Adintoviruses: A Proposed Animal-Tropic Family of Midsize Eukaryotic Linear dsDNA (MELD) Viruses

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Abstract

Polintons (also known as Mavericks) were initially identified as a widespread class of eukaryotic transposons named for their hallmark type B DNA polymerase and retrovirus-like integrase genes. It has since been recognized that many polintons encode possible capsid proteins and viral genome-packaging ATPases similar to those of a diverse range of double-stranded DNA (dsDNA) viruses. This supports the inference that at least some polintons are actually viruses capable of cell-to-cell spread. At present, there are no polinton-associated capsid protein genes annotated in public sequence databases. To rectify this deficiency, we used a data-mining approach to investigate the distribution and gene content of polinton-like elements and related DNA viruses in animal genomic and metagenomic sequence datasets. The results define a discrete family-like clade of viruses with two genus-level divisions. We propose the family name Adintoviridae, connoting similarities to adenovirus virion proteins and the presence of a retrovirus-like integrase gene. Although adintovirus-class PolB sequences were detected in datasets for fungi and various unicellular eukaryotes, sequences resembling adintovirus virion proteins and accessory genes appear to be restricted to animals. Degraded adintovirus sequences are endogenized into the germlines of a wide range of animals, including humans.
Introduction

Analyses based on conserved protein structural features have increasingly revealed commonalities between families of eukaryotic viruses with double-stranded DNA (dsDNA) genomes. A current model places a loosely defined group known as polinton-like viruses at the center of a network of evolutionary relationships (Koonin, Dolja et al. 2015, Koonin, Krupovic et al. 2015). Polintons (also known as Mavericks) are defined by the presence of a type B DNA polymerase (PolB) and a retrovirus-like integrase gene. Although polintons were initially suspected to be transposons, the observation that many of them encode predicted virion proteins supports the later proposal that most elements initially designated as polinton transposons are actually integrated proviruses that may remain capable of infectious cell-to-cell spread (Krupovic, Bamford et al. 2014, Krupovic and Koonin 2015).

Adenoviruses, poxviruses, and baculoviruses are familiar groups of animal-tropic viruses that encode genes distantly similar to polinton PolB and virion proteins (Koonin, Dolja et al. 2015). An emerging group of viruses known as virophages, which are named for their ability to parasitize megaviruses that infect unicellular eukaryotes, also encode polinton-like PolB and virion protein genes as well as, in some cases, retrovirus-like integrase genes. (Duponchel and Fischer 2019).

Although polintons have been widely recognized in animal genomics and transcriptomics datasets (Krupovic, Bamford et al. 2014), the proposed capsid genes of these elements are not currently annotated in public sequence databases. This has led to confusion. For instance, a recent study detected two “Maverick transposons” in insect cell cultures but failed to annotate the capsid genes that identify them as likely viruses (Geisler 2018). In another example, a set of classic polinton PolB gene fragments detected in mouse fecal samples appear in GenBank with annotations incorrectly indicating that they are parvovirus structural proteins (Williams, Che et al. 2018). A primary goal of this study is to develop a coherent classification system for animal-tropic viruses with polinton-like genes and to facilitate further discovery by rendering annotated examples of these viruses searchable in public databases.

Results

Classification of animal-associated contigs with polinton-like PolB genes

TBLASTN searches using the inferred virion maturational protease (Adenain) of an arbitrarily chosen Parasteatoda spider contig (AOMJ02256338) identified hundreds of >10kb contigs of interest in NCBI’s whole genome shotgun (WGS) and transcriptome shotgun assembly (TSA) databases, as well as in de novo assemblies of various datasets of interest from the Sequence Read Archive (SRA). In animal datasets, a great majority of the larger adenain-bearing elements were found to encode either an archetypal polinton-like PolB (pfam03175) or a divergent PolB <30% identical to the pfam03175 type. Graphical illustrations of the gene detection and annotation process are shown in Supplementary Fig. S1. Both PolB classes were found encode a distinctive N-terminal domain with predicted structural similarity to the ovarian tumor
superfamily of ubiquitin-specific proteases (OTU). Adenovirus PolB sequences lack the OTU domain. In this study, we refer to the OTU-pfam03175 PolB class as Alpha and the more divergent OTU-PolB class as Beta. In RepBase https://www.girinst.org/, polinton groups 1, 2, 3, 4, and 9 each contain both Alpha and Beta PolB genes. Alpha PolB genes have previously been binned with hybrid virophages, ungrouped polinton-like viruses, and Polintons group 2, while Beta PolB genes have been binned with ungrouped polinton-like viruses, plant and fungal mitochondrial plasmids, and Polintons group 1 (Moriyama, Terasawa et al. 2008, Yutin, Raoult et al. 2013, Yutin, Kapitonov et al. 2015, Yutin, Shevchenko et al. 2015).

In BLASTP searches, Alpha PolB sequences give strong hits (E-values ~1e-60) for an emerging family of bipartite parovirus-like viruses called bidnaviruses (Krupovic and Koonin 2014, Koonin, Dolja et al. 2020). Use of the DELTA-BLAST algorithm (Boratyn, Schaffer et al. 2012) yields stronger hits (E-values <1e-100) for adenoviruses. Beta PolB sequences typically do not yield bidnavirus hits in BLASTP searches and instead give moderate hits (E-value ~1e-15) for the PolB proteins of megaviruses (e.g., faustoviruses and Klosneuvirus) as well as various bacteriophages (Fig. 1). Neither of the two PolB classes detects known virophage PolB sequences in BLASTP or DELTA-BLAST searches.

In addition to Adenain and PolB, nearly all >10 kb contigs from the WGS and TSA surveys encode a retrovirus-like integrase (protein family rve) as well as a protein similar to a group of FtsK/HerA-type nucleosome triphosphatases (FtsK) that are thought to mediate the packaging of viral genomes into virions (Iyer, Makarova et al. 2004).

