

**Research Article** 

# Comparison of seven complete mitochondrial genomes from *Lamprologus* and *Neolamprologus* (Chordata, Teleostei, Perciformes) and the phylogenetic implications for Cichlidae

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Abstract

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**Copyright:** © Jiachen Wang et al. This is an open access article distributed under terms of the Creative Commons Attribution License (Attribution 4.0 International – CC BY 4.0). In this study, mitochondrial genomes (mitogenomes) of seven cichlid species (*Lamprologus kungweensis*, *L. meleagris*, *L. ornatipinnis*, *Neolamprologus brevis*, *N. caudopunctatus*, *N. leleupi*, and *N. similis*) are characterized for the first time. The newly sequenced mitogenomes contained 37 typical genes [13 protein-coding genes (PCGs), two ribosomal RNA genes (rRNAs) and 22 transfer RNA genes (tRNAs)]. The mitogenomes were 16,562 ~ 16,587 bp in length with an A + T composition of 52.1~58.8%. The cichlid mitogenomes had a comparable nucleotide composition, A + T content was higher than the G + C content. The AT-skews of most mitogenomes were inconspicuously positive and the GC-skews were negative, indicating higher occurrences of C than G. Most PCGs started with the conventional start codon, ATN. There was no essential difference in the codon usage patterns of these seven species. Using Ka/Ks, we found the fastest-evolving gene were *atp8*. But the results of p-distance indicated that the fastest-evolving gene was *nad6*. Phylogenetic analysis revealed that *L. meleagris* did not cluster with *Lamprologus* species, but with species from the genus *Neolamprologus*. The novel information obtained about these mitogenomes will contribute to elucidating the complex relationships among cichlid species.

Key words: Cichlidae, Lamprologus, mitogenome, Neolamprologus, phylogenetic analyses

## Introduction

Cichlids (Teleostei: Perciformes: Cichlidae) are widely distributed across the Neotropics, Africa, the Middle East, Madagascar, as well as southern India and Sri Lanka (Smith et al. 2008; López-Fernández et al. 2010). They stand out as one of the most species-diverse groups of acanthomorphs. Kullander (1998) divided the family Cichlidae into eight subfamilies: Astronotinae, Cichlasomatinae, Cichlinae, Etroplinae, Geophaginae, Heterochromidinae, Pseudocrenilabrinae, and Retroculinae. The ninth subfamily, the Ptychochrominae, was later

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recognized by Sparks and Smith (2004). Cichlids gained recognition as a prominent model species for the study of evolutionary biology due to the numerous species, diverse genetics, distinct evolutionary lineages, and significant ecological and morphological divergences (Kocher 2004; Schwarzer et al. 2015; Reis et al. 2016; Nam et al. 2021).

African cichlids (subfamily Pseudocrenilabrinae) boasted an abundant variety of more than 2000 species (Brawand et al. 2014; Astudillo-Clavijo et al. 2022). Biologists have long been fascinated by the diversity of cichlids in the East African cichlid radiation (EAR), which has promoted high levels of endemism in the Lakes Tanganyika, Malawi, and Victoria (Kornfield and Smith 2000). Lake Tanganyika is a deep tropical and large Rift Valley lake with an age of 9-12 million years (Irisarri et al. 2018). It has the most diverse species of cichlid fish in terms of morphology, ecology, and behavior, including several mouth-brooding and substrate-spawning lineages (Takahashi 2003; Salzburger 2009). The cichlid fauna of Lake Tanganyika is dominated by lamprologine cichlids, which colonized most lacustrine habitats, but most often inhabits the littoral zone (Sturmbauer et al. 2010). Although classified as a single tribe, lamprologine cichlids exhibit significant diversity in morphology, ecology, and behavior. Lamprologus kungweensis, Lamprologus meleagris, Lamprologus ornatipinnis, Neolamprologus brevis, Neolamprologus caudopunctatus, Neolamprologus leleupi, and Neolamprologus similis are among the smallest species within the lamprologine cichlids, small enough to live inside the empty shells of gastropod mollusks (Sturmbauer et al. 2010). These species are regarded as a highly valuable ornamental species in the aquatic trade industry due to their ease of maintenance and handling in aquariums (Nam et al. 2021).

The genera *Lamprologus* and *Neolamprologus* can be difficult to distinguish due to their similar morphology, ecology, and behavior. As discussed by Stiassny (1991), meristic and morphometric measurements, osteology, and dentition were insufficient to differentiate between the species, as many of these traits were homoplastic. Furthermore, there might be instances of ancient ancestral polymorphism, introgressive hybridization, or lack of diagnostic synapomorphic characters among certain species within these two genera, further complicating their classification (Sturmbauer et al. 2010; Gante et al. 2016). Therefore, additional method, like molecular analysis might be required for more accurate classification.

Mitochondria are organelles found in most eukaryotic cells that play a critical role in energy production (Hebert et al. 2010). The mitochondrial genome (mitogenome) of acanthomorph fishes is usually a circular, double-stranded molecule that ranges from 16 to 23 kbp in size. It typically contains 13 protein-coding genes (PCGs), two ribosomal RNA genes (rRNAs), 22 transfer RNA genes (tRNAs), and one control region (CR) (Iwasaki et al. 2013). Mitogenomes have the characteristics of high evolutionary rate, matrilineal inheritance, low molecular weight, simple structure, and ease of amplification, which makes them a reliable marker for studying phylogenetics (Ye et al. 2022; Wang et al. 2023). Mitogenome components, such as *nad2* or *rrnL*, are widely used for phylogenetic analyses (Sturmbauer et al. 2010; Schwarzer et al. 2015). Although partial mitochondrial sequences can offer some insights into evolutionary relationships, they are limited in their ability to provide a comprehensive understanding due to the absence of information such as gene rearrangement, genetic

code changes, replication, and transcriptional regulation patterns. Therefore, complete mitogenome sequences can be more beneficial as they can provide improved resolution and sensitivity for investigating evolutionary relationships (Li et al. 2019; Fiteha et al. 2023; Wang et al. 2023).

In this study, we report the complete mitogenome organizations and characteristics of seven species (*L. kungweensis*, *L. meleagris*, *L. ornatipinnis*, *N. brevis*, *N. caudopunctatus*, *N. leleupi*, and *N. similis*). We also performed a phylogenetic analysis of the seven complete mitogenomes obtained in this study with the published complete cichlid mitogenomes. We hope that our study can enable better comprehension of cichlid biodiversity and expand genetic resources for future cichlid comparisons.

## **Materials and methods**

#### Sample collection and DNA extraction

The seven species are commonly sold as ornamental fish and can be found in many pet markets. Specimens were obtained from the Qiqiaoweng pet market in Nanjing, Jiangsu province, China. The specimens were identified using morphological characteristics described in FishBase (https://www.fishbase.de/). No fish were sacrificed during this study. The fish were reared at the Laboratory of Animal Molecular Evolution, Nanjing Forestry University. Total genomic DNA was extracted from each fin using a FastPure Cell/Tissue DNA Isolation Mini Kit (Vazyme, Nanjing, China), and stored at -80 °C for future use.

### Genome sequencing, assembly, and annotation

Seven complete mitogenomes were sequenced on an Illumina platform (Personalbio Nanjin, China) using total genomic DNA. The genomic DNA was used to generate an Illumina library with an insert size of 400 bp. The clean data were then assembled in Geneious Prime 2022 software, using *Lamprologus signatus* (MZ427900.1) as a template. The mitogenomes were assembled and manually revised using DNAstar v. 7.1 (Madison, WI, USA).

Conservative domains were detected using BLAST (https://www.ncbi.nlm. nih.gov/Structure/cdd/wrpsb.cgi) and MITOS WebServer (http://mitos.bioinf. uni-leipzig.de/index.py) (Bernt et al. 2013). Maps of the mitogenomes were constructed using CGView (https://cgview.ca/) (Stothard and Wishart 2005). MEGA X was used for base composition analysis, relative synonymous codon usage (RSCU) analysis, pairwise relative genetic distance (p-distance) calculation, as well as non-synonymous (Ka) and synonymous substitutions (Ks) analysis (Fay and Wu 2003; Kumar et al. 2016). Composition skew values were calculated using the following formulas: "AT-skew = (A – T) / (A + T) GCskew = (G – C) / (G + C)" (Perna and Kocher 1995).

