

# Large-scale comparative genomics unravels great genomic diversity across the *Rickettsia* and *Ca. Megaira* genera and identifies Torix group as an evolutionarily distinct clade.

Helen R Davison<sup>1</sup>, Jack Pilgrim<sup>1</sup>, Nicky Wybouw<sup>2</sup>, Joseph Parker<sup>3</sup>, Stacy Pirro<sup>4</sup>, Simon Hunter-Barnett<sup>1</sup>, Paul M Campbell<sup>5</sup>, Frances Blow<sup>6</sup>, Alistair C Darby<sup>1</sup>, Gregory D D Hurst<sup>1</sup> and Stefanos Siozios<sup>1\*</sup>

## Affiliations

1. Department of Evolution, Ecology and Behaviour, Institute of Infection, Veterinary and Ecological sciences, University of Liverpool, Liverpool, L69 7ZB, United Kingdom

2. Terrestrial Ecology Unit, Department of Biology, Faculty of Sciences, Ghent University, Ghent, Belgium

3. Division of Biology and Biological Engineering, California Institute of Technology, 1200 E California Boulevard, Pasadena, CA 91125, USA

4. Iridian Genomes, Bethesda, MD, USA

5. School of Health and Life Sciences, Faculty of Biology Medicine and Health, the University of Manchester, Manchester, United Kingdom

6. Center for Genomics and Systems Biology, Department of Biology, New York University, New York, New York, USA

\* Corresponding author

## Abstract

*Rickettsia* are intracellular bacteria originally described as arthropod borne pathogens that are emerging as a diverse group of often biologically important, non-pathogenic symbionts of invertebrates and microeukaryotes. However, sparse genomic resources for symbiotic strains and for the sister genus (*Candidatus* Megaira) inhibit our understanding of *Rickettsia* evolution and biology. Here, we present the first closed genomes of *Ca. Megaira* from an alga (*Mesostigma viride*), and Torix *Rickettsia* from midge (*Culicoides impunctatus*) and bed bug (*Cimex lectularius*) hosts. Additionally, we sequenced and constructed draft genomes for *Ca. Megaira* from another alga (*Carteria cerasiformis*), Transitional group *Rickettsia* from tsetse fly (*Glossina morsitans submorsitans*), and Torix *Rickettsia* from a spider mite (*Bryobia graminum*). We further extract 22 draft genomes from arthropod genome sequencing projects, including 1 *Adalia*, 4 Transitional, 1 Spotted Fever, 7 Torix, 7 Belli and the first Rhyzobius and Meloidae *Rickettsia* group genomes. We used new and existing *Rickettsia* genomes to estimate the phylogeny and metabolic potential across groups and reveal transitions in genomic properties. These data reveal Torix as unique amongst currently described *Rickettsia*, with highly distinct and diverse accessory genomes. We confirm the presence of a third subclade of Torix, previously only known from gene marker sequences. Further, Torix share an intact pentose phosphate pathway with *Ca. Megaira*, not observed in other *Rickettsia*. Considering the distinctness and diversity of Torix, we propose that the group be named *Candidatus* Tisiphia. The wide host range of *Ca. Tisiphia* symbionts necessitates onward research to understand the biological and physiological bases of *Ca. Tisiphia*-host interactions.

## 42 Importance statement

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44 Members of the genus *Rickettsia* were originally identified as causative agents of mammalian vector-borne

45 disease. In the last 25 years we have recognised that many *Rickettsia* are arthropod symbionts, and sit

46 alongside a sister taxon, *Ca. Megaira*, which are symbiotic associates of microeukaryotes. The lack of

47 genomic information for symbiotic strains affects our ability to determine the evolutionary relationships

48 between strains and understand the biological underpinnings of the different symbioses. We clarify these

49 relationships by assembling 26 genomes of *Rickettsia* from understudied groups, and the first two *Ca.*

50 *Megaira*, from various insects and microeukaryotes. Of note, the accessory genome diversity and broad host

51 range of *Torix Rickettsia* parallels all other *Rickettsia* combined. This diversity, alongside the breadth of host

52 species, make the *Torix* clade an important hidden player in invertebrate biology and physiology. We argue

53 this clade should be given its own genus status, for which we propose *Ca. Tisiphia*.

## Introduction

Symbiotic bacteria are vital to the function of most living eukaryotes, including microeukaryotes, fungi, plants, and animals (Boettcher et al., 1996; Clay et al., 2005; Douglas, 2011; Fujishima & Kodama, 2012). The symbioses formed are often functionally important to the host with effects ranging from mutualistic to detrimental. Mutualistic symbionts may provide benefits through the biosynthesis of metabolites, or by protecting their hosts against pathogens and parasitoids (Hendry et al., 2014; Oliver et al., 2010). Meanwhile parasitic symbionts can be detrimental to the host due to resource exploitation or through reproductive manipulation that favours its own transmission over the host's (Engelstädter & Hurst, 2009; Leclair et al., 2017). Across these different symbiotic relationships, symbionts are often important determinants of host ecology and evolution.

The Rickettsiales (Alphaproteobacteria) represent an order of obligate intracellular bacteria that form symbioses with a variety of eukaryotes (Weinert et al., 2015). Within Rickettsiales, the family Rickettsiaceae represent a diverse collection of bacteria that infect a wide range of eukaryotic hosts and can act as symbionts, parasites, and pathogens. Perhaps the best-known clade of Rickettsiaceae is the genus *Rickettsia*, which was initially described as the cause of spotted fever and other rickettsioses in vertebrates that are transmitted by ticks, lice, fleas and mites (Angelakis & Raoult, 2017).

*Rickettsia* have been increasingly recognised as heritable arthropod symbionts. Since the first description of a maternally inherited male-killer in ladybirds (Werren et al., 1994), we now know that heritable *Rickettsia* are common in arthropods (Pilgrim et al., 2021; Weinert et al., 2009). Further, *Rickettsia*-host symbioses are diverse, with the symbiont capable of reproductive manipulation, nutritional and protective symbiosis, as well as influencing thermotolerance and pesticide susceptibility (Bodnar et al., 2018; Brumin et al., 2011; Chiel et al., 2009; Giorgini et al., 2010; Hurst et al., 1994; Kontsedalov et al., 2008; Łukasik et al., 2013).

Our understanding of the evolution and diversity of the genus *Rickettsia* and its allies has increased in recent years. Weinert et al. (2009) defined 13 different groups of *Rickettsia* with two early branching clades that appeared genetically distant from other members of the genus. The first of these was defined from a symbiont of *Hydra* and was named the Hydra group *Rickettsia*, which has since been assigned its own genus status, *Candidatus Megaira* (Schrallhammer et al., 2013). *Ca. Megaira* forms a sister clade to *Rickettsia* and is common in ciliate protists, amoebae, chlorophyte and streptophyte algae, and cnidarians (Lanzoni et al., 2019). Members of this clade are found in

hosts from aquatic, marine and soil habitats which include model organisms (e.g., *Paramecium*, *Volvox*) and economically important vertebrate parasites (e.g., *Ichthyophthirius multifiliis*, the ciliate that causes white spot disease in fish) (Lanzoni et al., 2019). Whilst symbioses between *Ca. Megaira* and microeukaryotes are pervasive, there is no complete publicly available genome and the impact of these symbioses on the host are poorly understood.

A second early branching clade was first described from *Torix tagoi* leeches and is commonly coined *Torix Rickettsia* (Kikuchi & Fukatsu, 2005). Symbionts in the *Torix* clade have since been found in a wide range of invertebrate hosts from midges to freshwater snails, and in a fish-parasitic amoeba (Pilgrim et al., 2021). The documented diversity of hosts is wider than other *Rickettsia* groups, which are to date only found in arthropods and their associated vertebrate or plant hosts (Weinert et al., 2009). *Torix* clade *Rickettsia* are known to be heritable symbionts, but their impact on host biology is poorly understood, despite the economic and medical importance of several hosts (inc. bed bugs, black flies, and biting midges). Rare studies have described the potential effects on the host, which include: larger body size in leeches (Kikuchi & Fukatsu, 2005); a small negative effect on growth rate and reproduction in bed bugs (Thongprem et al., 2020); and an association with parthenogenesis in *Empoasca* Leafhoppers (Aguin-Pombo et al., 2021).

Current data seems to suggest an emerging macroevolutionary scenario where the members of *Rickettsia*-*Megaira* clade originated as symbionts of microeukaryotes, before diversifying to infect invertebrate symbionts. The *Torix Rickettsia* retained a broad range of hosts from microeukaryotes to arthropods. The remaining members of the genus *Rickettsia* evolved to be arthropod heritable symbionts and vector-borne pathogens. However, a lack of genomic and functional information for symbiotic clades limits our understanding of evolutionary transitions within *Rickettsia* and its sister groups. No *Ca. Megaira* genome sequences are currently publicly available and of the 165 *Rickettsia* genome assemblies available on the NCBI (as of 29/04/21), only two derive from the *Torix* clade and these are both draft genomes. In addition, dedicated heritable symbiont clades of *Rickettsia*, such as the *Rhyzobius* group, have no available genomic data, and there is a single representative for the *Adalia* clade. Despite the likelihood that heritable symbiosis with microeukaryotes and invertebrates was the ancestral state for this group of intracellular bacteria, available genomic resources are heavily skewed towards pathogens of vertebrates.

In this study we establish a richer base of genomic information for heritable symbiont *Rickettsia* and *Ca. Megaira*, then use these resources to clarify the evolution of these groups. We broaden

116 available genomic data through a combination of targeted sequencing of strains without complete  
 117 genomes, and metagenomic assembly of *Rickettsia* strains from arthropod genome projects. We  
 118 report the first closed circular genome of a *Ca. Megaira* symbiont from a streptophyte alga  
 119 (*Mesostigma viride*) and provide a draft genome for a second *Ca. Megaira* from a chlorophyte  
 120 (*Carteria cerasiformis*). In addition, we present the first complete genomes of two Torix *Rickettsia*  
 121 from a midge (*Culicoides impunctatus*) and a bed bug (*Cimex lectularius*) as well as a draft genome  
 122 for *Rickettsia* from a tsetse fly (*Glossina morsitans submorsitans*, an important vector species), and  
 123 a new strain from a spider mite (*Bryobia graminum*). A metagenomic approach established a  
 124 further 22 draft genomes for insect symbiotic strains, including the first Rhyzobius and Meloidae  
 125 group draft genomes. We utilize these to carry out pangenomic, phylogenomic and metabolic  
 126 analyses of our extracted genome assemblies, with comparisons to existing *Rickettsia*.

