Paleozoic origin of insect large dsDNA viruses

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To understand how extant viruses interact with their hosts, we need a historical framework of their evolutionary association. Akin to retrovirus or hepadnavirus viral fossils present in eukaryotic genomes, baculoviruses are integrated in bracoonid wasp genomes and are transmitted by Mendelian inheritance. However, unlike viral genomic fossils, they have retained functional machineries homologous to those of large dsDNA viruses pathogenic to arthropods. Using a phylogenomic approach, we resolved the relationships between baculoviruses and their closest free relatives: baculoviruses and nudiviruses. The phylogeny showed that baculoviruses are nested within the nudivirus clade. Bracoviruses establish a bridge between the virus and animal worlds. Their inclusion in a virus phylogeny allowed us to relate free viruses to fossils. The ages of the wasps were used to calibrate the virus phylogeny. Bayesian analyses revealed that insect dsDNA viruses first evolved at ~310 Mya in the Paleozoic Era during the Carboniferous Period with the first insects. Furthermore, the virus diversification time frame during the Mesozoic Era appears linked to the diversification of insect orders; baculoviruses that infect larvae evolved at the same period as holometabolous insects. These results imply ancient coevolution by resource tracking between several insect dsDNA virus families and their hosts, dating back to 310 Mya.

Results

Insect dsDNA Virus Phylogeny. Analyses were conducted on 19 baculovirus genes for which homologs have been found in free insect dsDNA viruses (Table S1). Phylogenetic congruence analyses on unrooted trees showed that there was no conflicting signal between the different genes (SF Fig. S1, and Table S2). The phylogenomic analyses are therefore based on a concatenated alignment of 19 genes (7,238 aa), representing all of the data available for three integrated symbiotic baculoviruses, aligned to homologs of 10 free pathogenic viruses from the nudiviruses, the Baculoviridae, and the Hytrosaviridae as outgroups.

Bayesian inference and maximum likelihood phylogenetic analyses gave identical highly supported trees, differing only in branch lengths (Fig. 1). Interrelationships within and between the baculovirus and nudivirus clades are in accordance with, and better supported than, phylogenies based on diverse datasets (12, 17–19). Consistent with their chromosomal inheritance, the relationships within bracoviruses mirror the phylogeny of the wasps (20, 29). The most notable result from this tree is the paraphyly of the nudivirus clade, which includes the baculoviruses. Thus, this phylogeny demonstrates that the baculovirus lineage derives from two common ancestor and their hosts, dating back to 310 Mya. Previous phylogenetic studies clearly show that the nudiviruses and baculoviruses are sister groups (17, 18). The Hytrosaviridae are outside, although how distantly related remains uncertain. Whole-proteome clustering analysis placed them at the base of the DNA virus tree (18). However, based on gene content, they are thought to form a monophyletic group with the Baculoviridae, nudiviruses, and Nimaviridae (19). Within the Polydnaviridae, bracoviruses are associated with bracoonid wasps of the microgastroid complex (20). Parasitoid wasps have domesticated these viruses for transferring virulence genes (21–23), interfering with the immune response and development of their lepidopteran hosts (24, 25). The long-dated genuine viral nature of baculoviruses has recently been demonstrated with the discovery in wasp genomes of genes involved in bacovirus particle production (26–28). These genes, related to nudivirus and baculovirus core genes (27), establish a bridge between the phylogeny of free dsDNA viruses and geologic animal fossil data, thus allowing the exploration of deep evolutionary timescales.

Here, we performed phylogenomic analyses based on 19 bracovirus genes. Furthermore, we estimated the divergence times of the insect dsDNA virus families based on the age estimations of parasitoid wasp lineages (29). This work provides unique insights into the coevolutionary interactions between viruses and arthropods dating since the Paleozoic Era.

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within the nudiviruses and shares a common ancestor with the \textit{Heliothis zea} nudivirus (HzNV)--\textit{Penaeus monodon} nudivirus (PmNV) clade.

**Molecular Dating.** Molecular dating requires both a phylogenetic species tree and calibration anchors. The dual identity of bracoviruses as integrated wasp symbionts and viruses allowed us to relate geologic fossils to the insect dsDNA virus phylogeny. Bracoviruses are solely inherited vertically in the chromosomes of their associated braconid wasp. Each bracovirus and its wasps together act as a single genomic entity sharing the same evolutionary history. So their speciation events are contemporaneous (20). Thus, to calibrate the dsDNA virus tree, we used the age, phylogenetically estimated based on fossil amber calibration, of the wasps belonging to the bracovirus-bearing microgastrid complex (29). The time to the most recent common ancestor (TMRCA) of \textit{Chelonus inanitus} bracovirus (CiBV) and \textit{Cotesia congregata} bracovirus (CcBV) was set to 103.38 ± 4.41 Mya, corresponding to \textit{C. inanitus} (Cheloninae) and \textit{C. congregata} (Microgastrinae) lineage divergence times. Similarly, the TMRCA of CiBV and \textit{Toxoneuron nigriceps} bracovirus (TnBV) was set at ∼87 ± 5 Mya, the divergence times between the Microgastrinae and Cardiochilinae (Fig. S2). Because bracoviruses are now included in the insect dsDNA virus phylogeny, a Bayesian inference approach, using an autocorrelated lognormal relaxed clock and calibration priors based on the above age estimates, was performed to estimate the divergence times of the nudivirus and baculovirus dsDNA virus lineages.