#### **Phylogenetic analysis**

Phylogenetic analysis was conducted using the sequences of 13 PCGs and two rRNA genes from the complete mitogenomes of 105 species, including seven species from this study (Suppl. material 1). *Channa andrao* and *Hyphessobrycon sweglesi* were selected as outgroups, while the remaining specimens belonged to the Cichlidae family. Phylogenetic analysis was conducted using maximum likelihood (ML) and Bayesian inference (BI) methods with PhyloSuite v. 1.2.3 software package (Zhang et al. 2020; Xiang et al. 2023). All genes were aligned using MAFFT v. 7.313, and the best-fit substitution model and partitioning scheme were determined using ModelFinder. ML phylogenies were inferred using IQ-TREE with the Edge-linked partition model for 5000 ultrafast bootstraps (Minh et al. 2013; Nguyen et al. 2015). BI phylogenies were inferred using MrBayes v. 3.2.7a with a partition model (Ronquist et al. 2012). The analysis consisted of two parallel runs with 2,000,000 generations each, and the initial 25% of sampled data was discarded as burn-in. The trees were visualized and edited using iTOL v. 6 (Letunic and Bork 2021).

# **Results and discussion**

## Genome organization and composition

Seven complete mitogenomes covering two genera were obtained. L. kungweensis (16,587 bp), L. meleagris (16,582 bp), L. ornatipinnis (16,585 bp), N. brevis (16,586 bp), N. caudopunctatus (16,586 bp), and N. similis (16,580 bp) had similar lengths, while N. leleupi had the shortest length at 16,562 bp (Fig. 1) (accession numbers: OP805601.1, OP805600.1, OQ076695.1, OP930818.1, OP930816.1, OP930817.1, and OP930815.1). The seven mitogenomes possessed the typical gene composition found in most bony fish, including 13 PCGs, 22 tRNAs, two rRNAs, and a CR. Among these genes, 12 PCGs, 14 tRNA genes, and two rRNA genes, were located on the major strand (H-strand), while the remaining eight tRNA genes and a PCG were encoded on the minor strand (L-strand). The gene order of these mitogenomes was identical to that of previously published species L. signatus (MZ427900.1) and Neolamprologus brichardi (AP006014.1) (Nam et al. 2021). Seventeen intergenic regions of the same length were observed between the mitochondrial regions of species L. kungweensis, L. meleagris, L. ornatipinnis, N. brevis, with lengths ranging from 10 bp (between atp8 and atp6) to 35 bp (between trnN and trnC). However, N. caudopunctatus exhibited a 24 bp intergenic region between trnV and rrnL, and N. similis displayed a 38 bp intergenic region at the same location. The trnC and trnY of N. similis overlapped by 1 bp, whereas there was no overlap in this region in the other six species. In addition, N. leleupi had one more intergenic region (24 bp between cox1 and trnS2) than other species (Table 1).

## **Nucleotide composition**

The nucleotide composition of the seven newly sequenced *Lamprologus* and *Neolamprologus* mitogenomes were biased toward A and T (Table 2). The AT-skews exhibited inconspicuously positive values, while all GC-skews were mark-edly negative. The analysis revealed a clear preference for the utilization of C, along with a minor inclination towards A, across the entire genome (Table 2).

To determine the nucleotide composition of Cichlidae, the A + T content, ATskew, G + C content, and GC-skew of 103 complete mitogenomes (including 8 subfamilies Astronotinae, Cichlasomatinae, Cichlinae, Etroplinae, Geophaginae, Pseudocrenilabrinae, Ptychochrominae, and Retroculinae of the family Cichlidae) Jiachen Wang et al.: Comparison and phylogenetic insights of mitogenomes of cichlids



Figure 1. The gene maps of the seven newly sequenced mitogenomes. Different gene types are shown in different colors.

were calculated. The H-strand in the mitogenomes of 103 cichlid species showed a similar preference for A and T nucleotides. The 103 Cichlidae mitogenomes had a comparable nucleotide composition, A + T content (52.1 ~ 58.8%) were higher than the G + C content (41.1 ~ 47.8%) (Fig. 2). The GC-skew were negative ( $-0.351 \sim -0.221$ ), indicating a higher occurrence of C than G except for *Andinoacara rivulatus* (-0.019), *Pelvicachromis pulcher* (-0.005), and *Etroplus canarensis* (-0.002). The AT-skew were inconspicuously positive ( $0.002 \sim 0.076$ ), indicating a small difference in the content of A and T in the mitogenomes. This phenomenon is also observed in other published Teleostei genomes (Liu et al. 2020; Ruan et al. 2020; Xu et al. 2021a, 2021b; Wang et al. 2023). The A nucleotide composition is commonly used to indicate gene direction and replication orientation during transcription and replication (Wei et al. 2010a, 2010b).

#### **Protein-coding genes**

In the seven newly sequenced mitogenomes, PCG *nad6* was on the L-strand, while other PCGs were on the H-strand. The average A + T content of the PCGs ranged from 53.0% (*N. leleupi* and *N. brevis*) to 54.7% (*L. meleagris*). Six of them had the same 13 PCGs length of 11,466 bp, while the remaining species, *N. leleupi*, had a slightly shorter length of 11,421bp. The reason for this difference was that the *cox1* gene in *N. leleupi* had a mutation causing a premature stop codon compared to other species, resulting in a reduction of 45 base pairs in length (Tables 1, 2).

Most of the PCGs in the seven newly sequenced mitogenomes began with the start codon ATG, except for *cox1*, which started with GTG. Most PCGs terminated with the codon TAA or incomplete codon (TA- / T--), with the exception of *nad1*, which ended with TAG (Table 2). The cichlid species are relatively conservative in their use of start codons, and their preferences are generally consistent with those of the seven newly sequenced species with the only exception of the occurrence of a rare start codon ATC in the *cox1* and *nad3*. All the Cichlids share the stop codons with TAA, TAG, AGA, and incomplete codons (TA- / T--) (Fig. 3).