127

## Methods

### Genomic data collection and construction

We employed two different workflows to assemble genomes for *Ca. Megaira* and *Rickettsia* symbionts (Figure 1). A) Targeted sequencing and assembly of focal *Ca. Megaira* and *Torix Rickettsia*. B) Assembly from SRA deposits of *Ca. Megaira* from *Mesostigma viride* NIES296 and the 29 arthropods identified in Pilgrim et al (2021) that potentially harbour *Rickettsia*. These were analysed alongside previously assembled genomes from the genus *Rickettsia*, and the outgroup taxon *Orientia tsutsugamushi*.

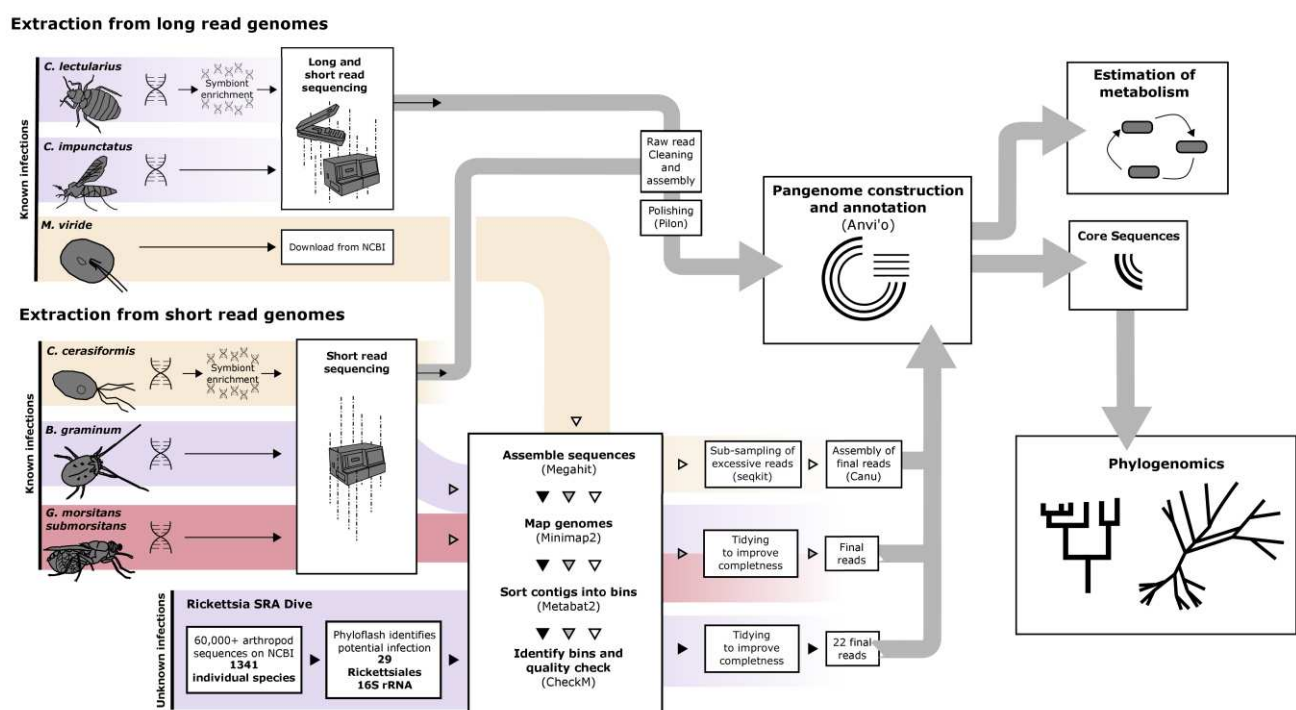


Figure 1. Workflow diagram for extraction, assembly and analyses performed in this study. Purple highlights *Torix Rickettsia* and orange highlights *Ca. Megaira* and red highlights *Transitional Rickettsia*. A full resolution version can be found here: <https://doi.org/10.6084/m9.figshare.15081975>.

DNA preparation, sequencing strategies and symbiont assembly methodologies varied between species. Methods are summarised in Figure 1 and detailed in supplementary material <https://doi.org/10.6084/m9.figshare.14865582>. The exact pipeline used to assemble genomes from Short Read Archive (SRA) data can be found here: <https://figshare.com/s/d1155765b523a6379443>.

### Sample collection for targeted genome assembly

*Cimex lectularius* were acquired from the 'S1' isofemale colony maintained at the University of Bayreuth described in Thongprem et al (2020). *Culicoides impunctatus* females were collected from

146 a wild population in Kinlochleven, Scotland (56° 42' 50.7"N 4° 57' 34.9"W) on the evenings of the  
147 2nd and 3rd September 2020 by aspiration. *Carteria cerasiformis* strain NIES 425 was obtained from  
148 the Microbial Culture Collection at the National Institute for Environmental Studies, Japan. The  
149 *Glossinia morsitans submorsitans* specimen Gms8 was collected in Burkina Faso in 2010 and  
150 *Rickettsia* infection was present alongside other symbionts as described in Doudoumis et al. (2017).  
151 The assembly itself is a result of later thesis work (Blow, 2017).

152 A *Bryobia* mite community was sampled from herbaceous vegetation in Turku, Finland. The  
153 Moomin isofemale line was established by isolating a single adult female and was maintained on  
154 detached leaves of *Phaseolus vulgaris* L. cv Speedy at 25 °C, 60 % RH, and a 16:8 light:dark  
155 photoperiod. The Moomin spider mite line was morphologically identified as *Bryobia graminum* by  
156 Prof Eddie A. Ueckermann (North-West University).

#### 157 *Previously published Rickettsia genomes*

158 A total of 86 published *Rickettsia* genomes, and one genome from *Orientia tsutsugamushi* were  
159 retrieved from the European Nucleotide Archive and assessed with CheckM v1.0.13 (Parks et al.,  
160 2015). Inclusion criteria for genomes were high completeness (CheckM > 90%), low contamination  
161 (CheckM < 2%) and low strain heterogeneity (Check M < 50%) except in the case of *Adalia* for which  
162 there is only one genome (87.6% completeness). Filtering identified 76 high quality *Rickettsia*  
163 genomes that were used in all subsequent analyses (S1  
164 <https://figshare.com/s/198c88c6e3ea5553192e>).

#### 165 *Genome content comparison and pangenome construction*

166 Anvi'o 7 (Eren et al., 2021) was used to construct a pangenome for *Rickettsia*. Included in this were  
167 the 22 MAGs retrieved from SRA data, 2 *Ca. Megaira* genomes and 4 targeted *Torix Rickettsia*  
168 genomes, and one transitional group *Rickettsia* genome acquired in this study. To these were  
169 added the 76 published and 1 *Orientia* described above, giving a total of 104 genomes. Individual  
170 Anvi'o genome databases were additionally annotated with HMMER, KofamKOALA, and NCBI COG  
171 profiles (Aramaki et al., 2020; Eddy, 2011; Galperin et al., 2021). For the pangenome itself,  
172 orthologs were identified with NCBI blast, mcl inflation was set to 2, and minbit was 0.5. Genomes  
173 were arranged according to cluster presence absence and average nucleotide sequence identity  
174 was calculated using pyANI (Pritchard et al., 2016). See  
175 <https://figshare.com/s/d1155765b523a6379443> for the exact code used in this section.



176 KofamKOALA annotation (Aramaki et al., 2020) in Anvi-o 7 was used to estimate completeness of  
 177 metabolic pathways. Then Pheatmap (Kolde, 2019) in R 3.4.4 (R Core Team, 2020) was used to  
 178 produce heatmaps of metabolic potential (figure 7). Annotations for function and *Rickettsia* group  
 179 were added *post hoc* in Inkscape.

180 The biotin operon found in the genome *Rhizobium Rickettsia* Oopac6 was identified from metabolic  
 181 prediction (figure 7). To confirm Oopac6 carries a complete biotin pathway that shares ancestry  
 182 with the existing *Rickettsia* biotin operon, Oopac6 biotin was compared to biotin pathways from  
 183 five other related symbionts: *Cardinium*, *Lawsonia*, *Buchnera aphidicola*, *Rickettsia buchneri*, and  
 184 *Wolbachia* (Seemann, 2014). Clinker (Gilchrist & Chooi, 2021) with default options was used to  
 185 compare and visualise the similarity of genes within the biotin operon region of all 6 bacteria.

186 All generated draft and complete reference genomes were annotated using the NCBI's Prokaryotic  
 187 Genome Annotation Pipeline (PGAP) (Tatusova et al., 2016). Secondary metabolite biosynthetic  
 188 gene clusters were identified using AntiSMASH version 6.0 (Blin et al., 2021) along with Norine  
 189 (Flissi et al., 2019) which searched for similarities to predicted non-ribosomal peptides.

190 Functional enrichment analyses between the main *Rickettsia* clade and the Torix – *Ca. Megaira*  
 191 clades were performed using the Anvi'o program `anvi-get-enriched-functions-per-pan-group` and  
 192 the "COG\_FUNCTION" as annotation source. A gene cluster presence - absence table was exported  
 193 using the command "`anvi-export-tables`". This was used to create an UpSet plot using the R package  
 194 ComplexUpset (Krassowski et al., 2020; Lex et al., 2014) to visualize unique and shared gene  
 195 clusters between different *Rickettsia* groups. A gene cluster was considered unique to a specified  
 196 *Rickettsia* group when it was present in at least one genome belonging to that group. Gene cluster  
 197 accumulation curves were performed for the pan-, core- and unique-genomes based on the same  
 198 presence-absence matrix using a custom-made R script (Siozios, 2021). In each case the cumulative  
 199 number of gene clusters were computed based on randomly sampled genomes using 100  
 200 permutations. The analysis was performed separately for each of the five major *Rickettsia* groups as  
 201 well as the complete *Rickettsia* dataset. Curves were plotted using the ggplot2 R package  
 202 (Wickham, 2016).

203 All information on extra genomes can be found at <https://doi.org/10.6084/m9.figshare.14865582>  
 204 and the code pipeline employed can be found at <https://figshare.com/s/d1155765b523a6379443>.

