# Ionotropic receptors as the driving force behind human synapse establishment

# Supplementary Material

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#### Abstract

The origin of nervous systems is a main theme in biology and its mechanisms are largely underlied by synaptic neurotransmission. The search for pan-neuronal genes has failed to explain synapses emergence since synaptic elements are present in multiple aneural organisms. We questioned how the interactions among these elements evolved and to what extent does it relate to our understanding of the nervous systems complexity. We infer the human neurotransmission gene network based on genes present in GABAergic, glutamatergic, serotonergic, dopaminergic, and cholinergic systems, and reconstruct the evolutionary scenario of synapse emergence. Our results outline the distribution of neurotransmitter systems and synaptic functions as taxa diverge. We find that the ionotropic receptors emergence in the chidarian last common ancestor was the driving force behind the advent of the anatomical synapse and the development of nervous systems. We suggest that after Placozoa origin there was a positive selection under a more restricted and specific activation of ionotropic receptors. This different selective pressure counterparting the broader amino acid activation in plants could be striking for the evolution of the nervous system.

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# Project structure

This is the title page

# Preprocessing

This topic refers mainly to data wrangling done before the actual analysis with the intent of making it simpler.

# Eukaryota species tree

We opted to use the TimeTree database in order to obtain an standardized Eukaryota species tree. However, some species were not present in it, so we devised a way to fill them in based on NCBI Taxonomy data.

# NCBI Taxonomy tree

First we preprocess NCBI Taxonomy data to leave only STRING eukaryotes, thus making the task easier.

#### Resources

Table 1: Lists all organisms in STRING v11.

	string_species									
#	# Col. name Col. type Used? Example Description									
1	taxid	character	yes	9606	NCBI Taxonomy identifier					
2	string_type	character	no	core	if the genome of this species is core or periphery					
3	string_name	character	yes	Homo sapiens	STRING species name					
4	$ncbi\_official\_name$	character	no	Homo sapiens	NCBI Taxonomy species name					

**Location:** data-raw/download/species.v11.0.txt

Source: stringdb-static.org/download/species.v11.0.txt

Table 2: Links outdated taxon IDs to corresponding new ones.

ncbi_merged_ids									
#	Col. name Col. type Used? Example Description								
1	taxid	character	yes	140100	id of node that has been merged				
2	${\it new\_taxid}$	character	yes	666	id of node that is the result of merging				

Location: data-raw/download/taxdump/merged.dmp

Source: ftp.ncbi.nlm.nih.gov/pub/taxonomy/taxdump.tar.gz

Table 3: Represents taxonomy nodes.

	ncbi_edgelist								
#	# Col. name Col. type Used? Example Description								
1	taxid	character	yes	2	node id in NCBI taxonomy database				
2	parent_taxid	character	yes	131567	parent node id in NCBI taxonomy database				
3	rank	character	no	superkingdom	rank of this node				
4			no		(too many unrelated fields)				

Location: data-raw/download/taxdump/nodes.dmp

Source: ftp.ncbi.nlm.nih.gov/pub/taxonomy/taxdump.tar.gz

Table 4: Links taxon IDs to actual species names.

	ncbi_taxon_names								
#	Col. name	Col. type	Used?	Example	Description				
1	taxid	character	yes	2	the id of node associated with this name				
2	name	character	yes	Monera	name itself				
3	unique_name	character	no	Monera  kacteria>	the unique variant of this name if name not unique				
4	$name\_class$	character	yes	scientific name	type of name				

Location: data-raw/download/taxdump/names.dmp

Source: ftp.ncbi.nlm.nih.gov/pub/taxonomy/taxdump.tar.gz

### Updating STRING taxon IDs

Some organisms taxon IDs are outdated in STRING. We must update them to work with the most recent NCBI Taxonomy data.

```
string_species %<>%
  left_join(ncbi_merged_ids) %>%
  mutate(new_taxid = coalesce(new_taxid, taxid))
```

# Creating tree graph

The first step is to create a directed graph representing the NCBI Taxonomy tree.

```
# leaving only "scientific name" rows
ncbi_taxon_names %<>%
filter(type == "scientific name") %>%
select(name, ncbi_name)

# finding Eukaryota taxid
eukaryota_taxon_id <- subset(ncbi_taxon_names, ncbi_name == "Eukaryota", "name", drop = TRUE)

# creating graph
g <- graph_from_data_frame(ncbi_edgelist[,2:1], directed = TRUE, vertices = ncbi_taxon_names)

# easing memory
rm(ncbi_edgelist, ncbi_merged_ids)</pre>
```

#### Traversing the graph

The second step is to traverse the graph from the Eukaryota root node to STRING species nodes. This automatically drops all non-eukaryotes and results in a species tree representing only STRING eukaryotes (476).

```
eukaryote_root <- V(g)[eukaryota_taxon_id]
eukaryote_leaves <- V(g)[string_species[["new_taxid"]]]

# not_found <- subset(string_species, !new_taxid %in% ncbi_taxon_names$name)
eukaryote_paths <- shortest_paths(g, from = eukaryote_root, to = eukaryote_leaves, mode = "out")$vpath
eukaryote_vertices <- eukaryote_paths %>% unlist %>% unique
eukaryote_tree <- induced_subgraph(g, eukaryote_vertices, impl = "create_from_scratch")</pre>
```

#### Saving

Saving ncbi\_tree and string\_eukaryotes for package use. These data files are documented by the package. We also create a plain text file 476\_ncbi\_eukaryotes.txt containing the updated names of all 476 STRING eukaryotes. This file will be queried against the TimeTree website.

```
ncbi_tree <- treeio::as.phylo(eukaryote_tree)

# plot(ncbi_tree %>% ape::ladderize(), type="cladogram")

string_eukaryotes <- string_species %>%
    filter(new_taxid %in% ncbi_tree$tip.label) %>%
    inner_join(ncbi_taxon_names, by = c("new_taxid" = "name"))

write(string_eukaryotes[["ncbi_name"]],"476_ncbi_eukaryotes.txt")

# usethis::use_data(ncbi_tree, overwrite = TRUE)
write.tree(ncbi_tree, "tree_ncbi.nwk")
usethis::use_data(string_eukaryotes, overwrite = TRUE)
```

```
## <U+2714> Setting active project to 'C:/R/neuro'
## <U+2714> Saving 'string_eukaryotes' to 'data/string_eukaryotes.rda'
```

#### **Duplicated Genera**

Some species from different kingdoms may share the same genus name. These genera must be noted down because one of the ways we fill in missing species is by looking at genera names. For taxid\_rank and ncbi\_taxon\_names see Table 3 and Table 4, respectively.

```
# keeping genera nodes
taxid_rank %<>% filter(rank == "genus")

# keeping scientific names
ncbi_taxon_names %<>%
filter(type == "scientific name") %>%
select(taxid, ncbi_name) %>%
inner_join(taxid_rank)

# extracting and saving duplicated values
duplicated_genera <- ncbi_taxon_names %>%
pull(ncbi_name) %>%
extract(duplicated(.)) %>%
write("duplicated_genera.txt")
```

### Hybrid tree

Once we have both the NCBI eukaryotes tree and the list of duplicated genera, we can start assembling the complete hybrid tree.

#### Resources

Besides downloading all TimeTree species data (Eukaryota\_species.nwk) we also need to manually query the website for the 476 STRING eukaryotes (476\_ncbi\_eukaryotes.txt). The file is called 476\_ncbi\_eukaryotes.txt because it contains updated NCBI Taxonomy names rather than STRING outdated names. This ensures better results.

timetree\_newick is the tree obtained by manually uploading 476\_ncbi\_eukaryotes.txt to the TimeTree website. tree 85k is the complete Eukaryota tree we have just downloaded.

```
# loading species names and taxon ids
load("../data/string_eukaryotes.rda")

# loading newick tree manually obtained from timetree
timetree_newick <- read.tree("download/timetree_335_eukaryotes.nwk")

# the following genera names are unreliable and should not be searched for
duplicated_genera <- scan("duplicated_genera.txt", what = "character")

# loading all TimeTree species data we have just download (85000 species)
tree_85k <- read.tree("download/Eukaryota_species.nwk")</pre>
```

### Unfound species with matching genera

Some of the 476 STRING eukaryotes are not present in the TimeTree database. However, sometimes TimeTree does contain tree data for closely related species (e.g. *Monosiga brevicollis* is not present, but *Monosiga ovata* is). Therefore, we can use these closely related species as proxies for the actual species. This is done by searching for genera names in the complete database (Eukaryota\_species.nwk). In the given *Monosiga brevicollis* example, we search for *Monosiga* in the complete database. We see that there is information for at least one other species of the *Monosiga* genus (in this case, *Monosiga ovata*), so we add *Monosiga brevicollis* as a sister branch to the found species.