Alignments of selected contigs back to parent read datasets showed that coverage depth fell to zero near the ends of some contigs. An example is shown graphically on page 6 of Supplementary Fig. S1. In some cases, such as a Mayetiola destructor (barley midge) read dataset, a single predominant apparently free-ended sequence could be assembled but the dataset also contained a range of lower-coverage variant reads near the termini, some of which extended into inverted terminal repeats (ITRs) and host genomic DNA sequences. The observation suggests that the integrase gene is functional and mediates integration events akin to those observed in virophages that encode rve integrases (Fischer and Hackl 2016).

Based on the similarities to adenoviruses and the presence of integrase and virus genome-packaging genes we suggest that this group of animal-associated elements could be referred to as “adintoviruses.” Graphical maps of reference Alpha and Beta adintoviruses are shown in Fig. 2. Maps of additional adintovirus genomes are shown in Supplementary Fig. S2.

HHpred searches confirmed the presence of ORFs with high-probability predicted structural similarity to the double-jellyroll major capsid proteins (hexons) and single-jellyroll vertex capsomers (pentons, also known as penton bases) of adenoviruses, virophages, megaviruses, or poxviruses (Supplementary Fig. S1). As expected, contigs with Beta PolB genes encode Hexon and Penton proteins that occupy discrete clusters that encompass the Terrapene Beta adintovirus cognates (Supplementary Fig. S3). Although most contigs with Alpha PolB genes encode Hexon and Penton proteins that cluster with the Mayetiola cognates, some Alpha PolB contigs unexpectedly encode virion proteins that are interspersed within the Terrapene cluster. Similar results were observed in analyses using traditional phylogenetic trees. The results suggest the
existence of distinct Alpha and Beta adintovirus lineages, but with some examples reflecting horizontal transfer of the virion protein module of the Beta PolB lineage into an Alpha PolB background. We have previously proposed a similar intra-family horizontal gene transfer scenario for some species of polyomaviruses (Buck, Van Doorslaer et al. 2016).

Other adintovirus genes

Adintoviruses encode three classes of proteins with predicted structures resembling known membrane-active proteins. A previously noted class (Yutin, Raoult et al. 2013) is similar to the phospholipase A2 (PLA2) domain of parvovirus VP1 virion proteins (see page 7 of Supplementary Fig S1). In parvoviruses, the domain is thought to be involved in membrane disruption during the infectious entry process. The PLA2-like genes, which are characteristic of Mayetiola-class (Alpha) virion protein modules, include a C-terminal domain similar to adenovirus virion core precursor protein ten (pX). We suggest the gene name PLA2X.

Beta adintoviruses, as well as Alpha PolB adintoviruses with Terrapene-class (Beta) virion protein modules, encode homologs of the C-terminal regulatory domain of gasdermins, a group of pore-forming proteins that serve as executioners in pyroptosis (a form of inflammatory programmed cell death) (Dubois, Sorgeloos et al. 2019). Like PLA2X, adintovirus gasdermin homologs typically encode a pX-like domain near the C-terminus. Apparent homologs of a membrane-active spider venom protein known as cupiennin were also observed in Beta-class virion protein modules. The pairing of hallmark Beta-class virion accessory genes (GasderminX, Cupiennin) with a subset of Alpha PolB adintoviruses (Supplementary Fig. S2) supports the hypothesis that some adintovirus species arose through horizontal gene transfer between the Alpha and Beta adintovirus lineages.

Some classes of predicted protein sequences were conserved among adintoviruses but did not show clear hits for known proteins in BLASTP or HHpred searches. We assigned these groups of adintovirus-conserved proteins of unknown function numbered “Adintoc” names.

Small DNA tumor viruses (adenoviruses, polyomaviruses, and papillomaviruses (Pipas 2019) encode proteins harboring conserved LXCXE motifs that that are known to engage cellular retioblastoma (Rb) and related tumor suppressor proteins (de Souza, Iyer et al. 2010). Adenovirus E1A, papillomavirus E7, polyomavirus LT, and parvovirus NS3 oncoproteins typically encode the Rb-binding motif just upstream of a consensus casein kinase 2 acceptor motif ((ST)XX(DE)). Some oncogenes, such as E1A, encode an additional conserved region ((DEN)(LMV)XX(LM)(FY)), referred to as CR1, that binds the groove containing the A and B cyclin folds within the Rb pocket domain (Pipas 1992, Gouw, Michael et al. 2018). In general, these predicted Rb-interacting motifs are adjacent to potential zinc- or iron-sulfur-binding motifs (typically, paired CXXC). Open reading frames encoding combinations of these short linear motifs were observed in adintovirus contigs. We refer to these predicted proteins, which typically occupy a region upstream of the PolB gene, as “Oncoid” genes, connoting their similarities to the known oncogenes of small DNA tumor viruses. Adintovirus homologs of anti-apoptotic proteins, such as Bcl2 and IAP, were also observed (Supplementary Figs. S1 and S2).
Distribution of adintovirus-like PolB sequences in eukaryotic WGS datasets

The conserved catalytic core PolB sequences of either the *Mayetiola* barley midge Alpha adintovirus or *Terrapene* box turtle Beta adintovirus were used separately as baits in TBLASTN searches of WGS databases for eukaryotes. Retrieved protein sequences were trimmed to 80% similarity and subjected to clustering with an alignment score threshold of 60 (Shannon, Markiel et al. 2003, Li and Godzik 2006, Huang, Niu et al. 2010, Fu, Niu et al. 2012, Zallot, Oberg et al. 2018). The clustering segregated away Beta adintovirus-like PolB sequences encoded by plant and fungal mitochondria (e.g., EU365401, AF061244). The filtered sequences were subjected to phylogenetic analyses (Fig. 3).

