EnsEMBL Compara Perl API Tutorial

By Cara Woodwark, Abel Ureta-Vidal and Javier Herrero

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WARNING: this is a 'test' version. By now this tutorial is 'warranty' work with branch-ensembl-26, and with ensembl databases release 26. As it is a 'test' version, you may find errors. Please email ensembl-dev@ebi.ac.uk, so that we can correct them. We will be extending/completing this tutorial in the near future.

Introduction

This tutorial is an introduction to the ensembl compara API. A knowledge of the ensembl core API is presumed, it is assumed that concepts and conventions presented in the ensembl core API tutorial have been assimilated by the user. The ensembl core API tutorial can be found at http://www.ensembl.org/Docs/linked_docs/ensembl_tutorial.pdf (in cvs, in ensembl/docs/tutorial/ensembl_tutorial.pdf) and should be read first as it provides a comprehensive guide to the ensembl environment.

A documentation about the compara database schema is available http://www.ensembl.org/???? (in cvs ensembl-compara/docs/docs/schema_doc.html), and while not necessary for this tutorial, an understanding of the database tables may help, as many of the Adaptor modules are table specific.

Obtaining the code

To use the ensembl compara API, you have the same requirement that when using the ensembl core API i.e. perl 5.6 or later, bioperl 1.2 or later, DBI, DBD::mysql and ensembl core code. Please refer to the ensembl core API tutorial that will tell you everything about these modules, how and where to get them.

You may start by creating a directory for storing the API in your home directory:

```
cd
mkdir src
mv src
```

In addition, you will need the ensembl compara code that is available by cvs from the ensembl cvs repository using the following cvs commands:

```
cvs -d :pserver:cvsuser@cvsro.sanger.ac.uk:/cvsroot/CVSmaster login
```

When prompted the password is 'CVS'.

```
\verb|cvs -d :pserver:cvsuser@cvsro.sanger.ac.uk:/cvsroot/CVSmaster co -r branch-ensembl-26| ensembl-compara| \\
```

This will check out ensembl-compara code for stable branch 26. Make sure the ensembl core code you have already checked out is on the same branch. Note that the branch that is checked out should correspond to the database version being used. Thus ensembl_compara_26_1 and e.g. homo-sapiens_core_26_35 and mus_musculus_core_26_33b should be used with the above ensembl branch 26 code.

Environment Variables

The following PERL5LIB environment variables should be set up:

- under tcsh/csh shell with

```
setenv PERL5LIB ${PERL5LIB}:{HOME}/src/bioperl-live: \
${HOME}/src/ensembl/modules:${HOME}/src/ensembl-compara/modules
```

- under bash shell with

```
export PERL5LIB=${PERL5LIB}:{HOME}/src/bioperl-live: \
${HOME}/src/ensembl/modules:${HOME}/src/ensembl-compara/modules
```

These presume that bioperl and ensembl are in a directory called src set up in your home directory.

Code Conventions (and unconventions)

Refer to the ensembl core tutorial for a good description of the coding conventions normally used in ensembl. Due to historical accidents, there may be exceptions to these rules in compara.

Connecting a ensembl compara database

There are two ways to connect to the EnsEMBL Compara database. The old way uses the Bio::EnsEMBL::Compara::DBSQL::DBAdaptor explicitely. The new one uses the Bio::EnsEMBL::Registry module which can read either a global or a specific configuration file.

Explicitely, using the Bio::EnsEMBL::Compara::DBSQL::DBAdaptor

Ensembl compara data as ensembl core data, is stored in a MySQL relational database. If you want to access a compara database, you will need to connect to it. This is done in exactly the same way as when connecting an ensembl core database, but using a Compara specific DBAdaptor.

As for a ensembl core connection, in addition to the parameters provided above, the optional port, driver and pass parameters can also be used to specify the TCP connection port, the type of database driver and the password respectively. These values have sensible defaults and can often be omitted.

