq-interval_strand -if Na-strand@value -equals minus -lbl "-"
```

The -pattern, -group, -block, and -subset commands provide a nested hierarchy of loop organizers for exploration of XML objects. Each pattern can contain multiple groups, each group can encompass multiple blocks, and each block can have multiple subsets.

Use of different argument names allows a linear representation of loop nesting, and provides sufficient flexibility to identify and extract arbitrary data from XML records in Entrez.

Sketching in pseudo code can clarify relative nesting levels. The extraction command:

```
xtract -pattern PubmedArticle \
  -block Author -element Initials,LastName \
  -block MeshHeading \
    -if QualifierName \
    -element DescriptorName \
    -subset QualifierName -element QualifierName
```

could be represented as a computer program in pseudo code by:

```
for each Pubmed record {
  for each Author {
    print Initials LastName
  }
  for each MeSH term {
    if Subheadings are present {
      print Term Name
      for each Subheading {
        print Subheading Name
      }
    }
  }
}
```

Extra arguments (-division, -branch, -section, and -unit) are held in reserve to provide additional levels of organization, should the need arise in the future for processing complex, deeply-nested XML data. The full set of commands, in order of rank, are:

```
-pattern
-division
-group
-branch
-block
-section
-subset
-unit
```

Starting xtract exploration with -block, and expanding with -group and -subset, leaves additional level names that can be used wherever needed without having to redesign the entire command.

# **Querying External Web Services**

The EDirect nquire function can be used to obtain data from an arbitrary URL. Queries are built up from command-line arguments. For example:

```
nquire -url "https://eutils.ncbi.nlm.nih.gov/entrez/eutils/esearch.fcgi" \
  -db pubmed -term insulin
```

reads the URL and then tag/value pairs to generate an E-utilities query:

```
https://eutils. ... .gov/entrez/eutils/esearch.fcgi?db=pubmed&term=insulin
```

Paths can be separated into components, which are combined with slashes, so:

```
-url https://eutils.ncbi.nlm.nih.gov entrez/eutils efetch.fcgi
```

is converted to:

```
https://eutils.ncbi.nlm.nih.gov/entrez/eutils/efetch.fcgi
```

Multiple values between tags are combined with commas. Thus:

```
-db nuccore -id U54469 V00328 -rettype fasta
```

is transformed into:

```
db=nuccore&id=U54469,V00328&rettype=fasta
```

A value that starts with a hyphen (or minus sign) can be distinguished from a tag by prefixing it with a backslash, so:

```
nquire -url http://api.geonames.org/countryCode -lat 41.796 -lng "\-87.577" will be sent as:
```

```
http://api.geonames.org/countryCode?lat=41.796&lng=-87.577
```

and will return "US" for coordinates within Chicago, which has a negative (western hemisphere) longitude value.

The -alias argument can read a file of shortcut keywords and URL aliases. The following aliases are always available:

```
ncbi_url https://www.ncbi.nlm.nih.gov
eutils_url https://eutils.ncbi.nlm.nih.gov/entrez/eutils
```

so the command:

```
nquire -url "(#eutils_url)" esearch.fcgi \
  -db gds -term "GSE22309 [ACCN] AND gse [ETYP]" -retmax 200
```

will run an ESearch query and return an eSearchResult XML object.

Raw XML with inconsistent line-wrapping and indentation can be reformatted for easier visual inspection of the data structure and content by piping it through:

```
xtract -format
```

#### **Automation**

# **Entrez Direct Commands Within Scripts**

Taking an adventurous plunge into the world of programming, a shell script can be written when each output line of one step needs to be processed independently, instead of output being piped in its entirety to the next command. (The simplest shell script is merely a copy of a set of commands that are typed into the terminal for execution.)

In scripts, variables can be set to the results of a command by enclosing the statements in backtick ("`") characters. The variable name is prefixed by a dollar sign ("\$") to use its value as an argument in another command. Comments start with a pound sign ("#") and are ignored. Quotation marks within quoted strings are entered by "escaping" with a backslash ("\"). Subroutines can be used to collect common code or simplify the organization of the script.

For example, executing a script file containing:

```
#!/bin/bash -norc
parse_fields() {
  echo "$1" |
 xtract -pattern Field \
    -pfx "[" -sfx "]" -element Name \
    -pfx "" -sfx "" -element FullName Description |
 sort -t $'\t' -k 2,2f | column -s $'\t' -t
dbs=`einfo -dbs | sort`
for db in $dbs
  eix=`einfo -db $db`
  flds=`parse_fields "$eix"`
  echo "$db"
  echo ""
  echo "$flds"
  echo ""
 sleep 1
done
```

will obtain the list of Entrez databases:

```
annotinfo
assembly
bioproject
...
```

and then return the abbreviations, names, and descriptions of indexed search fields, for each individual database:

```
mesh

[ALL] All Fields All terms from all searchable fields
[FILT] Filter Limits the records
[MESH] MeSH Terms MeSH Terms
[MHUI] MeSH Unique ID NLM MeSH Browser Unique ID
[MULT] Multi Multi
[PREV] Previous Indexing Previous Indexing
[TYPE] Record Type Record type
[REG] Registry Number Registry Number
[NOTE] Scope Note Scope Note
[ALSO] See Also See Also
[SUBS] Substance Name Substance Name
[WORD] Text Word Free text
[TN] Tree Number Unique number assigned to publication
```

The shell script command:

```
sleep 1
```

adds a one second delay between steps in a loop, and can be used to help prevent overuse of the Entrez servers by advanced scripts.

# Xargs/Sh Loop

Writing a script to loop through data can sometimes be avoided by creative use of the UNIX xargs and sh commands. Within the "sh -c" command string, the last name and initials arguments (passed in pairs by "xargs -n 2") are substituted at the "\$0" and "\$1" variables. All of the commands in the sh string are run separately on each name:

```
echo "Garber ED Casadaban MJ Mortimer RK" | xargs -n 2 sh -c 'esearch -db pubmed -query "$0 $1 [AUTH]" | xtract -pattern ENTREZ_DIRECT -lbl "$1 $0" -element Count'
```

This produces PubMed article counts for each author:

```
ED Garber 35
MJ Casadaban 46
RK Mortimer 85
```

# While Loop

A "while" loop can also be used to independently process lines of data. Given a file "organisms.txt" containing genus-species names, the UNIX "cat" command:

```
cat organisms.txt
```

#### writes the contents of the file:

```
Arabidopsis thaliana
Caenorhabditis elegans
Danio rerio
Drosophila melanogaster
Escherichia coli
Homo sapiens
Mus musculus
Saccharomyces cerevisiae
```

### This can be piped to a loop that reads one line at a time:

```
while read org
do
    esearch -db taxonomy -query "$org [LNGE] AND family [RANK]" < /dev/null |
    efetch -format docsum |
    xtract -pattern DocumentSummary -lbl "$org" \
        -element ScientificName Division
done</pre>
```

### looking up the taxonomic family name and BLAST division for each organism:

Arabidopsis thaliana	Brassicaceae	eudicots
Caenorhabditis elegans	Rhabditidae	nematodes
Danio rerio	Cyprinidae	bony fishes
Drosophila melanogaster	Drosophilidae	flies
Escherichia coli	Enterobacteriaceae	enterobacteria
Homo sapiens	Hominidae	primates
Mus musculus	Muridae	rodents
Saccharomyces cerevisiae	Saccharomycetaceae	ascomycetes

(The "< /dev/null" input redirection construct prevents esearch from "draining" the remaining lines from stdin.)

## For Loop

The same results can be obtained with organism names embedded in a "for" loop:

```
for org in \
   "Arabidopsis thaliana" \
   "Caenorhabditis elegans" \
   "Danio rerio" \
   "Drosophila melanogaster" \
   "Escherichia coli" \
   "Homo sapiens" \
   "Mus musculus" \
   "Saccharomyces cerevisiae"

do
   esearch -db taxonomy -query "$org [LNGE] AND family [RANK]" |
   efetch -format docsum |
   xtract -pattern DocumentSummary -lbl "$org" \
        -element ScientificName Division
done
```

### File Exploration

A for loop can also be used to explore the computer's file system:

```
for i in *
do
   if [ -f "$i" ]
   then
    echo $(basename "$i")
   fi
done
```

visiting each file within the current directory. The asterisk ("\*") character indicates all files, and can be replaced by any pattern (e.g., "\*.txt") to limit the file search. The if statement "-f" operator can be changed to "-d" to find directories instead of files, and "-s" selects files with size greater than zero.

### Processing in Groups

Because of technical limits in the Entrez link server, it may be necessary to perform an elink operation on a large set of records by using a function that splits unique identifiers or sequence accession numbers into smaller groups:

```
JoinIntoGroupsOf() {
   xargs -n "$@" echo |
   sed 's/ /,/g'
}
alias join-into-group-of='JoinIntoGroupsOf'
```

The following example will process sequence records in groups of 200 accessions at a time:

```
efetch -format acc |
join-into-groups-of 200 |
xargs -n 1 sh -c 'epost -db nuccore -format acc -id "$0" |
elink -target pubmed |
efetch -format abstract'
```

# **Examples**

Additional examples of using EDirect to answer ad hoc questions are shown in this section.

## **Author Frequency**

Who are the most prolific authors on rattlesnake phospholipase?