Two complete Alpha adintovirus-like contigs (NKLS02000104, NKLS02001728) were observed in assemblies of a PacBio-based WGS survey of bovine lung tissue. Sequences outside the inferred proviral inverted terminal repeats (ITRs) in the two sequences were highly diverse and mostly unidentifiable, but in a few reads the extra-proviral host sequences showed BLASTN similarity to genomic DNA sequences of various beetles, including *Tribolium castaneum* (a flour beetle that commonly infests cattle feed). Furthermore, the *Bos* lung-associated PolB sequences occupy phylogenetic clades comprised of insect-associated PolB sequences (Fig. 3). These observations suggest that the two Alpha adintovirus sequences in the bovine datasets are insect-derived environmental contaminants, rather than mammal-tropic viruses. Similarly, several Beta adintovirus-like contigs (e.g., AANG04004209) found in a housecat oral swab sample show close phylogenetic affinity for adintovirus sequences observed in salmon WGS datasets. In another example, integrated adintoviruses found in a genomic dataset for olive trees (*Olea europaea*) showed insect-like sequences outside the inferred ITRs and showed phylogenetic affinity with PolB sequences from insect WGS datasets. Other adintovirus-like sequences found in plant datasets resembled adintovirus PolB sequences associated with nematode datasets. It thus appears that adintovirus sequences in some datasets are derived from environmental sources, as opposed to a productive infection of the organism that was the target of the sequencing effort.

Although there are examples of apparent environmental contamination, most adintovirus sequences form discrete clades that recapitulate the phylogeny of the host organisms that were the subjects of the WGS surveys. For example, a distinct clade of Alpha adintovirus PolB sequences was observed in datasets for multiple related species of venomous snakes. Several distinct clades of Beta adintoviruses were observed in datasets for amphibians and reptiles, including the well-populated clade that houses the exemplar *Terrapene* adintovirus. The exemplar *Mayetiola* adintovirus likewise occupies a clade exclusively populated by sequences found in insect WGS datasets.

TBLASTN searches against *Terrapene* box turtle Beta adintovirus PolB and Hexon protein sequences both yielded weak hits (E-value ~1e-05) for a locus on human chromosome 7. An adintovirus GasderminX sequence was also detected at the locus. Alignments to *Terrapene* adintovirus protein sequences were used to assign pseudogene annotations (Fig. 4). The detected element is a highly disrupted endogenized Beta adintovirus. Homologous nucleotide sequences were detected in the genomes of primates, rodents, shrews, afrotherians, and xenarthrans but not in datasets for ungulates, carnivoires, bats, marsupials, or prototherians. Endogenized adintovirus sequences observed in amphibian and reptile genomes do not share recognizable nucleotide
similarity with placental mammal-endogenized adintovirus sequences. It is unclear whether a single adintovirus endogenization event affected an early placental mammal and the endogenized virus was then lost in non-shrew Laurasiatherians or whether multiple distinct endogenization events occurred in separate placental mammal lineages. Identification of extant examples of placental mammal adintoviruses could help resolve this question.

**Viruses with adinto-like genes in non-animal eukaryote datasets**

Eukaryotic viruses with midsize (10-50 kb) linear dsDNA genomes show a remarkable degree of genomic modularity (Koonin, Dolja et al. 2015, Yutin, Kapitonov et al. 2015, Yutin, Shevchenko et al. 2015). The apparently promiscuous horizontal gene transfer and lack of any single defining gene for these viruses makes the group taxonomically challenging. We propose the collective acronym MELD (midsize eukaryotic linear dsDNA) virus for the dizzyingly polyphyletic category. The name, which would encompass adenoviruses and adintoviruses, is intended to fill a gap between other operationally defined umbrella groups, such as circular Rep-encoding single-stranded DNA (CRESS) viruses, small DNA tumor viruses, nucleocytoplasmic large DNA viruses, and megaviruses.

Datasets for *Blastocystis hominis* (a diatom-related unicellular eukaryote that commonly inhabits the human gut) contain MELD virus sequences that unite Alpha adintovirus-like PolB and integrase genes with inferred virion proteins whose primary sequences are not recognizably similar to known virion proteins (Fig. 5). Gene identities for the *Blastocystis* virus were inferred based on HHpred results. Comparable MELD viruses were confirmed in rumen metagenomic datasets for sheep (Yutin, Kapitonov et al. 2015) and cattle, as well as in WGS datasets for green algae and fungi. In phylogenetic analyses, the PolB sequences of these viruses occupy long branches that are distant from animal-associated PolB clades (Fig. 3).

Contigs encoding Alpha adintovirus-like PolB and integrase genes were found in metagenomics datasets for bioreactor-cultured human feces, human urine samples, and human oral swab samples (Santiago-Rodriguez, Ly et al. 2015). This group of closely related sequences was only detected in datasets from a single laboratory and not in other human metagenomics surveys. Divergent variants of predicted proteins from the feces-associated virus were found in contigs from datasets for *Cyanophora paradoxa*, a species of glaucophyte algae (e.g., QPMI01000557), suggesting that the human feces-associated adintovirus-like sequences were derived from an environmental source.

Six MELD virus genomes assembled from a single *Powellomyces* SRA dataset unite sequences resembling adenovirus vertex fiber proteins with either a Beta adintovirus-like PolB (Fig. 5) or a surprising variety of non-PolB DNA replicases (Fig. 6). MELD virus genomes encoding genes similar to Alpha adintovirus virion proteins (E-values ~1e-7 to 1e-21) were assembled from datasets for *Capsaspora owczarzaki*, *Monosiga brevicollis*, and *Trichoplax H2* (unicellular eukaryotes that are thought to be closely related to animals). Aside from the abovementioned insect- and nematode-associated adintovirus sequences found in datasets for plants, adintovirus-like virion protein sequences were not detected in datasets for other non-animal eukaryotes. *Capsaspora* MELD virus 1 and the *Trichoplax* MELD virus both encode superfamily 1 helicase (S1H) genes instead of a PolB gene. Various megaviruses and bacteriophages encode similar
S1H genes, as does a MELD virus observed in *Physarum polycephalum* slime mold and *Powellomyces* MELD virus 4. Full-length S1H replicase genes of this class were not detected in animal WGS datasets, with the exception of seemingly endogenized degraded virus-like contigs in datasets for several coral and jellyfish species and a helitron-like element found in *Branchiostoma* lancelets (e.g., RDEB01009762, ABEP02037959).