When you search for a term in TimeTree, it uses a synonym list obtained from NCBI to try to resolve it. Sometimes TimeTree will resolve a searched term to a scientific name different from the one you searched for. The problem with this is that TimeTree does not make it obvious that it is returning a different term. The first step is to find out which species resolved to different names in the timetree\_335\_eukaryotes.nwk file:

```
# replacing timetree species underscores with spaces
timetree_newick[["tip.label"]] %<>% str_replace_all("_", " ")
taxid_indexes <- timetree_newick[["tip.label"]] %>% match(string_eukaryotes[["ncbi_name"]])
unmatched_names <- timetree_newick[["tip.label"]] %>% magrittr::extract(taxid_indexes %>% is.na)
print(unmatched_names)
## [1] "Cercospora fijiensis"
                                      "Arthroderma benhamiae"
## [3] "Macropus eugenii"
                                      "Ostreococcus lucimarinus"
## [5] "Oryza nivara"
# manually creating lookup table to be joined
ncbi_to_timetree <- tribble(</pre>
  ~timetree name,
                                   ~ncbi name,
  "Macropus eugenii",
species_dictionary <- string_eukaryotes %>% left_join(ncbi_to_timetree)
# coalescing NAs to ncbi name
species_dictionary %<>%
  mutate(timetree_name = coalesce(timetree_name, ncbi_name)) %>%
  mutate(timetree_name = ifelse(timetree_name %in% timetree_newick[["tip.label"]], timetree_name, NA))
```

Now we can start looking for unfound species genera in the complete tree data.

```
species_dictionary %<>%
  mutate(genus_search = coalesce(timetree_name, ncbi_name) %>%
strsplit(" ") %>%
   sapply("[", 1))
selected_genera <- species_dictionary[["genus_search"]] %>% unique
unreliable_genera <- intersect(selected_genera, duplicated_genera)</pre>
# this is actually really important
tree_85k[["node.label"]] <- NULL</pre>
tree_85k[["edge.length"]] <- NULL
# replacing timetree's underscores with spaces
tree_85k[["tip.label"]] %<>% str_replace_all("_", " ")
tree_85k[["tip.genus"]] <- sapply(strsplit(tree_85k[["tip.label"]]," "), "[", 1)</pre>
tree_85k_genera <- tree_85k[["tip.genus"]] %>% unique
# subtracting unreliable genera
tree_85k_genera %<>% setdiff(unreliable_genera)
tree_genus <- tree_85k %%% keep.tip(., tip.label[tip.genus %in% selected_genera])
tree_genus[["tip.genus"]] <- sapply(strsplit(tree_genus[["tip.label"]]," "), "[", 1)</pre>
# unfound species which genera are present in the 85k tree
unfound_species <- species_dictionary %>%
  filter(is.na(timetree_name) & genus_search %in% tree_85k_genera)
```

Once we figured out which species have proxy genera in the complete data, we can start filling them in as sister branches.

```
# for each unfound species which genus is present in the 85k tree,
for(i in 1:nrow(unfound_species)){
    # we search for all species of this genus ("sister species") in the 85k tree
    # this part is tricky because bind.tip rebuilds the tree from scratch
    # so we need to keep removing underscores. there are better ways to do this.
    tip_genus <- tree_genus[["tip.label"]] %% strsplit("[_ ]") %% sapply("[", 1)
    sister_species <- tree_genus[["tip.label"]] [tip.genus == unfound_species[[i, "genus_search"]]]
    # we obtain the sister_species" most recent common ancestor (MRCA)
    # c(.[1]) is a hack because the MRCA function only works with at least 2 nodes
    where <- getMRCA(tree_genus, sister_species %% c(.[1]))
    # and then add a leaf node linked to this MRCA
    tree_genus %<% bind.tip(tip.label = unfound_species[[i, "ncbi_name"]], where = where)
}

# for some reason bind.tip adds underscores to species names
    tree_genus[["tip.label"]] %<% str_replace_all("_", " ")

# keeping track of found species

found_species <- species_dictionary %>% filter(!is.na(timetree_name) | genus_search %in% tree_85k_genera)

# forced_name means it either was found in timetree or we forced it by looking at genera names

found_species %<>% mutate(forced_name = coalesce(timetree_name, ncbi_name))

# so we keep only found species in this tree we are building (timetree + forced by genera)

tree_genus %<>% keep.tip(found_species[["forced_name"]])

# which found_species rows correspond to each tip.label?

match_tiplabel_name <- match(tree_genus[["tip.label_name"]])

tree_genus %<>% list_modify(
# converting to ncbi taxids

tip.label = found_species[["new_taxid"]][match_tiplabel_name]

)
```

#### Species of unfound genera

In this part, we try to fill in the remaining missing species (those which genera were not found in TimeTree) by searching for their closest relatives (according to NCBI Taxonomy) that are present in the current tree. Once we find its two closest relatives, we can add the missing species as a branch from their LCA. This is a conservative approach.

```
graph_ncbi <- read.tree("tree_ncbi.nwk") %>% as.igraph.phylo(directed = TRUE)
# converting phylo to igraph
graph_genus <- as.igraph.phylo(tree_genus, directed = TRUE)
unfound_genera <- species_dictionary %>% filter(is.na(timetree_name) & !genus_search %in% tree_85k_genera)
tip_nodes <- V(graph_ncbi)[degree(graph_ncbi, mode = "out") == 0]</pre>
tip_distances <- graph_ncbi %>%
  distances(v = tip_nodes, to = tip_nodes, mode = "all") %>%
  as_tibble(rownames = "from") %>%
  pivot_longer(-from, names_to = "to", values_to = "distance")
tip_distances %<>% filter(distance > 0)
tip_distances %<>% inner_join(unfound_genera %>% select(from = new_taxid))
# we only want to find species already present in the genus_tree
tip_distances %<>% inner_join(found_species %>% select(to = new_taxid))
# we only want the two closest relatives
tip_distances %<>%
  group_by(from) %>%
  top_n(-2, distance) %>% # top 2 smallest distances
```

```
out_distances <- graph_genus %>% distances(mode = "out")
# we find the MRCA for its two closest relatives
unfound_genera_mrca <- tip_distances %>% group_by(from) %>% summarise(mrca = {
  # which rows have no infinite distances? the last one represents the MRCA
 mrca_row_index <- max(which(rowSums(is.infinite(out_distances[, to])) == 0))</pre>
 rownames(out_distances)[mrca_row_index]
graph_genus %<>% add_vertices(nrow(unfound_genera_mrca), color = "red", attr = list(name = unfound_genera_mrca[["from"]]))
edges_to_add <- V(graph_genus)[unfound_genera_mrca %>% select(mrca, from) %>% t %>% as.vector]$name
# connecting species leafs to the supposed MRCA
graph_genus %<>% add_edges(V(graph_genus)[edges_to_add])
# finally converting to phylo format
phylo_graph_genus <- treeio::as.phylo(graph_genus)</pre>
match_tiplabel_taxid <- match(phylo_graph_genus[["tip.label"]], species_dictionary[["new_taxid"]])</pre>
phylo_graph_genus %<>% list_modify(
  # adding tip.alias (this is not exported with write.tree)
tip.alias = species_dictionary[["string_name"]][match_tiplabel_taxid],
  tip.label = species_dictionary[["taxid"]][match_tiplabel_taxid]
phylo_graph_genus[["node.label"]] <- NULL</pre>
phylo_graph_genus[["edge.length"]] <- NULL
```

#### Ctenophora as sister to all animals

According to TimeTree, Ctenophora remains as a sister group to Cnidaria. We believe the most recent consensus in literature is to consider them a sister group to all animals. The following code block moves *Mnemiopsis leidyi*, the only ctenophore in our analysis, to the base of the metazoan lineage.

```
# moving ctenophora before porifera
mnemiopsis_taxid <- species_dictionary %>% filter(ncbi_name == "Mnemiopsis leidyi") %>% pull(taxid)
amphimedon_taxid <- species_dictionary %>% filter(ncbi_name == "Amphimedon queenslandica") %>% pull(taxid)

# reordering tip.labels
from_to <- c(
    "400682" = "27923", # amphimedon to mnemiopsis
    "10228" = "400682", # trichoplax to amphimedon
    "27923" = "10228" # mnemiopsis to trichoplax
)

modified_phylo <- phylo_graph_genus

modified_phylo[["tip.label"]] %<>% recode(!!!from_to)

write.tree(modified_phylo, "../data/hybrid_tree_modified.nwk")
```

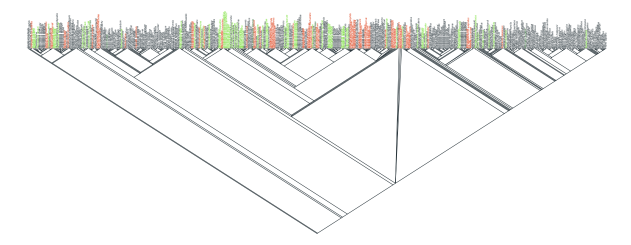


Figure 1: Complete 476 eukaryotes tree. Green species have been filled in by a genus proxy in TimeTree. Red species have been filled in by looking at NCBI Taxonomy.

#### Gene selection and annotation

The anchoring point for this study is basic annotation about genes and the pathways in which they participate. This section describes the process of structuring such data. In the end we will have a table to which all kinds of additional data will be left joined into.

# Neurotransmitter systems annotation

We start by querying the KEGG api for the pathways of interest. Resulting data is then pivoted to a wider format.

Table 5: All links between genes and pathways in KEGG.

link_pathway_entrez									
#	# Col. name Col. type Used? Example Description								
1	$entrez\_id$	character	yes	hsa:10411	NCBI Taxonomy identifier				
2	$pathway\_id$	character	yes	path:hsa04726	KEGG pathway ID				

Location: data-raw/download/link\_pathway\_entrez.tsv

Source: http://rest.kegg.jp/link/pathway/hsa

```
pivot_wider(
   id_cols = entrez_id,
   names_from = pathway_name,
   values_from = n,
   values_fr = list(n = length),
   values_fill = list(n = 0)
) %>%
   mutate(system_count = rowSums(select(., -entrez_id)))

# exporting for package use
usethis::use_data(gene_pathways, overwrite = TRUE)
```

```
## <U+2714> Setting active project to 'C:/R/neuro'
```

<sup>## &</sup>lt;U+2714> Saving 'gene\_pathways' to 'data/gene\_pathways.rda'

tail(gene_pathways)									
entrez_id	glutamatergic	cholinergic	serotonergic	gabaergic	dopaminergic	system_count			
805	0	0	0	0	1	1			
808	0	0	0	0	1	1			
810	0	0	0	0	1	1			
84152	0	0	0	0	1	1			
91860	0	0	0	0	1	1			
9575	0	0	0	0	1	1			

#### Base ID lookup table

Now we start building a base ID lookup table containing entrez gene IDs, STRING ensembl protein IDs, ensembl gene IDs, STRING protein names and entrez gene names. Every piece of data in subsequent analyses will be progressively joined to it.

