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Characterization of a New mtDNA Minicircle in a Chewing Louse, Geomydoecus aurei

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Campbell, Ashley L.; Bolic, Dino; Demastes, James W.; and Spradling, Theresa A., "Characterization of a New mtDNA Minicircle in a Chewing Louse, Geomydoecus aurei" (2020). Research in the Capitol. 3. https://scholarworks.uni.edu/rcapitol/2020/all/3

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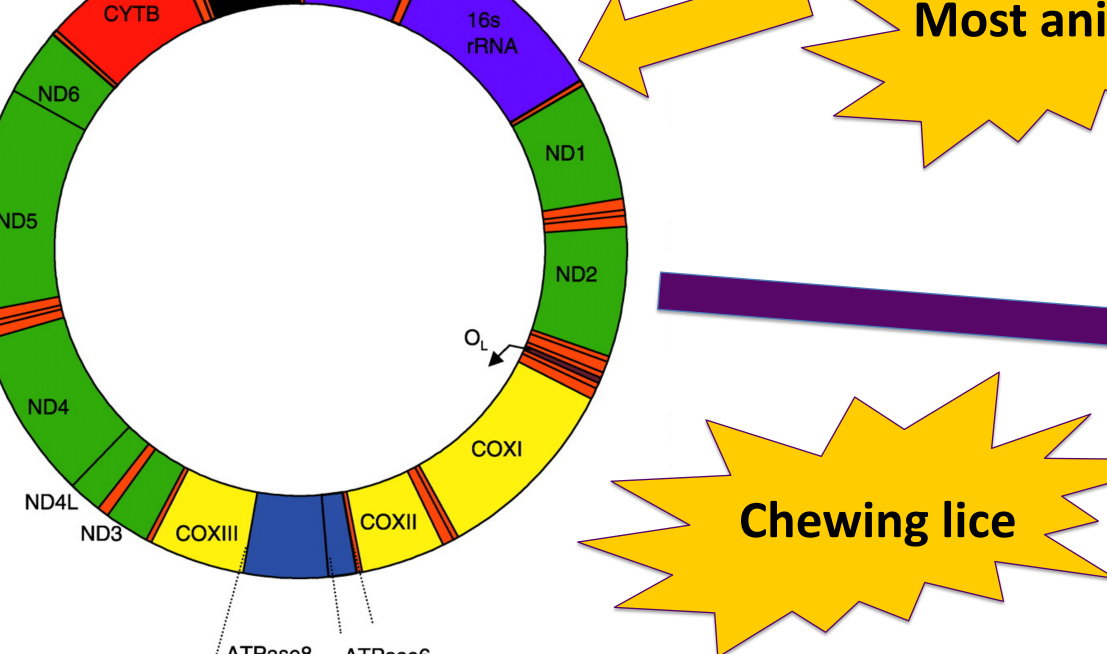
Ashley L. Campbell, Dino Bolic, James W. Demastes, and Theresa A. Spradling Department of Biology, University of Northern Iowa, Cedar Falls