## Phylogeny, Network, and recombination

The single-copy core of all 104 genomes was identified in Anvi'o 7 and is made up of 74 single-copy gene (SCG) clusters. Protein alignments from SCG were extracted and concatenated using the command "anvi-get-sequences-for-gene-clusters". Maximum likelihood phylogeny was constructed in IQ-TREE v2.1.2 (Nguyen et al., 2015). Additionally, 43 ribosomal proteins were identified through Anvi'o 7 to test phylogenomic relationships. These gene clusters were extracted from the pangenome and used for an independent phylogenetic analysis SUPPLEMENTARY FIG. The best model according to the Bayesian Information Criterion (BIC) was selected with Model Finder Plus (MFP) (Kalyaanamoorthy et al., 2017) as implemented in IQ-TREE; this was JTTDCMut+F+R6 for core gene clusters and JTTDCMut+F+R3 for ribosomal proteins. Both models were run with Ultrafast Bootstrapping (1000 UF bootstraps) (Hoang et al., 2018) with *Orientia tsutsugamushi* as the outgroup.

The taxonomic placement of Oopac6, Ppec13 and Dallo3 genomes within the Rhyzobius, Meloidae and Belli groups respectively were confirmed in a smaller phylogenetic analysis, performed as detailed in (Pilgrim et al. 2021) using reference MLST sequences (*gltA*, 16s rRNA, 17kDa, *COI*) from other previously identified *Rickettsia* profiles (S1 <https://figshare.com/s/198c88c6e3ea5553192e>). The selected models used in the concatenated partition scheme were as follows: 16S rRNA: TIM3e+I+G4; 17KDa: GTR+F+I+G4; *COI*: TPM3u+F+I+G4; *gltA*: K3Pu+F+I+G4a.

A nearest neighbour network was produced for core gene sets with default settings in Splitstree4 to further assess distances and relationships between *Rickettsia*, *Ca. Megaira* and *Torix* clades. All annotation was added post hoc in Inkscape. Furthermore, recombination signals were examined by applying the Pairwise Homoplasy Index (PHI) test to the DNA sequence of each core gene cluster extracted with Anvi'o-7. DNA sequences were aligned with MUSCLE (Edgar, 2004) and PHI scores calculated for each of the 74 core gene cluster with PhiPack (Bruen et al., 2006).

The taxonomic identity for new and newly expanded groups was established with GTDB-Tk (Chaumeil et al., 2020) to support the designation of new taxa through phylogenetic comparison of marker genes against an online reference database.

## Results and Discussion

We have expanded the available genomic data for several *Rickettsia* groups through a combination of draft and complete genome assembly. This includes an eight-fold increase in available *Torix*-

group genomes, and the first available genomes for Meloidae and Rhyzobius groups. We further report the first reference genomes for *Ca. Megaira*.

### Complete and closed reference genomes for *Torix Rickettsia* and *Ca. Megaira*

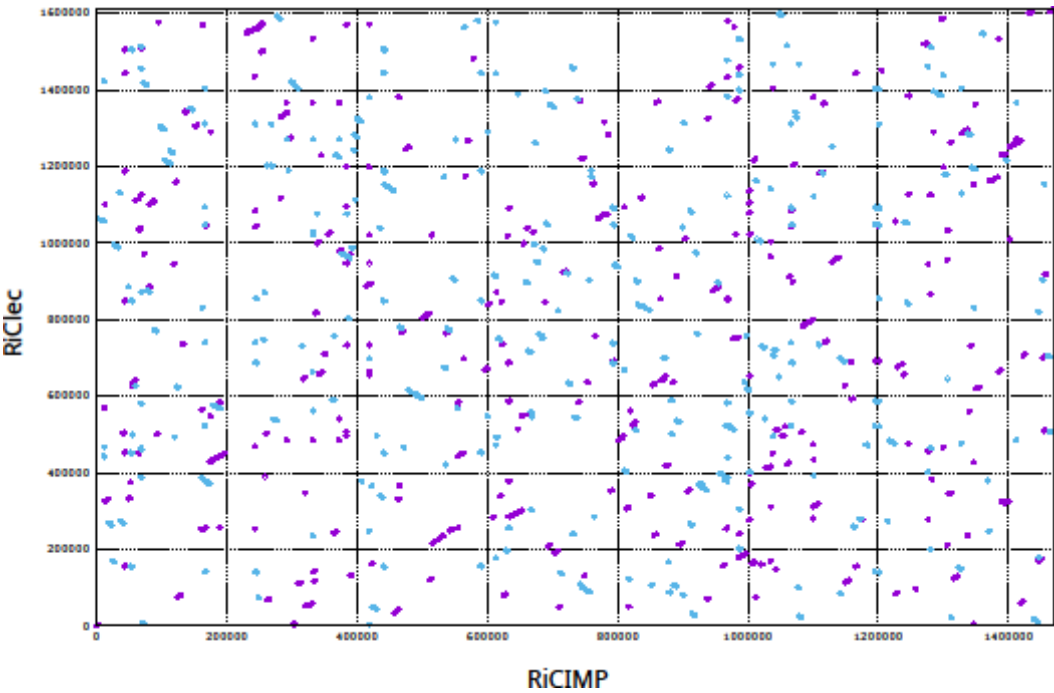
The use of long-read sequencing technologies produced the first complete genomes for two subclades of the *Torix* group (RiCimp-limoniae, RiClec-leech). Sequencing depth of the *Rickettsia* genomes from *C. impunctatus* (RiCimp) and *C. lectularius* (RiClec) were 18X and 52X respectively. The RiCimp genome provides the first evidence of plasmids in the *Torix* group (pRiCimp001 and pRiCimp002). In addition, we assembled the first complete closed reference genome of *Ca. Megaira* from *Mestostigma viride* (MegNEIS296) from previously published genome sequencing efforts.

General features of both genomes are consistent with previous genomic studies of the *Torix* group (Table 1). A single full set of rRNAs (16S, 5S and 23S) and a GC content of ~33% was observed. Notably, the two complete *Torix* group genomes show a distinct lack of synteny (see S2 <https://doi.org/10.6084/m9.figshare.14866263>), a genomic feature that is compatible with our phylogenetic analyses that placed these two lineages in different subclades (leech/limoniae) (figures 2 and 3). Of note within the closed reference genomes MegNEIS296 and RiCimp, is the presence of a putative non-ribosomal peptide synthetase (NRPS) and a hybrid non-ribosomal peptide/polyketide synthetase (NRPS/PKS) respectively (see S3 <https://doi.org/10.6084/m9.figshare.14865570>). Although, the exact products of these putative pathways are uncertain, *in silico* prediction by Norine suggests close similarity with both cytotoxic and antimicrobial peptides hinting at a potential defensive role (see S3 <https://doi.org/10.6084/m9.figshare.14865570>). A hybrid NRPS/PKS cluster has previously been reported in *Rickettsia buchneri* on a mobile genetic element, providing potential routes for horizontal transmission (Hagen et al., 2018). In addition, putative toxin-antitoxin systems similar to the one associated with cytoplasmic incompatibility in *Wolbachia* have recently been observed on the plasmid of *Rickettsia felis* in a parthenogenetic booklouse (Gillespie et al., 2015, 2018). Toxin-endotoxin systems are thought to be part of an extensive bacterial mobilome network associated with reproductive parasitism (Gillespie et al., 2018). A BLAST search found a very similar protein in Oopac6 to the putative large pLbAR toxin found in *R. felis* (88% aa identity), and a more distantly related protein in the *C. impunctatus* plasmid (25% aa identity).

265 Table 1. Summary of the complete *Ca. Megaira* and *Torix Rickettsia* genomes

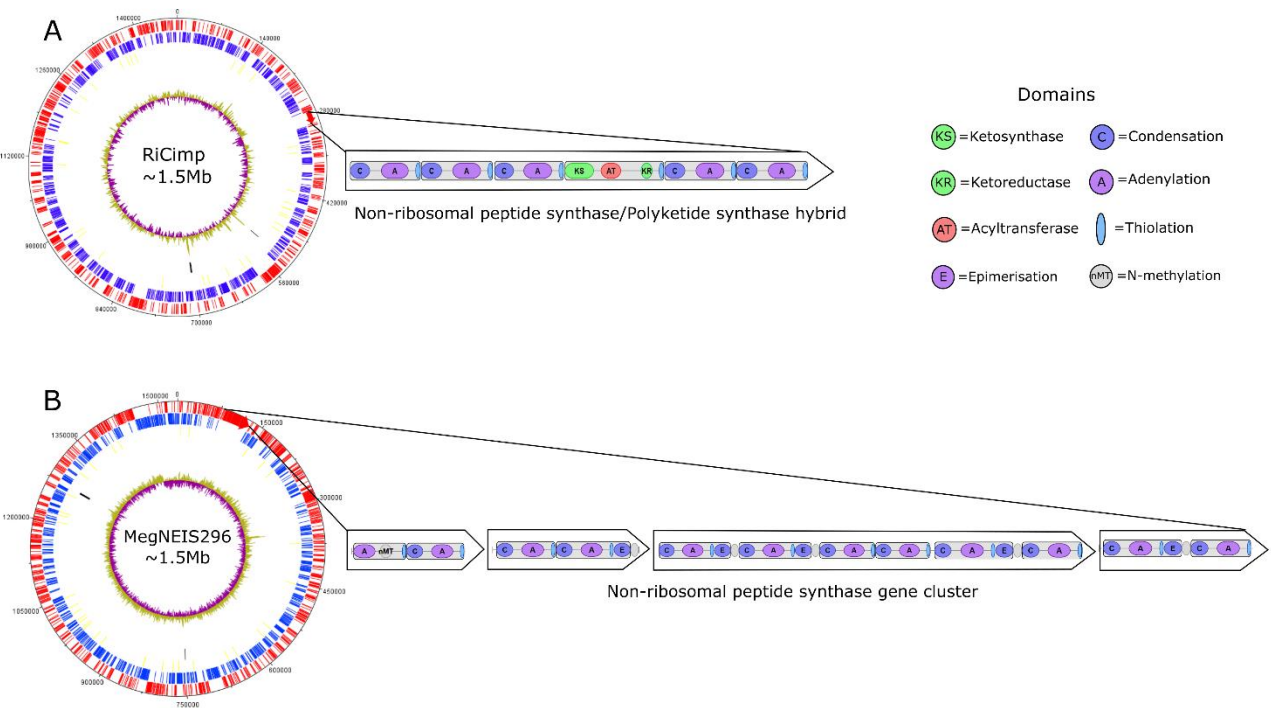
Group	<i>Ca. Megaira</i>	<i>Torix Rickettsia</i>	
Sub-group		Leech	Limoniae
Strain Name	MegNIES296	RiCimp	RiClec
Symbiont genome accession	CP084576-CP084577	CP084573-CP084575	CP084572
Host	Mesostigma viride NIES-296	Culicoides impunctatus	Cimex lectularius
Raw reads accession	SRR8439255, SRX5120346	SRR16018514, SRR16018513	SRR16018512, SRR16018511
Total nucleotides	1,532,409	1,566,468	1,611,726
Chromosome size (bp)	1,448,425	1,469,631	1, 611,726
Plasmids	1 (83,984 bp)	2 (77550bp + 19287bp)	None
GC content (%)	33.9	32.9	32.8
Number of CDS	1,359	1,397	1,544
Avg. CDS length (bp)	998	900	874
Coding density (%)	88.5	86	84
rRNAs	3	3	3
tRNAs	34	34	35

266



267

268 S2 Whole genome alignment between the complete *Torix limoniae* (RiCIMP) and *Torix Leech* (RiClec) genomes reveals complete lack  
269 of synteny. Magenta represents forward matches and blue reverse matches <https://doi.org/10.6084/m9.figshare.14866263>.