Age estimates and 95% highest posterior density (HPD) intervals were obtained for each node (virus tree in Fig. 2; Table 1). The TMRCA of nudiviruses and baculoviruses (node 1) is estimated at ∼310 Mya (Carboniferous, late Paleozoic Era). The TMRCA of the nudivirus (node 2) and the baculovirus (node 4) lineages are estimated at ∼222 Mya (Triassic, early Mesozoic Era) and ∼178 Mya (Jurassic, middle Mesozoic Era), respectively. Furthermore, the results outline that the diversification of
the nudiavirus and baculovirus lineages stretches back to the Mesozoic Era and occurred primarily during the Jurassic and the Cretaceous Periods.

To relate the evolution of the insect dsDNA viruses to their hosts, the host species tree (Fig. 2) was drawn from the literature (SI Text). This tree shows that the insect and crustacean lineages associated with the dsDNA viruses of this study appeared in the middle Paleozoic Era and the diversification of insect orders occurred during the second half of the Paleozoic Era. Diversification within insect orders occurred mainly during the Mesozoic Era after the Permian crisis. Within the insect lineage, the appearance of holometabolous insects at ~350 Mya is also highlighted on the host tree.

Discussion

Previous studies on braconid wasps have established the monophyly of the lineage bearing bracoviruses, the microgastroid complex (20, 29), suggesting the integration of an ancestral virus in the genome of the most recent common ancestor of these wasps. Recently, homologous viral machineries, related to those of nudiuviruses, were discovered to produce distantly related viruses. The phylogenetic tree is also in accordance with recent phylogenomic analyses clearly show the paraphyly of the nudiuviruses with the inclusion of the bracoviruses (27). The first objective of this study was to resolve the phylogenetic relationships between the bracoviruses and their free virus relatives, the nudiuviruses and baculoviruses. Our phylogenomic analyses show the paraphyly of the nudiuviruses with the inclusion of the baculoviruses (Fig. 1), thus closing the demonstration that bracoviruses derive from nudiuviruses. The phylogenetic tree is also in accordance with recent studies indicating that the baculoviruses and nudiuviruses are closely related but form distinct lineages and as such should be classified in distinct virus families (16, 17, 19, 30).

Interestingly, bracoviruses share a common ancestor with the HzNV-PmNV clade (Fig. 1), for which the highest genome content similarities (six unique genes) have been found (27). HzNV is able to develop particular infection strategies such as having a tropism for gonadal cells (31) and the establishment of latent infections (32). Both of these mechanisms could have been instrumental for the evolution of bracoviruses. The nudiuviral ancestor of the bracoviruses could have used this particular infection strategy to establish a persistent latent infection in hymenopteran gonad cells, leading to its integration in the wasp chromosome.

The parasitoid wasps obligatorily depend on the symbiotic bracoviruses for their parasitic success. These two entities share the same genetic inheritance and undergo specialized speciation, which led us to assume that the TMRCAs of both wasp and bracovirus lineages are contemporaneous. The additional peculiarity of bracoviruses is their phylogenetic relationships with other free dsDNA viruses. This link between the animal and the virus worlds allowed the estimation of baculovirus and nudiuvirus divergence times for inferring their macroevolutionary history. Our dataset is suitable for molecular dating approaches on a large timescale because it fills two important criteria relative to low evolutionary rates. First, unlike RNA viruses, large dsDNA virus genomes have relatively slow evolutionary rates (33–35), approaching those of bacteria and lower unicellular eukaryotes (36). They are therefore appropriate subjects for molecular dating, as shown in the case of hepadnaviruses, for which long-term evolution has been inferred from integrated genomic fossils (9). Similarly, we used the bracovirus sequences integrated in the wasp genomes as calibration anchors. The second criterion relates to the dataset itself. Unlike for the hepadnavirus example, here the bracovirus sequences are still functionally active. The gene set is involved in the viral machinery, produces the viral particles, and is conserved in baculoviruses, nudiuviruses, and bracoviruses. The functional homology even remains observable at the level of viral particle morphology because bracovirus and nudiuvirus nucleocapsids are very similar (37). Overall, this observation points to relatively slow evolutionary rates on a large timescale, which we assume to be similar between the free viruses and the eukaryotic viral symbionts. Given these assumptions, we inferred the age estimates of the dsDNA virus phylogeny from the eukaryotic calibration points.