Implicitely, using the Bio::EnsEMBL::Registry configuration file (recommended)

You will need to have a registry configuration file set up. By default, it takes the file defined by the ENSEMBL_REGISTRY environment variable or the file named .ensembl_init in your home directory if the former is not found. Additionally, it is possible to use a specific file (see perldoc Bio::EnsEMBL::Registry or later in this document for some examples on how to use a different file). An example of such file can be found in ensembl/modules/Bio/EnsEMBL/Utils/ensembl_init.example, and below you have a slightly modified copy of it.

```
# Example of configuration file used by Bio::EnsEMBL::Registry::load_all
# method to store/register all kind of Adaptors.
```

```
use Bio::EnsEMBL::Utils::ConfigRegistry;
use Bio::EnsEMBL::DBSQL::DBAdaptor;
use Bio::EnsEMBL::Compara::DBSQL::DBAdaptor;
my @aliases;
new Bio::EnsEMBL::DBSQL::DBAdaptor(-host => 'ensembldb.ensembl.org',
                                   -user => 'anonymous',
                                   -port => 3306,
                                   -species => 'Homo sapiens',
                                   -group => 'core',
                                   -dbname => 'homo_sapiens_core_26_35');
@aliases = ('H_Sapiens', 'homo sapiens', 'Homo_Sapiens', 'Homo_sapiens', 'Homo',
'homo', 'human');
Bio::EnsEMBL::Utils::ConfigRegistry->add_alias(-species => "Homo sapiens",
                                               -alias => \@aliases);
new Bio::EnsEMBL::DBSQL::DBAdaptor(-host => 'ensembldb.ensembl.org',
                                   -user => 'anonymous',
                                   -port => 3306,
                                   -species => 'Mus musculus',
                                   -group => 'core',
                                   -dbname => 'mus_musculus_core_26_33b');
@aliases = ('M_Musculus', 'mus musculus', 'Mus_Musculus', 'Mus_musculus', 'Mus', 'mus',
'mouse');
Bio::EnsEMBL::Utils::ConfigRegistry->add_alias(-species => "Mus musculus",
                                               -alias => \@aliases);
new Bio::EnsEMBL::DBSQL::DBAdaptor(-host => 'ensembldb.ensembl.org',
                                   -user => 'anonymous',
                                   -port => 3306,
                                   -species => 'Fugu rubripes',
                                   -group => 'core',
                                   -dbname => 'fugu_rubripes_core_26_2c');
@aliases = ('F_Rubripes', 'fugu rubripes', 'Fugu_Rubripes', 'Fugu_rubripes', 'Fugu',
'fugu');
Bio::EnsEMBL::Utils::ConfigRegistry->add_alias(-species => "Fugu rubripes",
                                                -alias => \@aliases);
new Bio::EnsEMBL::Compara::DBSQL::DBAdaptor(-host => 'ensembldb.ensembl.org',
                                             -user => 'anonymous',
                                            -port => 3306,
                                             -species => 'Compara26',
                                            -dbname => 'ensembl_compara_26_1');
@aliases = ('ensembl_compara_26_1', 'compara26');
Bio::EnsEMBL::Utils::ConfigRegistry->add_alias(-species => "Compara26",
                                               -alias => \@aliases);
1;
```

In this configuration file, you can list all the parameters needed to connect a compara

database. The compara database is a multi-species database that contains comparative genomic information on all ensembl species. One should then be able not only to connect to a compara database but also to every species ensembl core database. The use of the registry configuration file lets you the freedom to list connection parameters for all ensembl core databases you might need to access in relation to ensembl compara data (in our example, only 3 are mentioned, human, mouse and fugu). All this information is then in a single central place, easy to maintain (modify and update).

The access to database adaptor is done using either the main species alias (specified by the -species parameter) or one of the aliases specified (in the @aliases array). No need to remember the complete database name, one of the aliases will be enough.

WARNING: In previous version of this tutorial, an additional parameter disconnect_when_inactive => 1 was specified for all ensembl core databases. It is not needed anymore, as there is now a lazy connection in place i.e. connection will be established only at your first prepare statement and kept alive until you use a disconnect_if_idle (or a more drastic disconnect). If you want to use disconnect_when_inactive make sure you know what you are doing.