270  
271 *S3. The circular chromosomes of A) a Torix group Rickettsia (RiCimp) and B) a Ca. Megaira sp. (MegNEIS296). From outside to in, the*  
272 *circles represent: forward CDSs (Red), Reverse CDSs (blue), tRNAs (yellow) rRNAs (black), and GC content (green and magenta).*  
273 *Highlighted are the predicted modules formed by non-ribosomal peptide synthase genes (domains) that define individual amino acids*  
274 *in the synthesised peptide and show the catalytic domains within modules <https://doi.org/10.6084/m9.figshare.14865570>.*

275 Sequencing and *de novo* assembly of other *Rickettsia* and *Ca. Megaira* genomes.  
276 Our direct sequencing efforts enabled assembly of draft genomes for a second *Ca. Megaira* strain  
277 from the alga *Carteria cerasiformis*, and for *Rickettsia* associated with tsetse flies and *Bryobia* spider  
278 mites. The Transitional *Rickettsia* from a wild caught Tsetse fly, RiTSETSE, is a potentially chimeric  
279 assembly since we identified an excess of biallelic sites when the raw Illumina reads were mapped  
280 back to the assembly. It is also likely that RiTSETSE is not a heritable symbiont but comes from  
281 transient infection from a recent blood meal.

282 From the SRA accessions, the metagenomic pipeline extracted 29 full symbiont genomes for  
283 Rickettsiales across 24 host species. Five of 29 were identified as *Wolbachia* and discarded from  
284 further analysis, one was a *Rickettsia* discarded for low quality, and another was a previously  
285 assembled Torix *Rickettsia*, RiCNE (Pilgrim et al., 2017). Thus, 22 high quality *Rickettsia*  
286 metagenomes were obtained from 21 host species. One beetle (SRR6004191) carried coinfecting  
287 *Rickettsia* Lappe3 and Lappe4 (Table 2). The high-quality *Rickettsia* covered the Belli, Torix,  
288 Transitional, Rhyzobius, Meloidae and Spotted Fever Groups (Table 2 and S1  
289 <https://figshare.com/s/198c88c6e3ea5553192e>).

290 Beetles, particularly rove beetle (Staphylinidae) species, appear in this study as a possible hotspot  
 291 of *Rickettsia* infection. *Rickettsia* has historically been commonly associated with beetles, including  
 292 ladybird beetles (*Adalia bipunctata*), diving beetles (*Deronectes* sp.) and bark beetles (*Scolytinae*)  
 293 (Hurst et al., 1994; K  chler et al., 2009; Perlman et al., 2006; Weinert et al., 2009; Zchori-Fein et al.,  
 294 2006). Though a plausible and likely hotspot, this observation needs be approached with caution as  
 295 this could be an artefact of skewed sampling efforts.

296 All genome metadata and source information can be found here  
 297 <https://figshare.com/s/198c88c6e3ea5553192e>.

298

299 Table 2. Brief summary of draft genomes generated during the current project and their associated hosts. Full metadata can be found  
300 in S1 <https://figshare.com/s/198c88c6e3ea5553192e>.

Strain	Bacteria Biosample Accession	Group	Number of contigs	Total length (bp)	Host name	Order
<b>Blapp1</b>	SAMN21822536	Belli	171	1266633	<i>Bembidion lapponicum</i>	Coleoptera
<b>Btrans1</b>	SAMN21822537	Belli	241	1417452	<i>Bembidion nr. transversale</i> OSAC:DRMaddison DNA3205	Coleoptera
<b>Choog2</b>	SAMN21822538	Belli	16	1357829	<i>Columbicola hoogstraali</i>	Phthiraptera
<b>Cmasu2</b>	SAMN21822539	Transitional	196	1295004	<i>Ceroptres masudai</i>	Hymenoptera
<b>Dallo3</b>	SAMN21822540	Belli	196	990679	<i>Diachasma alloeum</i>	Hymenoptera
<b>Drufa1</b>	SAMN21822541	Belli	14	1364611	<i>Degeeriella rufa</i>	Phthiraptera
<b>Earac4</b>	SAMN21822542	Transitional	96	1350066	<i>Ecitomorpha arachnoides</i>	Coleoptera
<b>Econn1</b>	SAMN21822543	Transitional	238	1070326	<i>Eriopsis connexa</i>	Coleoptera
<b>Gbili3</b>	SAMN21822544	Torix limoniae	171	1188102	<i>Gnoriste bilineata</i>	Diptera
<b>Gdoso1</b>	SAMN21822545	Belli	34	1420758	<i>Graphium doson</i>	Lepidoptera
<b>Lappe3</b>	SAMN21822558	Torix limoniae	122	1368980	<i>Labidopullus appendiculatus</i>	Coleoptera
<b>Lappe4</b>	SAMN21822559	Torix limoniae	154	1332357	<i>Labidopullus appendiculatus</i>	Coleoptera
<b>MegCarte- ria</b>	SAMN21822546	Ca. Megaira	72	1298707	<i>Carteria cerasiformis</i>	Chlamydomonadales
<b>Ofont3</b>	SAMN21822560	Adalia	91	1529137	<i>Omalisus fontisbellaquei</i>	Coleoptera
<b>Oopac6</b>	SAMN21822548	Rhyzobius	181	1497231	<i>Oxypoda opaca</i>	Coleoptera
<b>Pante1</b>	SAMN21822549	Torix limoniae	70	1472610	<i>Pseudomimeceton antennatum</i>	Coleoptera
<b>Pfluc4</b>	SAMN21822550	Spotted Fever Group	7	1251895	<i>Proechinophthirus fluctus</i>	Phthiraptera
<b>Ppec13</b>	SAMN21822551	Belli	90	1426047	<i>Pyrocoelia pectoralis</i>	Coleoptera
<b>Psono2</b>	SAMN21822552	Torix limoniae	163	1492063	<i>Platyusa sonomae</i>	Coleoptera
<b>RiTSETSE</b>	SAMN21822553	Transitional	172	1451997	<i>Glossina morsitans submorsitans</i>	Diptera
<b>S2</b>	SAMN21822554	Torix limoniae	103	1251484	<i>Sericostoma</i>	Trichoptera
<b>Sanch3</b>	SAMN21822555	Belli	181	1487154	<i>Stiretrus anchorago</i>	Hemiptera
<b>Slati1</b>	SAMN21822556	Transitional	109	1301763	<i>Sceptobius lativentris</i>	Coleoptera
<b>Ssp4</b>	SAMN21822557	Torix limoniae	87	1231013	<i>Sericostoma sp.</i> HW- 2014	Trichoptera
<b>moomin</b>	SAMN21822560	Torix moomin	204	1137559	<i>Bryobia graminum</i>	Trombidiformes



## Phylogenomic analyses and taxonomic placement of newly assembled genomes

### *Phylogeny, network, and recombination*

The network and phylogeny illustrate the distance of Torix from *Ca. Megaira* and other *Rickettsia*, along with an extremely high level of within-group diversity in Torix compared to any other group (Figures 2 and 3). The phylogenies generated using core genomes are consistent with previously identified *Rickettsia* and host associations using more limited genetic markers. For instance, Pfluc4 from *Proechinophthirus fluctus* lice is grouped on the same branch as a previously sequenced *Rickettsia* from a different individual of *P. fluctus*. Four of 22 genomes from the SRA screen are identified as Transitional, 1 is in Spotted Fever Group, 1 is *Adalia*, 8 are *Belli* and 7 are *Torix limoniae*. Targeted sequences were confirmed as: *Torix limoniae* (RiCimp), *Torix leech* (RiClec), Transitional (RiTSETSE), *Ca. Megaira* (MegCarteria and MegNEIS296), and a new Torix clade, Moomin (Moomin). The new Torix include one double infection giving a total of 10 new genomes across 9 potential host species. The double infection is found within the rove beetle *Labidopullus appendiculatus*, forming two distinct lineages, Lappe3 and Lappe4 (Fig 2 and 3).

In addition, the pre-existing *Rickettsia helvetica*, which is typically cited as a member of the Spotted Fever group as a result of its first description in 1993 (Beati et al., 1993; Chisu et al., 2017), seems to form its own group in all trees and networks (figure 2, 3 and <https://doi.org/10.6084/m9.figshare.14865606>). We conclude from this that *Rickettsia helvetica* is most similar to Scapularis group *Rickettsia*, but because it does not fall into the same clade in any tree or network, it is likely that the strain belongs to a distinct lineage of tick-borne *Rickettsia*.