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|  |   |                       |                               |                       |                       | G/ TAG/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAG/TAG/ | G/ TAG/TAG/TAG/<br>TAG/TAA/TAG/TAG/                   | G/ TAG/TAG/TAG/<br>TAG/TAG/TAG/TAG/                          | G/ TAG/TAG/TAG/<br>TAG/TAG/TAG/TAG/   | G/ TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/<br>TA/TA/TA/   | G/ TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/                                | G/ TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/  | G/ TAG/TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/<br>TA/TA/TA/   | G/ TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/  | G/ TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/   | G/ TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/<br>TA/TA/TA/<br>TA/TAA/TAA/<br>TAA/TAA/   | G/ TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TAA/TA/TA/TA/<br>G/ TAA/TAA/TAA/TAA/   | G/ TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TA/TA/TA/TA/<br>TA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/   | G/ TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAA/TAA/TAG/<br>G/ TA/TA/TA/TA/TA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/  | G/ TAG/TAG/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/  | G/ TAG/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TA/TA/TA/TA/TA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>TAA/TAA/TAA/TAA/TAA/<br>TAA/TAA/   | G/ TAG/TAG/TAG/TAG/<br>TaG/TAA/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>TAG/TAG/TAG/<br>G/ TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/<br>TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/<br>TAA/TAA/TAA/TAA/<br>TAA/TAA/TA  | G/ TAG/TAG/TAG/TAG/<br>TaG/TAA/TAG/TAG/<br>Tag/TaA/TAG/TaG/<br>G/ TAA/TAA/TAG/TAG/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/TAA/<br>G/ TAA/TAA/TAA/TAA/TAA/TAA/<br>CA/TAA/TAA/TAA/TAA/TAA/TAA/TAA/<br>CA/TAA/TAA/TAA/TAA/TAA/TAA/TAA/<br>CA/TAA/TAA/TAA/TAA/TAA/TAA/TAA/TAA/TAA/T   | GI TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAA/TAG/TAG/<br>TAG/TAA/TAA/TAG/<br>GI TAA/TAA/TAA/TAA/<br>GI TAA/TAA/TAA/TAA/  |
|  |   |                       |                               |                       |                       | G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>TAG/ATG/ATG/      | G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>TAG/<br>TAG/        | G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG                | ATG/ATG/ATG/ TAG/<br>ATG/ATG/ATG/ TAG/  | G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG<br>6/ATG/ATG/<br>ATG/ATG/ATG/<br>TA/1<br>ATG/ATG/ATG/   | G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG<br>6/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/                           | 6/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/  | G/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/AT  | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/  | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG  | G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>G/ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>ATG/ATG/ATG/<br>S/GTG/GTG/CTG/<br>TAA/<br>GTG/GTG/GTG/   | ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/   | ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/   | S/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG   | S/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG   | 6/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG  | S/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG  | 6/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG   | 6/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG   |
| 0/0  | 0/0                                       | 0/0.                  |                               | 4/22/38               | 4/22/38               | 4/ 2// 38<br>0/0 ATG/ATC<br>0/0 ATG/I               | 4/ 2//38<br>0/0<br>0/0 ATG/ATC<br>3/3 3/3             | 4/ 2.// 38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>3/3<br>3/3 /1/-1 | 4/ 2// 38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>3/3<br>/-1/-1<br>/-1/-1  | 4/ 2// 38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>3/3<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/ | 4/ 2.// 38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>ATG//<br>3/3<br>/-1/-1<br>0/0 ATG/ATG<br>0/0 ATG/ATG                          | 4/ 2// 38<br>0/0 ATG/ATG<br>0/0 ATG/A<br>ATG/A<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>/ | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1<br>/-1/-1 ATG/ATG<br>0/0 ATG/ATG<br>1/1 1<br>1/1 1<br>1/1   | 4/ 2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>3/3<br>3/3<br>/-1/-1 ATG/ATC<br>0/0 ATG/ATC<br>0/0 ATG/ATC<br>1/1 1<br>1/1 5<br>5/35/35   | 4/ 2//38<br>0/0 ATG/ATG<br>0/0 ATG/A<br>ATG/A<br>/-1/-1<br>/-1/-1<br>/-1/-1<br>0/0 ATG/ATC<br>0/0<br>1/1<br>1/1<br>1/1<br>0/-1<br>5/35/35<br>0/-1  | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/1/-1<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG                         | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 ATG/ATC<br>/-1/-1 ATG/ATC<br>0/0 ATG/ATC<br>1/1 11<br>1/1 5/35/35 ATG//<br>1/1 GTG/GTC<br>0/-1 GTG/GTC<br>0/-1 GTG/GTC   | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>3/3 ATG/I<br>/-1/-1 ATG/ATC<br>/-1/-1 ATG/ATC<br>0/0 ATG/ATC<br>/-1/-1 CG/GTC<br>5/35/35 0/-1<br>1/1 GTG/CTC<br>0/-1 GTG/CTC<br>2/10 3/3 3/3  | 4/2//38 0/0 ATG/ATG ATG/A ATG/  | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>0/0 ATG/ATG<br>3/3 ATG/  | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 ATG/A<br>/-1/-1 ATG/A<br>0/0 ATG/A<br>1/1 ATG/A<br>0/0 ATG/A<br>0/0 ATG/A<br>0/0 ATG/ATG<br>1/1 GTG/GTC<br>0/-1 GTG/GTC<br>0/-1 GTG/GTC<br>0/-1 GTG/GTC<br>0/-1 ATG/ATG<br>1/1 ATG/ATG   | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 ATG/ATG<br>/-1/-1 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 CG1G/GTC<br>1/1 GTG/GTC<br>0/-1 GTG/GTC<br>24/0 24/0 ATG/ATG<br>1/1 ATG/ATC<br>1/1 ATG/ATC<br>1/1 ATG/ATC<br>10/-10/-10 ATG/ATG   | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 ATG/A<br>/-1/-1 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 CGG/GTG<br>1/1 GTG/GTC<br>2/1/1 GTG/GTC<br>2/1/1 GTG/GTC<br>2/1/1 ATG/ATC<br>1/1 ATG/ATC<br>1/1 ATG/ATC<br>1/1 ATG/ATC<br>1/1-1 ATG/ATC<br>1/1-1 ATG/ATC<br>1/-1/-1 ATG/ATC<br>1/-1/-1 ATG/ATC   | 4/2//38<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>3/3 AG/A<br>3/3 AG/A<br>/-1/-1 ATG/ATG<br>0/0 ATG/ATG<br>/-1/-1 ATG/ATG<br>5/35/35 ATG/A<br>1/1 GTG/C<br>2/0 ATG/ATG<br>0/0 ATG/ATG<br>1/1 GTG/C<br>2/1/1 GTG/C<br>2/1/1 ATG/ATG<br>1/1 ATG/ATG<br>1/1 ATG/ATG<br>1/1 ATG/ATG<br>1/1 ATG/ATG<br>0/0 ATG/ATG<br>1/1 ATG/ATG<br>0/0 ATG/ATG<br>1/1 ATG/ATG<br>0/0 ATG/ATG<br>0/0 ATG/ATG<br>1/1 ATG/ATG<br>0/0 ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/   |
| 0/0/0/0/0/0/0  |   | 0/0/0/0/0/0           | 22/22/22/22/24/22/            |                       | 0/0/0/0/0/0           | 0/0/0/0/0/0   | 6/6/0/0/0/0/0<br>0/0/0/0/0/0                          | 0/0/0/0/0/0/0<br>0/0/0/0/0/0<br>3/3/3/3/3/3/                 | 0/0/0/0/0/0/0<br>0/0/0/0/0/0/0<br>1-/1-/1-/1-/1-/1-/1-/1-/1-/1-/1-/1-/1-/1  | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | 0/0/0/0/0/0/0/0<br>0/0/0/0/0/0/0/0<br>5/8/8/8/8/8/8/8/8/8/8/8/8/8/8/8/8/8/8/8   | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/   | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/   | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/   | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/   | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | 0/0/0/0/0/0<br>0/0/0/0/0/0/0/0/0/0/0<br>3/3/3/3/3/3/3/3/3/3/3/3/3/3/3/3/3/3/3/   | 0/0/0/0/0/0<br>0/0/0/0/0/0/0/0/0/0/0/0/0/0   | 0/0/0/0/0/0<br>0/0/0/0/0/0/0/0/0/0/0/0/0/0  | 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  |
| 59/69/69/69<br>43/943/944/946<br>79/72/77/77   | 621621616                                 | 1.1.1.1               | 9/1671/1670/16 2<br>1652      | ALIALIALIA            | 4/ / 4/ / 4/ / 4      | 75/975/975/975                                      | -4/ 1-4/ 1-4/ 1-4/ 1-4/ 1-4/ 1-4/ 1-4/ 1              | -4/-4/-4/-4<br>75/975/975/975<br>0/70/70/70                  | -4/-4//4//4/<br>75/975/975/975<br>0/70/70/70<br>1/71/71/71<br>39/69/69/69   | 4/14/14/14<br>75/975/975/975<br>0/70/70/70<br>1/71/71/71<br>1/71/71/71<br>6/1046/1046/10<br>1046  | 4/4/14/14<br>75/975/975/975<br>0/70/70/70<br>1/71/71/71<br>1/71/71/71<br>6/1046/1046/10<br>1046                           | -4/-4//4//4<br>75/975/975/975<br>0/70/70/70/70<br>1/71/71/71<br>6/1046/1046/10<br>6/1046/1046/10<br>1046<br>2/72/72/72<br>59/69/69/69   | 4/14/14/14<br>75/975/975/975<br>0/70/70/70/70<br>1/71/71/71<br>9/69/69/69<br>6/1046/1046/10<br>1046<br>1046<br>1046<br>1046<br>1046<br>1046<br>1046<br>10   | -4/-4//4//4<br>75/975/975/975<br>0/70/70/70<br>1/71/71/7<br>1/71/71/7<br>6/1046/1046/10<br>1046<br>2/72/72/72<br>9/69/69/69<br>33/73/73/33<br>33/73/73/73<br>33/73/73/73                          | 75/975/975/975<br>75/975/975/975<br>1/71/71/71<br>1/71/71/71<br>6/1046/1046/10<br>6/1046/1046/10<br>1046<br>2/72/72/72<br>9/69/69/69<br>3/73/73/73<br>56/65/666<br>3   | 4/14/14/14<br>75/975/975/975<br>9/69/69/69/69<br>6/1046/1046/10<br>1046<br>5/1046/1046/10<br>1046<br>72/72/72<br>99/69/69<br>99/69/69<br>99/69/69<br>6/1596/15<br>1596<br>1596   | 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| 4/14/14/14   55/975/975/975/975   75/975/975/975/975   71/71/71/7   9/69/69/69   6/1046/10   1046   72/72/72/72   8/69/69/69   3/72/72/72   8/69/69/69   73/73/73/73   9/69/15   171/71/71   171/71/71   8/1596/159   19/69/159/1596/15   19/69/169116911   91/691169116911   88/168/168/168/168/168   84/784/784/784   | 4/14/14/14   55/975/975/975/975   75/975/975/975/975   71/71/71   1/71/71/71   6/1046/10   6/1046/10   6/1046/10   1046   72/72/72/72   8/65/66/66   3/73/73/73   73/73/73/73   8/1596/1596/15   19/691/691   19/1691/691   8/168/168/168   8/168/168/168   8/78/73/73/73   8/78/73/73/73   9/1691/691/691   8/78/78/784   8/168/168/168   8/78/784/784   8/78/782/72772  |
