Molecular evidence of *Chlamydia pecorum* and arthropod-associated Chlamydiaceae in an expanded range of marsupials

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The order *Chlamydiales* are biphasic intracellular bacterial pathogens infecting humans and domesticated animals. Wildlife infections have also been reported, with the most studied example being *Chlamydia pecorum* infections in the koala, an iconic Australian marsupial. In koalas, molecular evidence suggests that spill-over from *C. pecorum* infected livestock imported into Australia may have had a historical or contemporary role. Despite preliminary evidence that other native Australian marsupials also carry *C. pecorum*, their potential as reservoirs of this pathogen and other *Chlamydia*-related bacteria (CRBs) has been understudied. Mucosal epithelial samples collected from over 200 native Australian marsupials of different species and geographic regions across Australia were PCR screened for *Chlamydiales*. Previously described and genetically distinct *C. pecorum* genotypes and a range of 16S rRNA genotypes sharing similarity to different CRBs in the broader *Chlamydiales* order were present. One 16S rRNA *Chlamydiales* genotype recently described in Australian ticks that parasitise native Australian marsupials was also identified. This study provides further evidence that chlamydial infections are widespread in native fauna and that detailed investigations are required to understand the influence these infections have on host species conservation, but also whether infection spill-over plays a role in their epidemiology.

Evidence that infections by obligate intracellular bacterial pathogens in the order *Chlamydiales* are generally prevalent in wildlife is increasing, but their impact on the overall health of populations remains unclear. The koala (*Phascolarctos cinereus*), an iconic native Australian marsupial, is the most characterised example of the detrimental effects that chlamydial infections can have on the health of individuals and populations. Although asymptomatic infections are most common, *C. pecorum* infections in koalas can result in severe ocular, urinary tract and reproductive disease which, if left untreated, may progress to blindness, incontinence and infertility, respectively. The prevalence of koala *C. pecorum* infections varies from 10–90% in mainland populations, significantly contributing to the rapid decline of this threatened species in certain states of Australia. Koalas also carry unique *Chlamydia*-related bacteria (CRBs), although the role these organisms play in disease is yet to be defined.

The origin of *C. pecorum* infection in koalas is currently unclear. It is well known that domesticated livestock, such as sheep and cattle, are susceptible to *C. pecorum* infections. In Australia, genetically similar strains of livestock *C. pecorum* have been identified in koalas, in addition to koala-specific strains. This body of molecular evidence has led to the hypothesis that at least some *C. pecorum* infections in koalas may be the result of historical or contemporary pathogen spill-over from livestock sources, consistent with the significant deleterious

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Chlamydiales sequence results revealed the presence of one previously described and six novel genotypes belonging to the order Chlamydiales from multiple Chlamydiales organisms (data not shown). Nucleotide BLAST (nBLAST) analysis of the single sequence results that allowed for direct sequence analysis. The remaining samples resulted in mixed sequences, translating into a Chlamydiales ing of non-koala marsupial swabs from all regions sampled revealed 196/401 positive samples for DNA. MarsBar - HQ457465.1; Table 2). For the purpose of this investigation, this 16S rRNA sequence will be referred to as Chlamydia pecorum 16S rRNA genotype P787. In addition to this previously described 16S rRNA genotype, a second novel 16S rRNA Chlamydiapecorum sequence was also detected in three common brushtail possums, two from Tasmania and one from the East Coast (Table 2). This sequence, designated Chlamydia pecorum genotype 1, had 99% identity to the top BLAST hit C. pecorum strain P787 (CP004035.1) described above, yet differed from this strain by three SNPs. The phylogenetic positioning of these two genetically distinct sequences was confirmed following Bayesian phylogenetic reconstruction (Fig. 2).