## Core Phylogeny

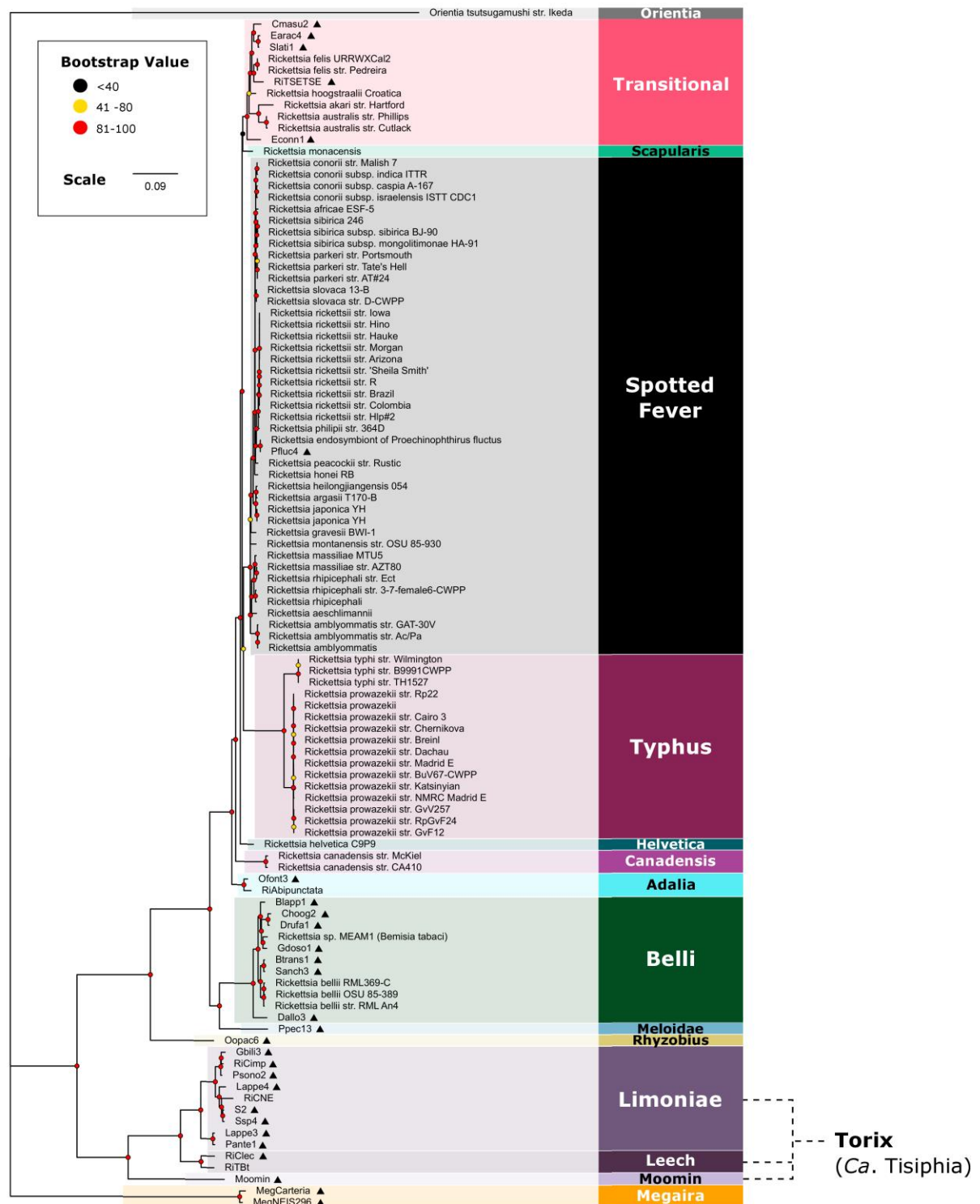
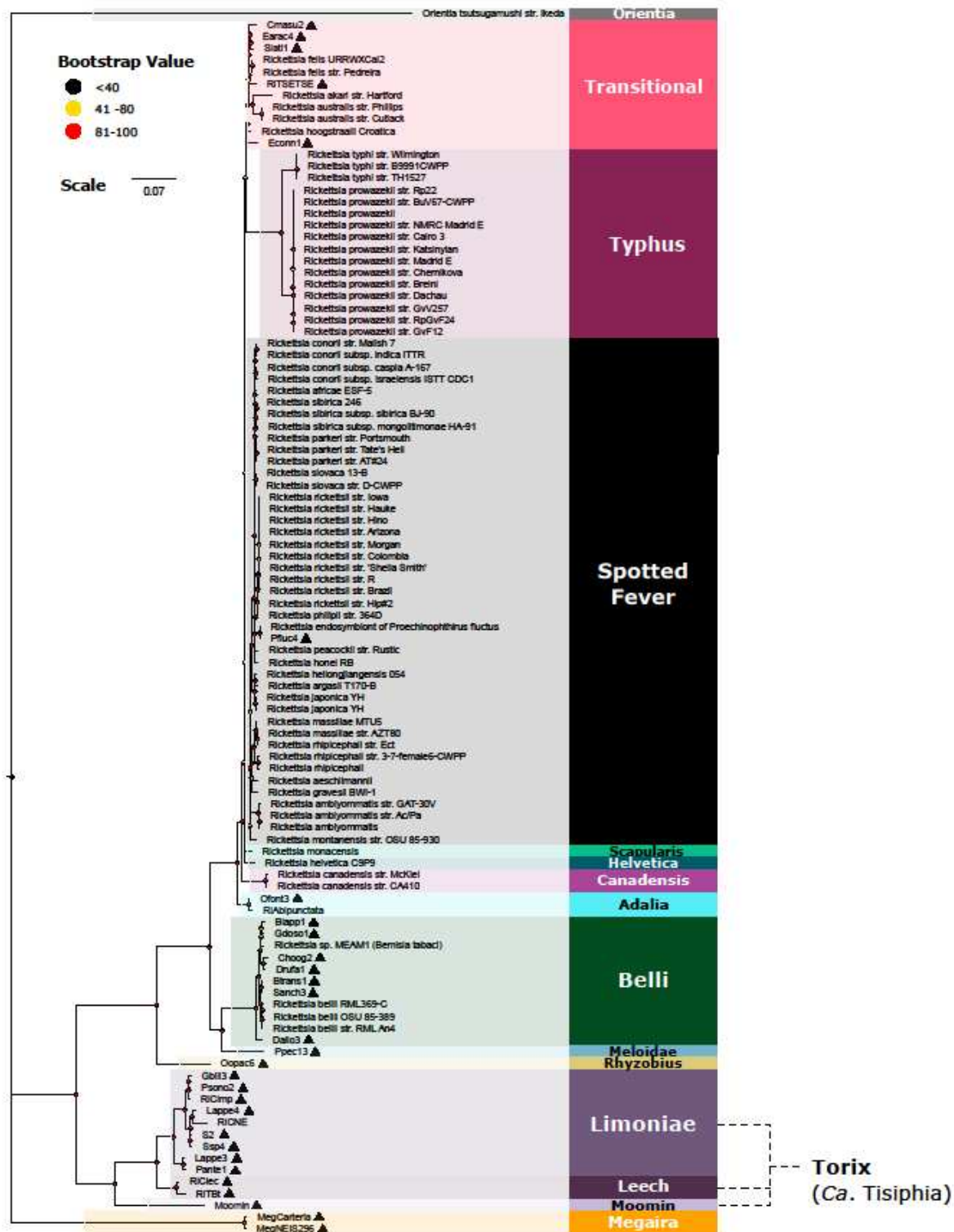


Figure 2. *Rickettsia* and *Ca. Megaira maximum* likelihood (ML) phylogeny constructed from 74 core gene clusters extracted from the pangenome. New genomes are indicated by ▲ and bootstrap values based on 1000 replicates are indicated with coloured circles. New complete genomes are: RiCimp, RiClec and MegNEIS296. A full resolution version can be found here: <https://doi.org/10.6084/m9.figshare.15081975>.

## Ribosomal Phylogeny



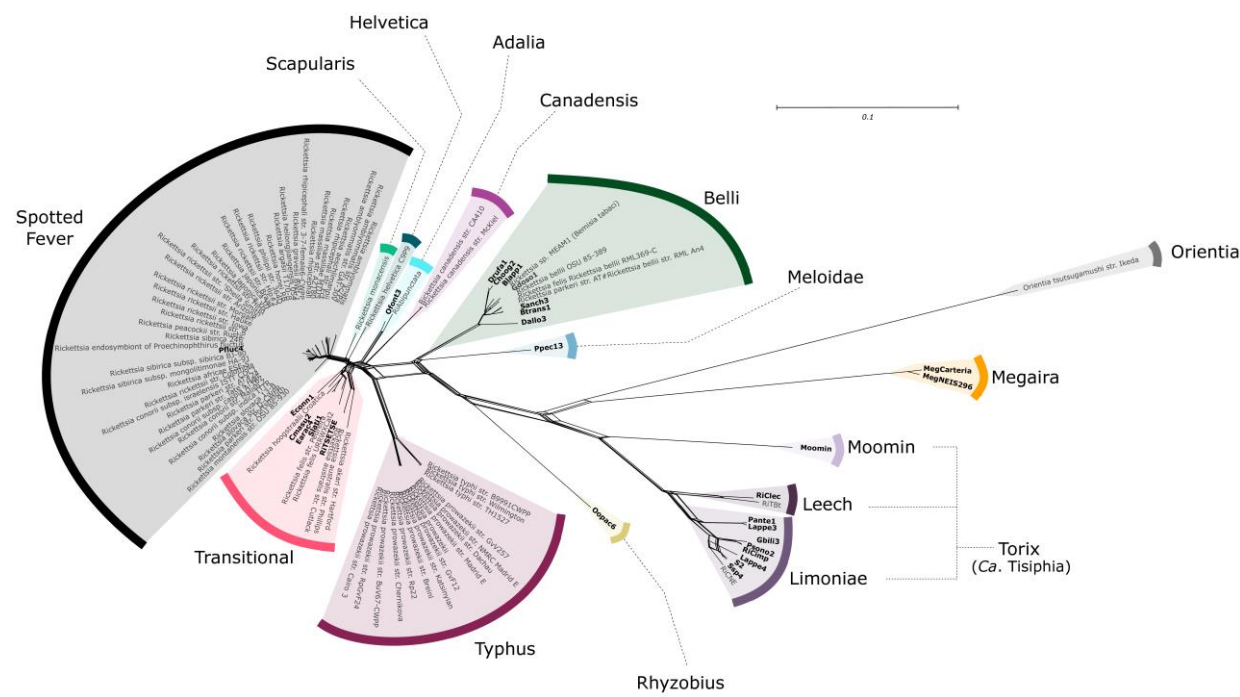
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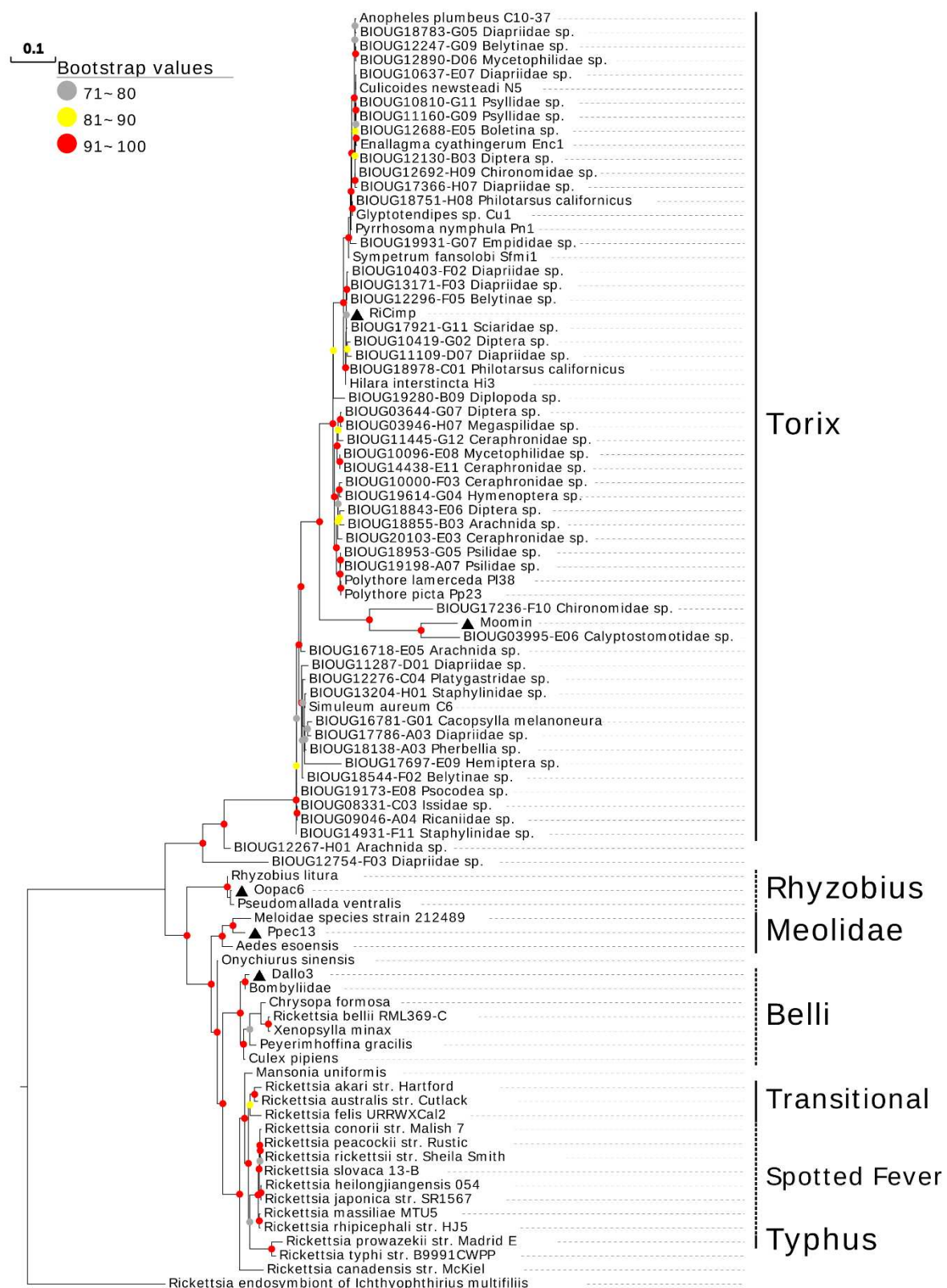
329