```
esearch -db pubmed -query \
   "crotalid venoms [MAJR] AND phospholipase [TIAB]" |
efetch -format xml |
xtract -pattern PubmedArticle \
```

```
-block Author -sep " " -tab "\n" -element LastName, Initials
sort-uniq-count-rank
This search produces:
```

```
74 Lomonte B
73 Gutiérrez JM
49 Soares AM
48 Marangoni S
43 Giglio JR
39 Bon C
```

### Publication Distribution

When were the most papers about Legionnaires disease published?

```
esearch -db pubmed -query "legionnaires disease [TITL]" |
efetch -format docsum |
xtract -pattern DocumentSummary -element PubDate |
cut -c 1-4 |
sort-uniq-count-rank
```

reports the number of selected papers per year:

```
173 1979
102 1980
96 1978
92 1981
66 1983
```

### **Treatment Locations**

What is the geographic distribution of sepsis treatment studies?

```
esearch -db pubmed -query \
   "sepsis/therapy [MESH] AND geographic locations [MESH]" |
efetch -format xml |
xtract -pattern PubmedArticle \
   -block MeshHeading -if DescriptorName@Type -equals Geographic \
   -tab "\n" -element DescriptorName |
sort-uniq-count-rank
```

returns the number of articles ranked by country (or region) of study:

```
567 United States
207 Spain
176 Great Britain
156 Germany
123 India
118 Europe
113 France
100 Taiwan
```

```
89 Japan83 Thailand75 Italy74 England
```

# Research History

What is the historic pattern of publication on diphtheria, pertussis, and tetanus?

```
#!/bin/bash
result=""
for disease in diphtheria pertussis tetanus
do
    current=`for (( yr = 2010; yr >= 1900; yr -= 10 ))
    do
        esearch -db pubmed -query "$disease [TITL] AND $yr:$((yr+9)) [PDAT]" |
        xtract -pattern ENTREZ_DIRECT -lbl "${yr}s" -element Count
    done`
    heading=`echo -e "${disease:0:4}" | tr [a-z] [A-Z]`
    current=`echo -e "Years\t$heading\n----\t---\n$current"`
    if [ -n "$result" ]
    then
        result=`join -t $'\t' <(echo "$result") <(echo "$current")`
    else
        result=$current
    fi
done
echo "$result"</pre>
```

gives per-decade counts of relevant papers for each disease:

```
DIPH
Years
                   PERT
                              TETA
          ----
____
                    ----
                              ----

      2010s
      577
      1708
      914

      2000s
      892
      1966
      1344

      1990s
      1150
      2661
      1615

1980s 780
                  1746
                             1485
1970s 749
                   698
                             1524
1960s 1152 635
                             2086
                             1540
1950s 1226 491
1940s 452
                   173
                             239
1930s 157
                   26
                             46
1920s 128
                   5
                             21
1910s 83
                   7
                             41
1900s 93
                             28
```

## Protein Homolog

Is there a mammalian equivalent of lycopene cyclase?

```
esearch -db protein -query \
  "lycopene beta cyclase [PROT] AND tomato [ORGN]" |
```

```
elink -related |
efetch -format gpc |
xtract -pattern INSDSeq -element INSDSeq_division |
sort-uniq-count-rank
```

In the resulting list of GenBank division codes:

```
905 BCT
856 ENV
609 PLN
197 CON
127 PAT
2 SYN
```

there are no similar sequences (protein neighbors) in the HUM, PRI, ROD, MAM, VRT, or INV divisions, so lycopene cyclase is not present in animals.

### **Longest Sequences**

What are the longest known insulin precursor molecules?

```
esearch -db protein -query "insulin [PROT]" |
efetch -format docsum |
xtract -pattern DocumentSummary -element Caption Slen Title |
grep -v receptor | sort -k 2,2nr | head -n 5 | cut -f 1 |
xargs -n 1 sh -c 'efetch -db protein -id "$0" -format gp > "$0".gpf'
```

Post-processing excludes the longer "insulin-like receptor" sequences and saves the GenPept results to individual files named by their sequence accessions:

```
EFN61235.gpf
EFN80340.gpf
EGW08477.gpf
EKC18433.gpf
ELK28555.gpf
```

using the right angle bracket (">") UNIX output redirection character.

### Archaea Enzyme

Which archaebacteria have chloramphenicol acetyltransferase?

```
esearch -db protein -query \
    "chloramphenicol acetyltransferase [PROT] AND archaea [ORGN]" |
efetch -format gpc |
xtract -pattern INSDSeq -element INSDSeq_organism INSDSeq_definition |
grep -i chloramphenicol | cut -f 1 | sort -f | uniq
```

produces a list of organism names:

```
Methanobrevibacter ruminantium
Methanobrevibacter smithii
Methanosarcina acetivorans
...
```

## Structural Similarity

What archaea structures are similar to snake venom phospholipase?

```
esearch -db structure -query "crotalus [ORGN] AND phospholipase A2" |
elink -related |
efilter -query "archaea [ORGN]" |
efetch -format docsum |
xtract -pattern DocumentSummary \
    -if PdbClass -equals Hydrolase \
    -element PdbDescr |
sort -f | uniq -i
```

This query uses geometric comparison (structure neighboring) to find proteins that are too divergent to be detected by sequence similarity with a BLAST search:

```
Crystal Structure Of Autoprocessed Form Of Tk-Subtilisin
Crystal Structure Of Ca2 Site Mutant Of Pro-S324a
Crystal Structure Of Ca3 Site Mutant Of Pro-S324a
```

## Taxonomy Search

Which organisms contain an annotated RefSeq genome MatK gene?

```
esearch -db nuccore -query "Matk [GENE] AND NC_0:NC_999999999 [PACC]" |
efetch -format docsum |
xtract -pattern DocumentSummary -element TaxId |
sort -n | uniq |
epost -db taxonomy |
efetch -format docsum |
xtract -pattern DocumentSummary -element ScientificName |
sort
```

The first query obtains taxonomy UIDs from nucleotide document summaries and uploads them for separate retrieval from the taxonomy database:

```
Acidosasa purpurea
Acorus americanus
...
Zingiber spectabile
Zygnema circumcarinatum
```

### **Chromosome Locations**

Where are mammalian calmodulin genes located?

```
esearch -db gene -query "calmodulin [PFN] AND mammalia [ORGN]" |
efetch -format docsum |
xtract -pattern DocumentSummary -MAP "(-)" -MAP MapLocation \
    -element Id Name "&MAP" ScientificName
```

The MAP variable is initialized with a literal dash to prevent missing data from shifting columns in the table:

801	CALM1	14q32.11	Homo sapiens
808	CALM3	19q13.32	Homo sapiens
805	CALM2	2p21	Homo sapiens
24242	Calm1	6q32	Rattus norvegicus
12313	Calm1	12 E	Mus musculus
326597	CALM	_	Bos taurus
50663	Calm2	6q12	Rattus norvegicus
24244	Calm3	1q21	Rattus norvegicus
12315	Calm3	7 9.15 cM	Mus musculus
12314	Calm2	17 E4	Mus musculus
617095	CALM1	-	Bos taurus
396838	CALM3	6	Sus scrofa

The -else command can also be used to insert placeholders for missing data:

```
esearch -db gene -query "calmodulin [PFN] AND mammalia [ORGN]" |
efetch -format docsum |
xtract -pattern DocumentSummary \
   -if MapLocation -element Id Name MapLocation ScientificName \
   -else -element Id Name -lbl "\-" -element ScientificName
```

The -def command can achieve the same result for missing elements:

```
esearch -db gene -query "calmodulin [PFN] AND mammalia [ORGN]" |
efetch -format docsum |
xtract -pattern DocumentSummary \
  -def "-" -element Id Name MapLocation ScientificName
```

### **Exon Counts**

How many exons are in each dystrophin transcript variant?