Below is a non exhaustive list of ensembl compara adaptors that are most often used

```
to fetch Bio::EnsEMBL::Compara::GenomeDB objects
GenomeDBAdaptor
DnaFragAdaptor
                              to fetch Bio::EnsEMBL::Compara::DnaFrag objects
                              to fetch Bio::EnsEMBL::Compara::GenomicAlignBlock objects
GenomicAlignBlockAdaptor
                              to fetch Bio::EnsEMBL::DnaDnaAlignFeature objects
DnaAlignFeatureAdaptor
                              (note that this adaptor return a ensembl core object)
                              to fetch Bio::EnsEMBL::Compara::SyntenyRegion objects
SyntenyAdaptor
                              to fetch Bio::EnsEMBL::Compara::Member objects
MemberAdaptor
                              to fetch Bio::EnsEMBL::Compara::Homology objects
HomologyAdaptor
                              to fetch Bio::EnsEMBL::Compara::Family objects
FamilyAdaptor
                              to fetch Bio::EnsEMBL::Compara::PeptideAlignFeature
PeptideAlignFeatureAdaptor
                              objects
```

Only some of these adaptors will be used for illustration as part of this tutorial through commented perl scripts code.

Whole Genome Alignments

The compara database contains a number of different types of whole genome alignments. A listing about what are these different types can be found in the ensembl-compara/docs/schema_doc.html document in method_link section.

The whole genome comparisons can be accessed through the API by 2 different ways using of the 2 different adaptors. Specifically, the DnaAlignFeatureAdaptor, which returns DnaDnaAlignFeatures objects (only used for pairwise alignment) and the GenomicAlignBlockAdaptor, which returns GenomicAlignBlock objects (can be used for pairwise and also multiple alignments).

DnaDnaAlignFeature objects (for pairwise alignments only)

Below it is a simple commented perl script to illustrate the use of DnaDnaAlignFeature objects.

```
use strict;
use Bio::EnsEMBL::Registry;
use Bio::EnsEMBL::Compara::DBSQL::DBAdaptor;
use Bio::AlignIO;
use Bio::LocatableSeq;
use Getopt::Long;
```

```
my $usage = "
  [--help]
                                this menu
                                (e.g. compara23) one of the compara database
   --dbname string
                                Bio::EnsEMBL::Registry aliases
   --seq_region string
                                (e.g. 22)
   --seq_region_start integer
                                (e.g. 50000000)
   --seq_region_end integer
                                (e.g. 50500000)
                                (e.g. human) the query species (i.e. {\tt a}
   --qy string
                                Bio::EnsEMBL::Registry alias) from which alignments
                                are queried and seq_region refer to
   --tg string
                                (e.g. mouse) the target sepcies (i.e. a
                                Bio::EnsEMBL::Registry alias) to which alignments are
                                queried
  [--alignment_type string]
                                (e.g. TRANSLATED_BLAT) type of alignment stored
                                (default: BLASTZ_NET)
  [--tsl]
                                print out a translated alignment
  [--00]
                                By default, the alignments are dumped so that the --qy
                                species sequence is always on forward strand. --oo is
                                mostly useful in association with -tsl option, when a
                                full translated alignment program has been used e.g
                                TRANSLATED_BLAT, and allow to obtain the right
                                translation phase. So the --qy species sequence might
                                be reverse complemented.
  [--ft string]
                                alignment format, available in bioperl Bio::AlignIO
                                (default: clustalw)
  [--ucl