S4. *Rickettsia* and *Ca. Megaira* maximum likelihood (ML) phylogeny constructed from 43 ribosomal protein gene clusters extracted from the pangenome. New genomes are indicated by ▲ and bootstrap values based on 1000 replicates are indicated with coloured circles. New complete genomes are: RiCimp, RiClec and MegNEIS296. <https://doi.org/10.6084/m9.figshare.14865606>



330 Figure 3. Nearest Neighbour Network, displaying the distances between the 74 core gene sets across all 104 *Rickettsia*, *Ca. Megaira*  
331 genomes, and the outgroup *Orientia*. New genomes are indicated with bold text. A full resolution version can be found here:  
332 <https://doi.org/10.6084/m9.figshare.15081975>.

333 We also report the first putative *Rhyzobius Rickettsia* genomes extracted from the staphylinid  
334 beetle *Oxypoda opaca* (Oopac6) and Meloidae *Rickettsia* from the firefly *Pyrocoelia pectoralis*  
335 (Ppec13). They have high completeness (S1 <https://figshare.com/s/198c88c6e3ea5553192e>), low  
336 pseudogenisation, and consistently group away from the other draft and completed genomes  
337 (Figures 2 and 3). MLST analyses demonstrate that these bacteria are most like the *Rhyzobius* and  
338 Meloidae groups described by Weinert *et al.* (2009) (see S5  
339 <https://doi.org/10.6084/m9.figshare.14865600>). The pangenome and metabolic profile of this draft  
340 genome suggests that Meloidae is a sister group to Belli and that *Rhyzobius Rickettsia* is  
341 superficially similar to Belli and Transitional groups. The *Rhyzobius*-group symbiont is  
342 phylogenetically distant from most *Rickettsia* and is potentially a sister clade linking Torix and the  
343 main *Rickettsia* clades. Further genome construction will help clarify this taxon and its relationship  
344 to the rest of *Rickettsia*.



345

346 S5 Phylogram of a maximum likelihood (ML) tree of 90 Rickettsia mutiocus profiles. The tree is based on 4 loci, 16S rRNA, 17KDa,  
347 gltA, and COI, under a partition model (2,781 bp total). <https://doi.org/10.6084/m9.figshare.14865600>



The sequencing data for the wasp, *Diachasma alloeum*, used here has previously been described to contain a pseudogenised nuclear insert of *Rickettsia* material, but not a complete *Rickettsia* genome (Tvedte et al., 2019). The construction of a full, non-pseudogenised genome with higher read depth than the insect contigs, low contamination (0.95%) and high completion (93.13%) suggests that these reads likely represent a viable *Rickettsia* infection in *D. alloeum*. However, these data do not exclude the presence of an additional nuclear insert. It is possible for a whole symbiont genome to be incorporated into the host's DNA (Hotopp et al., 2007), and there are recorded partial inserts of *Ca. Megaira* genomes in the *Volvox carteri* genome (Kawafune et al., 2015). The presence of both the insert and symbiont need confirmation through appropriate microscopy methods.

Recombination is low within the core genomes of *Rickettsia* and *Ca. Megaira*, but may occur between closely related clades that are not investigated here. Across all genomes, the PHI score is significant in 6 of the 74 core gene clusters, suggesting putative recombination events. However, it is reasonable to assume that most of these may be a result of systematic error due to the divergent evolutionary processes at work across *Rickettsia* genomes. Patterns of recombination can occur by chance rather than driven by evolution which cannot be differentiated by current phylogenetic methods (Murray et al., 2016). The function of each respective cluster can be found at <https://figshare.com/s/198c88c6e3ea5553192e>.

## Gene content and pangenome analysis

### Pangenome

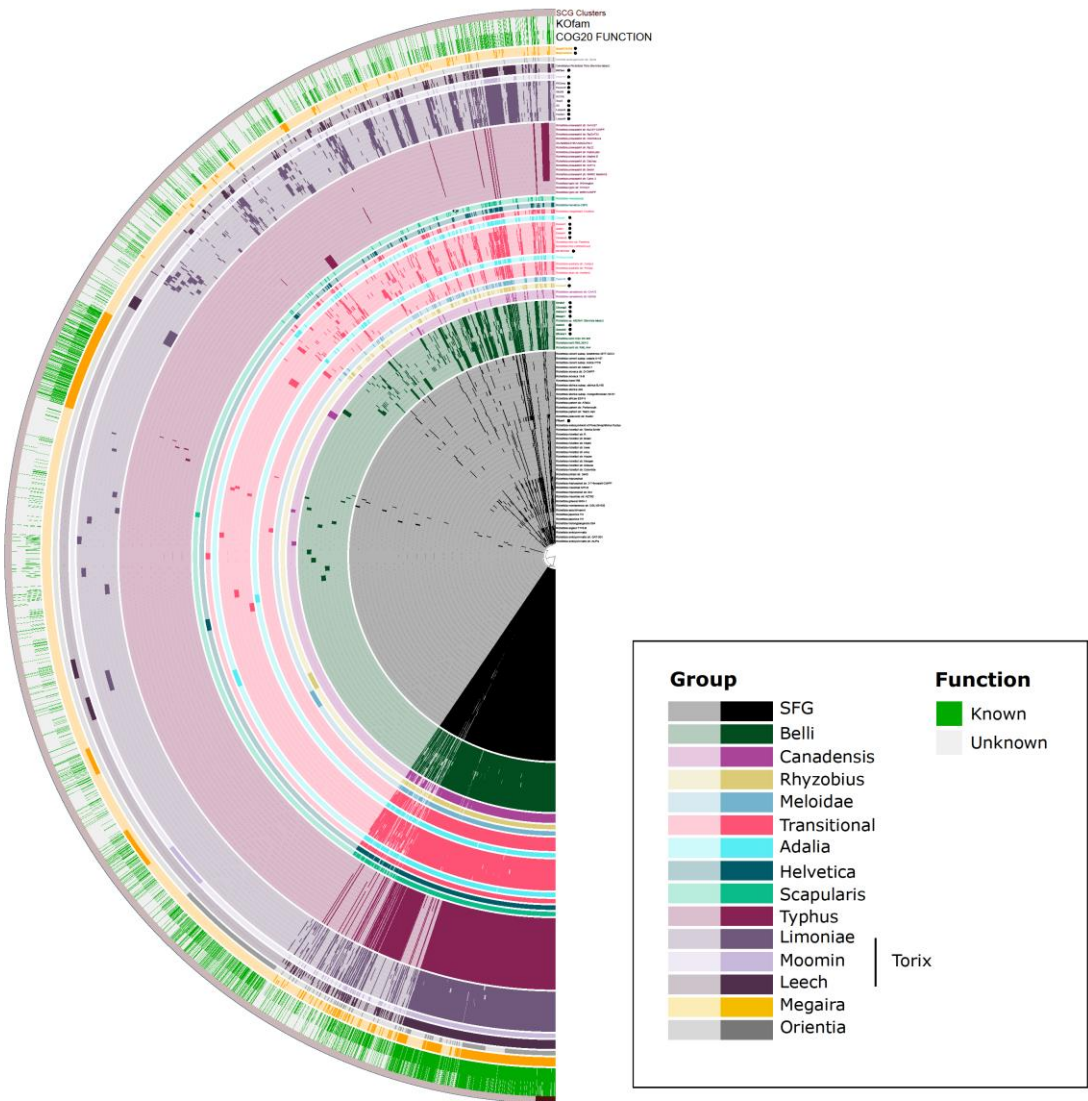
Across all 104 genomes used in the pangenome analysis (figure 2, full data in S6 <https://doi.org/10.6084/m9.figshare.14865576>), Anvi'o identified 208 core gene clusters of which 74 are represented by single-copy genes. Bacterial strains of the different *Rickettsia* groups, especially the neglected symbiotic Rickettsiaceae, seem to have large, open pangenomes an indication of rapid evolution. As expected, the more genomes that are included in analyses, the smaller the core genome extracted.