Table 6: Conversion dictionary from entrez ID to STRING's ensembl protein ID.

	link_entrez_string								
#	# Col. name Col. type Used? Example Description								
1	taxid	numeric	no	9606	NCBI Taxonomy ID				
2	$entrez\_id$	numeric	yes	7157	entrez gene ID				
3	$string\_id$	character	yes	9606.ENSP00000269305	STRING ID				

Location: data-raw/download/human.entrez\_2\_string.2018.tsv.gz

Source: https://string-db.org/mapping\_files/entrez/human.entrez\_2\_string.2018.tsv.gz

Table 7: Conversion dictionary from STRING ID to protein name.

	string_names								
# Col. name Col. type Used? Example Description									
1	taxid	numeric	no	9606	NCBI Taxonomy ID				
2	$string\_name$	character	yes	TP53	protein name				
3	$string\_id$	character	yes	9606.ENSP00000269305	STRING ID				

Location: data-raw/download/human.name\_2\_string.tsv.gz

Source: https://string-db.org/mapping\_files/STRING\_display\_names/human.name\_2\_string.tsv.gz

Table 8: Conversion dictionary from entrez ID to gene name.

	entrez_names								
#	# Col. name Col. type Used? Example Description								
1	taxid	numeric	no	9606	taxon ID				
2	$entrez\_id$	character	yes	7157	entrez gene ID				
3	$entrez\_name$	character	yes	TP53	gene name				
4			no		(too many unrelated fields)				

Location: data-raw/download/Homo\_sapiens.gene\_info.gz

Source: https://ftp.ncbi.nlm.nih.gov/gene/DATA/GENE\_INFO/Mammalia/Homo\_sapiens.gene\_info.gz

Table 9: Conversion dictionary from entrez ID to ensembl gene (ENSG) ID.

link_ensembl_entrez									
#	# Col. name Col. type Used? Example Description								
1	$entrez\_id$	character	yes	hsa:7157	entrez gene ID				
2	$ensembl\_id$	character	yes	ensembl: ENSG 00000141510	ensembl gene ID				

Location: data-raw/download/link\_ensembl\_entrez.tsv

Source: http://rest.genome.jp/link/ensembl/hsa

```
# removing all kegg prefixes (e.g. "hsa:")
link_ensembl_entrez %<>% mutate_all(str_split_n, ":", 2)

# joining all data
gene_ids <- gene_pathways %>%
select(entrez_id) %>%
left_join(link_ensembl_entrez) %>%
left_join(link_entrez_string) %>%
left_join(string_names) %>%
left_join(entrez_names)
```

Some STRING proteins couldn't be automatically resolved, so we perform it manually

gene\_ids[!complete.cases(gene\_ids),]

entrez_id	ensembl_id	string_id	string_name	entrez_name
100137049	ENSG00000243708	NA	NA	PLA2G4B
85358	ENSG00000251322	NA	NA	SHANK3
8681	ENSG00000168970	NA	NA	JMJD7-PLA2G4B
1139	ENSG00000175344	NA	NA	CHRNA7
107987478	NA	NA	NA	LOC107987478
107987479	NA	NA	NA	LOC107987479
1564	ENSG00000205702	NA	NA	CYP2D7
801	ENSG00000198668	NA	NA	CALM1
805	ENSG00000143933	NA	NA	CALM2
808	ENSG00000160014	NA	NA	CALM3

```
## <U+2714> Saving 'gene_ids' to 'data/gene_ids.rda'
```

# Neuroexclusivity

Neuroexclusivity data consists of gene expression collected from Gexe Expression Atlas and the KEGG pathways themselves.

# Expression neuroexclusivity

In this section we preprocess multiple wide .tsv files into a single long data.frame. We also create a template file for classifying tissues into nervous or non-nervous.

#### Resources

We start by searching Gene Expression Atlas for experiments that have human baseline expression data at the tissue level. For each experiment, TPM expression data is downloaded to the data-raw/download/gxa/directory. Found experiments:

- E-MTAB-513
- E-MTAB-2836
- E-MTAB-3358
- E-MTAB-3708
- E-MTAB-3716
- E-MTAB-4344
- E-MTAB-4840
- E-MTAB-5214

#### Reshaping data

We load and pivot all files to a long format.

```
gene_expression <- sapply(
  list.files("download/gxa/", full.names = T),
  read_tsv,
  comment = "#",
  simplify = FALSE,
  USE.NAMES = TRUE
)

# pivoting
gene_expression %<>%
  map_dfr(pivot_longer, cols = -(1:2), names_to = "tissue", values_to = "tpm") %>%
  na.omit %>%
  select(ensembl_id = `Gene ID`, tissue, tpm)
```

#### Cleaning

A lot of tissue information can be collapsed into a single level (e.g. "brain" and "brain fragment" can be considered the same tissue). The cleaning is performed and expression data is exported for analysis.

```
## <U+2714> Setting active project to 'C:/R/neuro'
## <U+2714> Saving 'gene_expression' to 'data/gene_expression.rda'
```

head(gene_expression)					
ensembl_id	tissue	tpm			
ENSG00000010379	adipose tissue	0.1			
ENSG00000010379	adrenal gland	0.1			
ENSG00000010379	cerebral cortex	5.0			
ENSG00000010379	colon	0.1			
ENSG00000010379	duodenum	5.0			
ENSG00000010379	fallopian tube	10.0			

# Tissue classification

For subsequent analyses, we need to distinguish if a tissue is part of the nervous system or not. This is done by hand. The first step is to write a temp file to data-raw/temp/temp\_tissue\_classification.tsv with all tissue names. This serves as a base for the completed data/neuroexclusivity\_classification\_tissue file.

```
gene_expression %>%
  select(tissue) %>%
unique %>%
arrange %>%
  arrange %>%
mutate(is_nervous = NA) %>%
  write_tsv("temp/temp_tissue_classification.tsv")
```

# Pathway neuroexclusivity

In this section we create a template file for classifying pathways into nervous or non-nervous.

#### Resources

For link\_pathway\_entrez see Table 5.

Table 10: KEGG pathway names.

	pathway_names					
#	Col. name	Col. type	Used?	Example	Description	
1	pathway_id	character	yes	path:hsa04726	KEGG pathway ID	
2	pathway_name	character	yes	Serotonergic synapse - Homo sapiens (human)	pathway name	

Location: data-raw/download/pathway\_names.tsv
Source: http://rest.kegg.jp/list/pathway/hsa

# Pathway classification

Just like tissues, we need to distinguish if a pathway is related to the nervous system or not. This is done by hand. The first step is to write a temp file to data-raw/temp/temp\_pathway\_classification.tsv with all pathway names. This serves as a base for the completed data/neuroexclusivity\_classification\_pathway.tsv file.

```
# removing species prefix "hsa:"
link_pathway_entrez[["entrez_id"]] %<>% str_split_n("\\:", 2)

selected_genes_pathways <- link_pathway_entrez %>% filter(entrez_id %in% gene_ids[["entrez_id"]])
unique_pathway_ids <- selected_genes_pathways %>% pull(pathway_id) %>% unique

pathway_names %<>% filter(pathway_id %in% unique_pathway_ids) %>%
    mutate(is_nervous = NA) %>%
    write_tsv("temp/temp_pathway_classification.tsv")
```

# Orthology data

This section refers to orthology data exported for geneplast use. Essentialy, we subset the global STRING mapping between proteins and orthologous groups into a smaller dataset containing only information about the orthogroups related to our selected genes.

Table 11: Orthologous groups (COGs, NOGs, KOGs) and their proteins.

	cogs						
#	Col. name	Col. type	Used?	Example	Description		
1	taxid.string_id	character	yes	9606.ENSP00000269305	STRING protein ID		
2	start_position	numeric	no	1	residue where orthogroup mapping starts		
3	end_position	numeric	no	393	residue where orthogroup mapping ends		
4	$\cos$ _id	character	yes	NOG08732	orthologous group ID		
5	protein_annotation	character	no	Cellular tumor antigen p53; []	protein description		

**Location:** data-raw/download/COG.mappings.v11.0.txt.gz

Source: https://stringdb-static.org/download/COG.mappings.v11.0.txt.gz

```
# spliting first column into taxid and string_id
cogs %<>% separate(taxid.string_id, into = c("taxid","string_id"), sep = "\\.", extra = "merge")

# subsetting cogs of interest
gene_cogs <- cogs %>% filter(string_id %in% gene_ids[["string_id"]]) %>% select(-taxid)

cogs %<>%
    # leave only eukaryotes
filter(taxid %in% string_eukaryotes[["taxid"]]) #%>%
    # leave only proteins which are part of cogs of interest
# (affects abundance metrics so we won't filter)
# filter(cog_id %in% cogs_of_interest[["cog_id"]])
```

```
# exporting for package use
usethis::use_data(cogs, overwrite = TRUE)

## <U+2714> Setting active project to 'C:/R/neuro'
## <U+2714> Saving 'cogs' to 'data/cogs.rda'

usethis::use_data(gene_cogs, overwrite = TRUE)

## <U+2714> Saving 'gene_cogs' to 'data/gene_cogs.rda'
```

#### Network

In this section we search the STRING API for our proteins of interest and recompute the combined interaction score.