In WGS searches for sequences resembling human adenovirus type 5 PolB, we did not detect any contigs resembling full-length viruses in non-animal datasets. The searches did reveal the complete ITR-bounded genome of a typical mastadenovirus in a dataset for *Dipodomys ordii* (a type of kangaroo rat) as well as apparently complete MELD viruses in datasets for *Hydra oligactis* (brown hydra) and *Anelosimus studiosus* (a type of tangle-web spider). Like known adenoviruses, the *Hydra* and *Anelosimus* MELD viruses do not encode integrase genes and their PolB genes do not encode detectable OTU domains.

**Proposal of a new virus group, bidnaparvoviruses**

BLASTP searches using Alpha adintovirus PolB sequences return high-likelihood matches (E-values <1e-80) for the PolB genes of viruses in the recently established family *Bidnaviridae* (Krupovic and Koonin 2014, Koonin, Dolja et al. 2020) (Fig. 1). Like adintoviruses, bidnavirus PolB genes encode an N-terminal OTU domain. Contigs with bidnavirus-like PolB genes were detected in datasets for the gut contents of African termites (*Cubitermes ugandensis*), dog (*Canis lupus familiaris*) feces, the silk glands of a false wolf spider (*Tengella perfuga*), and Tasmanian devil (*Sarcophilus harrisii*) feces (Fig. 7). The dog feces PolB sequence is 53% similar to a “structural protein” fragment of Fresh Meadows “densovirus” 3 reported in mouse (*Mus musculus*) feces (AWB14611) (Williams, Che et al. 2018).

In known bidnaviruses, the termini of each of the two genome segments have matching nucleotide sequences. To search for second segments, we probed assemblies for examples of other contigs with termini similar to the ITRs of the initially observed bidnavirus-like contigs. The datasets were also searched for possible second segments with sequences similar to the proteins of known bidnaviruses. Second segments were not detected, suggesting that the five new bidna-like viruses are monopartite. We propose that the apparently monopartite viruses be referred to as “bidnaparvoviruses,” highlighting their shared similarities with the two previously established families.

Searches for examples of parvovirus NS1-like sequences did not reveal clear examples outside of multicellular animal datasets. A marginal exception was a group of sequences found in datasets for *Abeoforma whisleri* and *Ichthyophonus hoferi*, two unicellular eukaryotes that are thought to be closely related to multicellular animals. The observations suggest an early-animal origin for parvoviruses that may have involved acquisition of genes from Alpha adintoviruses.

**Discussion**

We have identified a coherent family-like grouping of animal viruses that we call adintoviruses, connoting their hallmark adenovirus-like virion protein genes and retrovirus-like integrase genes.
Adintovirus sequences are detectable either as apparently free linear DNA molecules or as endogenized integrants in WGS datasets representing all eumetazoan phyla. Although sequences resembling the PolB proteins of Alpha and Beta adintoviruses can also be found in datasets for non-animal eukaryotes, the sequences of adintovirus virion proteins appear to be restricted to animals.

We imagine that the related Alpha and Beta adintovirus-like lineages might have infected early eukaryotes and the two lineages gradually co-evolved with major divisions of eukaryotes, including multicellular animals. In this model, the sequences of the virion protein genes presumably evolved more rapidly than the conserved catalytic core of PolB, resulting in distinctive mutually unrecognizable virion protein sequences specific to each major division of eukaryotes. The model suggests that adenoviruses could be thought of as a related sister lineage that also arose in or before the first animals. Although adenoviruses are currently only known to infect vertebrates, the idea that the lineage long predates the emergence of vertebrates is consistent with our identification of distantly adenovirus-like sequences in hydra and spider datasets (Figs. 5 and 6).

In non-animal eukaryote datasets, adintovirus-like PolB sequences can be found in a wide range of sequence contexts, ranging from elements with no obvious virion proteins to the genomes of megaviruses. Conversely, it appears that adintovirus-like PolB genes can readily be replaced with other types of DNA replicase genes (Fig. 6). This presumably reflects the previously proposed rampant horizontal gene transfer among virus lineages that infect unicellular eukaryotes (Koonin, Dolja et al. 2015). Although similar horizontal gene transfer events appear to have occurred between various animal-tropic virus families, including adintoviruses and paroviruses (Fig. 7), adomaviruses and polyomaviruses (Mizutani, Sayama et al. 2011, Dill, Camus et al. 2018, Welch, Tiszta et al. 2020) and papillomaviruses and polyomaviruses (Woolford, Rector et al. 2007), each of these cases appears to represent single ancient event. It may be that the evolution of distinct tissues and organs - or the development of cell-mediated immunity in multicellular animals - placed limits on the likelihood that different virus lineages can co-infect a single cell and productively recombine. From this view, the distinctive gene combinations seen in adintoviruses and adenoviruses might simply be bottlenecked examples of the much larger range of gene combinations observed in MELD viruses of unicellular eukaryotes (Yutin, Kapitonov et al. 2015, Yutin, Shevchenko et al. 2015).

It has generally been assumed that the functionally similar oncogenes found in adenoviruses, papillomaviruses, paroviruses, and polyomaviruses arose through convergent evolution or through horizontal gene transfer between virus families (de Souza, Iyer et al. 2010). Although small DNA tumor virus oncogenes show low overall sequence similarity, they can be roughly defined based on the presence of short linear motifs. Many adintoviruses encode candidate “Oncoid” proteins with these motifs. Bombyx silkworm bidnavirus NS3, which we have designated as a candidate Oncoid (Fig. 7), has previously been shown to be similar to a baculovirus protein of unknown function (Krupovic and Koonin 2014). We note that many of the proposed baculovirus homologs (e.g., YP_009506034) also share potential zinc-coordinating cysteine residues as well as a C-terminal LXCXE/CK2 site, qualifying the baculovirus proteins as candidate Oncoids as well. Surprisingly, candidate Oncoids were also observed in MELD viruses of unicellular eukaryotes (Figs. 5 and 6). The predicted Oncoid2 gene of Capsaspora
protist MELD virus 2 detects polyomavirus Large T oncogenes in DELTA-BLAST searches (E-value 4e-16). It is interesting to imagine that oncogenes in a broad range of animal DNA viruses might share an ancestry that pre-dates the emergence of multicellular animals.