Torix is a distinctly separate clade sharing less than 70% ANI similarity to any *Rickettsia* or *Ca. Megaira* genomes. It contains at least three groups that reflect its highly diverse niche in the environment (figure 5) (Jain et al., 2018; Pilgrim et al., 2021; Rodriguez-R et al., 2021). Torix has the most unique genes out of all the clades in this study followed by *Ca. Megaira* and Belli clades (figure 6). Rarefaction gene accumulation analysis suggest that Torix is the group where each additional

379 genome included increases the pangenome repertoire to the greatest extent (figure 7). Torix group  
 380 is thus more diverse in terms of genome content and size of the pangenome than other *Rickettsia*  
 381 groups.

382 *Rickettsia* lineages group together based on gene presence/absence and produce repeated patterns  
 383 of accessory genes that reliably occur within each group (figure 2). ANI scores are also strongest  
 384 within groups, while genomes tend to share lower similarity outside of their group (figure 4). This is  
 385 particularly apparent in Torix and *Ca. Megaira* which are divergent from the main *Rickettsia* clade  
 386 (figure 3 and 5).

387



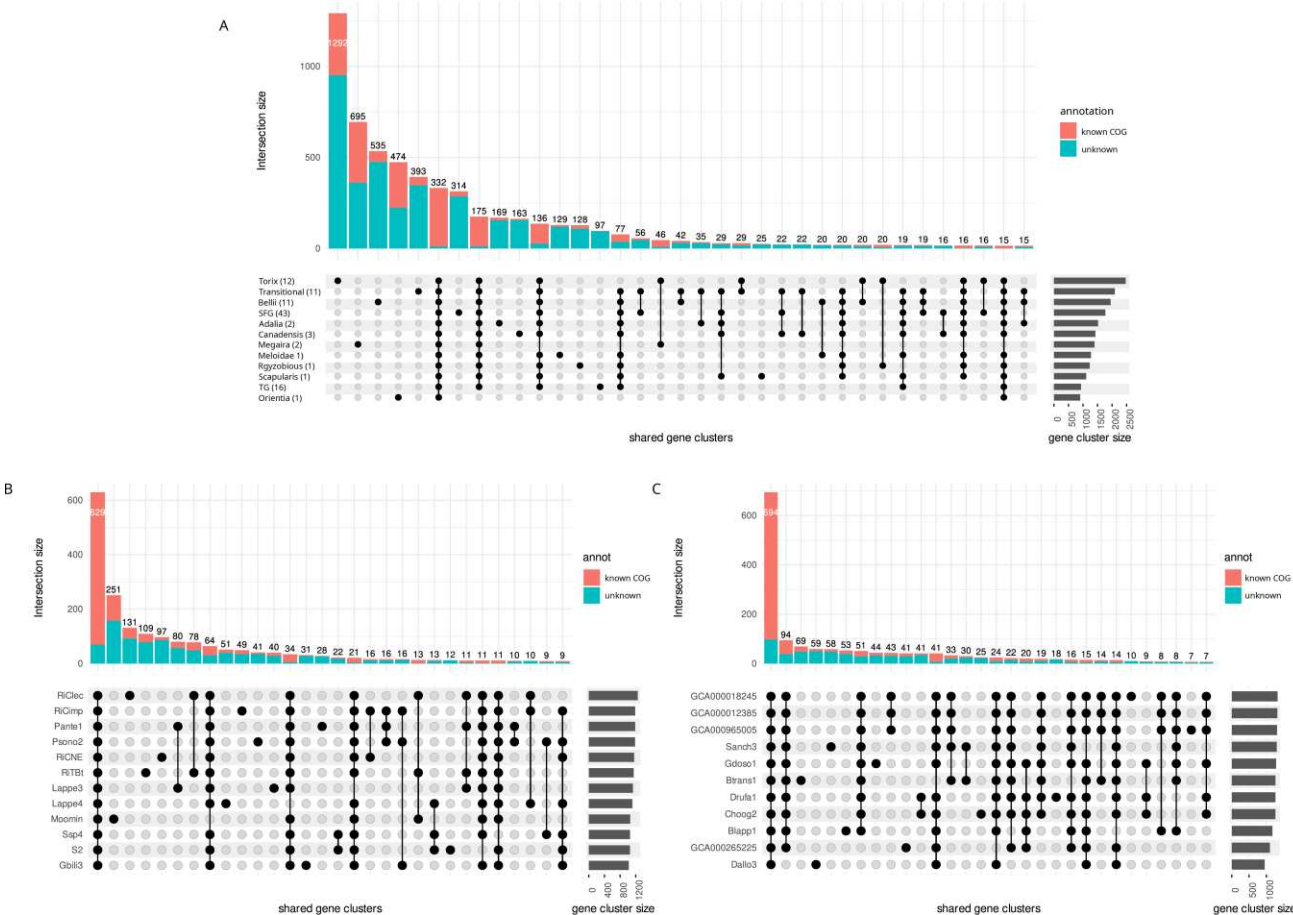
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389 Figure 4. Pangenome of all 104 genomes including *Rickettsia*, *Torix*, *Ca. Megaira* and the outgroup *Orientia*. New genomes are  
390 indicated by ●. Each genome displays gene cluster presence/absence and is organised by gene cluster frequency. Group identity was  
391 assigned from phylogeny. SFG is Spotted Fever Group. A full resolution version can be found here:  
392 <https://doi.org/10.6084/m9.figshare.15081975>.

393

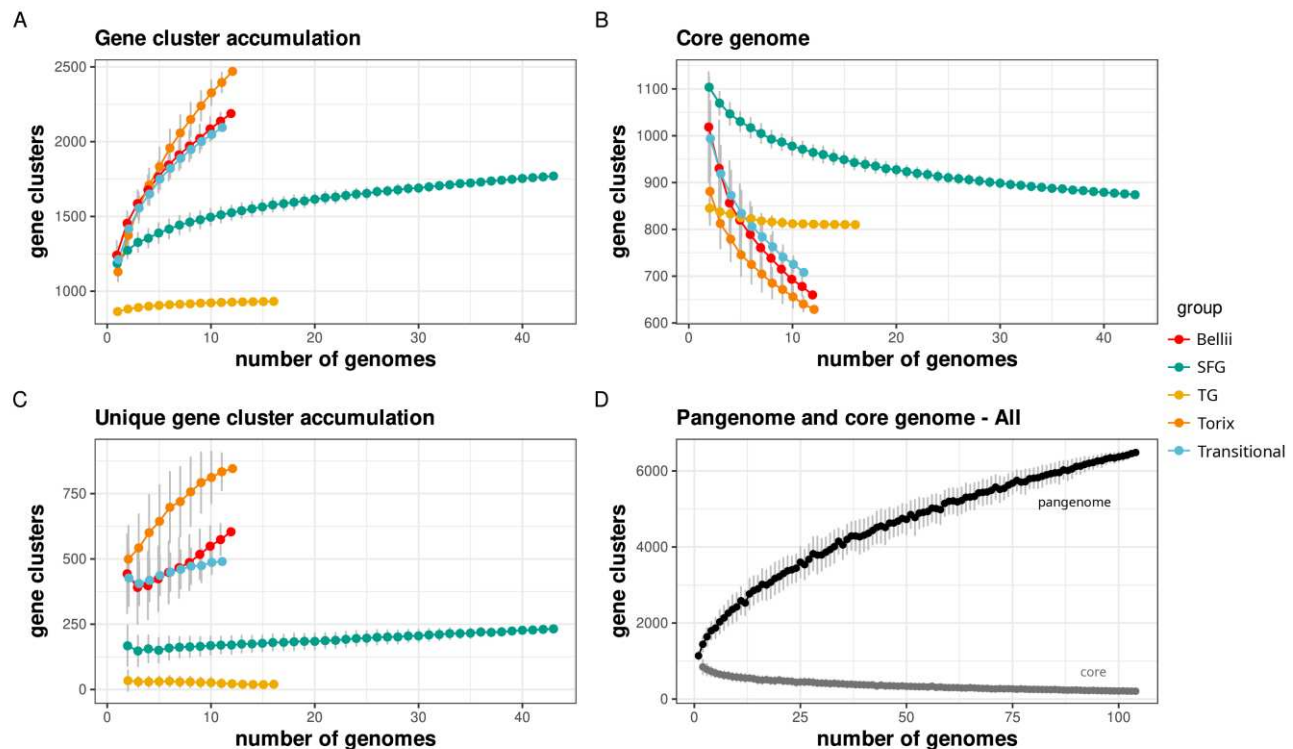






404

405 *Figure 6. Shared and unique gene clusters across A) All Rickettsia and Ca. Megaira genomes used in this study grouped by clade with*  
406 *Orientia as an outgroup B) all individual Torix genomes, and C) all individual Belli genomes. Horizontal grey bars to the right of each*  
407 *plot represent gene cluster size and vertical, coloured bars represent the size of intersections (the number of shared gene clusters)*  
408 *between genomes in descending order with known COG functions displayed in red and unknown in blue. Black dots mean the cluster*  
409 *is present and connected dots represent gene clusters that are present across groups. SFG is Spotted Fever Group and TG is Typhus*  
410 *Group. A full resolution version can be found here: <https://doi.org/10.6084/m9.figshare.15081975>.*



411

412 *Figure 7. Gene cluster accumulation curves for pangenome (A), core genome (B) and the unique genome (C) of the 5 largest Rickettsia*  
413 *groups as a function of the number of genomes sequenced. The pangenome and the core genome accumulation curves for the*  
414 *complete Rickettsia dataset is shown in panel D. Error bars represent  $\pm$  standard deviation based on 100 permutations. SFG is Spotted*  
415 *Fever Group and TG is Typhus Group. A full resolution version can be found here: <https://doi.org/10.6084/m9.figshare.15081975>.*

#### 416 *Gene content and metabolic analyses*

417 Rickettsial genomes extracted from SRA samples are generally congruent with the metabolic  
418 potential of their respective groups (Figure 8). Torix and *Ca. Megaira* have complete pentose  
419 phosphate pathways (PPP); a unique marker for these groups which seems to have been lost in the  
420 other *Rickettsia* clades. The PPP generates NADPH, precursors to amino acids, and is known to  
421 protect against oxidative injury in some bacteria (Christodoulou et al., 2018), as well as conversion  
422 of hexose monosaccharides into pentose used in nucleic acid and exopolysaccharide synthesis. The  
423 PPP has also been associated with establishing symbiosis between the Alphaproteobacteria  
424 *Sinorhizobium meliloti* and its plant host *Medicago sativa* (Hawkins et al., 2018). This pathway has  
425 previously been highlighted in Torix (Pilgrim et al., 2017) and its presence in all newly assembled  
426 Torix and *Ca. Megaira* draft genomes consolidates its importance as an identifying feature for these  
427 groups (Figure 8, S1 <https://figshare.com/s/198c88c6e3ea5553192e>). The PPP is likely an ancestral  
428 feature that was lost in the main *Rickettsia* clade.