# Retrieving network data

Querying the API endpoint for the STRING IDs we collected.

```
identifiers <- gene_ids %>% pull(string_id) %>% na.omit %>% pasteO(collapse="%Od")

if (!file.exists("download/string_ids.tsv")) {
    postForm(
        "http://string-db.org/api/tsv/get_string_ids"
        ,identifiers = identifiers
        ,echo_query = "1"
        ,species = "9606"
      ) %>%
        write("download/string_ids.tsv")
}
```

Table 12: STRING interaction network with channel specific scores.

	string_ids						
#	Col. name	Col. type	Used?	Example	Description		
1	queryItem	character	yes	ENSP00000258400	queried term		
2	queryIndex	numeric	yes	266	index of queried term		
3	$\operatorname{stringId}$	character	yes	9606.ENSP00000258400	STRING ID		
4	ncbiTaxonId	numeric	yes	9606	NCBI Taxonomy ID		
5	taxonName	character	yes	Homo sapiens	species name		
6	${\it preferredName}$	character	yes	HTR2B	common protein name		
7	annotation	character	yes	5-hydroxytryptamine receptor 2B; []	protein annotation		

**Location:** data-raw/download/string ids.tsv

Source: http://string-db.org/api/tsv/get\_string\_ids

Now we need to make sure that the API successfully resolves the protein IDs we searched for.

```
api_ids <- read_tsv("download/string_ids.tsv", comment = "", quote = "")

# removing taxid prefix
api_ids %<>% mutate(stringId = str_split_n(stringId, "\\.", 2))

# removing inexact matches (queried id is different from resolved id)
api_ids %<>% group_by(queryItem) %>% filter(queryItem == stringId)

# setequal must return true if ids matched exatcly
setequal(
gene_ids %>% pull(string_id) %>% na.omit,
api_ids %>% pull(stringId)
)
```

Once IDs are correct, we can query the network API endpoint to obtain the protein interaction edgelist.

```
# it is important to query this endpoint with the species prefix ("9606.")
identifiers <- api_ids %>% pull(stringId) %>% na.omit %>% { pasteO("9606.", ., collapse="%Od") }

if (!file.exists("download/string_edgelist.tsv")) {
    postForm(
        "http://string-db.org/api/tsv/network"
        ,identifiers = identifiers
        ,species = "9606"
        ) %>%
    write("download/string_edgelist.tsv")
}
```

Table 13: STRING interaction network with channel specific scores.

	string_edgelist						
#	Col. name	Col. type	Used?	Example	Description		
1	stringId_A	character	yes	ENSP00000215659	STRING ID (protein A)		
2	$stringId\_B$	character	yes	ENSP00000211287	STRING ID (protein B)		
3	$preferredName\_A$	character	yes	MAPK12	common protein name (protein A)		
4	$preferredName\_B$	character	yes	MAPK13	common protein name (protein B)		
5	ncbiTaxonId	numeric	yes	9606	NCBI Taxonomy ID		
6	score	numeric	yes	0.948	combined score		
7	nscore	numeric	yes	0	gene neighborhood score		
8	fscore	numeric	yes	0	gene fusion score		
9	pscore	numeric	yes	0.014223	phylogenetic profile score		
10	ascore	numeric	yes	0	coexpression score		
11	escore	numeric	yes	0.485	experimental score		
12	dscore	numeric	yes	0.9	database score		
13	tscore	numeric	yes	0.02772	textmining score		

Location: data-raw/download/string\_edgelist.tsv Source: http://string-db.org/api/tsv/network

#### Recomputing scores

From string-db.org:

"In STRING, each protein-protein interaction is annotated with one or more 'scores'. Importantly, these scores do not indicate the strength or the specificity of the interaction. Instead, they are indicators of confidence, i.e. how likely STRING judges an interaction to be true, given the available evidence. All scores rank from 0 to 1, with 1 being the highest possible confidence."

For the sake of this project, we will only use experimental and database scores with a combined value >= 0.7, a high confidence threshold according to the STRING database. The combined score is given by the following expression, as stated in von Mering C et al, 2005:

$$S = 1 - \prod_{i} (1 - S_i)$$

```
string_edgelist <- read_tsv("download/string_edgelist.tsv")

string_edgelist %<>%
    mutate(cs = combine_scores(., c("e","d"))) %>%
    filter(cs >= 0.7) %>%
    select(stringId_A, stringId_B)

# how many edgelist proteins are absent in gene_ids (should return 0)
setdiff(
    string_edgelist %$% c(stringId_A, stringId_B),
    gene_ids %>% pull(string_id)
)

# exporting for package use
usethis::use_data(string_edgelist, overwrite = TRUE)
```

# **Analysis**

Analysis

## Root inference

To estimate the evolutionary root of a given gene, i.e. the ancestor from which its genetic archetype (orthologous group) is vertically inherited, we use orthologous group annotation from the STRING database. The presence and absence of orthologous groups in the species of a cladogram are used to determine its most likely ancestor. Loading initial resources:

```
library(tidyverse)
library(magrittr)
library(geneplast)
library(ape)
library(xML)
library(rentrez)
library(neurotransmissionevolution)

data(
    cogs,
    gene_cogs,
    string_eukaryotes,
    package = "neurotransmissionevolution"
)

phyloTree <- read.tree("../data/hybrid_tree_modified.nwk") %% rotatePhyloTree("9606")</pre>
```

We perform some minor data formatting before feeding it to geneplast

```
# formating cogdata column names for geneplast
cogs %<>% rename(protein_id = string_id, ssp_id = taxid) %>% select(protein_id, ssp_id, cog_id)

# adding species names to taxid tree
phyloTree %<>% list_modify(
   tip.alias = string_eukaryotes %$% string_name[match(phyloTree[["tip.label"]], taxid)]
)
```

#### Geneplast

Geneplast's groot.preprocess function structures an ogr object on which groot will perform the rooting. We then retrieve the numeric root (groot.get("results")) for the cogs\_of\_interest, that is, orthologous groups pertaining to neurotransmission genes.

```
cogs_of_interest <- gene_cogs %>% pull(cog_id) %>% unique
ogr <- groot.preprocess(
  cogdata = cogs,</pre>
```

```
phyloTree = phyloTree,
    spid = "9606",
    cogids = cogs_of_interest
)

roots <- groot(ogr, nPermutations = 1) %>%
    groot.get("results") %>%
    rownames_to_column("cog_id") %>%
    select(cog_id, root = Root)

write_tsv(roots, "geneplast_roots.tsv")

# setwd("plots/roots/")
# groot.plot(ogr, plot.lcas = TRUE, width=10, height=20, cex.lab = 0.2, cex.nodes = 0.4)
# setwd("../../")
```

#### Clade names

Each root branches to a clade that diverged from humans some time in the past. It is nice to have these clades taxonomically named to ease our interpretation. Unlike NCBI Taxonomy, TimeTree's internal nodes are not named. Therefore, we query the NCBI Taxonomy API to try to find most clade names automatically. It is important to note that we are using a hybrid tree primarily built from TimeTree data. This means NCBI Taxonomy naming will not perfectly match clades in our tree. For instance, root #36 branches to a clade containing 38 species from the SAR supergroup, but also 1 species from the Haptista rank, namely *Emiliania huxleyi*. The Haptista group is a sister clade to SAR, so it might be the case that *Emiliania huxleyi* is actually correctly placed together with SAR species by TimeTree, given their evolutionary proximity. Resolving these naming conflicts is not trivial and falls out of our scope.

```
lineages <- entrez_fetch(</pre>
 db
          = string_eukaryotes[["new_taxid"]],
  rettype = "xml",
 retmode = "xml",
 parsed = TRUE
string_eukaryotes %<>% mutate(
             = ogr@tree$tip.group[taxid],
  lineage_txt = xpathSApply(lineages, "//Lineage", XML::xmlValue)
roots_names <- string_eukaryotes %>%
 mutate(lineage_split = strsplit(lineage_txt, "; ")) %>%
 group_by(root) %>%
  summarise(lineage = Reduce(intersect, lineage_split) %>% list,
            lineage_list = lineage_split %>% list) %>%
 mutate(downstream_diff = mapply(setdiff,
                                                  lineage, lead(lineage))) %>%
 mutate(upstream_diff = mapply(setdiff, downstream_diff, lag(lineage))) %>%
 mutate(clade_name = map_chr(upstream_diff, 1, .default = NA)) %>%
 mutate(collapse_depth = lineage %>% map_int(length) + 1) %>%
 group_by(root) %>%
    fallback_name is the collapsed lineage ranks
 mutate(fallback_name = lineage_list %>%
           flatten %>%
           map2_chr(collapse_depth, `[`) %>%
           table %>%
           sort(TRUE) %>%
           paste0(names(.), " (", .,")") %>% paste0(collapse="; ")) %>%
  mutate(clade_name = coalesce(clade_name, fallback_name)) %>%
  select(root, clade_name)
```

```
write_tsv(roots_names, "temp/temp_geneplast_clade_names.tsv")
```

# Phyletic patterns

Visualizing the presence/absence matrix according to inferred roots and species' clades

```
lca_names <- read_tsv("geneplast_clade_names.tsv") %>% rename("lca" = root)
lca_spp <- ogr@spbranches %>% rename("taxid" = ssp_id, "species" = ssp_name, "lca" = `9606`)
# Saving for use in abundance computation lca_spp %>%
  select(lca, taxid) %>%
cog_pam <- ogr@orthoct[,-1]</pre>
long_pam <- cog_pam %>%
  rownames_to_column("taxid") %>%
   pivot_longer(-taxid, names_to = "cog_id") %>%
   left_join(lca_spp) %>%
   left_join(lca_names) %>%
   left_join(roots) %>%
                     = fct_reorder(cog_id, root),
= fct_reorder(species, desc(lca)),
     cog_id
     clade_name = fct_reorder(clade_name, lca),
root = as_factor(root),
     clade_stripe = as.numeric(as_factor(lca)) %% 2 == 0
   group_by(cog_id) %>%
   mutate(spp_stripe = as.numeric(species) %% 2 == 0) %>%
   filter(value == 1) %>%
   group_by(taxid) %>%
   mutate(cog_stripe = as.numeric(cog_id) %% 2 == 0)
ggplot(long_pam, aes(x = cog_id, y = species)) +
  geom_tile(aes(fill = clade_stripe + 0.3 * xor(spp_stripe, cog_stripe))) +
  scale_fill_gradient(low = "#37474F", high = "#263238") +
  facet_grid(clade_name ~ fct_rev(root), scales = "free", space='free') +
   theme(
     text
     panel.spacing = unit(1, "pt"),
panel.grid.major.x = element_blank(),
     panel.grid.major.x = element_Drank(),

strip.background = element_rect(colour = "#FFFFFFF"),

strip.text.x = element_text(size = 6, angle = 90),

strip.text.y = element_text(size = 3, angle = 0, hjust = 0, lineheight = 3),

axis.text.x = element_text(size = 6, angle = 90, vjust = 0.5),
     legend.position = "none"
```