```
esearch -db gene -query "DMD [GENE] AND human [ORGN]" |
efetch -format docsum |
xtract -pattern DocumentSummary \
   -block GenomicInfoType -tab "\n" -element ChrAccVer, ChrStart, ChrStop |
```

This search returns the chromosome accession and the (0-based) gene start and stop positions:

```
NC 000023.11 33339608 31119221
```

These are then passed to efetch in (0-based) -chr\_start and -chr\_stop arguments:

```
xargs -n 3 sh -c 'efetch -db nuccore -format gbc \
  -id "$0" -chr_start "$1" -chr_stop "$2"' |
```

which converts them to (1-based) -seq\_start and -seq\_stop arguments and retrieves an INSDSeq XML subset record for the indicated region. That contains a number of alternatively-spliced dystrophin mRNA and CDS features.

Data extraction computes the number of intervals for each mRNA location (corresponding to individual exons or UTRs), and obtains the transcript sequence accession, transcript length, and product name from qualifiers:

```
xtract -insd complete mRNA "#INSDInterval" \
  transcript_id "%transcription" product |
```

Final processing sorts by number of exons:

```
grep -i dystrophin |
sed 's/dystrophin, transcript variant //g' |
sort -k 2,2nr -k 4,4nr
```

resulting in a table of exon counts and transcript lengths:

```
      NC_000023.11
      79
      NM_004010.3
      14083
      Dp427p2

      NC_000023.11
      79
      NM_000109.3
      14069
      Dp427c

      NC_000023.11
      79
      NM_004009.3
      14000
      Dp427p1

      NC_000023.11
      79
      NM_004006.2
      13993
      Dp427m

      NC_000023.11
      78
      XM_006724468.1
      13920
      X1

      NC_000023.11
      78
      XM_006724469.1
      13802
      X2

      NC_000023.11
      77
      XM_006724470.1
      13881
      X3
```

### Genome Range

What genes are in a given range on the human Y chromosome?

```
esearch -db gene -query "Homo sapiens [ORGN] AND Y [CHR]" |
efilter -status alive | efetch -format docsum |
xtract -pattern DocumentSummary -NAME Name -DESC Description \
   -block GenomicInfoType -if ChrLoc -equals Y \
   -min ChrStart,ChrStop -element "&NAME" "&DESC" |
sort -k 1,ln | cut -f 2- |
between-two-genes ASMT IL3RA
```

This query returns a table of gene names and descriptions, for the human "Y" chromosome, in the region between the ASMT and IL3RA genes:

```
IL3RA interleukin 3 receptor subunit alpha
LOC101928032 uncharacterized LOC101928032
LOC101928055 uncharacterized LOC101928055
SLC25A6 solute carrier family 25 member 6
LOC105373102 uncharacterized LOC105373102
LINC00106 long intergenic non-protein coding RNA 106
ASMTL-AS1 ASMTL antisense RNA 1
ASMTL acetylserotonin O-methyltransferase-like
P2RY8 purinergic receptor P2Y8
AKAP17A A-kinase anchoring protein 17A
acetylserotonin O-methyltransferase
```

(The "-if ChrLoc -equals Y" test is necessary because certain genes (e.g., IL9R) are present in the pseudoautosomal regions common to both X and Y chromosomes:

```
<GenomicInfo>
 <GenomicInfoType>
   <ChrLoc>Y</ChrLoc>
   <ChrAccVer>NC 000024.10</ChrAccVer>
   <ChrStart>57184100</ChrStart>
   <ChrStop>57197336</ChrStop>
    <ExonCount>10</ExonCount>
 </GenomicInfoType>
 <GenomicInfoType>
   <ChrLoc>X</ChrLoc>
    <ChrAccVer>NC_000023.11</ChrAccVer>
   <ChrStart>155997580</ChrStart>
   <ChrStop>156010816</ChrStop>
   <ExonCount>10</ExonCount>
  </GenomicInfoType>
</GenomicInfo>
```

with each gene copy annotated in its own GenomicInfoType block.)

### Gene Counts

How many genes are on each human chromosome?

returns a count of unique protein-coding genes per chromosome:

```
2067
       1
1268
       2
1071
       3
755
       4
       5
873
1034
      6
       7
935
690
      8
801
      9
739
      10
1288
      11
1027
      12
335
      13
607
      14
608
      15
```

```
862
      16
1181
       17
277
      18
1402
      19
545
      20
      21
248
445
      22
844
      X
71
13
```

The range construct cannot be used for Roman numerals, so the equivalent query on Saccharomyces cerevisiae would need to explicitly list all chromosomes:

### Complete Genomes

What complete genomes are available for Escherichia coli?

```
esearch -db assembly -query \
   "Escherichia coli [ORGN] AND representative [PROP]" |
elink -target nuccore -name assembly_nuccore_refseq |
efetch -format docsum |
xtract -pattern DocumentSummary -element AccessionVersion Slen Title |
sed 's/,.*//' |
sort -t $'\t' -k 2,2nr
```

This search finds genomic assemblies and sorts the results by sequence length, allowing complete genomes to be easily distinguished from smaller plasmids:

```
      NC_002695.1
      5498450
      Escherichia coli 0157:H7 str. Sakai chromosome

      NC_018658.1
      5273097
      Escherichia coli 0104:H4 str. 2011C-3493 ...

      NC_011751.1
      5202090
      Escherichia coli UMN026 chromosome

      NC_011750.1
      5132068
      Escherichia coli IAI39 chromosome

      NC_017634.1
      4747819
      Escherichia coli 083:H1 str. NRG 857C chromosome

      NC_000913.3
      4641652
      Escherichia coli str. K-12 substr. MG1655

      NC_017659.1
      147060
      Escherichia coli 083:H1 str. NRG 857C plasmid ...
```

The sed command removes extraneous text in the title (e.g., complete genome, complete sequence, primary assembly) after a comma.

A similar query for humans, additionally filtering out scaffolds, contigs, and plasmids:

```
esearch -db assembly -query "Homo sapiens [ORGN] AND representative [PROP]" | elink -target nuccore -name assembly_nuccore_refseq | efetch -format docsum | xtract -pattern DocumentSummary -element AccessionVersion Slen Title | sed 's/,.*//' | grep -v scaffold | grep -v contig | grep -v plasmid | sort
```

returns the assembled chromosome and mitochondrial sequence records:

```
NC 000001.11
                                248956422
                                                           Homo sapiens chromosome 1
 NC_000002.12 242193529 Homo sapiens chromosome 2
 NC 000003.12 198295559 Homo sapiens chromosome 3
 NC_000004.12 190214555 Homo sapiens chromosome 4
                                181538259 Homo sapiens chromosome 5
 NC_000005.10
 NC_000006.12 170805979 Homo sapiens chromosome 6
 NC 000007.14 159345973 Homo sapiens chromosome 7
 NC_000008.11 145138636 Homo sapiens chromosome 8
NC_000009.12 138394717 Homo sapiens chromosome 9
 NC_000010.11 133797422 Homo sapiens chromosome 10
 NC_000011.10 135086622 Homo sapiens chromosome 11
                                133275309 Homo sapiens chromosome 12
 NC_000012.12
NC_000013.11 114364328 Homo sapiens chromosome 13
                             107043718 Homo sapiens chromosome 14
 NC 000014.9

        NC_000014.9
        107043718
        Homo sapiens chromosome 14

        NC_000015.10
        101991189
        Homo sapiens chromosome 15

        NC_000016.10
        90338345
        Homo sapiens chromosome 16

        NC_000017.11
        83257441
        Homo sapiens chromosome 17

        NC_000018.10
        80373285
        Homo sapiens chromosome 18

        NC_000019.10
        58617616
        Homo sapiens chromosome 19

        NC_000020.11
        64444167
        Homo sapiens chromosome 20

        NC_000021.9
        46709983
        Homo sapiens chromosome 21

        NC_000022.11
        50818468
        Homo sapiens chromosome X

        NC_000023.11
        156040895
        Homo sapiens chromosome Y

        NC_012920.1
        16569
        Homo sapiens mitochondrion

                                                           Homo sapiens mitochondrion
 NC 012920.1
                                16569
```

This process can be automated to loop through a list of specified organisms:

```
for org in \
  "Agrobacterium tumefaciens" \
  "Bacillus anthracis" \
  "Escherichia coli" \
  "Neisseria gonorrhoeae" \
  "Pseudomonas aeruginosa" \
  "Shiqella flexneri" \
  "Streptococcus pneumoniae"
  esearch -db assembly -query "$org [ORGN]" |
  efilter -query "representative [PROP]"
  elink -target nuccore -name assembly_nuccore_refseq |
  efetch -format docsum |
 xtract -pattern DocumentSummary -element AccessionVersion Slen Title
  sed 's/,.*//'
  grep -v -i -e scaffold -e contig -e plasmid -e sequence -e patch |
  sort -t $'\t' -k 2,2nr
done
```

### which generates:

```
NC_011985.1 4005130 Agrobacterium radiobacter K84 chromosome 1
NC_011983.1 2650913 Agrobacterium radiobacter K84 chromosome 2
NC_005945.1 5228663 Bacillus anthracis str. Sterne chromosome
NC_003997.3 5227293 Bacillus anthracis str. Ames chromosome
```

```
NC_002695.1 5498450 Escherichia coli O157:H7 str. Sakai chromosome NC_018658.1 5273097 Escherichia coli O104:H4 str. 2011C-3493 ...
NC_011751.1 5202090 Escherichia coli UMN026 chromosome NC_011750.1 5132068 Escherichia coli IAI39 chromosome NC_017634.1 4747819 Escherichia coli O83:H1 str. NRG 857C chromosome NC_000913.3 4641652 Escherichia coli str. K-12 substr. MG1655 NC_002946.2 2153922 Neisseria gonorrhoeae FA 1090 chromosome NC_002516.2 6264404 Pseudomonas aeruginosa PAO1 chromosome NC_004337.2 4607202 Shigella flexneri 2a str. 301 chromosome NC_003028.3 2160842 Streptococcus pneumoniae TIGR4 chromosome NC_003098.1 2038615 Streptococcus pneumoniae R6 chromosome
```

### Amino Acid Composition

What is the amino acid composition of human titin?

produces a table of residue counts using the three-letter amino acid abbreviations:

```
Ala
      2084
      1640
Arq
Asn
      1111
    1720
Asp
Cys
      513
Gln
    942
Glu
    3193
Gly
      2066
His
      478
Ile
    2062
Leu
    2117
     2943
Lys
Met
      398
Phe
    908
    2517
Pro
Ser
      2463
      2546
Thr
Trp
      466
```

```
Tyr 999
Val 3184
```

### Amino Acid Substitutions

What are the missense products of green-sensitive opsin?