Adintoviruses encode a number of accessory genes that appear to be homologs of membrane-active proteins found in animal venom. These include bee and snake venom PLA2 and melittin, as well as a spider venom protein called cupiennin. Interestingly, venom PLA2 and melittin (which shows similarity to adenovirus protein X in HHpred searches) act in concert (Vogt, Patzer et al. 1970), suggesting the speculative hypothesis that these venom genes might have arisen from a captured viral PLA2X-like gene.

In unpublished work, our group used a standard baculovirus-based expression system (ThermoFisher) to generate a virus-like particle (VLP) vaccine against BK polyomavirus (BKV) (Peretti, Geoghegan et al. 2018). The project provided an inadvertent natural experiment. Recombinant baculoviruses were generated in Sf9 cells and bulk protein expression was performed using the Trichoplusia ni cell line High Five. BKV VLPs were purified according to previously reported methods (Cardone, Moyer et al. 2014) involving ultracentrifugation through density gradients, nuclease digestion, and size exclusion chromatography. Deep sequencing of DNA extracted from the purified VLP preparation shows high-depth coverage of Spodoptera adintovirus genomes alongside incomplete patchy coverage of endogenized Trichoplusia-specific homologs of the two Spodoptera viruses (Supplementary File S6). It appears that Sf9-derived adintoviruses infected the High Five cells and this led to the production of adintovirus virions that co-purified with the recombinant BKV VLPs. The results suggest that standard insect cell cultures could serve as a laboratory model for productive adintovirus infection.

A Beta adintovirus was detected in transcriptomic and WGS datasets for Mexican blind tetra cavefish (Astyanax mexicanus). Adintovirus transcripts were most abundant in head, kidney, and intestine samples and least abundant in muscle and whole embryo samples (Supplementary Table S2). Analysis of the WGS dataset showed that adintovirus DNA reads outnumbered reads for a single copy host gene (gamma tubulin, NW_019172896) by a factor of 25. At both an RNA and DNA level the Astyanax sequence showed a high degree of uniformity, suggesting a clonal infection. In contrast, pet store samples of a different species of tetra, Gymnocorymbus ternetzi (SRR2040422), showed such a complex range of adintovirus sequence variants that assembly of contigs representing complete viral genomes was challenging. These observations suggest that tetrars might serve as a tractable laboratory model for adintovirus infection.

Adintoviruses have a number of features that could make them useful as recombinant gene transfer vectors. Their genome size is substantially larger than commonly used retroviral and parvoviral vectors. In contrast to adenovirus- and baculovirus-based vector systems, adintovirus genomes are small enough to be manipulated entirely in the setting of standard plasmids. An intriguing feature of the adintovirus integrase gene is the presence of a predicted chromodomain that, in LTR retrotransposons, is believed to influence integration site specificity (Kordis 2005). This could theoretically offer an advantage over retroviral vectors, which show little integration site specificity. Another potential practical use for adintoviruses might be as biocontrol agents for pest organisms, such as Mayetiola destructor barley midges or chytrid fungi that parasitize amphibians.
An important implication of this study is that there may be additional unappreciated families of animal viruses hiding in plain sight in sequence databases. Adintoviruses may have been relatively easy quarry because they are able to integrate into host genomes, such that they are detectable in WGS datasets of randomly sampled animals that did not happen to be suffering from an active infection. In contrast to the hundreds of adintovirus-like contigs detected in our initial WGS survey, focused searches for adenoviruses (which do not encode integrases) detected only a single complete adenovirus genome. For future discovery efforts, it will be important to develop higher throughput methods using sensitive structure-guided searches to identify divergent new examples of viral hallmark genes in sequence datasets representing many individuals, including subjects suffering from disease. A key goal will be to understand which combinations of genes tend to co-occupy single contigs. Recently reported bioinformatics pipelines, such as Cenote-Taker (Tisza, Pastrana et al. 2020) and Mash Screen (Ondov, Starrett et al. 2019), should be useful for these purposes. Deposition of annotated viral genome sequences into publicly searchable databases will be critical for further expanding our understanding of the eukaryotic virome.

Materials and Methods

Detection and analysis of viral sequences

Adomavirus LO8 (Adenain) sequences were initially used for TBLASTN searches of the NCBI TSA and WGS databases. The relationship between adomavirus and adintovirus virion proteins is the subject of a separate manuscript (Welch, Tisza et al. 2020). The Adenain sequences of Nephila orb-weaver spider contig (GFKT014647032) or a Parasteatoda spider contig (AOMJ02256338) were arbitrarily chosen for further TBLASTN searches of eukaryotic datasets in TSA and WGS databases. Adenain-bearing contigs 4-50 kb in length were further searched (using CLC Genomics Workbench) for BLASTP-detectable PolB homologs. Contigs were inspected for the presence of nearly overlapping arrays of large (>100 AA) open reading frames. Contigs with inverted repeats flanking the ORF cluster were favored, but this was not a strict sorting criterion. Selected contigs of interest were initially annotated using DELTA-BLAST searches of GenBank nr or HHpred analyses of single or aligned protein sequences against PDB_mmCIF70, COG_KOG, Pfam-A, and NCBI_CD databases (Altschul, Madden et al. 1997, Altschul, Wootton et al. 2005, Soding 2005, Hildebrand, Remmert et al. 2009, Gerlt, Bouvier et al. 2015, Meier and Soding 2015, Zimmermann, Stephens et al. 2017). Protein sequences were extracted from the contigs using getORF http://bioinfo.nhri.org.tw/cgi-bin/emboss/getorf (Rice, Longden et al. 2000). Extracted protein sequences were clustered using EFI-EST https://efi.igb.illinois.edu/efi-est/ (Gerlt, Bouvier et al. 2015, Zallot, Oberg et al. 2018) and displayed using Cytoscape v3.7.1 (Shannon, Markiel et al. 2003). Multiple sequence alignments were constructed using MAFFT https://toolkit.tuebingen.mpg.de/#/tools/mafft. Contigs were annotated using Cenote-Taker (Tisza, Pastrana et al. 2020) with an iteratively refined library of conserved adintovirus protein sequences. Compiled protein sequences are provided as a zipped
set of fasta-format text files in Supplementary File S2. Maps were drawn using MacVector 17 software. Phylogenetic analyses were performed using MAFFT 7
https://mafft.cbrc.jp/alignment/server/ (Kuraku, Zmasek et al. 2013, Katoh, Rozewicki et al. 2019) and displayed using FigTree 1.4.4 http://tree.bio.ed.ac.uk/software/figtree/.