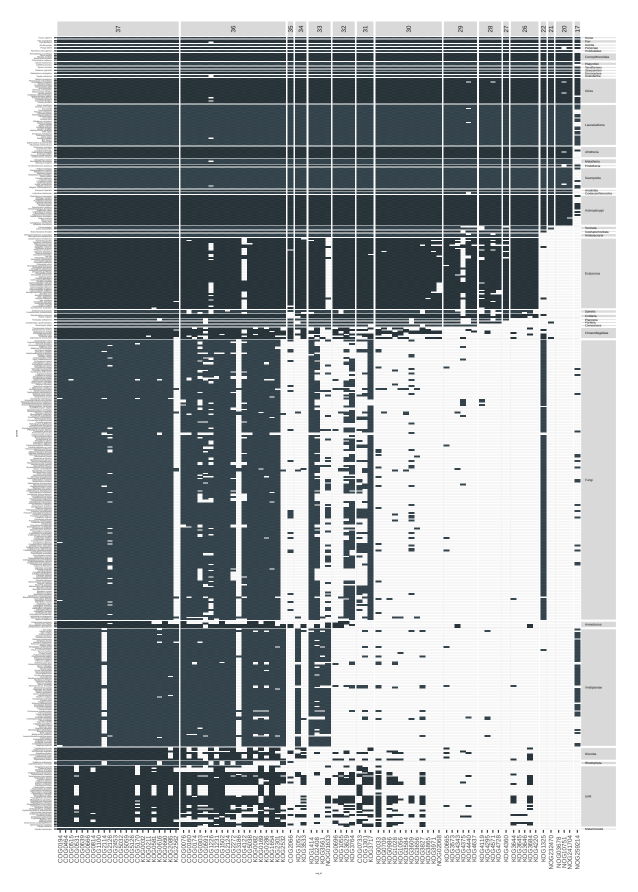


Figure 2: Presence of orthologous groups in species. The horizontal axis is grouped by COGs rooted at some specific LCA. The vertical axis is grouped by species' clades. A checkerboard pattern is superimposed to aid visual examination.

# Neuroexclusivity

We characterize genes' relevance to the nervous system by inspecting what proportion of its activity is related to nervous processes. We relied on tissue RNA-Seq data, as well as KEGG's pathways themselves. Loading resources.

```
library(tidyverse)
library(magrittr)

data(
   gene_ids
   ,gene_pathways
   ,gene_expression
   ,package = "neurotransmissionevolution"
)
```

# Expression neuroexclusivity

We start by averaging all gene\_expression collected from the Expression Atlas by tissue (tpm\_avg). The averaged expression is filtered for values greather than 0.5 TPM. This ensures further computations only account for tissues in which genes are actually expressed. Then, we add the manual tissue classification indicating which tissues are nervous or not (described in Preprocessing). The neuroexclusivity index for a gene is the sum of its tpm\_avg values in nervous tissues divided by the sum its values in all tissues.

```
tissue_classification <- read_tsv(
    file = "../data/neuroexclusivity_classification_tissue.tsv"
, col_types = "ci"
)

# Averaging TPM expression by tissue
avg_by_tissue <- gene_expression %>%
    group_by(ensembl_id, tissue) %>%
    summarise(tpm_avg = mean(tpm)) %>%
    filter(tpm_avg >= 0.5) %>%
    left_join(tissue_classification)

# Measuring expression neuroexclusivity
expression_neuroexclusivity <- avg_by_tissue %>%
    group_by(ensembl_id) %>%
    summarise(expression_neuroexclusivity = sum(tpm_avg[is_nervous == 1])/sum(tpm_avg)) %>%
    write_tsv("neuroexclusivity_expression.tsv")
```

#### Pathway neuroexclusivity

To find the pathway neuroexclusivity of a gene, we simply divide the count of nervous pathways by the count of all pathways it participates in.

```
pathway_classification <- read_tsv(
    file = "../data/neuroexclusivity_classification_pathway.tsv"
    ,col_types = "cci"
)

link_pathway_entrez <- read_tsv(
    file = "../data-raw/download/link_pathway_entrez.tsv"
    ,col_names = c("entrez_id", "pathway_id")
    ,col_types = "cc"
)

# Removing "hsa:" prefix
link_pathway_entrez[["entrez_id"]] %<>% str_split_n("\\:", 2)

# Pathway data related to our genes of interest
selected_genes_pathways <- link_pathway_entrez %>%
filter(entrez_id %in% gene_ids[["entrez_id"]]) %>%
left_join(pathway_classification) %>%
drop_na # Dropping general pathways

# Measuring pathway neuroexclusivity
```

```
pathway_neuroexclusivity <- selected_genes_pathways %>%
  group_by(entrez_id) %>%
  summarise(pathway_neuroexclusivity = sum(is_nervous)/length(is_nervous)) %>%
  write_tsv("neuroexclusivity_pathway.tsv")
```

#### Network

In this section we search the STRING API for our proteins of interest and recompute their combined interaction score.

### Graph data

Loading resources.

```
# Data manipulation
library(tidyverse)
library(igraph)
library(magrittr)
library(scatterpie)
library(UpSetR)
library(gridExtra)
library(patchwork)
library(neurotransmissionevolution)
data(
  gene_ids
  ,gene_pathways
  ,string_edgelist
  ,pathway_neuroexclusivity
  ,expression_neuroexclusivity
  ,package = "neurotransmissionevolution"
col_types = "ci")
                                                                             col_types = "ic")
expression_neuroexclusivity <- read_tsv("neuroexclusivity_pathway.tsv", col_types = "cn")

# Collapsize rigid ("neuroexclusivity_expression.tsv", col_types = "cn")
gene_annotation <- read_tsv(".../data/gene_annotation.tsv", col_types = "cc") %>%
  mutate(annotation = case_when(
    grepl("clearance", annotation) ~ "depletion"
,grepl("degradation", annotation) ~ "depletion"
    ,grepl("transport", annotation) ~ "synthesis"
     ,TRUE ~ annotation
```

Joining all gene data and creating the network graph object.

```
# If a gene has more than 1 COG, select the oldest one.

# This is unusual, but can happen in cases of gene fusion, for instance.

gene_cogs %<>%
    inner_join(cog_roots) %>%
    group_by(string_id) %>%
    filter(root == max(root)) %>%
    inner_join(clade_names)

# Gathering all gene info available

vertices <- gene_ids %>%
    na.omit %>%
    inner_join(gene_cogs) %>%
    inner_join(gene_pathways) %>%
    inner_join(gene_annotation) %>%
    inner_join(gene_annotation) %>%
    inner_join(gene_annotation) %>%
    inner_join(gene_annotation) %>%
    inner_join(gene_annotation) %>%
    inner_join(gene_annotation) %>%
```

```
inner_join(expression_neuroexclusivity) %>%
mutate(ne = pathway_neuroexclusivity >= 0.9) %>%
select(string_id, everything())

# Quick color hack to aid visualization
vertices %<>%
    unite(color, glutamatergic:dopaminergic, remove = F) %>%
    mutate(color = rainbow(color %>% n_distinct)[color %>% as.factor])

g <- graph_from_data_frame(string_edgelist, directed = F, vertices = vertices)

# Setting node sizes
V(g)$size <- V(g)$system_count %>% sqrt %>% multiply_by(5)
```

Generating the network layout (i.e. node coordinates).

```
if(file.exists("network_layout.tsv")) {
   layout <- read_tsv("network_layout.tsv", col_types = "dd") %>% as.matrix
} else {
   layout <- vivagraph(g, precompute_multiplier = 200, precompute_niter = 1000)
}

# inserting layout coordinates into graph object
V(g)$x <- layout[, 1]
# layout matrix comes vertically flipped
V(g)$y <- -layout[, 2]</pre>
```

Generating tidy edge coordinates for plotting.

```
# Recreating the vertices data.frame, now with layout coordinates (lazy)
vertices <- igraph::as_data_frame(g, what = "vertices") %>% rename(string_id = name)

# The edges data.frame will be used to draw lines with geom_line
edges <- string_edgelist %>%
    map(match, vertices[["string_id"]]) %>%
    map_dfr(~ vertices[.x,]) %>%
    select(x:y) %>%
    cbind(group = 1:nrow(string_edgelist))
```

