

Eutherian comparative genomic analysis protocol

Marko Premzl (✉ Marko.Premzl@alumni.anu.edu.au)

The Australian National University Alumni <https://orcid.org/0000-0002-3362-689X>

Method Article

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Abstract

The eutherian genomics momentum greatly advanced biological and medical sciences. Yet, **future revisions and updates of eutherian genomic sequence data sets** were expected, due to **potential genomic sequence errors** and incompleteness of genomic sequences. The **eutherian comparative genomic analysis protocol** was established as guidance in protection against **potential genomic sequence errors** in public eutherian genomic sequence assemblies. The protocol revised, updated and published **14 major eutherian gene data sets**, including **2615 complete coding sequences** deposited in European Nucleotide Archive as curated third party data gene data sets under accession numbers: [FR734011-FR734074](#), [HF564658-HF564785](#), [HF564786-HF564815](#), [HG328835-HG329089](#), [HG426065-HG426183](#), [HG931734-HG931849](#), [LM644135-LM644234](#), [LN874312-LN874522](#), [LT548096-LT548244](#), [LT631550-LT631670](#), [LT962964-LT963174](#), [LT990249-LT990597](#), [LR130242-LR130508](#) and [LR760818-LR761312](#).

Introduction

The eutherian genomics momentum greatly advanced biomedical research. For example, one major aim of initial sequencing and analysis of human genome was to **update and revise human genes**, as well as to uncover potential new drugs, drug targets, and molecular markers in medical diagnostics. However, **future revisions and updates of eutherian genomic sequence data sets** were expected, due to **potential genomic sequence errors** and incompleteness of reference genomic sequences. Specifically, the **potential genomic sequence errors** included **Sanger DNA sequencing method errors** (artefactual nucleotide deletions, insertions and substitutions) and **analytical and bioinformatical errors** (erroneous gene annotations, genomic sequence misassemblies). Thus, the **eutherian comparative genomic analysis protocol** [RRID:SCR_014401](#) was established as **guidance in protection against potential genomic sequence errors** in public eutherian genomic sequence assemblies ^{1,2,3,4,5,6,7,8,9,10,11,12,13,14,15}.

Reagents

Public biological databases:

[Ensembl genome browser](#)

[European Nucleotide Archive](#)

[National Center for Biotechnology Information](#)

Equipment

The present protocol required personal computer and internet connectivity.

Procedure

Eutherian comparative genomic analysis protocol

The eutherian comparative genomic analysis protocol [RRID:SCR_014401](#) integrated **gene annotations**, **phylogenetic analysis** and **protein molecular evolution analysis** into one framework of eutherian gene descriptions. The protocol included 3 original genomics and protein molecular evolution tests, including **tests of reliability of public eutherian genomic sequences** using genomic sequence redundancies, **tests of contiguity of public eutherian genomic sequences** using multiple pairwise genomic sequence alignments and **tests of protein molecular evolution** using relative synonymous codon usage statistics.

1. Gene annotations

The eutherian **gene annotations** included **gene identifications in public genomic sequence assemblies**, **analyses of gene features**, **tests of reliability of public eutherian genomic sequences** and **tests of contiguity of public eutherian genomic sequences**.

1.1. All analyses and manipulations of nucleotide and protein sequences used sequence alignment editor [BioEdit](#).

1.2. The eutherian reference genomic sequence data sets were accessible in National Center for Biotechnology Information's (NCBI) [GenBank](#), as well as in [Ensembl](#) genome browser.

1.3. The **identifications of potential coding sequences** used public eutherian reference genomic sequence assemblies and NCBI's [BLAST program including BLAST Genomes](#) and [Ensembl genome browser's BLAST or BLAT programs](#).

1.4. The **analyses of gene features** used **potential coding sequences** and direct evidence of eutherian gene annotations accessible in NCBI's [nr](#), [est_human](#), [est_mouse](#) and [est_others](#) databases.

1.5. The **tests of reliability of eutherian public genomic sequences** analysed potential coding sequences using good laboratory practice in Sanger DNA sequencing method. The first test steps analysed nucleotide sequence coverages of **potential coding sequences** using [NCBI's BLAST program](#) and processed Sanger DNA sequencing reads or traces accessible in [NCBI's Trace Archive](#). The second test steps discriminated **complete coding sequences** and **putative coding sequences**. Specifically, the tests described **potential coding sequences** as **complete coding sequences** only if consensus trace nucleotide sequence coverages were available for every nucleotide. Alternatively, if consensus trace nucleotide sequence coverages were not available for every nucleotide, the **potential coding sequences** were described as **putative coding sequences** that were not used in analyses. For example, the good laboratory practice in Sanger DNA sequencing method exacted that minimal consensus trace nucleotide sequence coverage included 2 identical trace nucleotide sequences.