| 69/69/69/69/69/69/<br>944/943/944/943/94<br>72/72/72/72/72/                          | 12/12/12/12/                              |                       | 1669/1670/1669/167<br>68/1652 | 74/74/74/74/          | 075/075/075/075/07    | 16/016/016/016/016                                  | 10L/0L/0L/0L/0L                                       | 16/07/07/07/07/07/07/07/07/07/07/07/07/07/                   | 16/07/07/07/07/07/07/07/07/07/07/07/07/07/  | 10/07/07/07/07/07/07/07/07/07/07/07/07/07   | 72/07/07/07/07/07/07/07/07/07/07/07/07/07/  | 72/07/07/07/07/07/07/07/07/07/07/07/07/07/  | 70/70/70/70/70/70/70/70/70/70/70/70/70/7  | 70/70/70/70/70/70/70/70/70/70/70/70/70/7  | 70/70/70/70/70/70/70/70/70/70/70/70/70/7   | 70/70/70/70/70/70/70/70/70/70/70/70/70/7   | 70/07/07/07/07/07/07/07/07/07/07/07/07/0   | 70/70/70/70/70/70/70/70/70/70/70/70/70/7   | 70/70/70/70/70/70/70/70/70/70/70/70/70/7  | 70/70/70/70/70/70/70/70/70/70/70/70/70/7  | 70/70/70/70/70/70/70/70/70/70/70/70/70/7   | 70/70/70/70/70/70/70/70/70/70/70/70/70/7   | 70/70/70/70/70/70/70/70/70/70/70/70/70/7  | 70/70/70/70/70/70/70/70/70/70/70/70/70/7  |
| 6<br>13/1015 944/9<br>35/1087 7<br>75/2777 1669/                                     | 35/1087 7<br>75/2777 1669/                | 75/2777 1669/         |                               | 49/2851 7             | 0/32/0                | 6/C/6 070C/H7                                       | 21/3899 77  | 57/3969 77   | evic.ve 0.200.042<br>77 3899<br>77 557/3969<br>76<br>557/3969<br>66<br>66<br>66<br>66<br>66<br>66<br>66<br>66<br>66 | 6%/27     02000 (42)       77     397/3969     7       7     7     66       35/4037     6     6       31/5083     1046/     1046/   | 7/3899     7       37/3899     7       57/3969     7       35/4037     6       31/5083     1046/       33/5155     7      | 8/L01/2     8/L01/2     8/L01/2       7     899     7       7     7     8/L01/2       8     1/L01/2     6       8     1/L01/2     6       8     1/L01/2     6       8     1/L01/2     1046/       7     23/L02     6       23/L02     23/L02     6  | F(L)     F(L) <th< td=""><td>24,5220     24,5220     7       37/3899     7       35/4037     6       35/4037     6       35/4033     1046/       31/5083     1046/       37/5255     6       37/5299     7       38/5400     6</td><td>SV/2020     SV/3999     T       37/3899     7     7       35/4037     6     7       35/4037     1046/     7       53/5155     7     7       53/5155     6     7       53/525     6     7       37/5299     6     38/5400       38/5469     7     7</td><td>24,3220 24,3220   37/3899 7   35/4037 6   35/4037 6   31/5083 1046/   31/5083 1046/   31/5299 7   38/5409 6   38/5409 6   38/5469 7   38/5469 1596/</td><td>247,302.0 247,302.0   37/3899 7   35/4037 6   35/4037 1046/   33/5155 7   33/5125 7   33/5225 6   37/5299 7   38/5469 7   20/7066 1596/   15/137 7</td><td>24,5220 24,5220   37/3899 7   37/3899 7   35/4037 1046/   31/5083 1046/   31/5259 6   37/5299 7   38/5400 6   38/5400 6   38/5400 6   38/5400 6   30/5299 7   31/5299 7   31/5299 7   31/5291 7   31/5293 15/513   31/5213 7</td><td>24/3020 24/3020 24/3020   37/3969 7   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    35/4037     1046/     7       53/5155     7     7       53/5155     6     7       53/525     6     7       37/5299     6     38/5400       38/5469     7     7 | 24,3220 24,3220   37/3899 7   35/4037 6   35/4037 6   31/5083 1046/   31/5083 1046/   31/5299 7   38/5409 6   38/5409 6   38/5469 7   38/5469 1596/  | 247,302.0 247,302.0   37/3899 7   35/4037 6   35/4037 1046/   33/5155 7   33/5125 7   33/5225 6   37/5299 7   38/5469 7   20/7066 1596/   15/137 7   | 24,5220 24,5220   37/3899 7   37/3899 7   35/4037 1046/   31/5083 1046/   31/5259 6   37/5299 7   38/5400 6   38/5400 6   38/5400 6   38/5400 6   30/5299 7   31/5299 7   31/5299 7   31/5291 7   31/5293 15/513   31/5213 7   | 24/3020 24/3020 24/3020   37/3969 7   35/4037 1046/   31/5083 1046/   31/5299 7   37/5299 6   38/5469 7   38/5469 1596/   30/7066 1596/   37/7009 691/6   | 24,5220 24,5220   37/3899 7   35/4037 6   35/4037 1046/   35/5155 7   53/5155 7   53/5155 7   35/525 6   37/5299 7   38/5469 7   38/5469 1596/   38/5400 6   38/7400 6   38/7400 6   38/7403 1596/   38/7403 691/6   37/7903 691/6  | 24,3220 24,3220   37/3899 7   35/4037 6   35/4037 1046/   53/5155 7   53/5155 7   53/5155 7   53/5155 7   53/5155 6   53/5155 7   53/5155 6   53/5155 6   53/5155 7   53/5155 6   53/5155 6   58/5469 1596/   7 7   58/5469 1596/   7 7   58/5469 1596/   7 7   7 7   7 7   7 7   51/7213 691/6   51/7983 7   30/8152 168/1  | 24/5020 24/5020   37/3899 7   35/4037 1046/   53/5155 7   53/5155 7   53/5155 6   37/5299 6   38/5400 6   38/5400 6   37/5299 7   38/5400 6   37/5299 6   37/7909 691/6   31/7909 691/6   31/7909 691/6   30/8152 168/1   30/8152 168/1  | 24/3020 24/3020   37/3899 7   37/3899 7   35/4037 1046/   53/5155 7   53/5155 7   53/5259 6   37/5299 7   38/5400 6   38/5400 6   38/5400 6   37/5299 7   38/5400 6   38/5400 6   38/5400 6   38/5400 6   38/5400 6   38/5400 6   31/7909 6   31/7909 6   30/8152 16   31/7909 6   31/7909 6   31/7909 6   31/7909 6   31/7909 6   31/7909 6   31/7909 6   31/7909 6   31/7909 6  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| /69/69/69/69/69<br>13/1012/1012/1013/101<br>35/1084/1085/108<br>76/2777/278/2775/277 | 35/1084/1085/108<br>76/2777/2778/2775/277 | 76/2777/2778/2775/277 |                               | 50/2851/2852/2849/285 | 25/3826/3827/3824/382 |   | 38/3899/3900/3892/386                                 | 38/3899/3900/3897/389<br>58/3969/3970/3967/396               | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>36/4037/4038/4035/403  | 38/3899/3900/3897/388<br>58/3969/3970/3967/396<br>36/4037/4038/4035/403<br>32/5083/5084/5081/508  | 38/3899/3900/3897/385<br>58/3969/3970/3967/395<br>36/4037/4038/4035/403<br>32/5083/5084/5081/505<br>54/5155/5156/5153/515 | 38/3899/3900/3897/388<br>58/3969/3970/3967/396<br>36/4037/4038/4035/405<br>32/5083/5084/5081/508<br>54/5155/5156/5153/515<br>24/5225/5226/5223/522  | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>36/4037/4038/4035/405<br>32/5083/5084/5081/508<br>54/5155/5156/5153/515<br>24/5225/5226/5223/522<br>38/529/5300/529/522/525  | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>36/4037/4038/4035/40<br>32/5083/5084/5081/50<br>54/5155/5156/5153/515<br>24/5225/5226/5223/522<br>39/5200/5299/5300/5297/525                     | 38/3899/3900/3897/388<br>58/3969/3970/3967/398<br>58/4037/4038/4035/402<br>32/5083/5084/5081/508<br>54/51 55/5156/51 53/515<br>54/51 55/5156/51 53/515<br>24/5225/5226/5223/52<br>39/5400/5470/5398/546<br>39/5470/5470/5398/546                   | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>36/4037/4038/4035/405<br>32/5083/5084/5081/508<br>54/5155/5156/5153/52<br>24/5225/5226/5223/522<br>38/5299/5300/5297/52<br>39/5470/5400/5308/546<br>59/5470/5468/546  | 38/3899/3900/3897/385<br>58/3969/3970/395/403<br>56/4037/4038/4035/403<br>54/5155/5156/5153/515<br>54/5155/5156/5153/51<br>54/5155/5156/5153/51<br>54/5299/5300/5297/525<br>39/5400/5400/5398/54(<br>56/7057/7067/7020/705<br>56/7138/7115/713   | 38/3899/3900/3897/38<br>58/3069/3970/3967/396<br>58/4037/4038/4035/403<br>32/5083/5084/5081/506<br>54/5155/5156/5153/515<br>54/5155/5156/5153/52<br>54/5155/5156/5153/52<br>54/5155/5156/5153/52<br>59/5400/5398/54<br>59/5400/5398/54<br>56/7067/7067/7020/705<br>56/7067/7020/715<br>37/7138/7138/7115/715   | 38/3899/3900/3897/38<br>58/3969/3970/3967/395<br>58/4037/4038/4035/405<br>32/5083/5084/5081/505<br>54/5155/5156/5153/515<br>24/525/525/525/523/52<br>98/5299/5300/5297/525<br>99/5400/5400/5398/546<br>59/5470/5470/5468/546<br>59/7067/7067/7020/706<br>56/7067/7020/708<br>13/7138/7138/713/715<br>13/7214/7214/71910/7887/790  | 88/3899/3900/3897/388<br>58/3969/3970/3967/398<br>36/4037/4038/4035/403<br>54/5155/5156/5153/5153/515<br>54/525/525/526/523/522<br>39/5400/529/5300/5297/522<br>39/5400/5400/5398/546<br>56/7057/706777067/7020/702<br>37/7138/7115/715<br>37/7138/7105777020/702<br>33/7984/7984/79847795  | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>58/4037/4038/4035/403<br>54/5155/5156/5153/515<br>54/5155/5156/5153/51<br>54/5155/5156/5153/52<br>39/5400/5400/5299/522<br>39/5400/5400/5298/54<br>39/5470/5470/5388/54<br>56/7057/7067/7020/70<br>56/7057/7067/7020/70<br>56/7057/7067/7067/7020/70<br>56/70577067/706777020/70<br>56/705770677706777020/70<br>56/705770677706777020770<br>57/713877138771157715<br>33779847798477961779<br>52/8153/8153/8153/8130/815   | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>58/4037/4038/4035/403<br>54/5155/5156/5153/5153/51<br>54/5155/5156/5153/5153/51<br>54/5155/5156/5153/52<br>99/5400/5400/5297/525<br>99/5400/5400/5398/54(<br>56/7057/7067/7020/705<br>59/5470/5470/5468/54<br>59/5470/5470/548/7115/715<br>13/7138/71138/7115/715<br>37/7138/7138/7115/715<br>59/7910/7910/7910/7887/796<br>33/7984/7984/7961/795<br>52/8153/8153/8130/882<br>52/8153/8153/8130/885   | 38/3899/3900/3897/38<br>58/3969/3970/3967/396<br>58/4037/4038/4035/403<br>54/5155/5156/5155/5155/5155/51<br>54/5155/5156/5155/5155/51<br>54/5155/5156/5153/52<br>99/5400/529/5300/5297/525<br>99/5400/5400/5398/54<br>56/7067/7067/706<br>77/138/7115/715<br>59/5470/5470/5468/54<br>59/5770/5470/548/7159<br>59/7910/7910/7910/7887779<br>09/7910/7910/7913/8130/815<br>52/8153/8153/8130/815<br>52/8827/8827/8827/8804/86<br>52/8827/8827/8827/8804/86  | 38/3899/3900/3897/388<br>58/3969/3970/3967/398<br>58/4037/4038/4035/403<br>54/5155/5156/5153/515<br>54/5155/5156/5153/52<br>54/5155/5156/5153/52<br>54/5155/5156/5153/52<br>54/5299/5300/5398/54<br>54/5299/5300/5398/54<br>54/7067/7057/706<br>55/7067/7067/702<br>52/7138/7138/7138/713<br>37/7138/7138/7138/7138<br>52/8153/8153/8130/815<br>26/8827/8827/9610/9587/966<br>31/9682/96822/96559/966   |
