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Detection and quantitation of mixed infected samples with *Agrotis* **specific baculoviruses**

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Four baculoviruses, namely Agrotis nucleopolyhedrovirus segetum А (AgseNPV-A), A. segetum (Agse) NPV-B, A.ipsilon (Agip) NPV and A. segetum granulovirus (AgseGV) from the genera Alpha- and Betabaculovirus, respectively, are known to infect larvae of the lepidopteran pests A. segetum and A. ipsilon. The potential as biocontrol agents against Agrotis species benefits of two abilities, namely to crossinfect both pests A. segetum and A. ipsilon and to coinfect single larvae in mixed portions. In order to obtain a detailed understanding of mixed infections, especially between Agrotis spp. NPV and GV, and the amount of virus progeny produced, reproducible quantitative methods are necessary. This has been already set up with a SybrGreenbased RT-qPCR assay with specific primers binding in the core gene polyhedrin, or granulin respectively. By this method, mixed infections of AgseNPV-B and AgseGV in portions of their corresponding median lethal concentrations have been examined in experiments with neonate larvae of Agrotis segetum.

These findings revealed first hints to viral interactions between AgseNPV-B and AgseGV, as given by their potential virus progeny per larva. Ongoing experiments will consider investigations of mixed infections in cell culture studies with varieties of *Agrotis* specific baculoviruses and will be used for viral quantitation and microscopic picturing of mixed infections.

The SybrGreen-based RT-qPCR assay, however, requires a single RT-qPCR detection for each single virus and is thusly time- and sample-consuming. This lacks the ability for being used in experiments with more than two viruses, and/or in experiments with cell cultures of *Agrotis ipsilon*. An alternative assay with highly-specific *TaqMan* probes improves the quantification by a simultaneous RT-qPCR of up to four viruses in one reaction.