Selected contigs for which SRA datasets were available were subjected to reference-guided re-assembly using Megahit 1.2.9 (Li, Liu et al. 2015, Li, Luo et al. 2016) and/or the map reads to reference function of CLC Genomics Workbench. Annotated maps were submitted to GenBank as third party annotation assemblies (TPA_asm). Graphical examples of the annotation process are depicted in Supplementary Fig. S1.

**Data Availability**

GenBank accession numbers for sequences deposited in association with this study are:
BK010888 BK010889 BK010890 BK010893 BK010894 BK010998 BK010999 BK011000 BK011001 BK011002 BK011003 BK011004 BK011005 BK011006 BK011007 BK011008 BK011009 BK011010 BK011011 BK011022 BK011023 BK011024 BK011025 BK011026 BK012042 BK012043 BK012044 BK012045 BK012046 BK012047 BK012048 BK012049 BK012050 BK012051 BK012052 BK012053 BK012054 BK012055 BK012056 BK012057 BK012058 BK012059 BK012060 BK012061 BK012062 BK012063 BK012064 BK012084 BK012085 BK012086.

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Altschul, S. F., T. L. Madden, A. A. Schaffer, J. Zhang, Z. Zhang, W. Miller and D. J. Lipman (1997). "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs." Nucleic Acids Res 25(17): 3389-3402.

Altschul, S. F., J. C. Wootten, E. M. Gertz, R. Agarwala, A. Morgulis, A. A. Schaffer and Y. K. Yu (2005). "Protein database searches using compositionally adjusted substitution matrices." FEBS J 272(20): 5101-5109.

Boratyn, G. M., A. A. Schaffer, R. Agarwala, S. F. Altschul, D. J. Lipman and T. L. Madden (2012). "Domain enhanced lookup time accelerated BLAST." Biol Direct 7: 12.

Buck, C. B., K. Van Doorslaer, A. Peretti, E. M. Geoghegan, M. J. Tisza, P. An, J. P. Katz, J. M. Pipas, A. A. McBride, A. C. Camus, A. J. McDermott, J. A. Dill, E. Delwart, T. F. Ng, K. Farkas, C. Austin, S. Kraberger, W. Davison, D. V. Pastrana and A. Varsani (2016). "The Ancient Evolutionary History of Polyomaviruses." PLoS Pathog 12(4): e1005574.

Cardone, G., A. L. Moyer, N. Cheng, C. D. Thompson, I. Dvoretzky, D. R. Lowy, J. T. Schiller, A. C. Steven, C. B. Buck and B. L. Trus (2014). "Maturation of the human papillomavirus 16 capsid." MBio 5(4): e01104-01114.

de Souza, R. F., L. M. Iyer and L. Aravind (2010). "Diversity and evolution of chromatin proteins encoded by DNA viruses." Biochim Biophys Acta 1799(3-4): 302-318.

Dill, J. A., A. C. Camus, J. H. Leary and T. F. F. Ng (2018). "Microscopic and Molecular Evidence of the First Elasmobranch Adomavirus, the Cause of Skin Disease in a Giant Guitarfish, Rhynchobatus djiddensis." MBio 9(3).

Dubois, H., F. Sorgeloos, S. T. Sarvestani, L. Martens, Y. Saeyes, J. M. Mackenzie, M. Lamkanfi, G. van Loo, I. Goodfellow and A. Wullaert (2019). "Nlrp3 inflammasome activation and Gasdermin D-driven pyroptosis are immunopathogenic upon gastrointestinal norovirus infection." PLoS Pathog 15(4): e1007709.

Duponchel, S. and M. G. Fischer (2019). "Viva lavidaviruses! Five features of virophages that parasitize giant DNA viruses." PLoS Pathog 15(3): e1007592.

Fischer, M. G. and T. Hackl (2016). "Host genome integration and giant virus-induced reactivation of the virophage mavirus." Nature 540(7632): 288-291.

Fu, L., B. Niu, Z. Zhu, S. Wu and W. Li (2012). "CD-HIT: accelerated for clustering the next-generation sequencing data." Bioinformatics 28(23): 3150-3152.

Geisler, C. (2018). "A new approach for detecting adventitious viruses shows Sf-rhabdovirus-negative Sf-RVN cells are suitable for safe biologicals production." BMC Biotechnol 18(1): 8.

Gerlt, J. A., J. T. Bouvier, D. B. Davidson, H. J. Imker, B. Sadkhin, B. R. Slater and K. L. Whalen (2015). "Enzyme Function Initiative-Enzyme Similarity Tool (EFI-EST): A web tool for generating protein sequence similarity networks." Biochim Biophys Acta 1854(8): 1019-1037.

Gouw, M., S. Michael, H. Samano-Sanchez, M. Kumar, A. Zeke, B. Lang, B. Bely, L. B. Chemes, N. E. Davey, Z. Deng, F. Diella, C. M. Gurfth, A. K. Huber, S. Kleinsorg, L. S. Schlegel, N. Palopoli, K. V. Roey, B. Altenberg, A. Remenyi, H. Dinkel and T. J. Gibson (2018). "The eukaryotic linear motif resource - 2018 update." Nucleic Acids Res 46(D1): D428-D434.