                                print out sequence in upper cases (default is lower
                                cases)
  [--limit integer]
                                (e.g. 2) limit the output to the number of alignments
                                specified
                                the Bio::EnsEMBL::Registry configuration file. If none
  [--reg_conf filepath]
                                given, the one set in ENSEMBL_REGISTRY will be used if
                                defined, if not ~/.ensembl_init will be used.
\n";
my $dbname;
my ($seq_region,$seq_region_start,$seq_region_end);
my ($qy_species,$tg_species);
my $help = 0;
my $alignment_type = "BLASTZ_NET";
my $limit;
my $reg_conf;
my $format = "clustalw";
my $translated = 0;
my \$uc = 0;
my $original_orientation = 0;
unless (scalar @ARGV) {
 print $usage;
  exit 0;
}
GetOptions('help' => \$help,
           'dbname=s' => \$dbname,
           'seq_region=s' => \$seq_region,
           'seq_region_start=i' => \$seq_region_start,
           'seq_region_end=i' => \$seq_region_end,
           'qy=s' => \$qy_species,
           'tg=s' => \$tg_species,
```

```
'alignment_type=s' => \$alignment_type,
          'tsl' => \$translated,
          'ft=s' => \$format,
          'uc' => \$uc,
          'oo' => \$original_orientation,
          'limit=i' => \$limit,
          'reg_conf=s' => \$reg_conf);
$|=1;
if ($help) {
 print $usage;
 exit 0;
# Setting up Bio::EnsEMBL::Regitry
# if $reg_conf is undef, ~/.ensembl_init will be loaded if it exists
Bio::EnsEMBL::Registry->load_all($reg_conf);
$format = lc $format;
# Getting the core SliceAdaptor for the query species
my $qy_sa = Bio::EnsEMBL::Registry->qet_adaptor($qy_species,'core','Slice');
# Fetching a Slice. In compara, all slices are 'toplevel' coordinate system.
my $qy_slice = $qy_sa->fetch_by_region('toplevel',$seq_region,
                                    $seq_region_start,$seq_region_end);
# Getting the core MetaContainer adaptor for the target species
my $tg_mc = Bio::EnsEMBL::Registry->get_adaptor($tg_species,'core','MetaContainer');
# Getting a Bio:: Species object and from it the Species genus (e.g. Mus
# musculus) of the target species, using the binomial call
my $tg_binomial = $tg_mc->get_Species->binomial;
# Getting the compara DnaAlignFeatureAdaptor to query the compara database
my $dafad = Bio::EnsEMBL::Registry->get_adaptor($dbname,'compara','DnaAlignFeature');
# Fetching DnaDnaAlignFeatures object (these are core objects) using the
# fetch_all_by_Slice. The 3rd argument that can specify the assembly version
# can be undef. The compara API will find for you the default assembly for
# the target species.
my $DnaDnaAlignFeatures =
$dafad->fetch_all_by_Slice($qy_slice,$tg_binomial,undef,$alignment_type,$limit);
# Go through each alignment to print out in the requested format
foreach my $ddaf (sort {$a->start <=> $b->start
```

```
|| $a->end <=> $b->end}
                @{$DnaDnaAlignFeatures}) {
  # if the original alignment strand orientation is requested
  # ($original_orientation is true) and effectively the alignment obtained
  # is reverse complement from the originally obtained by the alignment
  # program used (if $ddaf->strands_reversed is true), then reverse
  # complement the alignment.
  if ($original_orientation && $ddaf->strands_reversed) {
    $ddaf->reverse_complement;
 # Create a list of flags to be used in the get_SimpleAlign method call
 my @flags;
 push @flags, 'translated' if ($translated);
 push @flags, 'uc' if ($uc);
  # Get a Bio::SimpleAlign from the DnaDnaAlignFeature object
 my $sa = $ddaf->get_SimpleAlign(@flags);
  # Create a Bio::AlignIO with the requested output format
 my $alignIO = Bio::AlignIO->newFh(-interleaved => 0,
                                 -fh => \*STDOUT,
                                 -format => $format,
                                 -idlength => 20);
  # print out the alignment (Bio::SimpleAlign object) in the requested
  # output format through the Bio::AlignIO handler
 print $alignIO $sa;
exit 0;
```