Confirmation of Chlamydia pecorum in non-koala marsupials and molecular typing using a C. pecorum-specific MLST scheme. In order to confirm the detection of C. pecorum in non-koala marsupials and to gain insight into the genetic relationships of these strains to those from other hosts, amplification of seven highly conserved C. pecorum MLST housekeeping genes was performed on the eight C. pecorum positive non-koala marsupial samples, as previously described. Unfortunately, we were only successful in amplifying one of the seven C. pecorum MLST genes (gidA) from six of the non-koala marsupial samples. Analysis of these gidA gene sequence fragments and comparisons to previously described gidA sequences deposited in the C. pecorum PubMLST database revealed a 100% sequence match for the spotted tail quoll sample to C. pecorum gidA allele 31, an allele previously described in koalas. The squirrel glider sample was an exact match to C. pecorum gidA.

Results Chlamydiales PCR positivity in Australian marsupials. Pan-Chlamydiales 16S rRNA gene PCR screening of non-koala marsupial swabs from all regions sampled revealed 196/401 positive samples for Chlamydiales DNA, translating into a Chlamydiales positive rate of 48% for individual animals at one or more anatomical sites (111/231 animals: Table 1). A concurrent screening of 37 koala tests (57 swab samples) from the East Coast revealed 100% PCR positivity for Chlamydiales DNA.

To determine the identity of the Chlamydiales detected in the marsupials screened in this study, a pan-Chlamydiales PCR amplifying an 800 bp fragment of the 16S rRNA gene was targeted. PCR products for 96 non-koala and 43 koala samples were obtained and directly sequenced. Results revealed that 18.7% (18/96) and 58.1% (25/43) of the PCR products for non-koala marsupials and koalas, respectively, returned single sequence results that allowed for direct sequence analysis. The remaining samples resulted in mixed sequences, despite repeated PCR amplification and sequencing efforts, suggesting the majority of samples contained DNA from multiple Chlamydiales organisms (data not shown). Nucleotide BLAST (nBLAST) analysis of the single sequence results revealed the presence of one previously described and six novel genotypes belonging to the order Chlamydiales.

Chlamydiales prevalence and genotype diversity varied by geography and marsupial host species. Common brushtail possums harboured the most diverse range of Chlamydiales, carrying four of the seven genotypes, with the remaining marsupial species carrying two or less genotypes. Chlamydiales prevalence and diversity within non-koala marsupials was also noticeably larger at conjunctival sites (Table 1). PCR positivity of non-koala marsupials varied geographically, with Tasmania having the highest prevalence of 100%, followed by the East Coast and Northern Territory collection (26%; 41/154). In the minority of samples where we were able to resolve sequences, the distribution of 16S rRNA sequences across these different geographic locations varied. Only one genotype (Ca. Rhabdochlamydia porcellionis genotype 1) was detected in all three geographical locations (Fig. 1). The highest genotypic diversity was found in the East Coast collection where four of the seven different genotypes could be resolved. Three genotypes could each be resolved in the Tasmanian and Northern Territory cohorts (Tables 1 and 2).

Detection of C. pecorum 16S rRNA sequences in koalas and other marsupials. BLAST searching revealed the presence of two distinct C. pecorum 16S rRNA sequences in koalas and other marsupials in the East Coast sample set. The first of these partial sequences, detected in three common brushtail possums, one squirrel glider, one spotted tail quoll and 20 koalas was found to share 100% nucleotide identity to previously reported C. pecorum 16S rRNA sequences from cattle (P787–CP004035.1; E58–NR_102975.1) and koalas (MC/MarsBar - HQ457465.1; Table 2). For the purpose of this investigation, this 16S rRNA sequence will be referred to as Chlamydia pecorum 16S rRNA genotype P787. In addition to this previously described 16S rRNA genotype, a second novel 16S rRNA C. pecorum sequence was also detected in three common brushtail possums, two from Tasmania and one from the East Coast (Table 2). This sequence, designated Chlamydia pecorum genotype 1, had 99% identity to the top BLAST hit C. pecorum strain P787 (CP004035.1) described above, yet differed from this strain by three SNPs. The phylogenetic positioning of these two genetically distinct sequences was confirmed following Bayesian phylogenetic reconstruction (Fig. 2).
and Tasmania (Fig. 1) and, additionally, in five koalas from the East Coast sample collection. Possums, one short-eared possum and one northern brown bandicoot across the East Coast, Northern Territory DNA derived from ocular, nasal and cloacal swabs.*UGT includes cloaca and penile swabs; #Mixed sites are a pool of