```
ApplySNPs() {
  seq=""
  last=""
  while read rsid accn pos res
    if [ "$accn" != "$last" ]
    then
      insd=$(efetch -db protein -id "$accn" -format gbc < /dev/null)</pre>
      seq=$(echo $insd | xtract -pattern INSDSeq -element INSDSeq_sequence)
      last=$accn
    fi
    pos=$((pos+1))
    pfx=""
    sfx=""
    echo ">rs$rsid [$accn $res@$pos]"
    if [ $pos -gt 1 ]
      pfx=$(echo ${seq:0:$pos-1})
    fi
    if [ $pos -lt ${#seq} ]
    then
      sfx=$(echo ${seq:$pos})
    echo "$pfx$res$sfx" | fold -w 50
  done
}
esearch -db gene -query "CBD [GENE] AND human [ORGN]" |
elink -target snp |
efetch -format xml |
xtract -pattern Rs -RSID Rs@rsId \
  -block FxnSet -if @fxnClass -equals missense \
    -sep "." -element "&RSID" @protAcc,@protVer @aaPosition \
    -tab "\n" -element @residue |
sort -t $'\t' -k 2,2 -k 3,3n -k 4,4 | uniq | ApplySNPs
```

The query returns an intermediate table of non-synonymous amino acid substitutions (with 0-based location coordinates) derived from single nucleotide polymorphisms:

```
104894915 NP_000504.1 93 K
782122931 NP_000504.1 95 V
781899063 NP_000504.1 97 T
781807082 NP_000504.1 102 A
```

The rows are then processed to produce protein sequences with the individual residue substitutions in upper case:

```
>rs104894915 [NP_000504.1 K@94]
maqqwslqrlagrhpqdsyedstqssiftytnsnstrgpfegpnyhiapr
wvyhltsvwmifvviasvftnglvlaatmkfkklrhplnwilvKlavadl
aetviastisvvnqvygyfvlghpmcvlegytvslcgitglwslaiiswe
```

### 3'UTR Sequences

What are the 3' UTR sequences for lycopene cyclase?

```
ThreePrimeUTRs() {
  xtract -pattern INSDSeq -ACC INSDSeq_accession-version -SEQ INSDSeq_sequence \
    -group INSDFeature -if INSDFeature_key -equals CDS -PRD "(-)" \
      -block INSDQualifier -if INSDQualifier_name \
        -equals product -PRD INSDQualifier_value \
      -block INSDFeature -pfc "\n" -element "&ACC" -rst \
        -last INSDInterval_to -element "&SEQ" "&PRD" |
  while read acc pos seq prd
  do
    if [ $pos -lt ${#seq} ]
    then
      echo -e ">$acc 3'UTR: $((pos+1))..${#seq} $prd"
      echo "${seq:$pos}" | fold -w 50
    elif [ $pos -ge ${#seq} ]
      echo -e ">$acc NO 3'UTR"
    fi
  done
esearch -db nuccore -query "5.5.1.19 [ECNO]" |
efilter -molecule mrna -source refseg |
efetch -format gbc | ThreePrimeUTRs
```

prints the sequences immediately following the CDS stop codon:

## **Upstream Sequences**

What sequences are upstream of phenylalanine hydroxylase genes?

```
esearch -db nuccore -query "U49897 [ACCN]" | elink -target gene |
```

```
elink -target homologene |
elink -target gene |
efetch -format docsum |
xtract -pattern DocumentSummary -if GenomicInfoType -element Id \
   -block GenomicInfoType -element ChrAccVer ChrStart ChrStop |
awk -F '\t' -v 'OFS=\t' '{print $1, $2, $3+1, $4+1}'
```

obtains a series of homologous genes, converting the gene coordinates to 1-based positions suitable for retrieving sequence regions:

```
      5053
      NC_000012.12
      102917603
      102838326

      18478
      NC_000076.6
      87521795
      87584137

      38871
      NT_037436.4
      7760453
      7763166

      24616
      NC_005106.4
      28066639
      28129772

      378962
      NC_007115.6
      17420391
      17402704
```

Given a shell script named "upstream.sh":

```
#!/bin/bash -norc
bases=1500
if [ -n "$1" ]
then
 bases=$1
while read id accn start stop
  if [[ $start -eq 0 || $stop -eq 0 || $start -eq $stop ]]
    echo "Skipping $id due to ambiguous coordinates"
    continue
  fi
  if [ $start -gt $stop ]
  then
    stop=$(( start + bases ))
    start=$(( start + 1 ))
    strand=2
  else
    stop=$(( start - 1 ))
    start=$(( start - bases ))
    strand=1
  rslt=`efetch -db nuccore -id $accn -format fasta \
        -seq_start $start -seq_stop $stop -strand $strand < /dev/null`</pre>
  echo "$rslt"
done
```

the data lines can be piped through:

```
upstream.sh 500
```

to extract and print the 500 nucleotides immediately upstream of each gene. (Without the argument it will default to 1500 nucleotides.)

### **Author Combinations**

What are the authorship patterns among selected individuals?

The "coauthors.sh" script takes author name arguments to construct a custom data extraction command for analyzing research collaboration patterns:

```
#!/bin/bash -norc
if [ "$#" -lt 2 ]
 echo "Must supply at least two author names"
fi
query="xtract -pattern PubmedArticle -element MedlineCitation/PMID"
# append a -block statement for each author argument
for auth in "$@"
do
 query=`echo "$query -block Author -if LastName -equals \"$auth\"" \
              "-sep \" \" -element LastName, Initials"`
done
query=`echo "$query | sort -t \'\\t' -k 2f -k 1,1n"`
if [ -t 0 ]
  # stand-alone command, print constructed query for later use
 echo "$query"
  # dynamically execute query on XML data piped to script
 res=`eval "$query"`
 echo "$res"
```

If XML publication data are piped to the script, it will read the data and immediately execute the generated xtract query. Otherwise, if called as a stand-alone command, it will print the custom query instructions for later use.

Running the following command:

```
esearch -db pubmed -query "Casadaban MJ [AUTH] OR Berg CM [AUTH]" | efetch -format xml | ./coauthors.sh Casadaban Groisman Berg Garber | ./extract-fuse.pl pubmed > author_patterns.htm
```

first produces an internal result table of PMIDs grouped by author combination:

```
7635839 Casadaban MJ
9634770 Casadaban MJ
1827084 Casadaban MJ Groisman EA
2954879 Casadaban MJ Groisman EA
3020001 Casadaban MJ Groisman EA
3525518 Casadaban MJ Groisman EA
3542967 Casadaban MJ Groisman EA
6324195 Casadaban MJ Groisman EA
3301525 Casadaban MJ Groisman EA
Berg CM
```

### The sorted lines are then piped to the "extract-fuse.pl" script:

```
#!/usr/bin/perl
my $max = scalar @ARGV;
if ( $max < 1 ) {
 die "Need argument for database\n";
my $db = $ARGV[0];
my $thisline = "";
my $laststr = "";
my $str = "";
my $uid = "";
my $uidlist = "";
my $count = 0;
my $base = "https://www.ncbi.nlm.nih.gov";
my \ pfx = "";
while ($thisline = <STDIN>) {
  thisline =  s/\r//;
  thisline =  s/n//;
  if (thisline = (([^{t}]+)\t(.+))) {
   $uid = $1;
   \$str = \$2;
    if ( lc ($str) ne lc ($laststr) and $laststr ne "" ) {
     \frac{1}{2}
     print "(<a href=\"$base/$db/$uidlist\">";
     print " $count </a>) - $laststr\n";
     pfx = "";
     $count = 0;
     $uidlist = "";
    $laststr = $str;
    $uidlist .= "$pfx$uid";
   $pfx = ",";
    $count++;
}
```

```
if ( $laststr ne "" ) {
    $laststr =~ s/\t/, /g;
    print "(<a href=\"$base/$db/$uidlist\">";
    print " $count </a>) - $laststr\n";
}
```

which combines them into PubMed query URLs, one for each author pattern:

```
https://www.ncbi.nlm.nih.gov/pubmed/1827084,2954879,3020001,...
```

Those are then wrapped, along with a record count, in the appropriate HTML tags for web display. If the resulting file is opened with a browser, it presents an argument-order-dependent view of author collaboration:

```
( 55 ) - Berg CM
( 10 ) - Berg CM, Berg DE
( 1 ) - BERG CM, GARBER ED
( 6 ) - Berg DE, Berg CM
( 39 ) - Casadaban MJ
( 6 ) - Casadaban MJ, Groisman EA
( 1 ) - Casadaban MJ, Groisman EA, Berg CM
```

Clicking on a hyperlinked record count number opens the document summary or individual article page, so the actual publications can be examined.

### Indexed Fields

What date fields are indexed for PubMed?