So to pull out BLASTZ_NET_TIGHT alignments, let's say on part of ENCODE region ENm004 on human chromosome 22, between position 30184430 and position 30184485, against the mouse genome in clustalw format, we can use know the following command line,

Now on the same region, TRANSLATED_BLAT alignments against fugu in clustalw format, but at translation level now (-tsl) not nucleotide level, we can run the following command line.

By default, the alignments will dump with --qy species sequence on forward strand. To make sure that the alignment, you got is on the strand on which it was originally generated using the --oo option will check that and restore the right strandness. See below the difference in the translation level alignment obtained.

GenomicAlignBlock objects (pairwise/multiple alignments)

GenomicAlignBlocks are the new way to store and fetch genomic alignments. A GenomicAlignBlock contains several GenomicAlign objects. Every GenomicAlign object corresponds to a piece of genomic sequence aligned with the other GenomicAlign in the same GenomicAlignBlock. A GenomicAlign object is always related with other GenomicAlign objects and this relation is defined through the GenomicAlignBlock object. Therefore the usual way to fetch genomic alignments is by fetching GenomicAlignBlock objects. We have to start by getting the corresponding adaptor:

Getting the GenomicAlignBlock adaptor:

In order to fetch the right alignments we need to specify a couple of data: the type of alignment and the piece of genomic sequence in which we are looking for alignments. The type of alignment is a more tricky now: you need to specify both the alignment method and the set of genomes. In order to simply this task, you could use the new Bio::EnsEMBL::Compara::MethodLinkSpeciesSet object. The best way to use them is by fetching them from the database:

```
# Getting the GenomeDB adaptor:
```