| 69/69/69/<br>1013/1012/1013/1<br>1085/1084/1085/1<br>2776/2776/2                     | 1085/1084/1085/1<br>2776/2776/2776/2      | 2776/2776/2776/2      |                               | 2850/2850/2850/2      | 3825/3825/3825/3      | 3898/3898/3898/3                                    |   | 3968/3968/3968/3   | 3968/3968/3968/3<br>4036/4036/4036/4  | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5  | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5082/5<br>5154/5154/5154/5   | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5154/5<br>5224/5224/5224/5224/5   | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5224/5<br>5298/5298/5298/5  | 3968/3968/3968/3<br>4036/4036/4036/4036/4<br>5082/5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5<br>5298/5298/5298/5<br>5399/5399/5  | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5224/5<br>5298/5298/5399/5<br>5399/5399/5399/5<br>5469/5469/5469/5469/5  | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/<br>52298/5298/5<br>5399/5399/5<br>5469/5469/5469/5<br>7066/7066/7066/7  | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5224/<br>5298/5298/5298/5<br>5399/5399/5<br>5469/5469/5<br>7137/7137/7   | 3968/3968/3968/3968/3<br>4036/4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5224/5<br>5299/5399/5<br>5399/5399/5<br>5399/5399/5<br>7137/7137/7<br>7137/7137/7  | 3968/3968/3968/3<br>3968/3968/3968/3<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5224/5224/5<br>5299/5399/5<br>5399/5399/5<br>5399/5399/5<br>7066/7066/7066/7<br>7137/7137/7<br>7213/7213/7213/77  | 3968/3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5<br>5224/5224/5224/5224/<br>5239/5399/5<br>5469/5469/5469/5<br>5469/5469/5469/<br>7066/7066/7066/7<br>7213/7213/7<br>7909/7909/7909/7903/7<br>7983/7983/7983/7   | 3968/3968/3968/3968/3<br>4036/4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5224/5224/5224/5<br>5298/5298/5298/5<br>5399/5399/5<br>5469/5469/5469/5<br>7066/7066/7066/7<br>7066/7066/7066/7<br>7083/7983/7983/7<br>8152/8152/8152/8152/8  | 3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5224/5224/5224/5<br>5224/5224/5224/5224/  | 3968/3968/3968/3968/3<br>4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5224/5224/5224/5<br>5224/5224/5224/5224/  | 3968/3968/3968/3<br>4036/4036/4036/4036/4<br>5082/5082/5082/5<br>5154/5154/5<br>5224/5224/5224/5224/5<br>5299/5399/5<br>5399/5399/5<br>5399/5399/5<br>5399/5399/  |
| 014/1016   | 14/1016                                   | 2011/00               | 1 I 70                        | 76/2778               | 850/2852              |   | 128/3830  | 397/3899   | .28/3830<br>197/3899<br>167/3969  | 28/3830<br>197/3899<br>167/3969<br>136/4038   | 28/3830<br>197/3969<br>167/3969<br>136/4038<br>182/5084   | 28/3830<br>197/3899<br>167/3969<br>136/4038<br>136/4038<br>182/5084<br>55/5157  | 28/3830<br>197/3969<br>167/3969<br>136/4038<br>136/4038<br>135/5157<br>55/5157  | 28/3830<br>97/3969<br>167/3969<br>136/4038<br>136/4038<br>55/5157<br>255/5227<br>133/5335   | 28/3830<br>97/3969<br>67/3969<br>136/4038<br>136/4038<br>135/5157<br>55/5157<br>155/5157<br>133/5335<br>133/5335<br>199/5400   | 28/3830<br>97/3969<br>167/3969<br>136/4038<br>182/5084<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>53/5335<br>899/5400<br>899/5471   | 28/3830<br>97/3969<br>167/3969<br>136/4038<br>182/5084<br>55/5157<br>255/5227<br>255/5227<br>255/5227<br>233/5335<br>133/5335<br>133/5335<br>170/5471  | 28/3830<br>97/3969<br>(67/3969<br>(67/3969<br>(82/5084<br>55/5157<br>55/5157<br>(82/5084<br>(82/5084<br>(82/5084<br>(82/5084<br>(82/5084)<br>(15/5167)<br>(170/5471<br>(19/7141)<br>(19/7141)  | 28/3830<br>97/7219<br>67/3969<br>67/3969<br>136/4038<br>136/4038<br>55/5157<br>55/5157<br>55/5157<br>133/5335<br>133/5335<br>133/5335<br>197/7219<br>197/7219   | 28/3830<br>97/3899<br>67/3969<br>136/4038<br>182/5084<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>55/5157<br>133/5335<br>133/5335<br>133/5335<br>137/7141<br>19/7141<br>19/77219<br>97/7219  | 28/3830<br>97/3899<br>67/3969<br>136/4038<br>182/5084<br>55/5157<br>55/5157<br>25/5227<br>25/5227<br>133/5335<br>133/5335<br>133/5335<br>133/5335<br>133/5335<br>137/219<br>19/71219<br>97/7219<br>97/7219<br>888/7910   | 28/3830<br>97/3899<br>67/3969<br>186/4038<br>182/5084<br>55/5157<br>55/5157<br>25/5227<br>133/5335<br>133/5335<br>133/5335<br>199/5400<br>170/5471<br>19/7141<br>19/7141<br>19/71219<br>97/7219<br>97/7219<br>19/7219<br>21/8143   | 28/3830<br>97/3899<br>67/3969<br>67/3969<br>55/526<br>55/527<br>25/5227<br>25/5227<br>25/5227<br>25/5227<br>19/7471<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>21/8143<br>888/7910<br>63/7985<br>63/7985<br>838/7910   | 28/3830<br>97/3899<br>67/3969<br>67/3969<br>55/5227<br>55/5227<br>25/5227<br>133/5335<br>133/5335<br>199/5471<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/7141<br>19/ 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 | /3830/3831/3826<br>/3899/3900/3895<br>/3969/3970/3965<br>/4038/4038/4038<br>/5084/5085/508;<br>/5084/5085/508;<br>/5158/5158/515<br>/5227/528/523<br>/5335/5336/533;<br>/5472/5472/547/  | /3830/3831/3826<br>/3899/3900/38956<br>/3999/3970/3965<br>/4038/4039/4036<br>/5084/5085/5082<br>/5157/5158/5155<br>/5227/5228/5225<br>/5235/5336/533<br>/5472/5472/5472<br>/5472/5472/04<br>/7142/7111   | /3830/3831/3826<br>/3899/3900/3890/<br>/3969/3970/3965<br>/4038/4039/4036<br>/5084/5085/5082<br>/5157/5158/5156<br>/5157/5228/5228<br>/527/5228/5228<br>/5355/5336/533<br>//5472/5472/5477<br>/5472/5472/5477<br>/7142/7119<br>//7142/7119  | /3830/3831/3826<br>/3899/3900/38395<br>/3999/3970/3965<br>/4038/4039/4036<br>/5084/5085/5085<br>/5157/5158/5158<br>/5157/5158/5158<br>/5335/5336/5335<br>/5472/5471<br>/5472/5472/5471<br>/5472/5472/5471<br>/7142/7115<br>/7142/71158  | /3830/3831/3826<br>/3899/3900/3890/<br>/3999/3970/3965<br>/4038/4039/4036<br>/5084/5085/5085<br>/5157/5158/5155<br>/5157/5158/5155<br>/5335/5336/5335<br>/5401/5397<br>/5401/5397<br>/5472/5472/5470<br>/5472/5472/5470<br>/5472/7427115<br>/7911/7911/788   | /3830/3831/3826<br>/3899/3900/3896;<br>/3969/3970/3965<br>/5084/5085/5085<br>/5084/5085/5085<br>/5084/5085/5085<br>/5335/5336/5335<br>/5335/5335/5335<br>/5335/5335  | /3830/3831/3825<br>/3899/3900/3895/<br>/3096/3970/3965<br>/5084/5085/5085<br>/5084/5085/5085<br>/5084/5085/5085<br>/5084/5085/5085<br>/515/5158/5155<br>/5227/5225/5335<br>/5335/5335/5335<br>/5227/5226/704<br>/7142/7111<br>/7142/71115<br>/7142/71115<br>/7142/71115<br>/7142/71115<br>/7142/71115<br>/7142/71115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7142/7115<br>/7220/7220/715<br>/7142/7115<br>/7220/7220/715<br>/7142/7115<br>/7220/7220/715<br>/7220/7220/715<br>/7142/7115<br>/71175<br>/7220/7220/715<br>/7220/7220/715<br>/71175<br>/7220/7220/715<br>/7220/7220/715<br>/71175<br>/7220/7220/715<br>/71175<br>/7220/7220/715<br>/71175<br>/7220/7220/715<br>/71175<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/715<br>/71175<br>/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7220/715<br>/7220/7720/7220/715<br>/7220/7720/715<br>/7220/7720/7220/775<br>/7220/7720/7220/775<br>/7220/7720/7220/775<br>/7220/775<br>/7220/775<br>/7220/775<br>/7220/775<br>/7220/775<br>/7220/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/775<br>/7720/7750<br>/7720/775<br>/7720/7 | /3830/3831/3826<br>/3899/3900/3890/<br>/3999/3900/3895<br>/4038/4039/4036<br>/5084/5085/5085<br>/5084/5085/5085<br>/5157/5158/5155<br>/527/528/5235<br>/5335/5335/5335/5335<br>/5472/5175<br>/5472/5472/5477<br>/5472/5472/5477<br>/7142/71170<br>/7142/71170<br>/7142/71196<br>/7142/71198<br>/7986/7986/7986/7965<br>/7986/7986/7986/7965<br>/7986/7986/7986/7965<br>/7986/7986/7986/7965   |
| 70/70  |   | 1014/1013/1014/       | 1108/1107/1108/               | 17772/7772/7772       | 2851/2851/2851/       |   | 3829/3829/3829/                                       | 3829/3829/3829/<br>3898/3898/3898/                           | 3829/3829/3829/<br>3898/3898/3898/<br>3968/3968/3968/   | 3829/3829/3829/<br>3898/3898/3898/3898/<br>3968/3968/3968/<br>4037/4037/4037/   | 3829/3829/3829/<br>3898/3898/3898/<br>3968/3968/3968/<br>4037/4037/4037/  | 3829/3829/3829/<br>3898/3898/3898/3968/<br>3968/3968/3968/<br>4037/4037/4037/<br>5083/5083/5083/  | 3829/3829/3829/<br>3898/3898/3968/<br>3968/3968/3968/<br>4037/4037/4037/4037/<br>5083/5083/5083/<br>5156/5156/5156/<br>5226/5226/5226/  | 3829/3829/3829/<br>3898/3898/3968/<br>3968/3968/3968/<br>4037/4037/4037/<br>5083/5083/5083/<br>5156/5156/5156/<br>5226/5226/<br>5334/5334/5334/   | 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| цч   |   | >                     | Γ                             | nL2                   | ad1                   | 2   |   | nl<br>nQ   | nl<br>nQ<br>Mu  | rnl<br>rnQ<br>iad2  | mi<br>mQ<br>aad2<br>mW  | mn<br>mm<br>mad2<br>mw<br>mad2<br>mad2<br>mad2  | mi<br>mA<br>mM<br>mM<br>mad2<br>mA<br>mad2<br>mad2<br>mad2<br>mad2<br>mad2<br>mad2<br>mad2<br>mad2  | ml<br>ma<br>mm<br>ma<br>ad2<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn<br>mn   | nnl<br>mR<br>nnM<br>nad2<br>nnW<br>nnM<br>nnC<br>nnC<br>nnC  | ml m   | ml mind mnd mnd mnd mnd mnd mnd mnd mnd mnd m  | rnl<br>rnd<br>ad2<br>ad2<br>ad2<br>rnW<br>rnA<br>rnA<br>rnA<br>rnS2<br>sox1<br>rnS2  | nd<br>mn<br>ad2<br>ad2<br>ad2<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn<br>nn   | rnl<br>rnR<br>ad2<br>ad2<br>rnW<br>rnW<br>rnN<br>rnN<br>rnS<br>cox1<br>rnS<br>cox2<br>rnS<br>rnS  | trnl<br>trnl<br>had2<br>had2<br>trnW<br>trnM<br>trnA<br>trnN<br>trnS<br>cox1<br>trnS<br>cox2<br>trnS<br>cox2<br>trnK<br>trnK   | trnl<br>trnl<br>had2<br>had2<br>trnW<br>trnW<br>trnN<br>trnN<br>trnS2<br>sox1<br>trnS2<br>sox2<br>trnS2<br>sox2<br>trnS2<br>sox2<br>trnS2<br>sox2<br>trnS2<br>trnS2<br>trnS2<br>trnS2<br>trnS2<br>trnM   | rtnl<br>TrnQ<br>had2<br>had2<br>trnW<br>trnW<br>trnN<br>trnS2<br>sox1<br>trnS2<br>sox2<br>sox2<br>atp8<br>atp8  | trnl<br>trnl<br>had2<br>had2<br>had2<br>had2<br>trnW<br>trnW<br>trnN<br>trnS2<br>trnS2<br>trnS2<br>trnS2<br>trnS2<br>trnS2<br>trnS2<br>trnB<br>trnS2<br>trnB<br>trnB<br>trnB<br>trnB<br>trnS2<br>trnB<br>trnS2<br>trnA<br>trnA<br>trnA<br>trnA<br>trnA<br>trnA<br>trnA<br>trnA  |