Setting up reusable aesthetic parameters for the plots.

```
pie_colors <- c(
    "cholinergic" = "#D84315"
, "dopaminergic" = "#F98625"
, "gabaergic" = "#598625"
, "gabaergic" = "#568050"
, "serotonergic" = "#6A1B9A"
)
plot_pie_fill <- scale_fill_manual(values = pie_colors)

element_colors <- c(
    "depletion" = "#F40000"
, "excitability" = "#F6800"
, "receptor-associated" = "#B6E000"
, "ionotropic receptor" = "#43FF10"
, "metabotropic receptor" = "#3FFF0"
, "signaling" = "#0091EA"
, "g-protein" = "#0093f1"
, "synthsis" = "#A000FF"
, "vestile" = "#F600A"
#------ is_neuroexclusive ------
, "TRUE" = "#00BFC4"
, "FALSE" = "#587660"
)
# Color and size scales for neurotransmission functions
plot_scales <- list(
    scale_fill_manual(values = element_colors)
, scale_color_manual(values = element_colors %>% darken(0.25))
, scale_radius(range = c(1.75, 5.00), guide = FALSE)
)
systems <- names(pie_colors)
edge_color <- rgb(0.7, 0.7, 0.7, alpha = 0.3)</pre>
```

```
past_fill <- "#FFFFFF" # past nodes' fill color
past_color <- "#888888" # past nodes' border color</pre>
vertices %<>% mutate(
  shape = ifelse(ne, "square filled", "circle filled"),
color_node = ifelse(ne, "#000000", element_colors[annotation] %>% darken(0.2)),
color_pie = ifelse(ne, "#000000", NA),
# Some recurrent ggplot aesthetics
edge_aes <- aes(x = x, y = y, group = group)
text_aes <- aes(x = x, y = y, label = string_name)
pie_aes <- aes(x = x, y = y, group = string_id, r = size^(0.94) - 1.5)</pre>
# Fixing xy limits across all plots
xy_lim <- list(</pre>
  scale_x_continuous(limits = range(vertices[["x"]]) + c(-50, 50)),
scale_y_continuous(limits = range(vertices[["y"]]) + c(-50, 50))
# Emptying theme defaults
plot_theme <- list(coord_equal(), theme_void())</pre>
# Allowing more space for multiple network plots
diff_theme <- list(</pre>
   coord_equal(),
     theme_void(),
    theme(
                                                 = element_text(size = 8, hjust = 0.5)
        plot.title = element_text(size = 8, n)]

,legend.text = element_text(size = 6)
,legend.title = element_text(size = 8)
,legend.key.size = unit(1, "mm")
,legend.box.spacing = unit(-2, "mm")
,legend.box.margin = unit(c(0, 2, 0, 0), "mm")
,plot.margin = unit(c(0, 0, 0, 0), "mm")
roots <- vertices %>%
   arrange(-root) %>%
   distinct(root, clade_name) %%% set_names(root, clade_name)
```

```
plot_pies <- geom_scatterpie(</pre>
   ,mapping = pie_aes
  ,color = NA
fig1a <- ggplot() +
  plot_theme +</pre>
  plot_edges +
  plot_pies +
  plot_pie_fill +
  plot_text
## Figure 1B
#############
plot_nodes <- geom_point(</pre>
   ,mapping = aes(x, y, fill = annotation, color = annotation, size = size)
  ,shape = 21
,stroke = 0.5
fig1b <- ggplot() +
  plot_theme +
  plot_edges +
  plot_nodes +
  plot_scales +
# Plotting and saving
fig1a + fig1b
```

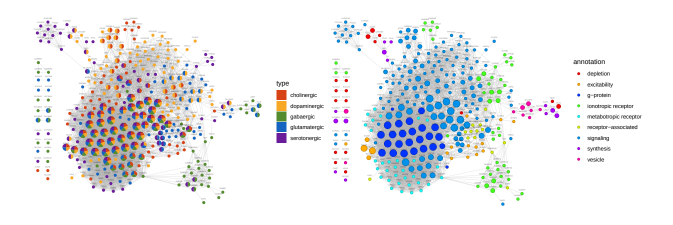


Figure 3: The human neurotransmission network with nodes colored by neurotransmitter systems (left) and neurotransmission functions (right).

```
ggsave("plots/fig1_raw.pdf", width = 14, height = 7, onefile = F, useDingbats = F)
```

```
# Retrieving the largest connected component
subgraphs <- decompose.graph(g)
lcc_index <- which.max(sapply(subgraphs, vcount))</pre>
```

```
<- subgraphs[[lcc_index]]</pre>
xy_hack <- data.frame(
                                = c("top", "bot")
= range(V(lcc)$x) + c(-75, 75)
  name
                                = range(V(1cc)$y) + c(-75, 75)
  ,pathway_neuroexclusivity
  ,expression_neuroexclusivity = 0
  ,stringsAsFactors
pajek_nodes <- lcc %>%
  igraph::as_data_frame("vertices") %>%
  bind_rows(xy_hack) %>%
  mutate(id = row_number(), y = -y)
pajek_edges <- igraph::as_data_frame(lcc, "edges")</pre>
write_tsv(
                = pajek_edges
 ,append
  ,col_names
  ,quote_escape = F
write("*nodes", "network_viacomplex.net", append = T)
write_tsv(
                = pajek_nodes %>% select(name, x, y)
  ,path
 ,append
  ,col_names
  ,quote_escape = F
write_tsv(
        = pajek_nodes %>% select(id, name, pathway_neuroexclusivity)
  ,path = "network_viacomplex_pathway.dat
        = pajek_nodes %>% select(id, name, expression_neuroexclusivity)
  ,path = "network_viacomplex_expression.dat
```

The process for generating Figures 3 and 4 (as well as Supplementary Figures 2 and 3) is roughly the same. It consists of finding what nodes have numeric roots in a given range. In our analysis, the largest root is numbered 37 and represents the furthest human common ancestor in the cladogram (the Human-Metamonada LCA, as seen in previous sections). Root #1 is represented by *Homo sapiens* itself.

The nodes we need to draw are either current\_nodes (roots in a specified numeric range), or past\_nodes (roots > such specified range). The edges we need to draw are all edges between both sets of nodes.

#### Figure 3A

```
# Finding which genes should be drawn
current_genes <- vertices %>% filter(root == 37)

# Finding which edges should be drawn
partial_ids <- current_genes %>% pull(string_id)
which_edges <- apply(string_edgelist, 1, function(r) all(r %in% partial_ids))
partial_edges <- edges[which_edges,]

plot_edges <- geom_path(
    data = partial_edges
    ,mapping = edge_aes
    ,color = edge_color
    ,size = 0.1
)

plot_text <- geom_text(
    data = current_genes</pre>
```

```
,mapping = text_aes
  ,vjust = 0
  ,nudge_y = 1.75
 ,alpha = 0.5
plot_current_pies <- geom_scatterpie(</pre>
 data = current_genes
  ,mapping = pie_aes
 ,color = NA
fig3a <- ggplot() +
 plot_edges +
 plot_scales +
 xy_lim +
 plot_current_pies +
 plot_pie_fill +
 plot_text +
 plot_theme
```

# Figure 3B

For Figure 3B, we want to see what nodes have numeric roots < 37 (Human-Metamonada LCA) and >= 26 (Human-Cnidaria LCA).

```
# Finding which genes should be drawn
current_genes <- vertices %>% filter(root < 37 & root >= 26)
past_genes <- vertices %>% filter(root == 37)
# Finding which edges should be drawn
partial_ids <- c(current_genes[["string_id"]], past_genes[["string_id"]])
which_edges <- apply(string_edgelist, 1, function(r) all(r %in% partial_ids))
partial_edges <- edges[which_edges,]
plot_edges <- geom_path(</pre>
   data = partial_edges
    ,mapping = edge_aes
   ,color = edge_color
,size = 0.1
plot_past <- geom_point(</pre>
   data = past_genes
  mapping = aes(x, y, size = size)
,fill = past_fill
,color = past_color
,shape = past_genes$shape
,stroke = 0.25
plot_text <- geom_text(</pre>
   data = current_genes
    ,mapping = text_aes
   ,size = 1
,vjust = 0
  ,nudge_y = 1.75
,alpha = 0.5
plot_current_pies <- geom_scatterpie(</pre>
  data = current_genes
   ,mapping = pie_aes
,cols = systems
,color = NA
fig3b <- ggplot() +
  plot_edges +
    plot_past +
   plot_current_pies +
    plot_pie_fill +
```

```
plot_text +
  plot_theme

# Plotting and saving
fig3a + fig3b
```

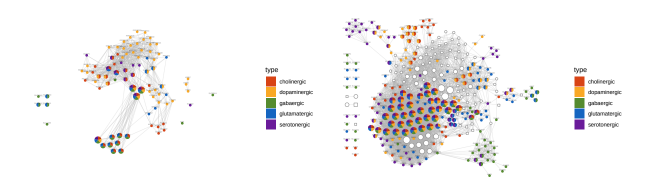


Figure 4: The human neurotransmission network with nodes colored by neurotransmitter systems and neurotransmission functions.

```
ggsave("plots/fig3_raw.pdf", width = 14, height = 7, onefile = F, useDingbats = F)
```

Additionally, we cumulatively count nodes by their categories (function and neuroexclusivity) and inferred root:

```
cumulative_emergence <- vertices %>%
    select(root, annotation, is_neuroexclusive = ne) %>%
    # Adding clade info
    right_join(clade_names) %>%
    # Pivoting from wide to long
    pivot_longer(annotation:is_neuroexclusive, values_ptypes = list(value = "character")) %>%
    # Counting nodes by category (name) for each root
    count(root, clade_name, name, value) %>%
    # Making absent counts explicit
    group_by(name) %>%
    complete(nesting(root, clade_name), name, value, fill = list(n = 0)) %>%
    # No reason to include NA observations in cumulative sum
    na.omit %>%
    # Cumulative sum node count at each root
    group_by(name, value) %>%
    mutate(cumulative_count = order_by(-root, cumsum(n)))
```

### Plotting such cumulative counts:

```
cumulative_emergence %<>% ungroup %>%
  # Creating ordered factors for plotting
mutate(
    clade_name = fct_reorder(clade_name, -root)
    ,value = fct_reorder(value, name)
)

ggplot(cumulative_emergence) +
  #---- Barplot -----
geom_bar(
    mapping = aes(clade_name, cumulative_count, group = value)
    ,stat = "sum"
    ,fill = "#999999"
    ,show.legend = F
) +
```

```
#---- Lines -----
geom_line(
    mapping = aes(clade_name, cumulative_count, group = value, color = value)
    ,size = 1
) +
#---- Styling -----
scale_color_manual(values = element_colors) +
facet_grid(name ~ .) +
theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust = 1))
```

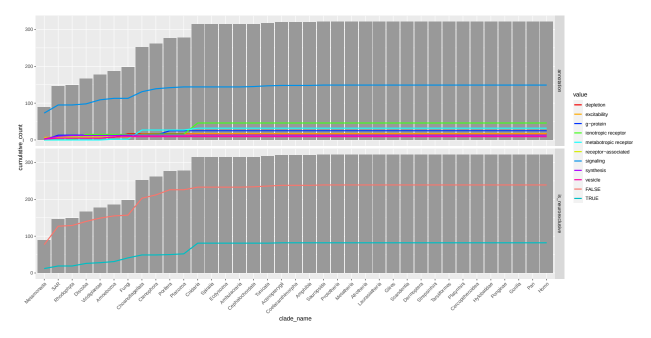


Figure 5: Cumulative node counts by categories at each root.