429 Glycolysis, gluconeogenesis and cofactor and vitamin metabolism are absent or incomplete across  
430 all *Rickettsia*, except the Rhyzobius group member, Oopac6 (Figure 8). Oopac6 has a complete



456 *hoogstraali*, *Degeeriella rufa*), one is a butterfly (*Graphium doson*), and one is a ground beetle  
 457 (*Bembidion lapponicum*). dTDP-L-rhamnose is an essential component of human pathogenic  
 458 bacteria like *Pseudomonas*, *Streptococcus* and *Enterococcus*, where it is used in cell wall  
 459 construction (van der Beek et al., 2019). This pathway has also been utilized in the synthesis of  
 460 plant cell walls (Jiang et al., 2021), may be involved in the moulting process of *Caenorhabditis*  
 461 *elegans* (Feng et al., 2016), and is a precursor to rhamnolipids that are used in quorum sensing  
 462 (Daniels et al., 2004). In the root symbiont *Azospirillum*, disruption of this pathway alters root  
 463 colonisation, lipopolysaccharide structure and exopolysaccharide production (Jofré et al., 2004). No  
 464 *Rickettsia* from typically pathogenic groups assessed in figure 8 has this pathway, and the hosts of  
 465 these four bacteria are not involved with human or mammalian disease. Presence in feather lice  
 466 provides little opportunity for this *Rickettsia* to be pathogenic as feather lice are chewers rather  
 467 than blood feeders, and Belli group *Rickettsia* more generally are rarely pathogenic. Further, this  
 468 association does not explain its presence in a butterfly and ground beetle; it is most likely that this  
 469 pathway, if functional, would be involved in establishing infection in the insect host or host-  
 470 symbiont recognition.

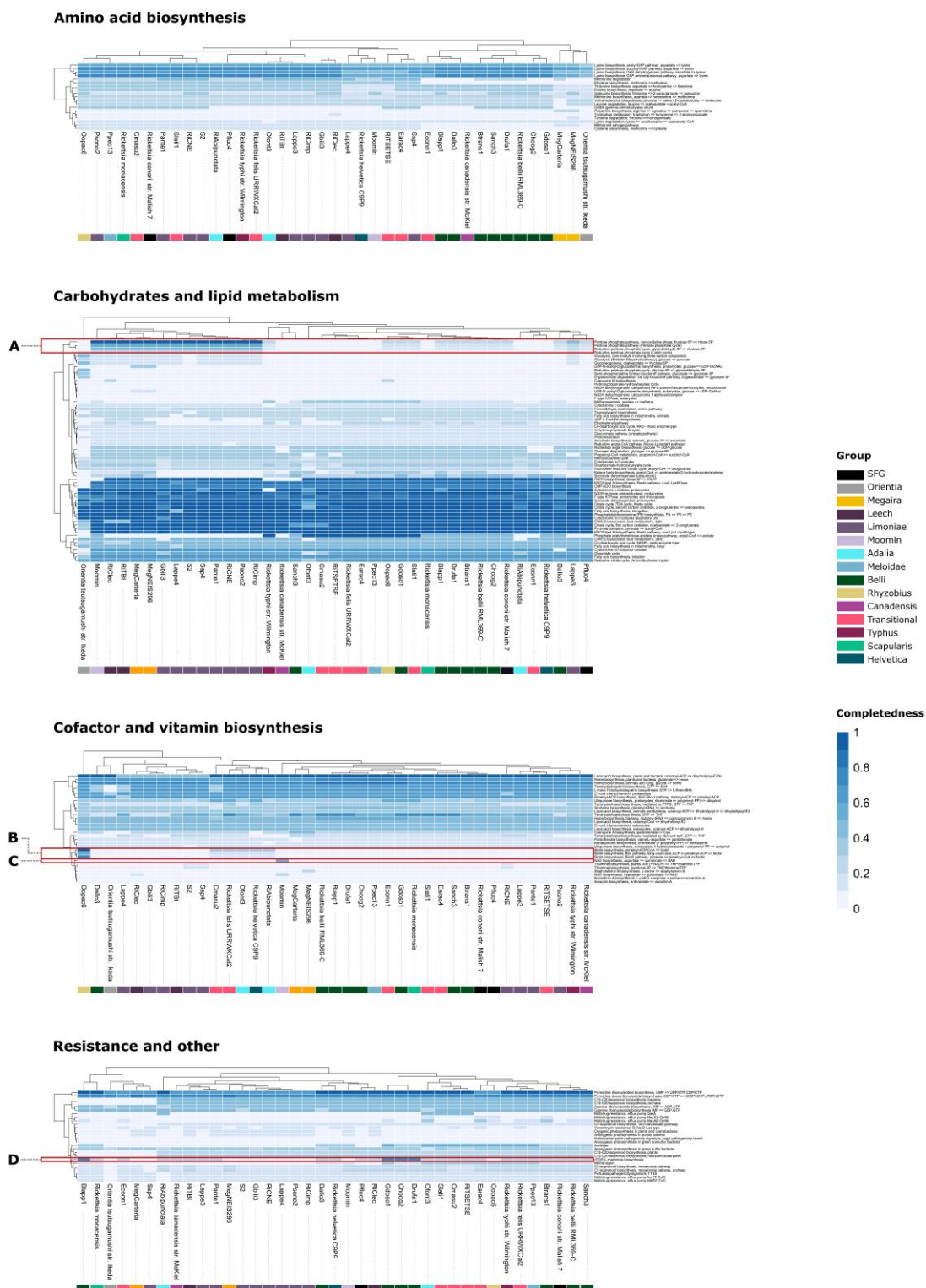


Figure 8. Heatmaps of predicted KEGG pathway completion estimated in Anvi'o 7, separated by function and produced with Pheatmap. Pathways of interest are highlighted: A) The pentose phosphate pathway only present in Torix and Ca. Megaira, B) the biotin pathway present only in the Rhyzobius Rickettsia Oopac6, C) NAD biosynthesis only present in Moomin Rickettsia, D) dTDP-L-rhamnose biosynthesis pathway in Gdoso1, Choog2, Drufa1, and Blapp1. SFG is Spotted Fever. A full resolution version can be found here: <https://doi.org/10.6084/m9.figshare.15081975>.



## Designation of *Ca. Tisiphia*

In all analyses, Torix consistently cluster away from the rest of *Rickettsia* as a sister taxon. Despite the relatively small number of Torix genomes, its within group diversity is greater than any divergence between previously described *Rickettsia* in any other group (figures 2, 3 and 5). Additionally, Torix shares characteristics with both *Ca. Megaira* and *Rickettsia*, but with many of its own unique features (figures 6 and 8). The distance of Torix from other *Rickettsia* and *Ca. Megaira* is confirmed in both the phylogenomic and metabolic function analyses to the extent that Torix should be separated from *Rickettsia* and assigned its own genus. This is supported by GTDB-Tk analysis which places all Torix genomes separate from *Rickettsia* (S1 <https://figshare.com/s/198c88c6e3ea5553192e>) alongside ANI percentage similarity scores less than 70% in all cases. To this end, we propose the name *Candidatus Tisiphia* after the fury Tisiphone, reflecting the genus *Ca. Megaira* being named after her sister Megaera.

## Conclusion

The bioinformatics approach has successfully extracted a substantial number of novel *Rickettsia* and *Ca. Megaira* genes from existing SRA data, including the first putative Rhyzobius *Rickettsia* and several *Ca. Tisiphia* (formerly Torix *Rickettsia*). Successful completion of two *Ca. Megaira* and two *Ca. Tisiphia* genomes provide solid reference points for the evolution of *Rickettsia* and its sister groups. From this, we can confirm the presence of a complete Pentose Phosphate Pathway in *Ca. Tisiphia* and *Ca. Megaira*, suggesting that this pathway was lost during *Rickettsia* evolution. We also describe the first Meloidae and Rhyzobius *Rickettsia* and show that Rhyzobius group *Rickettsia* has the potential to be a nutritional symbiont due to the presence of a complete biotin pathway. These new genomes provide a much-needed expansion of available data for symbiotic *Rickettsia* clades and clarification on the evolution of *Rickettsia* from *Ca. Megaira* and *Ca. Tisiphia*.

## Supporting information

All original genomes and raw readsets produced in this study can be accessed at Bioproject accession PRJNA763820 and all assemblies produced from previously published third party data can be accessed at Bioproject PRJNA767332.

Supplementary data and full resolution figures can be accessed on figshare here:

<https://doi.org/10.6084/m9.figshare.c.5518182.v1>

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 519 Culture Collection at the National Institute for Environmental Studies, Japan for use of the sample  
 520 *Carteria cerasiformis* NIES-425.

## 521 Contributions

522 Project concept: HRD, SS, JP and GH

523 Manuscript written by HRD, SS, JP and GDDH

524 SRA dive and metagenome assembly carried out by HRD with aid from SS.

525 Assembly of genome from SRA, pangenomics and phylogenomics carried out by HRD with advice from SS,  
 526 GH

527 Metabolic analysis carried out by HRD, JP and SS

528 Sequencing and assembly of bacteria from *Cimex lectularius* and *Culicoides impunctatus* genomes by SS  
 529 and JP.

530 Sequencing and assembly of symbionts from *Carteria* by SHB and SS, supervised by PC and GH.

531 Sequencing and construction of RiTSETSE carried out by FB as part of thesis work supervised by AD.

532 SP collected and sequenced staphylinid genomes that were released through NCBI by iridian genomics.

533 NW collected and sequenced the *Bryobia* Moomin strain and performed preliminary metagenomic analyses

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