| OfferForTot  |       | Pos                                       | ion   |   | a bita a la cina de la c | Cod                             | u                               | 2      |
|--|-------|---|---|---|--------------------------|---------------------------------|---------------------------------|--------|
| mm     1003/10031/10031/10031/1002/100301     1009/1009/1003/100301     000/00/00     Mickards       mm     1000/1010/10101/1001/10071/10078/1003     1009/1009/10036/1039/1039/1039/1039/1039/1039/1039/1039  | auao  | From                                      | P   | (da) azıs                                   | Intergenic nucleotides   | Start                           | Stop                            | orrand |
| madd     1000/100/100/100/100/100/100/100     000/00/00     ATG/ATG/ATG       madd     1000/100/100/100/100/100/100/100     177/1177/1177/1172/1177/1172/1177/1172/1170/1177/1170/1177/1170/1177/1170/1177/1170/1177/1172/1177/1172/1177/1177  | trnR  | 10031/10031/10031/10032/10032/10009/10031 | 10099/10099/10099/10100/10100/10077/10099       | 69/69/69/69/69/69/69                        | 0/0/0/0/0/0/0            |                                 |                                 | т      |
| mad     10390/10390/10391/10361/10361/10361     11770/11771/1171/11748/1176     1381/1381/1381/1381/1381/1381/1381/1381  | nad4l | 10100/10100/10100/10101/10101/10078/10100 | 10396/10396/10396/10397/10397/10396             | 297/297/297/297/297/297                     | 0/0/0/0/0/0/0            | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG | ТАА/ТАА/ТАА/ТАА/<br>ТАА/ТАА/ТАА | т      |
| tml11771/11771/1172/1172/1172/11749/1177111839/11839/11839/11830/113317373737373737373737373737373737373   | nad4  | 10390/10390/10390/10391/10391/10368/10390 | 11770/11770/11770/11770/11771/11748/11770       | 1381/1381/1381/1381/1381/13<br>81/1381      | L-/L-/L-/L-/L-/L-        | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG | Τ/Τ/Τ/Τ/Τ/Τ                     | т      |
| tmll11840/11840/11841/1181/11841/11841/11841/1184111966/11906/11907/11807/11907/11884/1196467/66/67/67/670/0/0/0/0/073/73/73/73/73/73/73/73/73tmll11911/11912/11912/1182/11889/119111983/11983/11983/11983/11984/11984/1191211983/11983/11983/11984/119120/0/0/0/0/073/73/73/73/73/73/73/73/73/73/73/73/73/7   | trnH  | 11771/11771/11772/11772/11772/11749/11771 | 11839/11839/11839/11840/11840/11817/11839       | 69/69/69/69/69/69/69                        | 0/0/0/0/0/0/0            |                                 |                                 | т      |
| tml111911/11910/11911/1192/1198/1198/1198/1198/1198/119  | trnS1 | 11840/11840/11840/11841/11841/11818/11840 | 11906/11905/11906/11907/11907/11884/11906       | 67/66/67/67/67/67/67                        | 0/0/0/0/0/0              |                                 |                                 | т      |
| nad5     11984/11983/1198/11965/11962/11984     13822/13822/13822/13823/1380/1382     1839/1839/1839/1839/1839/1839/1839/183/1363/163/163     AG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/A   | trnL1 | 11911/11910/11911/11912/11912/11889/11911 | 11983/11982/11983/11984/11984/11961/11983       | 73/73/73/73/73/73/73/73                     | 4/4/4/4/4/4              |                                 |                                 | т      |
| nad6     13819/13818/13820/13820/13820/13820/13820/13820/13819     14340/14361/1431/1431/1431/1431     522/522/522/522/522/522/522/522/522/522   | nad5  | 11984/11983/11984/11985/11985/11962/11984 | 13822/13821/13822/13823/13823/13800/13822       | 1839/1839/1839/1839/1839/18<br>39/1839      | 0/0/0/0/0/0/0            | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG | ТАА/ТАА/ТАА/ТАА/<br>ТАА/ТАА/ТАА | т      |
| tml     14341/14340/14341/14342/14319/14341     14409/14400/14410/14310/1430/1430     69/69/69/69/69/69/69/69     0/0/0/0/0/0     0/0/0/0/0/0     Display       cybb     14414/1141/1141/1141/1141/1141/1141/11  | nad6  | 13819/13818/13819/13820/13820/13797/13819 | 14340/14339/14340/14341/14341/14318/14340       | 522/522/522/522/522/522/522                 | -4/-4/-4/-4/-4/-4        | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG | ТАА/ТАА/ТАА/ТАА/<br>ТАА/ТАА/ТАА |        |
| cyth     14414/1413/1414/1415/14392/1414     15564/155371555/1555/1555/1555/1555/1555/1552/1555     1141/1141/1141/1141/11     4/4/4/4/4     ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/ATG/  | trnE  | 14341/14340/14341/14342/14342/14319/14341 | 14409/14408/14409/14410/14410/14410/14387/14409 | 69/69/69/69/69/69/69                        | 0/0/0/0/0/0/0            |                                 |                                 | _      |
| tmT     15555/15554/15556/15556/15556/15625/15525/15626/15625/15626/15627/15626/15627/15626/15627/15626/15627/15626/15627/15626/15627/15626/15627/15626/15627/15626/15627/15626/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15696/15697/15698/15697/15698/15697/15698/15697/15696/15697/15696/15697/15696/15697/15698/15698/15688/15698/15698/15698/156988/188978889889889898989898898988/1899/100 | cytb  | 14414/14413/14414/14415/14415/14414       | 15554/15553/15554/15555/15555/15532/15554       | 1141/1141/1141/1141/1141/1141/11<br>41/1141 | 4/4/4/4/4/4/4            | ATG/ATG/ATG/ATG/<br>ATG/ATG/ATG | Τ/Τ/Τ/Τ/Τ/Τ                     | т      |
| tmp     15627/15626/15627/15628/15637/15628/15635/15637/15635/15636/15637/15636/15637/15636/15637/15636/15637/15636/15637/15636/15637/15636/15637/15636/15637/15636/15637/15636/15637/15636/15586/15636/15586/15586/15586/15586/15580/15636/15580/15636/15580/15636/15586/15580/1558 | trnT  | 15555/15555/15555/15556/15556/15533/15555 | 15626/15625/15626/15627/15627/15604/15626       | 72/72/72/72/72/72                           | 0/0/0/0/0/0/0            |                                 |                                 | т      |
| CR 15697/15696/15697/15698/15697/15674/15697 16587/16582/16586/16586/16586/16586/16586/16586/16580 891/887/889/890/889/890/889/889/890/884 0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/0/  | trnP  | 15627/15626/15627/15628/15628/15605/15627 | 15696/15695/15696/15697/15696/15673/15696       | 70/70/70/20/69/69/20                        | 0/0/0/0/0/0/0            |                                 |                                 | _      |
|  | CR    | 15697/15696/15697/15698/15697/15674/15697 | 16587/16582/16585/16586/16586/16562/16580       | 891/887/889/889/890/889/884                 | 0/0/0/0/0/0/0            |                                 |                                 |        |