Overview of marsupial species, number and location included in this study and their corresponding

Table 1. Detection of other CRB 16S rRNA sequences in Australian marsupials.

| Marsupial Species | # of Chlamydiaceae PCR Positive Individuals | Total # of Individuals | PCR positive sites | Ocular | UGT | Rectal | Mixed |
|-------------------|------------------------------------------|------------------------|------------------|--------|-----|--------|-------|
| **Northern Territory** | | | | | | | |
| Northern quoll (Dasyurus hallucatus) | 21 | 103 | 5/62 | 9/59 | 2/3 | 9/41 |
| Northern brown bandicoot (Isoodon macrourus) | 13 | 37 | n/a | n/a | n/a | 13/37 |
| Common brushtail possum (Trichosurus vulpecula) | 7 | 13 | n/a | n/a | n/a | 7/13 |
| Fawn antechinus (Antechinus bellus) | 0 | 1 | 0/1 | n/a | 0/1 | n/a |
| **Total number of marsupial species = 4** | 41 | 154 | 5/63 | 9/59 | 2/4 | 29/91 |
| **East Coast** | | | | | | | |
| Ring-tailed possum (Pseudocheirus peregrinus) | 10 | 13 | 4/13 | 9/13 | n/a | n/a |
| Common brushtail possum (Trichosurus vulpecula) | 24 | 26 | 22/28 | 20/25 | n/a | n/a |
| Short-eared possum (Trichosurus caninus) | 3 | 3 | 3/3 | 3/3 | n/a | n/a |
| Spotted tail quoll (Dasyurus maculatus) | 5 | 5 | 7/10 | 6/6 | 1/1 | n/a |
| Eastern grey kangaroo (Macropus giganteus) | 3 | 4 | 1/4 | 3/4 | n/a | n/a |
| Swamp wallaby (Macropus rufogriseus) | 1 | 1 | 1/1 | 1/1 | n/a | n/a |
| Long-nosed bandicoot (Perameles nasuta) | 1 | 1 | 1/1 | 1/1 | n/a | n/a |
| Squirrel glider (Petaurus norfolcensis) | 1 | 2 | 1/2 | 1/2 | n/a | n/a |
| **Total number of marsupial species = 8** | 48 | 55 | 40/62 | 44/55 | 1/1 | n/a |
| **Tasmania** | | | | | | | |
| Common brushtail possum (Trichosaurus vulpecula) | 22 | 22 | 44/44 | 22/22 | n/a | n/a |
| **Total number of marsupial species = 1** | 22 | 22 | 44/44 | 22/22 | n/a | n/a |
| **Totals** | | | | | | | |
| **Overall total number of marsupial species = 11** | 111 | 231 | 89 | 75 | 3 | 29 |

Table 1. Overview of marsupial species, number and location included in this study and their corresponding individual and tissue site PCR positivity. *UGT includes cloaca and penile swabs; *Mixed sites are a pool of DNA derived from ocular, nasal and cloacal swabs.

allele 22, which has previously been described in both sheep and cattle. However, the four common brushtail possum samples were all identified to have novel C. pecorum allele 12 sequences, each encompassing one to two unique SNPs (Fig. 3), when compared to gidA alleles derived from sheep and cattle.