```
einfo -db pubmed |
xtract -pattern Field \
  -if IsDate -equals Y -and IsHidden -equals N \
    -pfx "[" -sfx "]" -element Name \
    -pfx "" -sfx "" -element FullName |
sort -k 2f | expand
```

This produces a list of field abbreviations and names filtered by index type:

```
[CDAT] Date - Completion

[CRDT] Date - Create

[EDAT] Date - Entrez

[MHDA] Date - MeSH

[MDAT] Date - Modification

[PDAT] Date - Publication
```

## Digital Object Identifiers

How are digital object identifiers obtained from PubMed articles?

```
esearch -db pubmed -query "Rowley JD [AUTH]" |
efetch -format xml |
xtract -head '<html><body>' -tail '</body></html>' \
    -pattern PubmedArticle -PMID MedlineCitation/PMID \
    -block ArticleId -if @IdType -equals doi \
    -tab '\n' -pfx '<a href="http://dx.doi.org/' \
    -sep '">' -sfx '</a>' -encode ArticleId, "&PMID"
```

extracts the DOIs and constructs the appropriate URL references:

```
<html><body>
<a href="http://dx.doi.org/10.1038/leu.2013.340">24496283</a>
<a href="http://dx.doi.org/10.1073/pnas.1310656110">23818607</a>
<a href="http://dx.doi.org/10.1073/pnas.1310144110">23798388</a>
```

These intermediate lines are then piped through:

```
xtract -format
```

to produce a minimal HTML document with clickable links:

## Phrase Searching

Can phrase searching be simulated in Entrez?

The "entrez-phrase-search" script included with EDirect takes advantage of the fact that some short phrases are indexed in certain Entrez fields. Given an input phrase, the script generates overlapping pairs of adjacent words, separately queries on each pair to determine which are present in the pubmed title or abstract index, and keeps those that appear in at least 10 articles. Independent phrases are separated by a plus ("+") sign.

For example, running the following command:

```
entrez-phrase-search -db pubmed -field WORD \
   selective serotonin reuptake inhibitor + monoamine oxidase inhibitor
```

will generate word pairs from each phrase and run a query on each pair. The individual term counts are:

```
11343 selective serotonin
11892 serotonin reuptake
6714 reuptake inhibitor
21722 monoamine oxidase
3680 oxidase inhibitor
```

The combined query will return a search result with 36 articles, and these can then be retrieved by piping to efetch. The script in its current form will not match phrases with plurals (e.g., serotonin reuptake inhibitors) or hyphens (e.g., monoamine-oxidase inhibitor).

### Gene-Protein Links

What proteins are produced by a given gene?

Given a query in the gene database, the following commands:

```
esearch -db gene -query "beta galactosidase [PFN]" |
elink -target protein -name gene_protein_refseq -cmd neighbor |
xtract -pattern LinkSet -element Id
```

will show the gene ID in the first column and linked RefSeq protein UIDs in subsequent columns.

Piping the results to a Perl script named "gene-protein-links.pl" will read the identifiers and run separate efetch queries on the gene and protein databases:

```
#!/usr/bin/perl
while ($line = <STDIN>) {
  chomp ($line);
  @uids = split( /\t/, $line);
  $gene = $uids [0];
  $proteins = join (',', @uids [1..$#uids]);
  $symbol = $data = '';
  $cmd = "efetch -format docsum -db gene -id $gene | ";
  $cmd .= "xtract -pattern DocumentSummary -element Name CommonName";
  open (CMD, "$cmd|");
  while (<CMD>) {
    symbol .= s_;
  }
  close CMD;
  if ($proteins ne "") {
    $cmd = "efetch -format docsum -db protein -id $proteins | ";
    $cmd .= "xtract -pattern DocumentSummary -element Caption Slen Title";
    open (CMD, "$cmd|");
```

```
while (<CMD>) {
    $data .= $_;
}
close CMD;
}
print "$symbol$data\n";
}
```

printing the gene symbol and organism common name, followed by the protein accessions, lengths, and titles:

```
GLB1
                human
NP 001129074
                546
                       beta-galactosidase isoform c preproprotein ...
                beta-galactosidase isoform c preproprotein
beta-galactosidase isoform b [Homo sapiens]
NP_001073279
NP 000395
                beta-galactosidase isoform a preproprotein ...
Glb1
                house mouse
NP_033882
                647 beta-galactosidase preproprotein [Mus musculus]
Glb1
               Norway rat
NP_001101662 647 beta-galactosidase precursor [Rattus norvegicus]
```

### **Bulk Downloads**

How can the entire set of GenBank records for mammals be obtained?

```
ftp-ls ftp.ncbi.nlm.nih.gov genbank |
grep ".seq.gz" |
grep -e gbmam -e gbpri -e gbrod |
while read file
do
   ftp-cp ftp.ncbi.nlm.nih.gov genbank "$file"
   gzcat "$file"
   rm "$file"
done
```

will use the ftp-ls and ftp-cp scripts (included with the EDirect software) to retrieve and print GenBank flatfiles for human, primate, rodent, and other mammals:

```
GBMAM1.SEQ Genetic Sequence Data Bank
February 15 2015

NCBI-GenBank Flat File Release 206.0

Other Mammalian Sequences (Part 1)

20709 loci, 155323216 bases, from 20709 reported sequences

LOCUS AB000170 2732 bp mRNA linear MAM ...

DEFINITION Sus scrofa mRNA for endopeptidase 24.16, complete cds.
```

```
ACCESSION AB000170
VERSION AB000170.1 GI:1783121
KEYWORDS endopeptidase 24.16 type M3; endopeptidase 24.16 type M1.
SOURCE
          Sus scrofa (pig)
  ORGANISM Sus scrofa
           Eukaryota; Metazoa; Chordata; Craniata; Vertebrata;
            Euteleostomi; Mammalia; Eutheria; Laurasiatheria;
            Cetartiodactyla; Suina; Suidae; Sus.
REFERENCE
 AUTHORS Kato, A., Sugiura, N., Saruta, Y., Hosoiri, T., Yasue, H. and
           Hirose,S.
  TITLE
           Targeting of endopeptidase 24.16 to different subcellular
           compartments by alternative promoter usage
  JOURNAL J. Biol. Chem. 272 (24), 15313-15322 (1997)
   PUBMED 9182559
REFERENCE 2 (bases 1 to 2732)
  AUTHORS Hirose, S.
  TITLE
         Direct Submission
  JOURNAL Submitted (27-DEC-1996) Shigehisa Hirose, Tokyo Institute of
           Technology, Department of Biological Sciences; 4259
            Nagatsuta-cho, Midori-ku, Yokohama, Kanagawa 226-8501, Japan
FEATURES
                    Location/Qualifiers
     source
                     1..2732
                     /organism="Sus scrofa"
                     /mol type="mRNA"
                     /db_xref="taxon:9823"
                     /tissue_type="Liver"
                     /dev_stage="Adult"
                     /note="porcine"
     mRNA
                     1..2732
                     /note="corresponding to exon1,5-16 of this gene;
                     endopeptidase 24.16 type 1"
     CDS
                     175..2289
                     /standard_name="endopeptidase 24.16"
                     /note="oligopeptidase M :neurolysin :sBAP(soluble
                     angiotensin-binding protein) :MEP(microsomal
                     metalloendopeptidase)"
                     /codon start=1
                     /product="endopeptidase 24.16 type M1"
                     /protein id="BAA19060.1"
                     /db xref="GI:1783122"
```

For systems with Aspera Connect installed, the asp-ls and asp-cp scripts can be used for faster retrieval:

```
asp-ls genbank |
grep ".seq.gz" |
grep -e gbmam -e gbpri -e gbrod |
while read file
do
   asp-cp genbank "$file"
   gzcat "$file"
```

```
rm "$file"
done
```

# **Appendices**

# Setting Contact Address and Script Name

EDirect automatically obtains the user's e-mail address from the system, to have someone to notify in case a runaway script causes problems with an Entrez server, but if another contact address is desired (e.g., that of a system administrator or software developer) it can be explicitly set at the beginning of a pipeline or script:

```
econtact -email author_email_address -tool name_of_script
```

That way the NCBI has information on who to contact if an infinite loop in a script accidentally abuses NCBI resources. (For convenience, the preferred e-mail address and software tool name can also be set in all E-utilities-calling operations.)

# Command-Line Arguments

Arguments for the EDirect functions are listed below:

Use esearch to start a new Entrez search on indexed terms:

