

neurogenes synteny project

1.

Paper notes - neural genes presence/absences

Moroz et al., 2014 [1]

- Table 34S: -structure of figure shows that it is from bilaterian perspective (closer to bilateria more rectangles filled in)
- remember there are also random absences in cnidaria; the mirror of bilateria, but all the proteins are characterized from bilateria
- no examples where ctenophores or sponges don't have something present in fungi, capsaspora, monosiga
- Suppl Table 12as - it's possible didn't use Amphimedon genome, but Amphimedon was covered by the other papers and I crossreferenced them.

Mentioned in text:

Not in ctenophores:

neurogenin

NeuroD

Achaete-scute

REST

HOX

Otx

-not that much overlap in genes looked at by Riesgo vs Moroz

Ryan et al., 2013 [2]

netrin, slit, unc-5 (axon guidance) not in Mnemiopsis or Amphimedon

-used genomes, since based on Alie and Manuel 2010

Supplementary Table S17: Presence and absence of post-synaptic genes - pretty much Alie and Manuel 2010

Supplementary Table S19: Presence and absence of Dopamine / Norepinephrine /Epinephrine Biosynthetic Pathway components

-are seqs of the animals in S17 genomes? Unsure, but all animals in table have genomes (and the Mle seqs are from the genome)

-AMPA iGluR and NMDA iGluR included as iGluR

Alie and Manuel, 2010 [3]

-used genomes

-Ryan built on Fig. 1. Cross ref with current data to make sure have everything.

-Only use Monosiga, Trichoplax, Amphimedon, Nematostella, Hydra, Homo

Capitella (3 absences), Drosophila (2 absences), Homo very similar with few differences

Unicellular animals mostly missing everything (except B-cat and PMCA). Start with Monosiga which has more things. B-cat and PMCA are ancient - interesting?

AMPA and NMDAR collapsed into iGluR in table; presence of one of these trumped absence of the other
PKC alpha-beta-gamma = PKC on table

Table

Table abbreviations

DBH - dopamine-B-hydroxylase

DDC - DOPA decarboxylase

TH - tyrosine hydroxylase

TPH - tryptophan hydroxylase

PAH - phenylalanine hydroxylase

GAD - glutamate decarboxylase

Qdpr - quinoid dihydropteridine reductase,

Slc18A2 = Homo sapiens solute carrier family 18 member 2,

Pnmt = phenylethanolamine N-methyltransferase

Missing domains

Piccolo - Pleurobrachia - missing ZF (Moroz et al., 2014)

Erbin - Pleurobrachia - missing PDZ (Moroz et al., 2014)

Species names written to the broadest level - eg. *Monosiga brevicollis* in Riesgo et al but only *Monosiga* in Moroz, so put *Monosiga* only

Many entries have NA but if combine:

Salpingoeca + *Monosiga* = Choanoflagellida

Pleurobrachia + Mnemiopsis = Ctenophora

Amphimedon + Oscarella = Porifera

Nematostella + Hydra = Cnidaria

Get only 4 entries that have an NA.

(*What about 0/1s (conflicting info?)*) >> decided to transform 0/1s into NA

Loss_Status: P1C0: present in Porifera, absent in Ctenophora - 6 instances

C1P0: present in Ctenophora, absent in Porifera - 3 instances

T0: absent in Trichoplax but present in Ctenophora or Porifera - 5 instances

There is only 2 instances where Capsaspora has a 1 while choanoflagellates have 0: GABAR and DDC. Don't use column in second iteration

'Fungi' is very vaguely defined - don't use column in second iteration

Stopped at Delta catenin

Make new table where all 0/1s or missing_domains (i.e. not 0,1,NA) into NA

Create a new table where species for Ctenophora, Porifera combined: <https://stackoverflow.com/questions/14563531/combine-column-to-remove-nas>

Synteny programs/papers

ghost locus hypothesis:

Ramos et al., 2012 [4]

<https://www.sciencedirect.com/science/article/pii/S0960982212009888>

-first ghost locus paper - parahox, Amphimedon + Trichoplax

Fortunato et al, 2014 [5] - sycon Parahox

<https://www.nature.com/articles/nature13881>

-second ghost locus paper?

Ferrier 2015 [6] (review)

<https://academic.oup.com/bfg/article/15/5/333/1741867>

Reviews:

Liu et al 2018 [7]

<https://bmcbioinformatics.biomedcentral.com/articles/10.1186/s12859-018-2026-4>

-most synteny programs designed with assumption working on complete high quality genomes.

-need N50 of at least 1Mb for robust synteny analysis

-Fig. 5 - Below 0.5 Mb error rate skyrockets - looks like close to 20-30% for ~100-200 kbs N50s -This can represent a major source of systemic bias in my analysis

Scaffold N50s of my genomes

Mnemiopsis (from orig [2]): 187 kb

Pleurobachia (Suppl Table 5S from [1]): 20.607 kb

Amphimedon orig, Ensembl (https://metazoa.ensembl.org/Amphimedon_queenslandica/Info/Annotation/):

120 kb >> likely improved, looking for Aqu2.1 (or is this only a re-annotation not reassembly?)