There are two ways to fetch GenomicAlignBlocks. One is uses Bio::EnsEMBL::Slice objects while the second one is based on Bio::EnsEMBL::Compara::DnaFrag objects for specifying the piece of genomic sequence in which we are looking for alignments.

```
# Getting the Slice adaptor:
```

);

Here is an example script with all of this:

```
use strict;
use Bio::EnsEMBL::Registry;
use Bio::EnsEMBL::Utils::Exception qw(throw);
use Bio::SimpleAlign;
use Bio::AlignIO;
use Bio::LocatableSeq;
use Getopt::Long;

my $usage = qq{
perl DumpMultiAlign.pl
   Getting help:
    [--help]

General configuration:
```

```
[--reg_conf registry_configuration_file]
        the Bio::EnsEMBL::Registry configuration file. If none given,
        the one set in ENSEMBL_REGISTRY will be used if defined, if not
        ~/.ensembl_init will be used.
    [--dbname compara_db_name]
        the name of compara DB in the registry_configuration_file or any
        of its aliases. Uses "compara" by default.
  For the query slice:
    [--species species]
       Query species. Default is "human"
    [--coord_system coordinates_name]
       Query coordinate system. Default is "chromosome"
    --seq_region region_name
        Query region name, i.e. the chromosome name
    --seq_region_start start
    --seq_region_end end
  For the alignments:
    [--alignment_type method_link_name]
        The type of alignment. Default is "BLASTZ_NET"
    [--set_of_species species1:species2:species3:...]
        The list of species used to get those alignments. Default is
        "human:mouse". The names should correspond to the name of the
        core database in the registry_configuration_file or any of its
        aliases
  Ouput:
    [--output_format clustalw|fasta|...]
        The type of output you want. "clustalw" is the default.
    [--output_file filename]
        The name of the output file. By default the output is the
        standard output
};
my $reg_conf;
my $dbname = "compara";
my $species = "human";
my $coord_system = "chromosome";
my $seq_region = "14";
my $seq_region_start = 75000000;
my $seq_region_end = 75010000;
my $alignment_type = "BLASTZ_NET";
my $set_of_species = "human:mouse";
my $output_file = undef;
my $output_format = "clustalw";
my $help;
GetOptions(
    "help" \Rightarrow \$help,
    "reg_conf=s" => \$reg_conf,
    "dbname=s" => \$dbname,
    "species=s" => \$species,
    "coord_system=s" => \$coord_system,
    "seq_region=s" => \$seq_region,
    "seq_region_start=i" => \$seq_region_start,
    "seq_region_end=i" => \$seq_region_end,
    "alignment_type=s" => \$alignment_type,
    "set_of_species=s" => \$set_of_species,
    "output_format=s" => \$output_format,
```

```
"output_file=s" => \$output_file,
 );
# Print Help and exit
if ($help) {
 print $usage;
  exit(0);
if ($output_file) {
  open(STDOUT, ">$output_file") or die("Cannot open $output_file");
# Configure the Bio::EnsEMBL::Registry
# Uses $req_conf if supllied. Uses ENV{ENSMEBL_REGISTRY} instead if defined.
# Uses ~/.ensembl_init if all the previous fail.
Bio::EnsEMBL::Registry->load_all($reg_conf);
# Getting all the Bio::EnsEMBL::Compara::GenomeDB objects
my $genome_dbs;
my $genome_db_adaptor = Bio::EnsEMBL::Registry->get_adaptor($dbname, 'compara',
       'GenomeDB');
throw("Registry configuration file has no data for connecting to <$dbname>")
       if (!$genome_db_adaptor);
foreach my $this_species (split(":", $set_of_species)) {
  my $this_meta_container_adaptor = Bio::EnsEMBL::Registry->qet_adaptor(
      $this_species, 'core', 'MetaContainer');
  throw("Registry configuration file has no data for connecting to <$this_species>")
      if (!$this_meta_container_adaptor);
 my $this_binomial_id = $this_meta_container_adaptor->get_Species->binomial;
  # Fetch Bio::EnsEMBL::Compara::GenomeDB object
  my $genome_db = $genome_db_adaptor->fetch_by_name_assembly($this_binomial_id);
  # Add Bio::EnsEMBL::Compara::GenomeDB object to the list
 push(@$genome_dbs, $genome_db);
# Getting Bio::EnsEMBL::Compara::MethodLinkSpeciesSet obejct
my $method_link_species_set_adaptor = Bio::EnsEMBL::Registry->get_adaptor(
       $dbname, 'compara', 'MethodLinkSpeciesSet');
my $method_link_species_set =
       $method_link_species_set_adaptor->fetch_by_method_link_type_GenomeDBs(
              $alignment_type, $genome_dbs);
throw("The database do not contain any $alignment_type data for $set_of_species!")
       if (!$method_link_species_set);
# Fetching the query Slice:
my $slice_adaptor = Bio::EnsEMBL::Registry->get_adaptor($species, 'core', 'Slice');
throw("Registry configuration file has no data for connecting to <$species>")
       if (!$slice_adaptor);
my $query_slice = $slice_adaptor->fetch_by_region('toplevel', $seq_region,
$seq_region_start, $seq_region_end);
throw("No Slice can be created with coordinates $seq_region:$seq_region_start-".
       "$seq_region_end") if (!$query_slice);
# Fetching all the GenomicAlignBlock corresponding to this Slice:
my $genomic_align_block_adaptor = Bio::EnsEMBL::Registry->get_adaptor(
    $dbname, 'compara', 'GenomicAlignBlock');
my $genomic_align_blocks =
```

```
$genomic_align_block_adaptor->fetch_all_by_MethodLinkSpeciesSet_Slice(
        $method_link_species_set, $query_slice);
my $all_aligns;
# Create a Bio::SimpleAlign object from every GenomicAlignBlock
foreach my $this_genomic_align_block (@$genomic_align_blocks) {
  my $simple_align = Bio::SimpleAlign->new();
  $simple_align->id("GAB#".$this_genomic_align_block->dbID);
  $simple_align->score($this_genomic_align_block->score);
  my $all_genomic_aligns = $this_genomic_align_block->get_all_GenomicAligns;
  # Create a Bio::LocatableSeq object from every GenomicAlign
  foreach my $this_genomic_align (@$all_genomic_aligns) {
   my $seq_name = $this_genomic_align->dnafrag->genome_db->name;
   seq_n = - s/(.) w* (.) w*/$1$2/;
   $seq_name .= $this_genomic_align->dnafrag->name;
   my $aligned_sequence = $this_genomic_align->aligned_sequence;
   my $seq = Bio::LocatableSeq->new(
           -SEO
                 => $aligned_sequence,
           -START => $this_genomic_align->dnafrag_start,
           -END => $this_genomic_align->dnafrag_end,
                  => $seq_name,
           -ID
           -STRAND => $this_genomic_align->dnafrag_strand
       );
    # Add this Bio::LocatableSeq to the Bio::SimpleAlign
    $simple_align->add_seq($seq);
 push(@$all_aligns, $simple_align);
# print all the genomic alignments using a Bio::AlignIO object
my $alignIO = Bio::AlignIO->newFh(
       -interleaved => 0,
       -fh => \*STDOUT,
       -format => $output_format,
       -idlength => 10
foreach my $this_align (@$all_aligns) {
 print $alignIO $this_align;
exit:
```