```
Query Specification
  -db
                 Database name
  -query
                 Query string
Document Order
  -sort
            Result presentation order
Date Constraint
  -days Number of days in the past
-datetype Date field abbreviation
-mindate Start of date range
-maxdate End of date range
Spell Check
  -spell
                 Correct misspellings in query
Miscellaneous Arguments
                   Alias for query step
  -label
```

The elink function looks up related articles or associated records:

```
Destination Database
```

```
-related Neighbors in same database
-target Links in different database
```

-name Link name (e.g., pubmed\_protein\_refseq)

Direct Record Selection

-db Database name

-id Unique identifier(s)

Advanced Control

-cmd Command type (returns eLinkResult XML)
-mode "ref" uses LinkOut provider's web site

-holding Name of LinkOut provider

Batch Processing

-batch Bypass Entrez history mechanism

Miscellaneous Arguments

-label Alias for query step

Use efilter to restrict search or link results by indexed terms:

Query Specification

-query Query string

Document Order

-sort Result presentation order

Date Constraint

-days
 -datetype
 -mindate
 -maxdate
 Number of days in the past
 Date field abbreviation
 Start of date range
 End of date range

Spell Check

-spell Correct misspellings in query

Publication Filters

preprint, review, structured

Sequence Filters

-feature gene, mrna, cds, mat\_peptide, ...

-location mitochondrion, chloroplast, plasmid, plastid

-molecule genomic, mrna, trna, rrna, ncrna

-organism animals, archaea, bacteria, eukaryotes, fungi,

human, insects, mammals, plants, prokaryotes,

protists, rodents, viruses

-source genbank, insd, pdb, pir, refseq, swissprot, tpa

Gene Filters

-status alive

-type coding, pseudo

Miscellaneous Arguments

-label Alias for query step

#### The record retrieval function is efetch:

Format Selection

-format Format of record or report -mode text, xml, asn.1, json

Direct Record Selection

-db Database name

-id Unique identifier or accession number

Sequence Range

-seq\_start First sequence position to retrieve -seq\_stop Last sequence position to retrieve

-strand Strand of DNA to retrieve

Gene Range

-chr\_start Sequence range from 0-based coordinates -chr\_stop in gene docsum GenomicInfoType object

Miscellaneous

-complexity 0 = default, 1 = bioseq, 3 = nuc-prot set -extend Extend sequence retrieval in both directions

-extrafeat Bit flag specifying extra features

### The xtract function is used for processing XML data:

Processing

-cleanup Fix non-ASCII spaces
-compress Compress runs of spaces
-plain Delete Unicode accents
-relaxed Allow PubMed mixed content
-strict Remove HTML highlight tags

#### Data Source

Read XML from file instead of stdin -input

#### Exploration Argument Hierarchy

Name of record within set -pattern Use of different argument -group -block names allows command-line -subset control of nested looping

#### Exploration Constructs

Object DateCreated Book/AuthorList Parent/Child Heterogeneous "PubmedArticleSet/\*"

Nested "\*/Taxon"

"\*\*/Gene-commentary" Recursive

#### Conditional Execution

-if Element [@attribute] required

-unless Skip if element matches All tests must pass -and -or Any passing test suffices

-else Execute if conditional test failed

-position Must be at [first|last] location in list

#### String Constraints

-equals String must match exactly -contains Substring must be present -starts-with Substring must be at beginning -ends-with Substring must be at end

String must not match -is-not

#### Numeric Constraints

Greater than -qt

Greater than or equal to -ge

-lt Less than

-le Less than or equal to

Equal to -eq Not equal to -ne

#### Format Customization

-ret	Override line break between patterns
-tab	Replace tab character between fields
-sep	Separator between group members
-pfx	Prefix to print before group
-sfx	Suffix to print after group

-clr	Clear queued tab separator
-pfc	Preface combines -clr and -pfx
-rst	Reset -sep, -pfx, and -sfx

-def Default placeholder for missing fields

-lbl Insert arbitrary text

#### Element Selection

-element Print all items that match tag name
 -first Only print value of first item
 -last Only print value of last item
 -NAME Record value in named variable

#### -element Constructs

Tag Caption

Group Initials, LastName
Parent/Child MedlineCitation/PMID

Attribute DescriptorName@MajorTopicYN Recursive "\*\*/Gene-commentary\_accession"

Object Count "#Author"

Item Length "%Title"

Element Depth "^PMID"

Variable "&NAME"

#### Special -element Operations

Parent Index "+"
XML Subtree "\*"
Children "\$"
Attributes "@"

### Numeric Processing

-num Count -len Length -sum Sum -min Minimum Maximum -max -inc Increment Decrement -dec -sub Difference Average -avq -dev Deviation

#### String Processing

-encode URL-encode <, >, &, ", and ' characters

-title Capitalize initial letters of words

#### Phrase Processing

terms
 partition phrase at spaces
 words
 split at punctuation marks
 pairs
 Adjacent informative words
 letters
 separate individual letters
 indices
 Experimental index generation

#### Sequence Coordinates

-0-based Zero-Based -1-based One-Based -ucsc-based Half-Open

#### Command Generator

-insd Generate INSDSeq extraction commands

#### -insd Argument Order

Descriptors INSDSeq\_sequence INSDSeq\_definition INSDSeq\_division

Flags [complete|partial]

Feature(s) CDS, mRNA

Qualifiers INSDFeature\_key "#INSDInterval" gene product

#### Miscellaneous

-head
-tail
-hid
-tail
-hid
-tail
-hid
-tail
-hid
-tail
-hid
-tail
-t

#### Reformatting

-format [copy|compact|flush|indent|expand]

#### Modification

-filter Object

[retain|remove|encode|decode|shrink|expand|accent]
 [content|cdata|comment|object|attributes|container]

#### Validation

-verify Report XML data integrity problems

#### Summary

-outline Display outline of XML structure -synopsis Display count of unique XML paths

#### Documentation

```
-examples Examples of EDirect and xtract usage
-extras Batch and local processing examples
```

The einfo function returns information on Entrez indexed fields:

```
Database Selection
```

```
-db Database name
-dbs Get all database names

Data Summaries

-fields Print field names
-links Print link names
```

Several additional functions are provided by EDirect:

```
epost
  -db
            Database name
  -id
           Unique identifier(s) or accession number(s)
  -format uid or acc
 -input Read from file instead of stdin-label Alias for query step
eproxy
 -alias File of aliases
  -pipe
          Read aliases from stdin
econtact
  -email Contact person's address
           Name of script or program
  -tool
nquire
            Uses HTTP GET instead of POST
  -get
            Base URL for external search
```

In addition, -email and -tool are available in all E-utilities-calling functions to override default values, -http get will force the use of GET instead of POST, -alias will specify a file of shortcut keywords and query strings or URL sections, and -help will print the list of arguments for each function.

For debugging, -silent will suppress link failure retry messages, -verbose will display the <ENTREZ\_DIRECT> field values at each step, -debug will print the internal URL query and XML results of each step, and -base will specify a particular server for quality assurance testing.

# **EFetch Formats**

EFetch -format and -mode values for each database are shown below:

-db	-format	-mode	Report Type
(all)			
	docsum docsum	igon	DocumentSummarySet XML
	full	json	DocumentSummarySet JSON Same as native except for mesh
	uid		Unique Identifier List
	url		Entrez URL
	xml		Same as -format full -mode xml
bioproject			
	native		BioProject Report
	native	xml	RecordSet XML
biosample			
	native		BioSample Report
	native	xml	BioSampleSet XML
biosystems			
	native	xml	Sys-set XML
gds			
	native	xml	RecordSet XML
	summary		Summary
gene			
	gene_table		Gene Table
	native	1	Gene Report
	native native	asn.1 xml	Entrezgene ASN.1 Entrezgene-Set XML
	tabular	XIIII	Tabular Report
la			<del>-</del>
homologene	alignmentscores		Alignment Scores
	fasta		FASTA
	homologene		Homologene Report
	native		Homologene List
	native	asn.1	HG-Entry ASN.1
	native	xml	Entrez-Homologene-Set XML
mesh			
	full		Full Record
	native	<sup>7</sup>	MeSH Report
	native	xml	RecordSet XML
nlmcatalog			- 11 p
	native		Full Record

	native	xml	NLMCatalogRecordSet XML
pmc	medline		MEDLINE
	native	xml	pmc-articleset XML
pubmed			
-	abstract		Abstract
	medline		MEDLINE
	native	asn.1	Pubmed-entry ASN.1
	native	xml	PubmedArticleSet XML
(sequences)			
	acc		Accession Number
	est		EST Report
	fasta	-	FASTA
	fasta	xml	TinySeq XML
	fasta_cds_aa fasta_cds_na		FASTA of CDS Products FASTA of Coding Regions
	ft		Feature Table
	gb		GenBank Flatfile
	gb	xml	GBSet XML
	gbc	xml	INSDSet XML
	gbwithparts		GenBank with Contig Sequences
	gene_fasta		FASTA of Gene
	gp		GenPept Flatfile
	gp	xml	GBSet XML
	gpc	xml	INSDSet XML
	gss		GSS Report
	ipg	-	Identical Protein Report
	ipg	xml	IPGReportSet XML
	native	text	Seq-entry ASN.1
	native seqid	xml	Bioseq-set XML Seq-id ASN.1
	seqia		seq-1d ASN.1
snp			_
	chr		Chromosome Report
	docset fasta		Summary FASTA
	flt		Flat File
	native	asn.1	Rs ASN.1
	native	xml	ExchangeSet XML
	rsr	211111	RS Cluster Report
	ssexemplar		SS Exemplar List
ara			
sra	native	xml	EXPERIMENT_PACKAGE_SET XML
	runinfo	xml	SraRunInfo XML
structure	mmdb		Ncbi-mime-asn1 strucseq ASN.1
	native		MMDB Report
	native	xml	RecordSet XML
		22111±	1.00010000 1111111

taxonomy

native Taxonomy List native xml TaxaSet XML

### ESearch Sort Order

ESearch -sort values for several databases are listed below:

-db -sort \_\_\_ \_\_\_

gene

Chromosome Gene Weight

Name Relevance

geoprofiles

Default Order Deviation Mean Value Outliers

Subgroup Effect

 ${\tt pubmed}$ 

First Author Journal Last Author Pub Date

Recently Added Relevance Title

(sequences)

Accession
Date Modified
Date Released
Default Order
Organism Name
Taxonomy ID

snp

Chromosome Base Position

Default Order Heterozygosity

Organism SNP\_ID

Success Rate

### **ELink Commands**

ELink -cmd options produce results as LinkSet XML:

-cmd	Result
neighbor	Neighbors or links
neighbor_score	Neighbors with computed similarity scores
acheck	All links available
ncheck	Existence of neighbors
lcheck	Existence of external links (LinkOuts)
llinks	Non-library LinkOut providers
llinkslib	All LinkOut providers
prlinks	Primary LinkOut provider, or URL for single UID with -mode ref

## Elnfo Data

EInfo field data contains status flags for several term list index properties:

### **UNIX Utilities**

Several useful classes of UNIX text processing filters, with selected arguments, are presented below:

Process by contents:

```
sort Sorts lines of text

-f Ignore case
-n Numeric comparison
-r Reverse result order

-k Field key (start,stop or first)
-u Unique lines with identical keys
```

```
Ignore leading blanks
   -b
         Stable sort
   -s
   -t
         Specify field separator
         Removes repeated lines
 uniq
   -C
         Count occurrences
   -i
         Ignore case
   -f
         Ignore first n fields
    -s
         Ignore first n characters
         Only output repeated lines
   -d
         Only output non-repeated lines
   -u
         Matches patterns using regular expressions
 grep
   -i
         Ignore case
         Invert search
    -v
         Search expression as a word
   -w
         Search expression as whole line
   -x
         Specify individual pattern
   -е
         Only count number of matches
   -c
         Print line numbers
    -n
Regular expressions:
  Characters
         Any single character (except newline)
         Alphabetic [A-Za-z], numeric [0-9], or underscore (_)
   \w
   \s
         Whitespace (space or tab)
         Escapes special characters
   []
         Matches any enclosed characters
  Positions
         Beginning of line
         End of line
         Word boundary
   \b
 Repeat Matches
         0 or 1
         0 or more
         1 or more
```

### Modify contents:

Exactly n

{n}

- sed Replaces text strings
  - -e Specify individual expression
- tr Translates characters
  - -d Delete character
- rev Reverses characters on line

#### Format contents:

- column Aligns columns by content width
  - -s Specify field separator
  - -t Create table
- expand Aligns columns to specified positions
  - -t Tab positions
- fold Wraps lines at a specific width
  - -w Line width

### Filter by position:

- cut Removes parts of lines
  - -c Characters to keep
  - -f Fields to keep
  - -d Specify field separator
  - -s Suppress lines with no delimiters
- head Prints first lines
  - -n Number of lines
- tail Prints last lines
  - -n Number of lines

#### Miscellaneous:

- wc Counts words, lines, or characters
  - -c Characters
  - -l Lines
  - -w Words
- xargs Constructs arguments
  - -n Number of words per batch

### File compression:

```
Archive files
tar
       Create archive
  - C
  -f
       Name of output file
       Compress archive with gzip
  -z
qzip
       Compress file
  -k
      Keep original file
  -9
       Best compression
unzip
      Decompress .zip archive
       Pipe to stdout
  -p
       Decompress .gz archive and pipe to stdout
gzcat
```

Directory conventions and arguments for file navigation commands are shown below:

```
cd
       Changes directory
 /
       Root
       Home
       Current
      Parent
       Previous
ls
       Lists file names
 -1
      One entry per line
 -a
      Show files beginning with dot (.)
      List in long format
 -1
 -R
      Recursively explore subdirectories
 -S
      Sort files by size
       Sort by most recently modified
 -t
       Prints working directory path
```

Additional documentation with detailed explanations and examples can be obtained by typing "man" followed by a command name.

# Terminal Keyboard Shortcuts

Control and escape sequences can be used within a terminal session to navigate through the command history and to move the cursor for editing the command currently being entered:

#### Command history:

```
Ctrl-n Next command Ctrl-p Previous command
```

#### Move cursor forward:

```
Ctrl-e To end of line
Ctrl-f By one character
Esc-f By one argument
```

#### Move cursor backward:

```
Ctrl-a To beginning of line
Ctrl-b By one character
Esc-b By one argument
```

#### Delete:

Del	Previous character
Ctrl-d	Next character
Ctrl-k	To end of line
Ctrl-u	Entire line
Ctrl-w	Previous word
Esc-Del	Previous argument
Esc-d	Next argument

### Autocomplete:

```
Tab Completes directory or file names
```

### Program control:

```
Ctrl-c Quit running program
^x^y Run last command replacing x with y
```

(Note that Control sequences are typed by holding down Control, hitting the other key, and releasing Control, while Escape sequences are typed by hitting Escape and then hitting the other key.)

### Release Notes

# EDirect Version 6.80: June 8, 2017

- Modified download instructions to use edirect.tar.gz archive.
- The ftp-cp script can now read from stdin without the need for xargs.
- Rerunning ftp-cp or asp-cp only attempts to download missing files.

# EDirect Version 6.70: May 8, 2017

- Added asp-cp script for faster download of NCBI ftp files using Aspera Connect.
- Xtract -strict and -relaxed handle empty HTML tag variants (e.g., <b/> and <sup/>).

## EDirect Version 6.60: April 25, 2017

- Xtract -strict replaces -degloss to remove HTML <i>, <b, <u>, <sup> and <sub> tags from XML contents.
- Xtract -relaxed allows HTML tags in XML contents, to support current PubMed ftp release files.
- Xtract -plain removes Unicode accents.
- The setup.sh script prints an error message if it cannot fetch missing Perl modules.

### EDirect Version 6.50: March 6, 2017

• Xtract -degloss replaces -html to remove HTML <i>, <b, <u>, <sup> and <sub> tags.

### EDirect Version 6.40: March 1, 2017

- Epost detects accession.version input for sequence databases and sets -format acc.
- Xtract -html [remove|encode] converts <i> and <b> tags embedded in XML contents.

### EDirect Version 6.30: February 13, 2017

- Efetch -format docsum skips GI-less sequences without summaries.
- Xtract local indexing commands moved to -extras documentation.

## EDirect Version 6.20: January 30, 2017

• Xtract -limit and -index allow extraction of selected records from XML file.

### EDirect Version 6.10: January 19, 2017

- Added run-ncbi-converter script for processing ASN.1 release files.
- Xtract -format flush option added.
- Removed obsolete accession-dot-version conversion code.

# EDirect Version 6.00: December 27, 2016

- Efetch -format docsum removes eSummaryResult wrapper.
- Fixed content truncation bug when Xtract encounters very long sequences.

## EDirect Version 5.90: December 21, 2016

- Efetch and Elink readied for switch to accession-dot-version sequence identifier.
- Xtract -insd recognizes INSDInterval\_iscomp@value and other boolean attributes.
- Xtract adds experimental phrase processing commands for word index preparation.

### EDirect Version 5.80: December 12, 2016

- Efilter adds shortcuts for -db gene (e.g., -status alive, -type coding).
- Xtract numeric conditional tests can use an element name for the second argument (e.g., -if ChrStop -lt ChrStart finds minus strand genes).

### EDirect Version 5.70: November 30, 2016

• Xtract -format takes an optional [compact|indent|expand] argument. Processing compact XML is about 15% faster than indent form. Using expand places each attribute on a separate line for ease of reading.

### EDirect Version 5.60: November 22, 2016

- Fixed bug in -datetype argument for Esearch and Efilter.
- Added optional argument to filter-stop-words script to indicate replacement.

### EDirect Version 5.50: November 16, 2016

- Efetch -id allows non-numeric accessions only for sequence databases.
- Xtract element selection no longer considers fields in recursive sub-objects.
- Xtract introduces a double-star "\*\*/Object" construct to flatten recursive child objects for linear exploration.
- Xtract conditional tests ignore empty self-closing tags.
- Xtract -else simplifies insertion of a placeholder to indicate missing data.

## EDirect Version 5.40: November 7, 2016

• Added filter-stop-words and xy-plot scripts.

### EDirect Version 5.30: October 31, 2016

- Added support for ecitmatch utility.
- Added amino-acid-composition and between-two-genes scripts.
- The sort-uniq-count and sort-uniq-count-rank scripts take an optional argument (e.g., -n for numeric comparisons, -r to reverse order).

## EDirect Version 5.20: October 26, 2016

- Setup script no longer modifies the user's configuration file to update the PATH
  variable. Instead, it now prints customized instructions for the user to execute. The
  user may choose to run these commands, but is free to edit the .bash\_profile file
  manually.
- Xtract deprecates -match and -avoid functions and the Element: Value conditional shortcut.
- Xtract -if and -unless commands use compound statements for conditional execution (e.g., -if Element -equals Value).

- Colon now separates namespace prefix from element name in xtract arguments (e.g., -block jats:abstract). Colon at start of element name matches any namespace prefix.
- Xtract -insd uses a dash as placeholder for missing field. Experimental -insdx command is deprecated.
- Precompiled versions of xtract are now provided for Darwin, Linux, and CYGWIN\_NT platforms. The appropriate executable is downloaded by the setup script.

# EDirect Version 5.10: October 13, 2016

• Xtract adds -0-based, -1-based, and -ucsc numeric extraction/conversion commands for sequence positions from several Entrez databases.

### EDirect Version 5.00: September 26, 2016

- Efetch -format fasta removes blank lines between records.
- Xtract -insdx uses a dash to indicate a missing field.
- Xtract -insd no longer has blank lines between records.
- Xtract -input allows reading XML data from a file.

## EDirect Version 4.90: September 14, 2016

- Epost -input allows reading from an input file instead of using data piped through stdin.
- Efilter now supports the -sort argument.
- Xtract -filter can recover information in XML comments and CDATA blocks.

## EDirect Version 4.80: August 9, 2016

• Xtract -insd controlled vocabularies updated.

### EDirect Version 4.70: August 4, 2016

- Einfo -db request can also display -fields and -links data summaries.
- Einfo -dbs prints database names instead of eInfoResult XML.

## EDirect Version 4.60: July 18, 2016

- Elink -cmd acheck returns information on all available links for a record.
- Efilter -pub structured limits to articles with structured abstracts.

# EDirect Version 4.50: July 1, 2016

- Esearch and Efilter detect and report -query phrase quotation errors.
- Efilter -pub shortcut adds last\_week, last\_month, and last\_year choices.
- Efetch sets -strand 2 for minus strand if -seq\_start > -seq\_stop or if -chr\_start > chr\_stop.

### EDirect Version 4.40: June 21, 2016

- Transitioning to use of https for access to NCBI services.
- Epost -db assembly -format acc uses [ASAC] field instead of [ACCN].

### EDirect Version 4.30: June 13, 2016

- Efilter -pub preprint limits results to ahead-of-print articles.
- Xtract -pattern Parent/\* construct can now process catenated XML files.

### EDirect Version 4.20: May 24, 2016

- Xtract command-line argument parsing improvements.
- Nquire -get supersedes -http get.

### EDirect Version 4.10: May 3, 2016

• Xtract -format removes multi-line XML comments and CDATA blocks.

## EDirect Version 4.00: April 4, 2016

- Esearch adds -spell to correct known misspellings of biological terms in the query string.
- Efilter adds -spell to correct query misspellings, and -pub, -feature, -location, molecule, -organism, and -source shortcuts. Run efilter -help to see the choices available for each argument.

## EDirect Version 3.90: March 21, 2016

• Code optimizations for increased Xtract speed.

### EDirect Version 3.80: February 29, 2016

• Xtract can distribute its work among available processor cores for additional speed.

## EDirect Version 3.70: February 8, 2016

• Xtract performance improvements.

## EDirect Version 3.60: January 11, 2016

• The setup.sh configuration script now downloads a precompiled Xtract executable for selected platforms.

## EDirect Version 3.50: December 27, 2015

• Xtract reports error for element:value construct outside of -match or -avoid arguments.

### EDirect Version 3.40: December 20, 2015

• Xtract -insd supports extraction from multiple features (e.g., CDS,mRNA).

### EDirect Version 3.30: December 3, 2015

• Efetch -format docsum can accept a single sequence accession number in the -id argument.

### EDirect Version 3.20: November 30, 2015

• Xtract supports -match conditional execution on values recorded in variables.

### EDirect Version 3.10: November 18, 2015

• Efetch adds -chr\_start and -chr\_stop arguments to specify sequence range from 0-based coordinates in gene docsum GenomicInfoType object.

### EDirect Version 3.00: October 30, 2015

- Xtract rewritten in the Go programming language for speed. The setup.sh configuration script installs an older Perl version (2.99) if a local Go compiler is unavailable.
- Efetch -format docsum only decodes HTML entity numbers in select situations.

## EDirect Version 2.90: October 15, 2015

• Xtract warns on use of deprecated arguments -present, -absent, and -trim, in preparation for release of much faster version.

### EDirect Version 2.80: September 9, 2015

• Xtract uses the "\*/Child" construct for nested exploration into recursive structures, replacing the -trim argument.

## EDirect Version 2.70: July 14, 2015

 Added entrez-phrase-search script to query on adjacent word pairs indexed in specific fields.

### EDirect Version 2.60: June 23, 2015

 $\bullet \quad X tract \ -match \ and \ -avoid \ support \ "Parent/Child" \ construct \ for \ BLAST \ XML.$ 

# EDirect Version 2.50: April 9, 2015

• Xtract capitalized -Pattern handles recursively-defined top-level objects.

### EDirect Version 2.40: March 25, 2015

• EDirect programs use the http\_proxy environment variable to work behind firewalls.

### EDirect Version 2.30: March 11, 2015

- Cleaned up logic in setup.sh configuration script.
- EPost -format acc works properly on protein accessions.

### EDirect Version 2.20: March 4, 2015

• Xtract -match and -avoid recognize "@attribute" without element or value.

## EDirect Version 2.10: February 3, 2015

 Added ftp-ls and ftp-cp scripts for convenient access to the NCBI anonymous ftp server.

### EDirect Version 2.00: August 28, 2014

• Introduced copy-and-paste installation commands with setup.sh configuration script.

### EDirect Version 1.90: August 8, 2014

- Xtract -format combines multiple XML results into a single valid object.
- Improved suppression of 0-count failure messages with -silent flag in scripts.

# EDirect Version 1.80: July 15, 2014

• EPost -format acc accepts accessions in an -id argument on the command line.

## EDirect Version 1.70: April 23, 2014

• EFetch -format docsum decodes HTML entity numbers embedded in the text.

# EDirect Version 1.60: April 3, 2014

• Minor enhancements to xtract -insd.

## EDirect Version 1.50: March 29, 2014

- Esearch -sort specifies the order of results when records are retrieved.
- Xtract exploration arguments (e.g., -block) now work on self-closing tags with data in attributes.

## EDirect Version 1.40: March 17, 2014

• Xtract -format repairs XML line-wrapping and indentation.

• Implemented -help flag to display the list of command-line arguments for each function.

### EDirect Version 1.30: March 3, 2014

• Xtract -insd partial logic was corrected to examine both 5' and 3' partial flags, and the location indicator recognizes "+" or "complete" and "-" or "partial".

### EDirect Version 1.20: February 26, 2014

Xtract -insd detects if it is part of an EDirect sequence record query, and
dynamically executes the extraction request for specific qualifier values. When run
in isolation it generates extraction instructions that can be incorporated (with
modifications, if necessary) into other queries.

### EDirect Version 1.10: February 10, 2014

- ESummary was replaced by "efetch -format docsum" to provide a single command for all document retrieval. The esummary command will continue to work for those who prefer it, and to avoid breaking existing scripts.
- Xtract processes each -pattern object immediately upon receipt, eliminating the need for using xargs and sh to split document retrieval into smaller units.

## EDirect Version 1.00: February 6, 2014

• Initial public release.

## For More Information

## **Announcement Mailing List**

NCBI posts general announcements regarding the E-utilities to the <u>utilities-announce</u> <u>announcement mailing list</u>. This mailing list is an announcement list only; individual subscribers may **not** send mail to the list. Also, the list of subscribers is private and is not shared or used in any other way except for providing announcements to list members. The list receives about one posting per month. Please subscribe at the above link.

### Getting Help

Please refer to the PubMed and Entrez help documents for more information about search queries, database indexing, field limitations and database content.

Suggestions, comments, and questions specifically relating to the EUtility programs may be sent to <a href="mailto:eutilities@ncbi.nlm.nih.gov">eutilities@ncbi.nlm.nih.gov</a>.