**Figure 2.** A + T content vs AT-skew and G + C content vs GC-skew in the 103 mitogenomes of family Cichlidae. Values are calculated on H-strands for full-length mitogenomes.

| Table 2. Base | compositions | of the | complete | genomes, | PCGs, | rRNAs, | tRNAs, | and | CRs | of the | seven | newly | sequence | ed: |
|---------------|--------------|--------|----------|----------|-------|--------|--------|-----|-----|--------|-------|-------|----------|-----|
| mitogenomes.  |              |        |          |          |       |        |        |     |     |        |       |       |          |     |

|                                  | Whole g | genome |       |        | PC     | Gs   | tR    | lAs  | rRi   | NAs  | CR   |      |
|----------------------------------|---------|--------|-------|--------|--------|------|-------|------|-------|------|------|------|
| Species                          | Size    | AT     | AT -  | GC -   | Size   | AT   | Size  | AT   | Size  | AT   | Size | AT   |
|                                  | (bp)    | (%)    | SKEW  | SKEW   | (bp)   | (%)  | (bp)  | (%)  | (bp)  | (%)  | (bp) | (%)  |
| Lamprologus<br>kungweensis       | 16,587  | 54.1   | 0.002 | -0.300 | 11,466 | 53.5 | 1,554 | 54.7 | 2,613 | 54.1 | 891  | 62.9 |
| Lamprologus<br>meleagris         | 16,582  | 55.1   | 0.002 | -0.300 | 11,466 | 54.7 | 1,553 | 55.8 | 2,613 | 54.1 | 887  | 63.6 |
| Lamprologus<br>ornatipinnis      | 16,585  | 53.9   | 0.006 | -0.304 | 11,466 | 53.2 | 1,554 | 55.4 | 2,613 | 53.5 | 889  | 62.7 |
| Neolamprologus<br>brevis         | 16,586  | 53.6   | 0.011 | -0.311 | 11,466 | 53.0 | 1,554 | 54.9 | 2,614 | 53.0 | 889  | 63.8 |
| Neolamprologus<br>caudopunctatus | 16,586  | 53.9   | 0.002 | -0.299 | 11,466 | 53.2 | 1,552 | 55.4 | 2,613 | 53.2 | 890  | 63.0 |
| Neolamprologus<br>Ieleupi        | 16,562  | 53.7   | 0.017 | -0.318 | 11,421 | 53.0 | 1,553 | 54.7 | 2,612 | 53.0 | 889  | 62.5 |
| Neolamprologus<br>similis        | 16,580  | 54.1   | 0.010 | -0.311 | 11,466 | 53.5 | 1,554 | 54.9 | 2,598 | 53.9 | 884  | 63.0 |