Hildebrand, A., M. Remmert, A. Biegert and J. Soding (2009). "Fast and accurate automatic structure prediction with HHpred." Proteins 77 Suppl 9: 128-132.
Huang, Y., B. Niu, Y. Gao, L. Fu and W. Li (2010). "CD-HIT Suite: a web server for clustering and comparing biological sequences." Bioinformatics 26(5): 680-682.

Iyer, L. M., K. S. Makarova, E. V. Koonin and L. Aravind (2004). "Comparative genomics of the FtsK-HerA superfamily of pumping ATPases: implications for the origins of chromosome segregation, cell division and viral capsid packaging." Nucleic Acids Res 32(17): 5260-5279.

Katoh, K., J. Rozewicki and K. D. Yamada (2019). "MAFFT online service: multiple sequence alignment, interactive sequence choice and visualization." Brief Bioinform 20(4): 1160-1166.

Koonin, E. V., V. V. Dolja and M. Krupovic (2015). "Origins and evolution of viruses of eukaryotes: The ultimate modularity." Virology 479-480: 2-25.

Koonin, E. V., V. V. Dolja, M. Krupovic, A. Varsani, Y. I. Wolf, N. Yutin, F. M. Zerbini and J. H. Kuhn (2020). "Global Organization and Proposed Megataxonomy of the Virus World." Microbiol Mol Biol Rev 84(2).

Koonin, E. V., M. Krupovic and N. Yutin (2015). "Evolution of double-stranded DNA viruses of eukaryotes: from bacteriophages to transposons to giant viruses." Ann NY Acad Sci 1341: 10-24.

Kordis, D. (2005). "A genomic perspective on the chromodomain-containing retrotransposons: Chromoviruses." Gene 347(2): 161-173.

Krupovic, M., D. H. Bamford and E. V. Koonin (2014). "Conservation of major and minor jelly-roll capsid proteins in Polinton (Maverick) transposons suggests that they are bona fide viruses." Biol Direct 9: 6.

Krupovic, M. and E. V. Koonin (2014). "Evolution of eukaryotic single-stranded DNA viruses of the Bidnaviridae family from genes of four other groups of widely different viruses." Sci Rep 4: 5347.

Krupovic, M. and E. V. Koonin (2015). "Polintons: a hotbed of eukaryotic virus, transposon and plasmid evolution." Nat Rev Microbiol 13(2): 105-115.

Kuraku, S., C. M. Zmasek, O. Nishimura and K. Katoh (2013). "aLeaves facilitates on-demand exploration of metazoan gene family trees on MAFFT sequence alignment server with enhanced interactivity." Nucleic Acids Res 41(Web Server issue): W22-28.

Li, D., C. M. Liu, R. Luo, K. Sadakane and T. W. Lam (2015). "MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph." Bioinformatics 31(10): 1674-1676.

Li, D., R. Luo, C. M. Liu, C. M. Leung, H. F. Ting, K. Sadakane, H. Yamashita and T. W. Lam (2016). "MEGAHIT v1.0: A fast and scalable metagenome assembler driven by advanced methodologies and community practices." Methods 102: 3-11.

Li, W. and A. Godzik (2006). "Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences." Bioinformatics 22(13): 1658-1659.

Meier, A. and J. Soding (2015). "Automatic Prediction of Protein 3D Structures by Probabilistic Multi-template Homology Modeling." PLoS Comput Biol 11(10): e1004343.

Mizutani, T., Y. Sayama, A. Nakanishi, H. Ochiai, K. Sakai, K. Wakabayashi, N. Tanaka, E. Miura, M. Oba, I. Kurane, M. Saijo, S. Morikawa and S. Ono (2011). "Novel DNA virus isolated from samples showing endothelial cell necrosis in the Japanese eel, Anguilla japonica." Virology 412(1): 179-187.

Moriyama, T., K. Terasawa, M. Fujiwara and N. Sato (2008). "Purification and characterization of organellar DNA polymerases in the red alga Cyanidioschyzon merolae." FEBS J 275(11): 2899-2918.
Ondov, B. D., G. J. Starrett, A. Sappington, A. Kostic, S. Koren, C. B. Buck and A. M. Phillippy (2019). "Mash Screen: high-throughput sequence containment estimation for genome discovery." Genome Biol 20(1): 232.

Peretti, A., E. M. Geoghegan, D. V. Pastrana, S. Smola, P. Feld, M. Sauter, S. Lohse, M. Ramesh, E. S. Lim, D. Wang, C. Borgogna, P. C. FitzGerald, V. Bliskovsky, G. J. Starrett, E. K. Law, R. S. Harris, J. K. Killian, J. Zhu, M. Pineda, P. S. Meltzer, R. Boldorini, M. Gariglio and C. B. Buck (2018). "Characterization of BK Polyomaviruses from Kidney Transplant Recipients Suggests a Role for APOBEC3 in Driving In-Host Virus Evolution." Cell Host Microbe 23(5): 628-635 e627.

Pipas, J. M. (1992). "Common and unique features of T antigens encoded by the polyomavirus group." J Virol 66(7): 276-277.

Pipas, J. M. (2019). "DNA Tumor Viruses and Their Contributions to Molecular Biology." J Virol 93(9).

Rice, P., I. Longden and A. Bleasby (2000). "EMBOSS: the European Molecular Biology Open Software Suite." Trends Genet 16(6): 276-277.

Santiago-Rodriguez, T. M., M. Ly, M. C. Daigneault, I. H. Brown, J. A. McDonald, N. Bonilla, E. A. Vercoe and D. T. Pride (2015). "Chemostat culture systems support diverse bacteriophage communities from human feces." Microbiome 3: 58.

