

Tackling the metagenomics of aphids microbial symbionts : are conifer-feeding aphids looking for new tenants

OBJECTIFS

The association of aphids with *Buchnera aphidicola*, established 150 Mya, is one of the most persistent obligate symbioses known to date. However, recent studies have shown that in a few species of the genus *Cinara*, this relationship has evolved into a "ménage à trois", in which a former facultative endosymbiont, *Serratia symbiotica*, performs some of the functions usually fulfilled by *Buchnera*. The aim of the project is to elucidate the long-term evolution and maintenance of symbiotic associations in conifer-feeding aphids (Aphididae : Lachnidae : *Cinara*) and determine the contribution of bacterial symbionts in shaping the distribution patterns of this highly diversified aphid genus.

ACTIONS

Investigate the composition of the endosymbiotic bacteria of *Cinara* across a large sampling (400 specimens collected worldwide) using deep sequencing of 16S rRNA bacterial genes.

Optimize a protocol to sequence the genomes of *Cinara*'s obligate endosymbionts.

Establish correlations between endosymbiont community evolution and aphids host-plant shifts and environmental transitions using phylogenetic comparative methods.

RESULTATS

Most *Cinara* species host both *Serratia* and *Buchnera* but, in several clades, endosymbionts related to *Sodalis*, *Erwinia* or an unnamed member of the Enterobacteriaceae have replaced *Serratia*. These findings confirm that the species of this genus require a co-resident symbiont to complement *Buchnera* in its nutritional role. Results further demonstrate that co-symbiont replacement is not associated with the adaptation of aphids to new ecological conditions.

PERSPECTIVES

It shows that symbiont replacement has repeatedly occurred during the diversification of *Cinara*. The succession of essential symbionts does not result from the adaptation of the hosts to changing ecological conditions. It might therefore be driven by factors intrinsic to endosymbiosis such as rapid genome deterioration or the competitive displacement of symbionts providing benefits to the host.

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