The results show that the baculoviruses and nudiuviruses are rooted at 310 Mya, deep in the Paleozoic Era in the Carboniferous Period, suggesting that the ancestors of these viruses had already infected the first insects, which appeared during the Devonian and Carboniferous Periods. Currently, this is the oldest age estimate to have been determined for any group of viruses, going far beyond the 19 Ma estimated for avian hepadnaviruses (DNA viruses) (9) or the 100 Ma established for mammalian retroviruses (RNA viruses) (5).

In the late Carboniferous Period, when the nudiuvirus and baculovirus families diversified, insects were also diversifying into new orders (Fig. 2). The phylogenies of both the baculovirus and nudiuvirus lineages are clearly not the mirror images of that of their hosts. This lack of cophylogeny shows that the diversification of both virus families and host orders occurred independently. On the macroevolutionary timescale, this finding suggests that these viruses may have colonized new insect hosts and also crustaceans many times during their evolution. However, the time frame of nudiuvirus and baculovirus diversification, starting at 222 Mya and 178 Mya respectively (Table 1), does not strongly support the notion of total independence of that of their hosts. Both virus families diversified during the Mesozoic Era, after the major Permian extinction, which is contemporary with the major diversification of insect orders and families (38).

Remarkably, at the late Carboniferous Period, the baculovirus lineage appeared shortly after the appearance of holometabolous insects, which are insects with larval stages undergoing complete metamorphosis (Fig. 2). These insects form a monophyletic clade, including (among others) the orders Diptera, Lepidoptera, Coleoptera, and Hymenoptera (39). Extant baculoviruses have only been characterized in holometabolous insect hosts from the orders Diptera, Hymenoptera, and Lepidoptera. Baculovirus phylogenetic studies have shown the ancient co-evolution between baculoviruses and insects of these three orders (12), although the lack of dating analyses did not allow inference of how long the relationship could have lasted. Our data suggest that the baculovirus ancestor was already specialized to infect the new susceptible host niche that represents the larval stage of holometabolous insects.

In contrast to baculoviruses, nudiuviruses appear to be a more generalist group of viruses. Although the pathological data are sparse, they are known to affect adult as well as larval stages with more varied infection strategies. The lack of environmentally resistant occlusion bodies in nudiuviruses has been proposed to
lead to the evolution of close association with their hosts, often involving viral persistence or latency (42). Nudviruses have a broad host spectrum (Fig. 2), encompassing many insect orders but also crustaceans like PmNV (17, 19). The derived phylogenetic position of PmNV may imply a large host shift between insects and crustaceans, which could date as far back as 125 Mya.

Relating the large-scale evolutionary history of insect dsDNA viruses to that of their hosts showed that the ancestors of the baculovirus and nudivirus lineages infected the already-evolved ancestors of their extant hosts, suggesting a coevolutionary scenario of colonization by resource tracking whereby the resources for these viruses are the insects (43). When new insect species evolved, providing in part an evolutionarily modified form of the ecological niche required for the virus replication cycle, virus individuals, adapted to the ancestral host, colonized the new host. This ecological speciation mechanism eventually produced new virus species. On the macroevolutionary timescale, with the extinction of many host and virus phylogenetic lineages, cophylogenetic divergence should not be expected. However, closely related virus species should infect closely related insects, as shown in baculoviruses, for which particular phylogenetic lineages are specialized to infect particular insect orders (12). When more data become available for nudviruses, similar lineage specializations are likely to be revealed. This coevolutionary process does not exclude large host shifts nor that dsDNA viruses evolve by coevolution with their hosts on the microevolutionary timescale. Indeed, most dsDNA viruses have a strong specificity toward their hosts and therefore have only a small probability of infecting a new host species during any short interval (44). The insect dsDNA virus phylogeny bears witness to the ancestral and in-timate coevolutionary relationships between viruses and their insect hosts and shows that the expansion of their host range has been driven by punctuated host-shift events through the ages. This result is in line with recent findings on the evolution of foamy retroviruses, which have coevolved with mammals for over 100 Ma (5), suggesting that common macroevolutionary scenarios affect viruses regardless of their replication modes or origin.