1.6. The **tests of contiguity of public eutherian genomic sequences** included multiple pairwise genomic sequence alignments. The tests used public eutherian reference genomic sequences encoding **complete coding sequences** and [mVISTA's program AVID](#). In eutherian genomic sequences, the tests analysed

translated exon numbers, as well as their chimerisms and relative orders and orientations. The tests of contiguity of eutherian public genomic sequences did not use masking of transposable elements in public eutherian reference genomic sequence assemblies.

1.7. The **curated eutherian gene collections** were deposited in [European Nucleotide Archive](#) as third party data gene data sets. The revised and updated eutherian gene classifications and nomenclatures used [guidelines of human gene nomenclature](#) and [guidelines of mouse gene nomenclature](#).

2. Phylogenetic analysis

The **phylogenetic analysis** included **protein and nucleotide sequence alignments**, **calculations of phylogenetic trees** and **calculations of pairwise nucleotide sequence identities**.

2.1. The **complete coding sequences** were translated using [BioEdit](#) and then aligned at amino acid level using [ClustalW](#) in **protein amino acid sequence alignments**. The **protein amino acid sequence alignments** were manually corrected, and **nucleotide sequence alignments** were prepared accordingly using [BioEdit](#).

2.2. The **calculations of phylogenetic trees** used **nucleotide sequence alignments** and [MEGA program](#).

2.3. Using **nucleotide sequence alignments**, the **pairwise nucleotide sequence identities of eutherian complete coding sequences** were calculated using [BioEdit](#). The statistical analyses using Microsoft Office Excel statistical functions included calculations of average pairwise nucleotide sequence identities (\bar{a}) and their average absolute deviations (\bar{a}_{ad}), as well as largest (a_{max}) and smallest (a_{min}) pairwise nucleotide sequence identities.

3. Protein molecular evolution analysis

The **protein molecular evolution analysis** included **analyses of protein amino acid sequence features** and **tests of protein molecular evolution** using relative synonymous codon usage statistics.

3.1. The **protein amino acid sequence features** were annotated manually, including analyses of common cysteine amino acid residue patterns among eutherian major protein clusters.

3.2. The **tests of protein molecular evolution** using relative synonymous codon usage statistics integrated patterns of nucleotide sequence similarities with protein primary structures. Using **nucleotide sequence alignments**, the [MEGA](#) calculated relative synonymous codon usage statistics as ratios between observed and expected amino acid codon counts ($R = \text{Counts} / \text{Expected counts}$). The amino acid codons including $R \leq 0.7$ were described as not preferable amino acid codons. In reference protein amino acid sequences, the tests described invariant amino acid sites (invariant alignment positions), forward amino acid sites (variant alignment positions that did not include amino acid codons with $R \leq 0.7$) and compensatory amino acid sites (variant alignment positions that included amino acid codons with $R \leq 0.7$).

Troubleshooting

Time Taken

Anticipated Results

The **eutherian comparative genomic analysis protocol** [RRID:SCR_014401](#) published **2615 complete coding sequences** that included: 64 interferon- γ -inducible GTPase genes ¹, 255 ribonuclease A genes ², 119 Mas-related G protein-coupled receptor genes ³, 116 lysozyme genes ⁴, 128 adenohipophysis cystine-knot genes ⁵, 30 D-dopachrome tautomerase and macrophage migration inhibitory factor genes ⁵, 100 growth hormone genes ⁶, 211 tumor necrosis factor ligand genes ⁸, 149 globin genes ⁹, 121 kallikrein genes ¹⁰, 211 adiponectin genes ¹¹, 349 connexin genes ¹³, 267 fibroblast growth factor genes ¹⁴, and 495 interferon genes ¹⁵. These **curated eutherian third party data gene data sets** were applicable in gene annotations and genome analyses – including differential gene expansion analyses and initial descriptions of human genes, phylogenetic analyses and protein molecular evolution analyses.

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Author: Marko Premzl PhD, The Australian National University Alumni (WFH), 4 Kninski trg Sq., Zagreb, Croatia

E-mail contact: Marko.Premzl@alumni.anu.edu.au

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