**Detection of arthropod-associated Chlamydiaceae 16S rRNA sequences in Australian marsupials.** The most geographically distributed CRB genotype (Ca. Rhabdochlamydia porcellionis genotype 1) detected in this study was 100% identical to a recently described novel Ca. Rhabdochlamydia porcellionis genotype (KX774317) detected in marsupial-feeding tick species removed from koalas. This genotype is previously described as having 98% BLAST similarity to Ca. Rhabdochlamydia porcellionis (AY223862.1) and branches as a distinct lineage (Fig. 4). Ca. Rhabdochlamydia porcellionis genotype 1 was detected in two common brushtail possums, one short-eared possum and one northern brown bandicoot across the East Coast, Northern Territory and Tasmania (Fig. 1) and, additionally, in five koalas from the East Coast sample collection.

Beyond this previously described arthropod-associated Chlamydiaceae, a novel Ca. Rhabdochlamydia crassificans strain was also detected in two northern quolls exclusive to the Northern Territory (Fig. 1). Ca. Rhabdochlamydia crassificans strain 1 exhibited 99% BLAST identity to the closest match Ca. Rhabdochlamydia crassificans (AY928092.1) and branches as a distinct lineage (Fig. 4). Ca. Rhabdochlamydia crassificans genotype 1 was also detected in two common brushtail possums, exclusively to the Tasmanian region. This partial 16S rRNA genotype, Ca. Rhabdochlamydia porcellionis genotype 2, demonstrated 93% BLAST identity to Ca. Rhabdochlamydia porcellionis (HF933203.1) (Fig. 1, Table 2) as well as previously described novel Ca. Rhabdochlamydia genotypes identified in ticks.

**Detection of other CRB 16S rRNA sequences in Australian marsupials.** A partial 16S rRNA gene sequence potentially representing a novel genus within the family Chlamydiaceae was also detected in a northern quoll from the Northern Territory (Fig. 1). Fine-detailed phylogenetic analysis of this 16S rRNA sequence against other sequences belonging to species in the family Chlamydiaceae suggests that this novel sequence sits between the Chlamydia and Ca. Amphibiichlamydia genera (Fig. 2, Table 2). This genotype, Chlamydia-like genotype 1, exhibited 92% identity to Chlamydia pneumoniae (LN847058.1) 16S rRNA from a western barred bandicoot, as well as strains detected in koalas (CP001713.1), African clawed frogs (AF139200.1) and other western barred bandicoots (DQ444323.1).

To conclude the descriptions, we also detected an unusual partial 16S rRNA sequence in a sample from a spotted tail quoll from the East Coast, sharing 99% identity and two single nucleotide differences (Fig. 1) to
a previously deposited sequence from the novel chlamydial pathogen of fish, Ca. Similichlamydia latridicola (KC686679.1).

**Discussion**

Koalas, an iconic Australian marsupial, and globally distributed domesticated animals such as sheep and cattle, are well-known hosts of *C. pecorum*. In the current study, a pan-*Chlamydiales* order specific 16S rRNA gene PCR screening strategy provided molecular evidence that a range of other Australian marsupials also carry *C. pecorum*. A distinct range of novel bacteria from the order *Chlamydiales* were also found, some of which have been previously reported in ticks, including those found parasitising Australian marsupials.

In this study, *C. pecorum* 16S rRNA genotype P787 and *C. pecorum* genotype 1 accounted for 44.4% of the partial 16S rRNA *Chlamydiales* genotypes obtained, suggesting that *C. pecorum* is more abundant among non-koala marsupials than previously thought. In this study the greatest diversity and abundance of *C. pecorum* was found in common brushtail possums, suggesting that some marsupial host species apart from koalas, may be more susceptible to *C. pecorum* infection than others. Possums have previously been identified as a carrier of *C. pecorum*. While we need to treat the geographic distribution data with caution since we were only able to resolve sequences for the minority of PCR-positive samples we detected, it is notable that the majority of *C. pecorum* positive samples were detected in the East Coast. This region strongly overlaps with the host range of koalas that are endemically infected with *C. pecorum* and, indeed, the majority of animal hospitals that were used for sampling in this study commonly receive high numbers of koalas into care with chlamydiosis. Notably, animals sampled in the Northern Territory where koalas are not found, were free of *C. pecorum*. Inconsistent with this trend was the more surprising detection of *C. pecorum* positive samples from Tasmanian brushtail possums. Wild koalas do not inhabit Tasmania and, to the author’s knowledge, this report represents the first reported detection of chlamydial pathogens in Tasmanian fauna. Tasmania was last connected to mainland Australia during the Pleistocene (~25–8,000 years ago) suggesting *C. pecorum* has been present in this region for many millions of years.