- Slightly confused with scaffold N50 stats in [8] suppl Table S2.3.2

Oscarella carmela (Suppl from [9]): 5.897 kb!! :O

Data/downloads

Pleurobachia genome: https://www.ncbi.nlm.nih.gov/assembly/GCA_000695325.1

P.bachei_draft_genome_v.1.1

Organism: Pleurobrachia bachei (ctenophores)

Submitter: University of Florida

Date: 2014/05/21

Assembly level: Scaffold

Genome representation: full

RefSeq category: representative genome

GenBank assembly accession: GCA_000695325.1 (latest)

RefSeq assembly accession: n/a

IDs: 180401 [UID] 1073948 [GenBank]

-paper said deposited at Moroz's website; links don't work on website.

Mnemiopsis leidyi: https://www.ncbi.nlm.nih.gov/assembly/GCA_000226015.1/

MneLei_Aug2011

Organism name: Mnemiopsis leidyi (sea walnut)

BioSample: SAMN02953801 BioProject: PRJNA64405

Submitter: National Human Genome Research Institute, National Institutes of Health

Date: 2011/09/19

Assembly level: Scaffold

Genome representation: full

RefSeq category: representative genome

GenBank assembly accession: GCA_000226015.1 (latest)

RefSeq assembly accession: n/a

RefSeq assembly and GenBank assembly identical: n/a

WGS Project: AGCP01

Assembly method: Phusion v. 1.02

Genome coverage: 12xSequencing technology: 454 GS-FLX Titanium; Illumina GA IIX

IDs: 304208 [UID] 304208 [GenBank]

Amphimedon queenslandica: v1.0: https://www.ncbi.nlm.nih.gov/assembly/GCF_000090795.1

Organism: Amphimedon queenslandica (sponges)

Submitter: US DOE Joint Genome Institute (JGI-PGF)

Date: 2010/05/28

Assembly level: Scaffold

Genome representation: full

RefSeq category: representative genome
GenBank assembly accession: GCA_000090795.1 (latest)
RefSeq assembly accession: GCF_000090795.1 (latest)
IDs: 293608 [UID] 111438 [GenBank] 293608 [RefSeq]

Oscarella carmela: <http://www.compagen.org/datasets.html> OCAR not *Oscarella* sp.

-added .fna and gzipped

-*Oscarella carmela* (this assembly) renamed *Oscarella pearsei*; *Oscarella* sp. in Compagen redescribed *Oscarella carmela*. See <https://journals.plos.org/plosone/article?id=10.1371/journal.pone.0183002> - old *O. carmela* papers based on two species (*carmela* + *pearsei*). Both in compagen, but although there is press release compagen didn't change their names in db. *Can't find annotation file*.

Papers+Links

Riesgo et al., 2014

<https://academic.oup.com/mbe/article/31/5/1102/993377>

Neural genes Fig

<https://academic.oup.com/view-large/figure/74385341/msu057f3p.jpeg>

Moroz et al., 2014

<https://www.nature.com/articles/nature13400>

Table 34S: neural genes

<https://media.nature.com/original/nature-assets/nature/journal/v510/n7503/extref/nature13400-s1.pdf>

Ryan et al, 2013

<http://science.sciencemag.org/content/342/6164/1242592>

Suppl Mat:

<http://science.sciencemag.org/content/sci/suppl/2013/12/11/342.6164.1242592.DC1/Ryan.SM.pdf>

Alie and Manuel, 2010

<https://bmcevolbiol.biomedcentral.com/articles/10.1186/1471-2148-10-34>

Srivastava et al., 2010

Suppl.S8.9 - neural genes <https://media.nature.com/original/nature-assets/nature/journal/v466/n7307/extref/nature09201-s1.pdf>

Nichols et al., 2012

<https://www.pnas.org/content/109/32/13046>

Suppl

<https://www.pnas.org/content/pnas/suppl/2012/07/25/1120685109.DCSupplemental/sapp.pdf>

2. Should we be expecting these genes in these animals?

Why should they use similar genes?

How misguided is this approach? What is the true question implied by this approach?

References

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3. Alié A, Manuel M. The backbone of the post-synaptic density originated in a unicellular ancestor of

choanoflagellates and metazoans. *BMC Evolutionary Biology*. 2010;10: 34. doi:10.1186/1471-2148-10-34

4. Ramos OM, Barker D, Ferrier DE. Ghost loci imply hox and parahox existence in the last common ancestor of animals. *Current biology*. Elsevier; 2012;22: 1951–1956.

5. Fortunato SA, Adamski M, Ramos OM, Leininger S, Liu J, Ferrier DE, et al. Calcisponges have a parahox gene and dynamic expression of dispersed nk homeobox genes. *Nature*. Nature Publishing Group; 2014;514: 620.

6. Ferrier DE. The origin of the hox/parahox genes, the ghost locus hypothesis and the complexity of the first animal. *Briefings in functional genomics*. Oxford University Press; 2015;15: 333–341.

7. Liu D, Hunt M, Tsai IJ. Inferring synteny between genome assemblies: A systematic evaluation. *BMC bioinformatics*. BioMed Central; 2018;19: 26.

8. Srivastava M, Simakov O, Chapman J, Fahey B, Gauthier ME, Mitros T, et al. The amphimedon queenslandica genome and the evolution of animal complexity. *Nature*. Nature Publishing Group; 2010;466: 720.

9. Nichols SA, Roberts BW, Richter DJ, Fairclough SR, King N. Origin of metazoan cadherin diversity and the antiquity of the classical cadherin/ β -catenin complex. *Proceedings of the National Academy of Sciences*. National Acad Sciences; 2012;109: 13046–13051.