Background

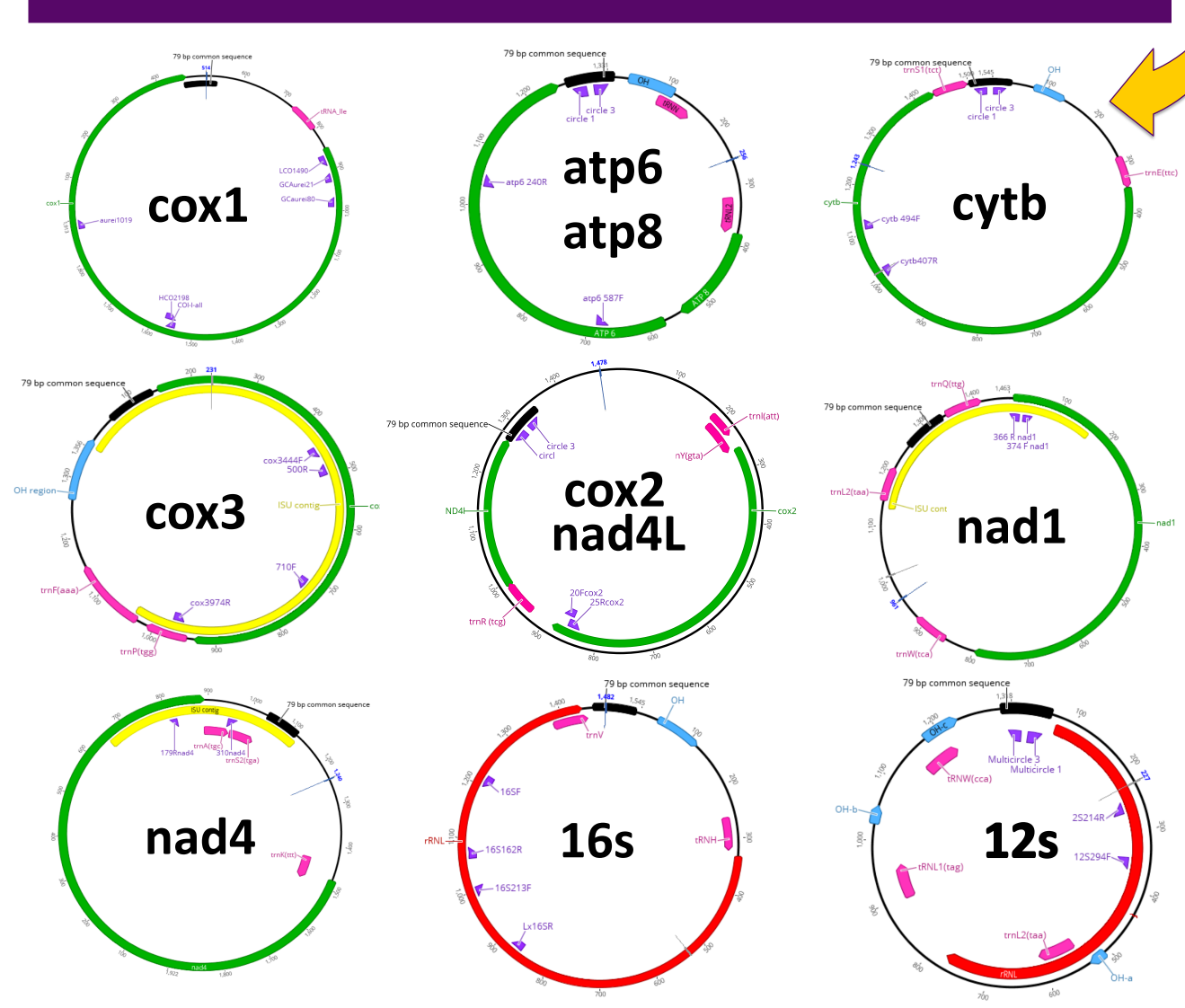
Nad2 Minicircle of Geomydoecus aurei

Typical Animal mtDNA Genome 128 rRNA 168 rRNA ND1 Most animals



- In most animals, the mitochondrial genome is very conserved with 13 protein coding genes, 22 tRNAs, and two rRNAs in a single circular chromosome (Cameron et al., 2011).
- Many species of lice are unique in the sense that they contain multiple minicircles of fragmented mtDNA instead of a single circular chromosome.

Characterized Minicircles in *G. aurei*



- trnS2

 had 2

 trnN

 trnN

 trnC

 trnC

 trnC
- The newly characterized *nad2* minicircle contains the nad2 gene and three putative tRNAs: *trnN*, *trnS2*, and *trnC*.
- The *nad2* gene is 1077 bp in length.
- The *nad2* minicircle contains the conserved 79-bp region, shown above in black, that has been identified in all *G. aurei* minicircles characterized to date.
- The *nad2* minicircle was identified and characterized using PCR, cloning reactions, and Sanger sequencing.

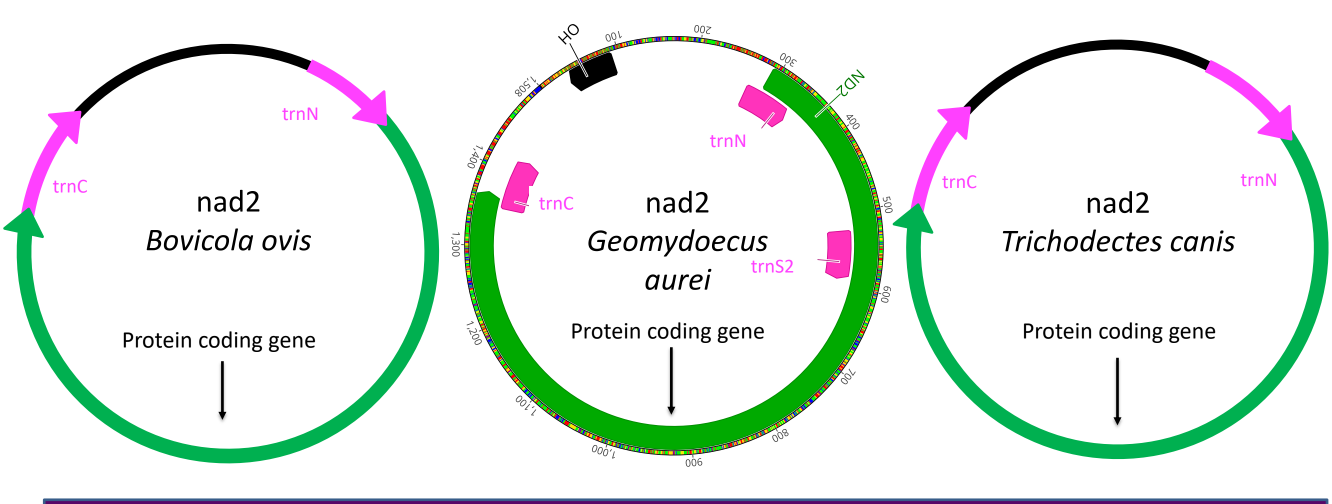
Literature

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Conclusions

Results

- The 79-bp DNA sequence found on this circle is identical to sequence found on nine other *G. aurei* minichromosomes, suggesting functional importance.
- The majority of the tRNAs identified in minicircles containing protein coding genes have been found nearly adjacent to the coding region or to very minimally overlap with the coding sequence.
- The gene order of the *G. aurei nad*2 minichromosome (*trnN-nad2-trnC*) is seen in other mammalian louse species (Song et al, 2018), including *B. ovis*, and *T. canis* (shown below).
- The position of *trnS2* within the interior of the *nad2* gene is unusual and requires further investigation. Another DNA sequence that is a likely *tRNS2* gene is found on the *G. aurei* nad4 minichromosome in a position also seen in other louse species. *tRNS2* has not been associated with *nad2* in other lice.



Next Steps

- Two protein coding genes, *nad3* and *nad5* have been identified and are in the process of being characterized and mapped.
- There are four tRNAs (trnD, trnG, trnM, and trnT) expected to be found upon the mapping of the remaining minicircles.
- Discovery of the remaining gene, nad6, has been problematic in this and other studies of many insects.
- Additional PCR, cloning reactions, and Sanger sequencing will be performed in an effort to identify *nad6* and map the mitochondrial genome of *G. aurei* to completion.

Acknowledgements

- Dr. Alan Orr Research Award
- NSF Award (DEB-1445708)
- UNI Intercollegiate Academics Travel Fund