

Use of molecular markers to identify cryptic species or complexes of sibling species in invasive or potentially invasive conifer pest insects

Géraldine Roux-Morabito, Alain Roques, Marie-Anne Auger-Rozenberg

▶ To cite this version:

Géraldine Roux-Morabito, Alain Roques, Marie-Anne Auger-Rozenberg. Use of molecular markers to identify cryptic species or complexes of sibling species in invasive or potentially invasive conifer pest insects. 23. International Congress of Entomology, Jul 2008, Durban, South Africa. 1 p. hal-02821372

HAL Id: hal-02821372 https://hal.inrae.fr/hal-02821372v1

Submitted on 6 Jun 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Use of molecular markers to identify cryptic species or complexes of sibling species in invasive or potentially invasive conifer pest insects

Géraldine Roux-Morabito, Alain Roques, Marie-Anne Auger-Rozenberg

INRA, Orleans, France

Introduction: Molecular genetic approaches are powerful tools for studying evolutionary aspects of biological invasions, especially for the identification of cryptic species or divergent lineages which can hinder a clear biological characterization of invasive species. Phytophagous insects show several examples of complexes of sibling species as well as successful invasion histories. We will present two case studies of phytophagous invaders. The first case will consider species which have been extensively transported globally for decades along with the seed trade (*Megastigmus* spp. seed chalcids), and the second pine pitch weevil of unknown origin, *Hylobitelus xiaoi*, which is presently in outbreak and is expanding in southern China. We will present data showing tests of the efficiency of molecular techniques for assessing whether the insect species correspond to cryptic native species or invasive species.

Material and methods: We use a phylogeographic approach by sequencing mitochondrial DNA genes (COI and/or COII). Individuals from different localities and from different host species were sampled in both the invaded and the presumed native range of the species.

Results and discussion: The study revealed misidentifications in *Megastigmus* species which have been previously recorded either as invasive or native species. It validated the use of molecular tools to detect cryptic species in this genus. These results also highlighted the importance of studying genetics in native populations to confirm or invalidate species identification only based on morphological traits. It should allow both a clear identification of native lineages and the tracking of the source population for expansion in *Hylobitelus*.