While only preliminary in nature, a combination of partial 16S rRNA *Chlamydiales* and *C. pecorum gidA* gene sequencing suggests that the *C. pecorum* strains detected may be genetically divergent and may differ from those previously detected in koalas and Australian livestock. This was most apparent in the gidA sequences, where several previously described and novel gidA haplotypes were detected, suggesting the possibility of marsupial host specific strains and no favourable association with koalas or livestock based on the *C. pecorum gidA* data available to date. Unfortunately, we were not able to gain any further insight into the genetic diversity of these strains since we failed to amplify any of the remaining house-keeping genes utilised in previously described chlamydial MLST schemes (Jelocnik et al., 2013). While it is not entirely clear why this was the case, we expect it was due to the presence of only very low levels of *C. pecorum* DNA in these samples, combined with varying sensitivities of each of the individual PCR assays included in this scheme. Further confirmation of strain diversity and insight into the relationships of these strains is not possible without detailed genomics studies to compare these non-koala Australian marsupial *C. pecorum* strains to those from koalas, sheep, and cattle.

In the absence of such data, a partial answer regarding the origin and impact of these infections in Australian marsupials may lie in the disease presentation of *Chlamydia* positive animals in this study, as well as the ecological niche that these animals occupy. All 231 individual non-koala marsupials screened in this study were void of any classical signs of chlamydiosis despite several being *C. pecorum* positive at either ocular or urogenital sites. However, the comparison koalas infected with the same strain (based on partial 16S rRNA gene sequence), ranged from being asymptomatic to presenting with severe chlamydiosis (data not shown). Previously, ocular and urogenital

### Table 2. Abundance, identity, SNP differences and non-koala marsupial host information of *Chlamydiales* genotypes identified in this study. *Also detected in East Coast koalas.*