Visualizing nodes with roots  $\leq$  30 (Human-Porifera LCA) and  $\geq$  26 (Human-Cnidaria LCA) at every distinct root.

```
plot_size <- scale_radius(range = c(0.5, 1.3), guide = FALSE)

fig4 <- roots[roots >= 26 & roots <= 30] %>%
    imap(- {
    # Finding which genes should be drawn
    current_genes <- vertices %>% filter(root == .x)
    past_genes <- vertices %>% filter(root > .x)

# Finding which edges should be drawn
    partial_ids <- c(current_genes[["string_id"]], past_genes[["string_id"]])
    which_edges <- apply(string_edgelist, 1, function(r) all(r %in% partial_ids))
    partial_edges <- edges[which_edges,]

plot_edges <- geom_path(
    data = partial_edges
    ,mapping = edge_aes
    ,color = edge_color
    ,size = 0.1
)

plot_past <- geom_point(
    data = past_genes
    ,mapping = aes(x, y, size = size)
    ,fill = past_fill
    ,color = past_color
    ,shape = past_genes$shape</pre>
```

```
plot_text <- geom_text(</pre>
     data = current_genes
     ,mapping = text_aes
     ,size = 0.8
,vjust = -0.5
     ,nudge_y = 1
     ,alpha
  plot_current_nodes <- geom_point(</pre>
     data = current_genes
     ,mapping = aes(x, y, fill = annotation, size = size)
     ,color = current_genes$color_node
,shape = current_genes$shape
  remove_legend <- guides(fill = "none", colour = "none")</pre>
  ggplot() +
    ggtitle(paste(.y, "LCA")) +
diff_theme +
    plot_edges +
    plot_current_nodes +
    plot_scales +
    plot_size +
plot_text +
     remove_legend
fig4 <- invoke(grid.arrange, fig4, ncol = 5)</pre>
```

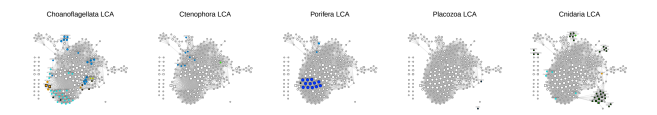


Figure 6: The human neurotransmission network with nodes rooted between roots 30 (Human-Choanoflagellata LCA) and 26 (Human-Chidaria LCA).

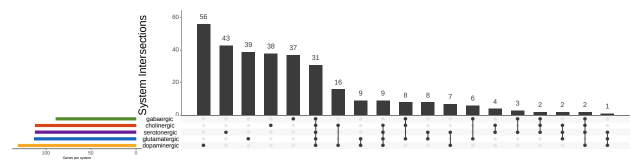
```
ggsave(
   "plots/fig4_raw.pdf"
   ,plot = fig4
   ,width = 9*0.8
   ,height = 5*0.8
   ,onefile = F
    ,useDingbats = F
)
```

# Set diagrams

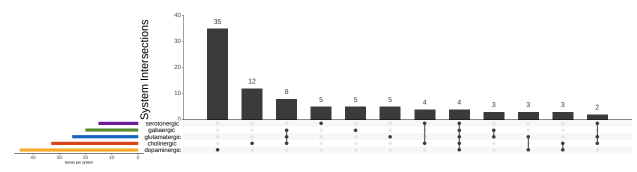
Given the dificulties of joining ggplot and base plots, the set diagrams have to be plotted by themselves:

```
# We have to manually find the correct order of colors
# Because UpSetR does not understand named vectors
get_colors <- function(df) {
    ordered_systems <- df %>%
    select(systems) %>%
    colSums %>%
```

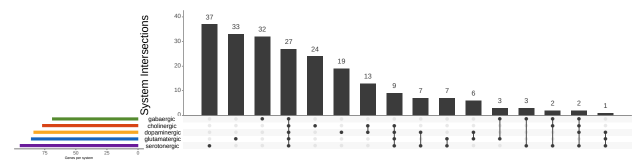
```
extract(. > 0) %>%
              extract(order(., names(.), decreasing = T))
      pie_colors[names(ordered_systems)]
 upset(
        select(vertices, systems)
                                                    = c(0.7, 0.3)
= "freq"
       ,mb.ratio
      ,mainbar.y.label = "System Intersections"
,sets.x.label = "Genes per system"
                                                                = upset_texts
       ,point.size
        ,sets.bar.color = get_colors(vertices)
dev.print(pdf, "plots/fig1a_set_raw.pdf", width = 18, height = 10, onefile = F, useDingbats = F)
 fig3a_set <- vertices %>% filter(root == 37) %>% select(systems)
 upset(
         fig3a_set
    - c(0.7, 0.3)
- green gr
                                                               = upset_texts
dev.print(pdf, "plots/fig3a_set_raw.pdf", width = 16, height = 8, onefile = F, useDingbats = F)
# Figure 3B set diagram
fig3b_set <- vertices %>% filter(root < 37 & root >= 26) %>% select(systems)
 upset(
       ,order.by = "freq"
,mainbar.y.label = "System Intersections"
,sets.x.label = "Genes per system"
                                                                = upset_texts
       ,point.size
                                                               = 3.5
        ,sets.bar.color = get_colors(fig3b_set)
dev.print(pdf, "plots/fig3b_set_raw.pdf", width = 16, height = 8, onefile = F, useDingbats = F)
```



(a) Set diagram for Figure 1A



(b) Set diagram for Figure 3A



(c) Set diagram for Figure 3B

Figure 7: Set diagrams

# Supplementary Figures 2 and 3

Supplementary figures 2 and 3 help us see what nodes have been rooted at each LCA. Nodes rooted at previous LCAs are painted white.

```
plot_edges <- geom_path(</pre>
     data = partial_edges
,mapping = edge_aes
    ,color = edge_color
,size = 0.1
  plot_past <- geom_point(</pre>
    lot_past <- geom_point(
    data = past_genes
,mapping = aes(x, y, size = size)
,fill = past_fill
,color = past_color
,shape = past_genes$shape
,stroke = 0.25</pre>
  plot_text <- geom_text(</pre>
     data = current_genes
     ,mapping = text_aes
     ,size = 1
,vjust = 0
    ,nudge_y = 1.75
,alpha = 0.5
  base <- ggplot() +</pre>
     ggtitle(paste0(.y, " LCA (#", .x, ")")) +
     diff_theme +
     plot_edges +
     plot_past +
   plot_current_pies <- geom_scatterpie(</pre>
      data = current_genes
     ,mapping = pie_aes
     ,color = NA
   system_plots[[as.character(.x)]] <<- base +</pre>
    plot_current_pies +
     plot_pie_fill +
     plot_text
   ## Supplementary Figure 3
  plot_current_nodes <- geom_point(</pre>
      data = current_genes
     mapping = aes(x, y, fill = annotation, size = size)
,color = current_genes$color_node
,shape = current_genes$shape
,stroke = 0.25
  legend_hack <- guides(fill = guide_legend(override.aes = list(shape = 21)))</pre>
   function_plots[[as.character(.x)]] <<- base +</pre>
     plot_current_nodes +
     plot_scales +
     plot_size +
     plot_text +
     legend_hack
ggsave(
   ,arrangeGrob(grobs = rev(system_plots), ncol = 3)
              = 210
= 297
   ,height
   ,useDingbats = F
ggsave(
```

# Abundance

Abundance is the average number of proteins in neurotransmission orthogroups present in a species. This abundance is decomposed by orthogroup function. The orthogroup function is considered to be the function of its humans proteins in neurotransmission, as described in previous sections. In other words: "what would be the average number of"X"-related proteins in neurotransmission COGs in a species?" Loading initial resources:

```
library(magrittr)
library(neurotransmissionevolution)
data(
  ,gene_ids
  ,gene_cogs
  ,string_eukaryotes
  ,package = "neurotransmissionevolution"
                                                                                  col_types = "ci")
col_types = "ic")
col_types = "ici")
cog_roots
                               <- read_tsv("geneplast_roots.tsv",
                               <- read_tsv("geneplast_clade_names.tsv",
clade_names
clade_taxids
                               <- read_tsv("geneplast_clade_taxids.tsv",</pre>
gene_annotation <- read_tsv("../data/gene_annotation.tsv", col_types = "cc") %>%
  mutate(annotation = case_when(
     grepl("clearance", annotation) ~ "depletion"
,grepl("degradation", annotation) ~ "depletion"
     ,grepl("transport", annotation) ~ "synthesis"
     ,TRUE ~ annotation
```

We start by setting up reusable data frames with useful metrics.

```
# If a gene has more than 1 COG, select the oldest one.
# This is unusual, but can happen in cases of gene fusion, for instance.
gene_cogs %x>%
inner_join(cog_roots) %>%
group_by(string_id) %>%
filter(root == max(root))

# The function of a COG is the function of its proteins
cog_annotation <- gene_ids %>%
inner_join(gene_cogs) %>%
inner_join(gene_cogs) %>%
distinct(cog_id, annotation)

# Number of proteins in a COG in every species
cog_size_by_taxid <- cogs %>%
filter(cog_id %;n% gene_cogs[["cog_id"]]) %>%
count(taxid, cog_id, name = "cog_size")

# Number of COGs in every species
cog_n_by_taxid <- cog_size_by_taxid %>%
count(taxid, name = "cog_n")

# Number of proteins in every species
protein_n_by_taxid <- cog_size_by_taxid %>%

# Number of proteins in every species
protein_n_by_taxid <- cog_size_by_taxid %>%

# Number of proteins in every species
```

```
count(taxid, wt = cog_size, name = "protein_n")

# Mapping species to clade info
ordered_species <- string_eukaryotes %>%
select(taxid, ncbi_name) %>%
left_join(clade_taxids) %>%
left_join(clade_names, by = c("lca" = "root")) %>%
mutate(
    lca = as_factor(lca) %>% fct_rev
    ,ncbi_name = fct_reorder(ncbi_name, -taxid_order)
    ,clade_name = fct_reorder(clade_name, -taxid_order)
)

# Plotting colors
annotation_colors <- c(
    "depletion" = "#F40000"
    ,"excitability" = "#FFABO0"
    ,"receptor-associated" = "#B6EEO0"
    ,"ionotropic receptor" = "#43FFIC"
    ,"metabotropic receptor" = "#43FFIC"
    ,"signaling" = "#0091EA"
    ,"g-protein" = "#0033ff"
    ,"synthesis" = "#AAOOFF"
    ,"vescicle" = "#FFOOAA"
)</pre>
```

