

Evolutionary history of Cytoplasmic Polyadenylation Element-Binding Proteins

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LIST OF ABBREVIATIONS

<Insert text>	<Insert text>
CPEB	Cytoplasmic Polyadenylation Element-Binding Protein
RBP	RNA-Binding Protein
RRM	RNA Recognition Motif
RBD	RNA-Binding Domain (region of CPEB containing RRM)
ZZ	Zinc (binding domain)
poly(A)	Polyadenosine

1 INTRODUCTION: BACKGROUND INFORMATION AND SCIENTIFIC RATIONALE

1.1 BACKGROUND INFORMATION

Cytoplasmic Polyadenylation Element-Binding Proteins (CPEBs) are pivotal regulators of gene expression during gametogenesis and the initial stages of embryonic development of bilaterians. These RNA-binding proteins recognize sequences (consensus UUUUA₁₋₂U) in the 3'untranslated region of specific mRNAs and recruit cytoplasmic poly(A)-polymerases that increase the poly(A)-tail length of substrate mRNAs and consequently their translation. CPEBs have not been studied in outside of bilaterians, but a recent study identified them as one of "25 groups of metazoan-specific genes that are essential across the Animal Kingdom" (Paps and Holland, 2018).

1.2 RATIONALE

CPEB orthologs corresponding to both CPEB1 and CPEB2 subgroups have been found in many bilaterian groups, but detailed phylogenetic analyses that have included non-bilaterian animals as well as closely related non-animal lineages have yet to be performed. The results of such analyses could shine light on the origin and evolution of cytoplasmic polyadenylation.

1.3 HYPOTHESES

Orthologs of CPEB1 and CPEB2 are present in the genomes of all animals and absent from the genomes of non-animals. The most identical proteins in non-animal genomes will be RNA-binding proteins with essential functions shared throughout Eukaryota. Metazoan proteins with sequence similarity with CPEBs, which are not CPEB orthologs or orthologs of proteins present in non-metazoan genomes, share biochemical and biological functions with CPEBs.

1.4 OBJECTIVES

1. To validate the presence or absence of CPEB orthologs in genomes of non-bilaterian animals.
2. To identify non-animal proteins that share a most-recent common genic ancestor with CPEBs.
3. To document the evolutionary history of CPEBs and related genes.

2 STUDY DESIGN AND ENDPOINTS

- 1) **Identify putative CPEBs within genomes:** Starting with the RBD (tandem RRM and ZZ domain) of human CPEB1 (NP_001275748.1; AA 234-479) and CPEB4 (AAH36899.1; AA 66-309) protein sequences (Afroz et al., 2014), perform TBLASTN searches against *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe* (Fungi), *Arabidopsis thaliana*, *Capsaspora oewazarki* (Filasterea), *Salpingoeca*

rosetta (Choanoflagellata), *Monosiga brevicollis* (Choanoflagellata), *Amphimedon queenslandica* (Porifera), *Mnemiopsis leidyi* (Ctenophora), *Nematostella vectensis* (Cnidaria), *Trichoplax adhaerens* (Placozoa), *Drosophila melanogaster* (Arthropoda), *Capitella teleta* (Annelida), *Schmidtea mediterranea* (Platyhelminth), *Caenorhabditis elegans* (Nematoda), as well as *Mus musculus* and *Danio rerio* (Vertebrata) gene models to identify organisms with CPEB orthologs.

- 2) **Validate orthology:** Perform reciprocal best BLASTP searches to validate orthology of human CPEB1 and CPEB4 with hits.
- 3) **Generate a databases of CPEB orthologous sequences:** A database of full-length sequence putative CPEBs and a database of CPEB RBDs will be built using the sequences identify at the end of (2) above.
- 4) **Align putative CPEBs:** Putative CPEB RBDs from the database generated in (3) will be aligned with MAFFT using default parameters.
- 5) **Identify proteins that share most-recent genic ancestor with CPEBs**
 - a. For species without CPEB orthologs, identify 1-3 sequences most similar to CPEB RBDs
 - b. Identify human sequences that are most identical to sequences from (3a).
 - c. Identify 1-3 genes that are not CPEB orthologs (according to 1 and 2, above) but share the highest sequence similarity to CPEBs in *H. sapiens*, *D. melanogaster*, *S. mediterranea*, *N. vectensis*, *M. leidyi*, *A. queenslandica*, *T. adhaerens* (i.e. genomes which are expected to have CPEBs)
- 6) **Generate a database of CPEB-like sequences:** A database of of RBDs from non-CPEB sequences will be built for genes identified in (5a-c).
- 7) Infer phylogenetic relationships among CPEBs and non-CPEB sequences (with similarity to CPEBs) by estimating gene trees of RBD sequence databases from step (3) and step (6).
 - a. IQTREE
 - b. RAxML with 25 starting parsimony trees and 25 random starting trees;
 - c. MrBayes
- 8) Choose best tree for main figure

3 WORK COMPLETED SO FAR W DATES

<sampling plan>

<statistical tests>

<inference criteria (p-values, bayes factors, model fit indices>

<criteria for accepting or rejecting hypotheses>

<description of or link to repo with custom scripts>

<command lines>

4 LITERATURE REFERENCES

Paps, J & Holland, PW, 2018, 'Reconstruction of the ancestral metazoan genome reveals an increase in genomic novelty'. *Nature Communications*, vol 9.

Afroz T, Skrisovska L, Belloc E, Guillén-Boixet J, Méndez R, Allain FH. A fly trap mechanism provides sequence-specific RNA recognition by CPEB proteins. *Genes Dev.* 2014;28(13):1498-1514. doi:10.1101/gad.241133.114

5 PHYLOTOCOL AMENDMENT HISTORY

Version	Date	Significant Revisions