| Genotype                  | Closest BLAST match and % identity (# detected) | # of SNPs/16S rRNA length (bp) | Region, Marsupial and Anatomical Site detected |
|---------------------------|-------------------------------------------------|---------------------------------|------------------------------------------------|
| *Chlamydia pecorum* 16S rRNA genotype P787 | *Chlamydia pecorum* CP004035.1 100% (5)          | 0/744                           | East Coast, common brushtail possum, ocular; East Coast, common brushtail possum, ocular; East Coast, common brushtail possum, ocular; East Coast, common brushtail possum, ocular; East Coast, spotted tail quoll, urogenital. |
| *Chlamydia pecorum* GT1  | *Chlamydia pecorum* CP004035.1 99% (3)           | 3/600                           | East Coast, common brushtail possum, ocular; Tasmania, common brushtail possum, ocular; Tasmania, common brushtail possum, ocular. |
| Ca. Rhabdochlamydia porcellonis GT1 | Ca. Rhabdochlamydia porcellonis AY23862.1 99% (4) | 16/759                          | Tasmania, common brushtail possum, ocular; Tasmania, common brushtail possum, ocular; East Coast, short eared possum, ocular; Northern Territory, northern brown bandicoot, mixed. |
| Ca. Rhabdochlamydia porcellonis GT 2 | Ca. Rhabdochlamydia porcellonis HF933203.1 93% (2) | 30/357                          | Tasmania, common brushtail possum, urogenital; Tasmania, common brushtail possum, ocular. |
| Ca. Rhabdochlamydia crassificans GT1 | Ca. Rhabdochlamydia crassificans AY28092.1 99% (2) | 7/736                           | Northern Territory, northern quoll, mixed; Northern Territory, northern quoll, urogenital. |
| *Chlamydia-like GT1*     | *Chlamydia pneumoniae* LN847058.1 92% (1)       | 56/734                          | Northern Territory, northern quoll, rectal. |
| Ca. Similichlamydia latridicola GT1 | Ca. Similichlamydia latridicola KC686679.1 99% (1) | 2/745                           | East Coast, spotted tail quoll, ocular. |
Chlamydiosis was observed in approximately 20% of *C. pecorum* negative non-koala marsupials and *C. pecorum* positive test results could only be obtained from three quarters of animals showing classical signs of disease. It is also interesting to note that in this and the previous study, only arboreal marsupials such as possums, gliders (and koalas) were thus far found to be more often *C. pecorum* positive than ground-based marsupials, such as bandicoots which are more often *C. pneumoniae* positive. For example, in our study, possums sampled from Tasmania had 100% *Chlamydiales* PCR positivity with *C. pecorum* strains identified twice. The quolls and bandicoots, predominantly sampled from the Northern Territory, had 27% *Chlamydiales* PCR positivity and no *C. pecorum* was identified; the *Chlamydia*-like genotype 1 identified from the quoll was, in fact, more similar to *C. pneumoniae*. While clearly too early yet to draw formal conclusions, we speculate that (i) *C. pecorum* infections are primarily asymptomatic in Australian marsupials and that (ii) ecological niche may play a role in predisposing individuals to infection. As to why koalas experience severe chlamydiosis, further comparative genomics analysis of koala and non-koala marsupial *C. pecorum* may reveal undetected *C. pecorum* strain diversity or other pathogenic agents, including other *Chlamydiales* that may predispose koalas to disease in a way that other marsupials are not.

Beyond *C. pecorum*, novel *Chlamydiales* were highly abundant amongst non-koala marsupials and made up approximately 50% of the genotypes retrieved in this screen (Table 2, Fig. 1). "Uncultured *Chlamydiales*" were previously described as the most abundant taxa in non-koala marsupials, even in smaller screens, with the impact of these infections on animal health still unknown. Compared to the previous studies, the majority of the novel CRBs identified from marsupials in this study, however, are thought to primarily be limited to infections of arthropods. Indeed, the predominant CRB sequence detected was a partial *Ca. Rhabdochlamydia porcellionis* 16S rRNA gene sequence (Genotype 1) in a range of marsupials that was 100% identical to a genotype recently described in engorged marsupial feeding tick species *Ixodes tasmani* and *Ixodes holocyclus* removed from koalas. In addition to this genotype being present in four non-koala marsupials distributed across all three geographic regions sampled, five of the 37 koalas we screened were also positive for this partial 16S rRNA gene sequence. Curiously, this genotype was found in marsupials in all three regions included in this study, however, are thought to primarily be limited to infections of arthropods. Indeed, the predominant CRB sequence detected was a partial *Ca. Rhabdochlamydia porcellionis* 16S rRNA gene sequence (Genotype 1) in a range of marsupials that was 100% identical to a genotype recently described in engorged marsupial feeding tick species *Ixodes tasmani* and *Ixodes holocyclus* removed from koalas. In addition to this genotype being present in four non-koala marsupials distributed across all three geographic regions sampled, five of the 37 koalas we screened were also positive for this partial 16S rRNA gene sequence. Curiously, this genotype was found in marsupials in all three regions included in this study, however, we only found this genotype in ticks removed from koalas in Queensland, a sampling region included in the East Coast. Although the sample sizes were smaller, ticks removed from other marsupial species did not carry this genotype. If this observation is true, this is surprising given that *I. tasmani*, the common marsupial tick is the most widely distributed *Ixodes* species and is found throughout mainland Australian as well as Tasmania. The fact that marsupials and their parasitising arthropods share the same CRBs could implicate ticks as a route of transmission for these novel *Chlamydiales*. This suggestion was also made by a recent study investigating the occurrence of CRBs...
Figure 2. Phylogenetic relationships of Chlamydiaceae genotypes identified in Australian marsupials. Bayesian tree incorporating representative 16S rRNA sequences of each species of the genus Chlamydia from GenBank, as well as the three partial 16S rRNA genotypes identified in this study. Tree was built using 18 sequences of 588 bp under the HKY85 evolutionary model, posterior probability exceeding 0.75 is shown at internal nodes.