Abundance is finally computed as the number of proteins pertaining to a function divided by the number of neurotransmission COGs in a species.

```
abundance_by_function <- cog_size_by_taxid %>%
  inner_join(cog_annotation) %>%
  count(taxid, annotation, wt = cog_size, name = "annotation_count") %>%
  left_join(cog_n_by_taxid) %>%
  left_join(protein_n_by_taxid) %>%
  mutate(
    abundance = annotation_count / cog_n
    ,abundance_prop = annotation_count / protein_n
)
```

#### Plotting:

```
abundance_by_function %<>% left_join(ordered_species)
metazoa_line <- geom_vline(</pre>
  xintercept = "Mnemiopsis leidyi"
            = "#FF0000'
  ,linetype = "11"
 ,alpha
tick_function <- function(skip_head = 1, skip_tail = 1) {</pre>
    breaks <- seq(x[2], 0, length.out = 5) \%\% head(-skip_head) \%\% tail(-skip_tail)
    ceiling(breaks/0.25)*0.25
ggplot(abundance_by_function) +
 ggtitle("Number of proteins in a function divided by number of all neural COGs in species") +
  geom_bar(aes(x = ncbi_name, y = abundance, fill = annotation, color = annotation), stat = "identity") +
  scale_y_continuous("Average protein count in neural COGs", breaks = tick_function(), minor_breaks = NULL) +
  facet_grid(annotation ~ clade_name, scales = "free", space = "free_x") +
  scale_fill_manual(values = annotation_colors %>% darken(0.1)) +
  scale_color_manual(values = annotation_colors %>% darken(0.2)) +
  theme (
     panel.spacing
    ,strip.background.x = element_blank()
    ,strip.background.y = element_rect(fill="#E0E0E0")
,panel.grid.major.x = element_blank()
    panel.grid.major.y = element_line(colour = "#F5F5F5", size = 0.25)
```

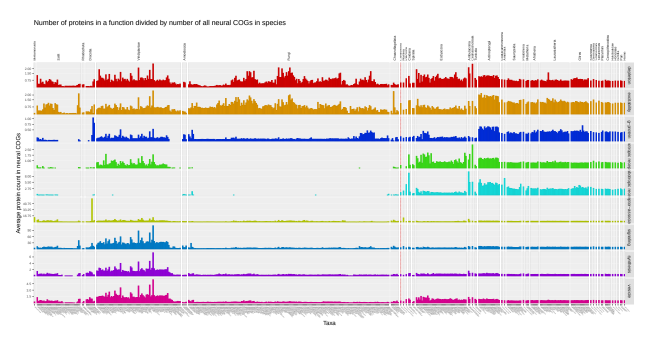


Figure 8: Abundance values by species. Species are ordered like in Supplementary Figure S1.

```
ggsave("plots/fig5_raw.pdf", width = 16, height = 8)
ggplot(abundance_by_function) +
  ggtitle("Number of proteins in a function divided by number of all neural proteins in species") +
  metazoa_line +
  geom_bar(aes(x = ncbi_name, y = abundance_prop, fill = annotation, color = annotation), stat = "identity") +
facet_grid(annotation ~ clade_name, scales = "free", space = "free_x") +
scale_y_continuous( minor_breaks = NULL) +
  scale_fill_manual(values = annotation_colors %>% darken(0.1)) +
   scale_color_manual(values = annotation_colors %>% darken(0.2)) +
  theme(
      panel.spacing
     ,strip.background.x = element_blank()
     ,strip.background.y = element_rect(fill="#E0E0E0")
     ,panel.grid.major.x = element_blank()
     panel.grid.major.y = element_line(colour = "#F5F5F5", size = 0.25)
,panel.background = element_rect(fill = '#EEEEEE', colour = '#E0E0E0')
,strip.text.x = element_text(size = 6, angle = 90, hjust = 0, vjust = 0.5)
,strip.text.y = element_text(size = 8, vjust = 0.5)
                                = element_text(size = 2, angle = -45, vjust = 0, hjust = 0)
                                = element_text(size = 6)
                                = element_line(size = 0.1)
     ,legend.position
```

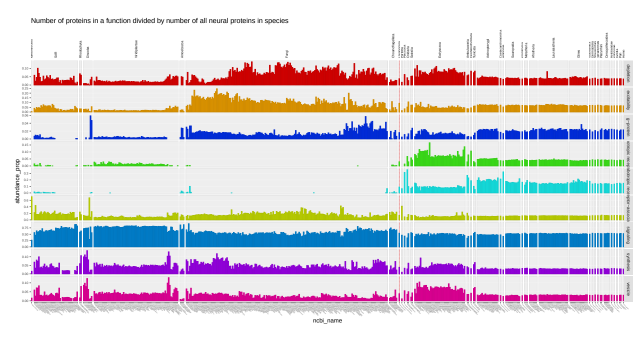


Figure 9: Abundance values by species. Species are ordered like in Supplementary Figure S1.

```
ggsave("plots/fig5_prop_raw.pdf", width = 16, height = 8)
```

Abundances averaged by clades.

```
ggplot(abundance_by_function) +
  geom_bar(aes(x = clade_name, y = abundance, fill = annotation), stat="summary", fun = "mean") +
  scale_fill_manual(values = annotation_colors, guide = "none") +
  facet_grid(annotation ~ ., scales = "free") +
  theme(
    panel.spacing = unit(1, "pt")
    ,strip.text.y = element_text(angle = 0, hjust = 0)
    ,axis.text.x = element_text(size = 5, angle = -45, vjust = 0, hjust = 0)
    ,axis.text.y = element_text(size = 5)
)
```

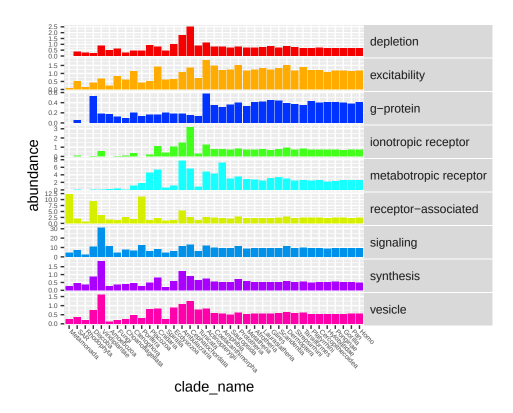


Figure 10: Abundances averaged by clades.

Breaking the data even further into the number of proteins in each neurotransmission COG.

```
cog_annotation_collapsed <- cog_annotation %>%
  group_by(cog_id) %>%
   summarise(annotation = paste(annotation, collapse = "/"))
# Simply joining cog annotation and sizes
protein_count_by_cog <- cog_size_by_taxid %>%
   inner_join(cog_annotation_collapsed) %>%
   left_join(ordered_species) %>%
  arrange(annotation) %>%
  mutate(cog_id = fct_inorder(cog_id))
ggplot(protein_count_by_cog) +
  metazoa_line +
   geom_bar(aes(x = ncbi_name, y = cog_size, fill = annotation), stat = "identity") +
   scale_fill_manual(values = annotation_colors %>% darken(0.2), guide = "none") + scale_y_continuous(breaks = tick_function(3, 1), minor_breaks = NULL) +
   facet_grid(cog_id ~ ., scales = "free_y") +
       panel.spacing
      ,panel.grid.major.x = element_blank()
      ,panel.grid.major.x = element_blank()
,panel.grid.major.y = element_line(size = 0.1, linetype = "dashed")
,strip.text.y = element_text(size = 4, angle = 0, hjust = 0)
,axis.text.x = element_text(size = 1.5, angle = -45, vjust = 0, hjust = 0)
,axis.text.y = element_text(size = 4)
                                  = element_line(size = 0.1)
      ,axis.ticks
```

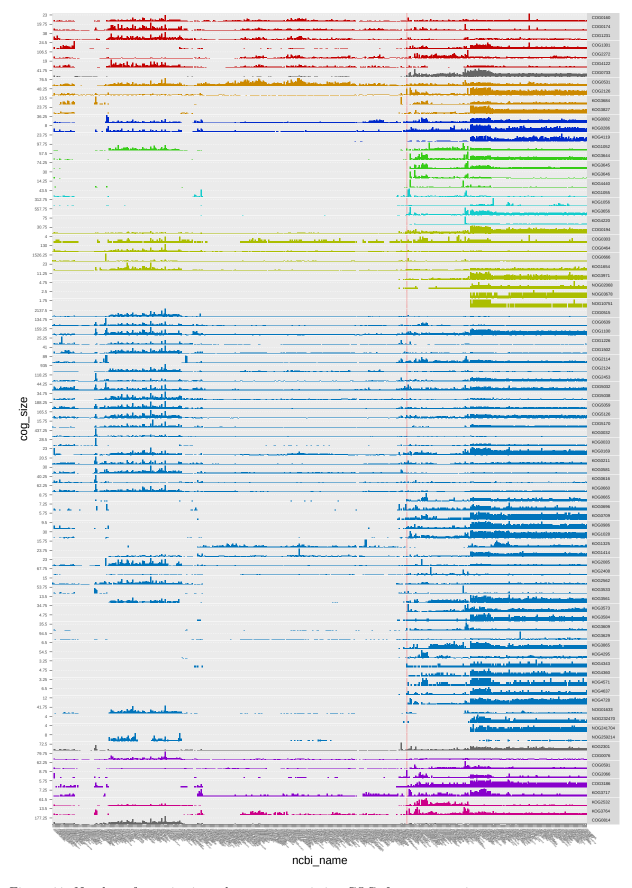


Figure 11: Number of proteins in each neurotransmission COG, for every species.