Our analysis is the largest timescale macroevolutionary scenario, supported with the oldest age estimates, reported for any extant virus families. Our findings broaden the general scope of paleovirology studies (45) and anchor the timeline of virus macro-evolution to very early stages of the history of viruses in the origin of cellular life. Ultimately, it should be possible to extend the molecular dating to the whole phylogeny of large dsDNA virus lineages involved in virus coevolution to test hypotheses on the role of viruses in the origin of life, the ancestry of their extant hosts, suggesting a coevolutionary scenario of colonization by resource tracking whereby the resources for these viruses are the insects (43). When new insect species evolved, providing in part an evolutionarily modified form of the ecological niche required for the virus replication cycle, virus individuals, adapted to the ancestral host, colonized the new host. This ecological speciation mechanism eventually produced new virus species. On the macroevolutionary timescale, with the extinction of many host and virus phylogenetic lineages, cophylogenetic divergence should not be expected. However, closely related virus species should infect closely related insects, as shown in baculoviruses, for which particular phylogenetic lineages are specialized to infect particular insect orders (12). When more data become available for nudviruses, similar lineage specializations are likely to be revealed. This coevolutionary process does not exclude large host shifts nor that dsDNA viruses evolve by coevolution with their hosts on the microevolutionary timescale. Indeed, most dsDNA viruses have a strong specificity toward their hosts and therefore have only a small probability of infecting a new host species during any short interval (44). The insect dsDNA virus phylogeny bears witness to the ancestral and intimate coevolutionary relationships between viruses and their insect hosts and shows that the expansion of their host range has been driven by punctuated host-shift events through the ages. This result is in line with recent findings on the evolution of foamy retroviruses, which have coevolved with mammals for over 100 Ma (5), suggesting that common macroevolutionary scenarios affect viruses regardless of their replication modes or origin.

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Materials and Methods

Sequence Data. Amino acid sequences of CiBV, CcBV, and TnBV nudivirus-related genes were retrieved from Bénzer et al. (27). BLASTp analyses (46) were used to identify homologs (e-value <0.01) from nudivirus and baculovirus genomes. When available, homologous sequences from hytrosaviruses (salivary gland hypertrophy virus) were used as outgroups.

Four species were used for the nudivirus family. Among these species, three nudivirus genome sequences have been fully sequenced: GbNV (47), Gryllus bimaculatus nudivirus (GbNV) (48), and Oryctes rhinoceros nudivirus (OrNV) (16) and one has been partially sequenced (PmNV). Four species representative of each genus of the baculovirus family were chosen: Culex nigripalpis nucleopolyhedrovirus (CnNPV) (49), Neodiprion lecontei nucleopolyhedrovirus (NeNPV) (50), Autographa california multiple nucleopolyhedrovirus (AcMNPV) (51), and Cydia pomonella granulovirus (CpGV) (52). Two species represent the hytrosavirus family: Musca domestica salivary gland hypertrophy virus (MsSGHV) (53) and Glossina pallidipes salivary gland hypertrophy virus (GsPShV) (54).

The dataset comprises 19 nudivirus-related genes: 13 belong to the 20 core genes conserved by all nudviruses and baculoviruses, 1 gene (odv-e66) is only found in some nudviruses and exclusively in baculoviruses infecting Lepidoptera, and 5 genes are found only in nudviruses (Table S1). Multiple amino acid alignments were obtained for each of the 19 genes with the program MAFFT (55), then they were trimmed with the program Gblocks (56) to only keep conserved domains before concatenation.

Phylogenetic Analyses. Phylogenetic analyses of the concatenated multiple amino acid alignment were performed by using both maximum likelihood and Bayesian inference. The appropriate substitution model and model parameters for each gene were selected using the Akaike information criterion (Blosum62 with the +I, +G, and +F parameters). Maximum likelihood analysis was performed with RAxML (58), and support for node in ML tree was obtained from 1,000 bootstrap iterations.

Mixed-model Bayesian phylogenetic analysis was performed with MrBayes (59). ProtTest and Bayesian information criteria were used to select appropriate substitution models and model parameters for each gene (Table S3). MrBayes analyses were run across four Monte Carlo Markov chains (MCMC) for 1 million generations, sampling every 500 generations. The consensus tree was obtained after a burn-in of 500 generations, and the value of average SD of split frequencies was used as a proof of stationarity if this value was under 0.01.

Molecular Dating Analysis. To test whether the combined dataset evolved in a clocklike fashion, we compared the InL of an unconstrained tree versus a tree constrained to evolve clockwise with the AAML program from the PAML package (60). Divergence times were then estimated by using a Bayesian inference approach implemented in BEAST (61). The analysis was performed on the combined dataset, preliminarily converted into codon-based sequence, and without outgroup because of the uncertainty concerning the sister-group relationships between the hytrosaviruses and the ingroup (16, 18). We assumed an uncorrelated lognormal relaxed clock and free mean substitution rates. MRMCAs of bracoonid wasp lineages bearing bracoviruses were used as priors to calibrate the insect dsDNA phylogeny under a normal distribution and a Yule process prior. Six independent MCMCs were run for 100 million generations sampling every 1,000 generations, using a burn-in of 1,000. The results of the six MCMCs were combined with the log-combiner program of the BEAST package, and age estimates were analyzed (effective sample size >200) with the Tracer program (62).

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