Figure 3. Phylogenetic relationships of C. pecorum gidA alleles identified in Australian marsupials. Bayesian tree incorporating six representative partial gidA alleles from PubMLST, as well as the six partial gidA alleles identified in this study. Marsupials identified with 16S rRNA as Chlamydia pecorum 16S rRNA genotype P787 are coloured green and those identified as Chlamydia pecorum genotype 1 are coloured blue. Tree was built using seven sequences of 474 bp under the HKY85 evolutionary model, posterior probability exceeding 0.75 is shown at internal nodes.
in ticks and skin biopsies from suspected tick bite individuals in Finland. Multiple sequences obtained from skin biopsies were found to be most closely related to CRBs detected from ticks species found in Europe, and in some cases, Finnish ticks. Further investigation into the pathogenesis and biology of novel CRBs is obviously warranted, especially given their presence in potential arthropod vectors.

Conclusions
This body of work further reinforces that chlamydial infections, including the previously described koala pathogen, C. pecorum, and other CRBs are widespread in Australian wildlife with some hosts more commonly infected than others. It appears that these infections do not cause traditional chlamydial diseases, certainly not like the debilitating chlamydial diseases that are observed in koalas, although longitudinal health studies would be required to ascertain this further. In the case of C. pecorum, what answers this holds about the origin and evolution of this pathogen in Australian marsupials is difficult to assess without further insight into the genetic differences that may exist between marsupial C. pecorum genotypes identified in this study with koala, sheep and cattle strains. Interestingly, the identification of a arthropod-associated Ca. Rhabdochlamydia porcellionis genotype in various marsupials, including koalas and koala-feeding tick species also suggests that arthropods are not the only host to this organism and further investigation into the pathogenesis and transmission of this and related CRBs is required.

Materials and Methods
Sampling. This study utilised a combination of (i) retrospective screening of wildlife samples collected as a part of other investigations and (ii) prospective sampling studies to investigate the prevalence of chlamydial infections in Australian marsupials from different geographic regions across Australia. Samples were obtained by swabbing the mucosal epithelia of conjunctiva, urogenital tract, rectal and penile tissues using Copan rayon tipped, aluminium-shafted applicator swabs (Interpath Services, Heidelberg West, Australia) for prospective sampling and both Copan rayon tipped, aluminium-shafted applicator swabs and Copan FLOQSwabs (Interpath Services) for retrospective sampling. Cumulatively, 401 swabs were opportunistically collected from 231 individual non-koala marsupials, comprised of 11 species across four Australian regions. The marsupial species, number of each marsupial and swab site are presented in Table 1. As a sample comparison set, 57 swabs from 37 koalas were opportunistically screened, as well.

All experimental protocols for the sampling of marsupials in this study were approved by the University of the Sunshine Coast Animal Ethics Committee (ANS1539). All methods were carried out in accordance with the...
Retrospective sampling. Northern Territory. A total of 217 DNA extracts from 154 individual non-koala marsupials spanning four species were included from the Northern Territory (Table 1). DNA extracts of 91 mixed mucosal epithelial sites (eye, nose, cloaca) from 91 individuals, representing four marsupial species were collected as part of a large-scale study of small mammal health in the Northern Territory Australia26. Another 126 DNA extracts of ocular, urogenital and rectal mucosal epithelial sites from 63 individuals, representing two marsupial species were collected as part of a large-scale study of a reintroduction program of the northern quoll.