RSCU was calculated to identify the predominant synonymous codon (Grantham et al. 1980). The comparative analysis based on RSCU of all PCG codons showed that the codon usage patterns of these seven species were similar (Fig. 4). Genes encoding IIe and Leu2 had high frequency, while those encoding Cys, Met, and Ser1 were infrequent.



#### **Evolutionary analyses**

The selection pressure was analyzed by calculating the ratio of Ka/Ks across *Lamprologus* and *Neolamprologus* for each aligned PCG (Fig. 5) (Yang and Nielsen 2002). It was found that *atp8* showed the largest Ka/Ks value among the 13 PCGs, which suggested more amino acid variety in the biomolecule. This suggests that the atp8 gene might have evolved faster than other PCGs due to slight selection pressure (Hassanin et al. 2005). The faster evolution of the atp8 gene could result in greater amino acid diversity, indicating its potential as an effective marker for population classification. The Ka/Ks values for all PCGs were lower than 1, suggesting that purifying selection was likely the main driver of mitochondrial PCG evolution (Hurst et al. 2002).



**Figure 5.** Ka/Ks values for the 13 PCGs. Pale pink box plots, five species of gnus *Neolamprologus*; orange box plots, four species of *Lamprologus*; blue box plots, nine species of *Lamprologus* and *Neolamprologus*. The band inside the box represents the median; upper and lower hinges correspond to the 25<sup>th</sup> and 75<sup>th</sup> percentiles; circles, to outliers.

Besides the Ka/Ks analysis, an assessment of the degree of divergence in *Lamprologus* and *Neolamprologus* was conducted by analyzing the overall p-distance between nucleotides of 13 PCGs + two rRNA genes (Fig. 6). The results of p-distance indicated that the fastest-evolving gene was *nad6*, which was inconsistent with the results of Ka/Ks value. However, the difference in this gene might be not comparable with the selection since this force is acting in a contemporary period.

#### Ribosomal RNA genes, transfer RNA genes, and control regions

The size of the *rrnS* genes were between 943 bp (*L. meleagris*, *N. brevis*, and *N. caudopunctatus*) and 946 bp (*N. similis*), while the size of the *rrnL* genes in seven species ranged between 1,652 bp (*N. similis*) to 1,671 bp (*N. brevis*) (Table 1). The two rRNA genes located between *trnF* and *trnL2*, with *trnV* separating them. The A + T content of rRNAs ranged from 53.0% ~ 54.1% (Table 2).

The sizes of the tRNA genes ranged from 66 bp (*trnY* of *N. caudopunctatus*) to 74 bp (*trnK*). The combined length of the 22 tRNA genes varied between 1,552 bp (*N. caudopunctatus*) and 1,554 bp (*L. kungweensis*, *L. ornatipinnis*, and *N. similis*). The A + T contents of tRNA genes ranged from 54.7% to 55.8% among the seven species analyzed in this study (Table 2).

As with other fish mitogenomes, the CRs were discovered to exist between *trnF* and *trnP* in all seven species. The sizes of the CRs ranged from 884 bp (*N. similis*) to 891 bp (*L. kungweensis*). The A + T contents of PCGs, tRNAs, and rRNAs sequences were found to be similar to that of the entire mitogenomes, whereas CR sequences had a higher A + T content (62.5% ~ 63.8%) (Table 2).

#### **Phylogenetic analysis**

To elucidate the phylogenetic inter-relationships within the family Cichlidae and genera *Lamprologus* and *Neolamprologus*, concatenated nucleotide sequences of 13 PCGs + two rRNAs from 103 cichlid species were obtained. Additionally,



**Figure 6.** Genetic p-distances for nucleotide sequences for 13 PCGs and 2 rRNAs. Pale pink box plots, five species of *Neolamprologus*; orange box plots, four species of *Lamprologus*; blue box plots, 9 species of genera *Lamprologus* and *Neolamprologus*.

*Channa andrao*, and *Hyphessobrycon sweglesi* from two other families were used as outgroups. It was found that BI and ML analysis generated the same topology structure on most nodes (Fig. 7).

Specifically, the seven complete mitogenomes covered two genera in this study have good clustering in phylogenetic trees, and within the family Cichlidae, the subfamily Etroplinae and Ptychochrominae were monophyletic across analyses. They diverged with species in other subfamilies early in the evolutionary history of cichlid fishes. This result was similar to a previous molecular phylogenetic study (Astudillo-Clavijo et al. 2022). Thirty-one species from subfamily Astronotinae, Cichlasomatinae, Cichlinae, Geophaginae, and Retroculinae were clustered into one branch, indicating these five subfamilies were closely related. Moreover, 67 Pseudocrenilabrinae species also formed a monophyletic clade. Pseudocrenilabrinae tribes and their interrelationships were for the most part well supported as reported by Astudillo-Clavijo et al. (2022). Due to the addition of seven newly sequenced mitogenomes, three pairs of sisters (N. brichardi + N. leleupi, N. caudopunctatus + L. meleagris, and L. signatus + L. kungweensis) were newly identified, as shown in Fig. 7. Lamprologus meleagris did not cluster with Lamprologus species, but with species from the genus Neolamprologus. Previous studies have identified such taxonomic issues in the genera Lamprologus and Neolamprologus (Schelly et al. 2006; Sturmbauer et al. 2010). Sturmbauer et al. (2010) think a viable way might be to re-assign the genus name Lamprologus to most Neolamprologus species. Our results also support this scenario. However, the species from the genera Lamprologus and Neolamprologus used in this study were limited, making it impossible to perform a more detailed analysis. Therefore, to better understand the relationships between members of these two genera, it will be beneficial to include more species in future studies.

In conclusion, our study increased the database of mitogenome in Cichlidae, and showed that mitogenome sequences are efficient molecular markers for studying the phylogenetic relationships within Cichlidae. However, there is a lack of analyses in nuclear genes. In the future study, we will further improve these deficiencies.



**Figure 7.** 13 PCGs-based phylogenetic tree of 103 cichlid species and two outgroups. Numbers at nodes represent the posterior probability and bootstrap values for BI and ML analysis, respectively. "-" indicates this clade not supported by BI or ML analysis.

## **Additional information**

#### **Conflict of interest**

The authors have declared that no competing interests exist.

#### **Ethical statement**

No ethical statement was reported.

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#### Author contributions

Conceptualization: WZ, HL. Data curation: JW. Formal analysis: JT, JW. Funding acquisition: WZ, JW, HL. Methodology: HL. Project administration: HL, WZ. Resources: WZ, HL, KH. Software: JT, JW. Supervision: HL, KH. Validation: HL. Visualization: JW, JT. Writing - original draft: JW.

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## **Data availability**

All of the data that support the findings of this study are available in the main text or Supplementary Information.

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## **Supplementary material 1**

#### Summary of the mitochondrial genomes used for phylogenetic analysis

Authors: Jiachen Wang, Jingzhe Tai, Wenwen Zhang, Ke He, Hong Lan, Hongyi Liu Data type: xlsx

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