Orthologues and Protein clusters

NB: This following is very much a draft at this stage with some piece of code to give examples, but not much comments.

Member objects

```
# get the MemberAdaptor
my $ma = Bio::EnsEMBL::Registry->get_adaptor($dbname,'compara','Member');
# fetch a Member
my $member = $ma->fetch_by_source_stable_id('ENSEMBLGENE','ENSG000000004059');
# print out some information about the Member
print join " ", map { $member->$_ } qw(chr_name chr_start chr_end description
```

```
source_name taxon_id taxon),"\n";
```

chr_name, chr_start, chr_end and description are self-explanatory.

source_name tells about the origin of the Member entry, and can be either

ENSEMBLPEP, derived from ensembl translation,

- or ENSEMBLGENE, derived from an ensembl gene,
- or SWISSPROT, derived from a Uniprot/Swissprot entry,
- or SPTREMBL, derived from a Uniprot/SP-TrEMBL entry.

taxon_id e.g. 9606 correspond to the NCBI taxonomy identifier (see <a href="http://www.ncbi.nlm.nih.gov/Taxonomy/taxonom

taxon returns a Bio::EnsEMBL::Compara::Taxon object that inherits itself from Bio::Species, so from this object you can get additional information about the species.

```
my $taxon = $member->taxon;
print join "; ", map { $taxon->$_ } qw(common_name genus species binomial
classification),"\n";
```

respectively for these method calls and in the case of human species, you will obtain

human; Homo; sapiens; Homo sapiens; sapiens Homo Hominidae Catarrhini Primates Eutheria Mammalia Euteleostomi Vertebrata Craniata Chordata Metazoa Eukaryota

Homology objects

```
# first you have to get a Member object. In case of homology is a gene, in
# case of family it can be a gene or a protein
my $ma = Bio::EnsEMBL::Registry->get_adaptor($dbname,'compara','Member');
my $member = $ma->fetch_by_source_stable_id('ENSEMBLGENE','ENSG00000004059');
# then you get the homologies where the member is involved
my $ha = Bio::EnsEMBL::Registry->get_adaptor($dbname,'compara','Homology');
my $homologies = $ha->fetch_by_Member($member);
fetch_by_Member_Homology_source (fetch_by_Member_MethodLink)
# That will return an array reference with all homologies (orthologues, and
# in some cases paralogues) against other species.
# Then for each homology, you get all the Members implicated
foreach my $homology (@{$homologies}) {
# You will find different kind of description
# UBRH, MBRH, MBRH, RHS, YoungParalogues
# see ensembl-compara/docs/docs/schema doc.html for more details
print $homology->description," ", $homology->subtype,"\n";
```

And if they are defined dN and dS related values

print join " ", map { \$homology->\$_ } qw(dn ds n s lnl threshold_on_ds),"\n";

```
# each homology relation have only 2 members, you should find there
# the initial member used in the first fetching
for each my $member_attribute (@{$homology->get_all_Member_Attribute})
# for each Member, you get information on the Member specifically and in
# relation to the homology relation via Attribute object
   my ($member, $attribute) = @{$member_attribute};
   print join " ", map { $member->$_ } qw(stable_id taxon_id),"\n";
   print join " ", map { $attribute->$_ } qw(perc_id perc_pos perc_cov),"\n";
# You can even retrieve the HSP alignment between the 2 proteins,
# HSP used to build the homology releationship at the peptide level
 my $sa = $homology->get_SimpleAlign();
 my $alignIO = Bio::AlignIO->newFh(-interleaved => 0,
                                  -fh \Rightarrow \*STDOUT,
                                  -format => "clustalw",
                                  -idlength => 20);
 print $alignIO $sa;
# or at the nucleotide level. You will need to make you have a connection to
# the corresponding core databases through the Bio::EnsEMBL::Registry
 $sa = $homology->get_SimpleAlign('cdna');
 my $alignIO = Bio::AlignIO->newFh(-interleaved => 0,
                                  -fh => \*STDOUT,
                                  -format => "phylip",
                                  -idlength => 20);
 print $alignIO $sa;
}
```

Family objects

You can obtain them in the same way as Homology objects

```
my $ma = Bio::EnsEMBL::Registry->get_adaptor($dbname,'compara','Member');
my $member = $ma->fetch_by_source_stable_id('ENSEMBLGENE','ENSG000000004059');

my $fa = Bio::EnsEMBL::Registry->get_adaptor($dbname,'compara','Family');
my $families = $fa->fetch_by_Member($member);

foreach my $family (@{$families}) {
  print join " ", map { $family->$_ } qw(description description_score),"\n";

  for each my $member_attribute (@{$family->get_all_Member_Attribute})
    my ($member, $attribute) = @{$member_attribute};
   print $member->stable_id," ",$member->taxon_id,"\n";
  }

my $sa = $family->get_SimpleAlign();
my $alignIO = Bio::AlignIO->newFh(-interleaved => 0,
```

```
-fh => \*STDOUT,
-format => "phylip",
-idlength => 20);

print $alignIO $sa;

$sa = $family->get_SimpleAlign('cdna');
my $alignIO = Bio::AlignIO->newFh(-interleaved => 0,
-fh => \*STDOUT,
-format => "phylip",
-idlength => 20);

print $alignIO $sa;
```

Further help

For additional information or help mail ensemb-dev@ebi.ac.uk. You will need to subscribe to this mailing list to use it (see how to subscribe in http://www.ensembl.org/Docs/Lists/).