Prospective Sampling. East Coast. One hundred and eighteen swabs were opportunistically taken from various mucosal epithelial sites (eye/s, urogenital tract, rectum and penis) of 55 non-koala marsupials representing eight marsupial species (Table 1) presenting to collaborating Australian veterinarian and wildlife care centres in New South Wales and Queensland. A total of 57 swabs from 37 individual koalas were taken from eye and urogenital mucosal epithelial sites of 9 and 28 koalas presenting to Wildlife Rehabilitation Centre at Eumundi, Queensland and Port Macquarie Koala Hospital, New South Wales, respectively.

Tasmania. Sixty six swabs were collected from the left and right eye and the urogenital tract mucosal epithelium of 22 deceased, freeze-thawed common brush-tail possums at the University of Tasmania (Table 1). The possums originated from commercial hunter-killed stock, collected from mixed agricultural-woodlands on private properties in Northern Tasmania.

DNA extraction. DNA extraction from swabs obtained from prospective sampling was performed using the QIAamp DNA mini kit (QIAGEN, Victoria, Australia). Swabs were vigorously shaken in TE (Tris-EDTA pH 8) buffer, followed by DNA extraction using the ‘DNA purification from tissues’ protocol as per the manufacturer’s instructions. DNA was stored at −20 °C until further use. DNA extractions performed on retrospective samples used the SIGMA Genelute bacterial DNA extraction kit in accordance with the manufacturer’s instructions.

Order specific 16S Chlamydiaceae PCR. To screen marsupials for Chlamydiaceae, an 800 bp fragment of the 16S rRNA gene partially covering the Chlamydiaceae signature sequence was amplified. PCR reactions consisted of 25 µl volumes containing 1.5 µl of each primer and 4 µl of template DNA, products were purified and sequenced following previously described methods11.

Chlamydia pecorum MLST PCR scheme. Samples with 16S rRNA sequences with a closest BLAST match to Chlamydia pecorum were subjected to C. pecorum MLST. The seven conserved C. pecorum specific housekeeping genes from the C. pecorum MLST scheme was amplified following previously described methods8.

Sequence analysis and phylogenetic parameters. 16S rRNA and gidA PCR products was purified using the Roche High Pure PCR Product Purification Kit (Roche, New South Wales, Australia) following the manufacturer’s instructions. Purified PCR products were sequenced by Macrogen Inc. (Seoul, Korea). Chromatograms of forward and reverse sequences were aligned in the Geneious R9.1.3 software package27. A consensus sequence was derived from each alignment and trimmed to the maximum length possible, gidA consensus sequences were trimmed to 474bp9. 16S rRNA gene consensus sequences, GenBank representatives of the closest BLAST match for each of the Chlamydiaceae, Ca. Rhabdochlamydiaceae and Ca. Parilichlamydiaceae families and an appropriate outgroup were then aligned in Geneious R9.1.3 using the MUSCLE plugin28 with default parameters. The alignments comprised of 18, 7 and 7 16S rDNA sequences trimmed to a length of 588, 368 and 701 bp, respectfully. For gidA, 12 sequences including consensus sequences, representatives of the closest PubMLST match19 from all possible host species and an appropriate outgroup were aligned in Geneious R9.1.3 using the MUSCLE plugin with default parameters. Bayesian phylogenies were constructed in Geneious R9.1.3 using the MrBayes plugin19 under the HKY85 substitution model. Run parameters included four Markov chain Monte Carlo (MCMC) chains with a million generations, sampled every 3,000 generations, and with the first 100,000 trees discarded as ‘burn-in’.

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Author Contributions
D.B. performed the majority of D.N.A. extractions, all molecular work, data analysis and was primarily responsible for preparation of the manuscript. W.H. and J.W. assisted with sample collection and D.N.A. extractions. M.J. assisted with molecular typing. A.G., A.R., S.F., S.C., J.C. and C.F. assisted with sample collection. A.P. and P.T. conceived the study and assisted with drafting of the manuscript. All authors have read and contributed to the manuscript.

Additional Information
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