Shannon, P., A. Markiel, O. Ozier, N. S. Baliga, J. T. Wang, D. Ramage, N. Amin, B. Schwikowski and T. Ideker (2003). "Cytoscape: a software environment for integrated models of biomolecular interaction networks." Genome Res 13(11): 2498-2504.

Soding, J. (2005). "Protein homology detection by HMM-HMM comparison." Bioinformatics 21(7): 951-960.

Tisza, M. J., D. V. Pastrana, N. L. Welch, B. Stewart, A. Peretti, G. J. Starrett, Y. S. Pang, S. R. Krishnamurthy, P. A. Pesavento, D. H. McDermott, P. M. Murphy, J. L. Whited, B. Miller, J. Brenchley, S. P. Rosshart, B. Rehermann, J. Doorbar, B. A. Ta'al a, O. Pletnikova, J. C. Troncoso, S. M. Resnick, B. Bolduc, M. B. Sullivan, A. Varsani, A. M. Segall and C. B. Buck (2020). "Discovery of several thousand highly diverse circular DNA viruses." Elife 9: e51971.

Vogt, W., P. Patzer, L. Lege, H. D. Oldagi and G. Wille (1970). "Synergism between phospholipase A and various peptides and SH-reagents in causing haemolysis." Naunyn Schmiedebergs Arch Pharmakol 265(5): 442-454.

Welch, N. L., M. J. Tisza, G. J. Starrett, A. K. Belford, D. V. Pastrana, Y.-Y. S. Pang, J. T. Schiller, P. An, P. G. Cantalupo, J. M. Pipas, S. Koda, K. Subramaniam, T. B. Waltzek, C. Bian, Q. Shi, Z. Ruan, T. F. F. Ng and C. B. Buck (2020). "Identification of Adomavirus Virion Proteins." bioRxiv: 341131.

Williams, S. H., X. Che, J. A. Garcia, J. D. Klena, B. Lee, D. Muller, W. Ulrich, R. M. Corrigan, S. Nichol, K. Jain and W. I. Lipkin (2018). "Viral Diversity of House Mice in New York City." MBio 9(2): e0135417.

Wooldford, L., A. Rector, M. Van Ranst, A. Ducki, M. D. Bennett, P. K. Nicholls, K. S. Warren, R. A. Swan, G. E. Wilcox and A. J. O'Hara (2007). "A novel virus detected in papillomas and carcinomas of the endangered western barred bandicoot (Perameles bougainville) exhibits genomic features of both the Papillomaviridae and Polyomaviridae." J Virol 81(24): 13280-13290.

Yutin, N., V. V. Kapitonov and E. V. Koonin (2015). "A new family of hybrid virophages from an animal gut metagenome." Biol Direct 10: 19.
Yutin, N., D. Raoult and E. V. Koonin (2013). "Virophages, polintons, and transpovirons: a complex evolutionary network of diverse selfish genetic elements with different reproduction strategies." Virol J 10: 158.

Yutin, N., S. Shevchenko, V. Kapitonov, M. Krupovic and E. V. Koonin (2015). "A novel group of diverse Polinton-like viruses discovered by metagenome analysis." BMC Biol 13: 95.

Zallot, R., N. O. Oberg and J. A. Gerlt (2018). "Democratized' genomic enzymology web tools for functional assignment." Curr Opin Chem Biol 47: 77-85.

Zimmermann, L., A. Stephens, S. Z. Nam, D. Rau, J. Kübler, M. Lozajic, F. Gabler, J. Söding, A. N. Lupas and V. Alva (2017). "A Completely Reimplemented MPI Bioinformatics Toolkit with a New HHpred Server at its Core." J Mol Biol S0022-2836(17): 30587-30589.
Figure Legends

Figure 1: PolB BLASTP relationships. PolB protein sequences were subjected to all-against-all sequence similarity network analysis with a BLASTP E-value cutoff of 1e-06. Supplementary File S1 presents an interactive version of Fig. 1 that can be viewed using Cytoscape software https://cytoscape.org. Supplementary File S2 contains sequence compilations for PolB and other proteins.

Figure 2: Genome maps of two representative adintoviruses. See main text for descriptions of gene names. Supplementary Table S1 presents accession numbers and full Linnaean designations of animal hosts.

Figure 3: Phylogenetic trees comprised of WGS hits for Alpha or Beta adintovirus PolB sequences (left and right panels, respectively). Hits from insect datasets are colored yellow, hits from tetrapod datasets are red and fungus-associated hits are blue. All other types of eukaryotes are represented by black lines. Annotated branches show two Alpha adintovirus sequences associated with bovine (Bos) lung samples clustering with adintovirus sequences from insect datasets, suggesting an environmental insect source. In contrast, exemplar Mayetiola and Terrapene adintoviruses cluster with sequences found in other insect or terrestrial vertebrate datasets, respectively. Similarly, adintovirus PolB-like sequences from Powellomyces and Rhizophagus fungi cluster with sequences from other types of fungi. Supplementary Files S3 and S4 are Nexus-format versions of the figure that can be viewed interactively using FigTree software http://tree.bio.ed.ac.uk/software/figtree/

Figure 4: An endogenized Beta adintovirus relic found on human chromosome 7. Degraded pseudogenes interrupted by nonsense and frameshift mutations were reconstructed based on alignments to the protein sequences of Terrapene box turtle adintovirus. Question marks indicate that the reconstructed gene does not yield hits in BLAST searches of GenBank’s viruses taxon. Tentative gene assignments are based on synteny with the Terrapene adintovirus. The reconstructed Hexon, GasderminX, and PolB protein sequences yield DELTA-BLAST hits with E-values of 1e-21, 4e-05, and 3e-25, respectively. Supplementary File S5 presents an annotated GenBank-format nucleotide map of the human chromosome 7 endogenized adintovirus.

Figure 5: MELD viruses (and related elements) with adintovirus-like PolB genes. Greek letters indicate genes similar to adintoviruses in BLASTP or DELTA-BLAST searches. “Ad-” indicates similarity to adenovirus sequences. Abbreviations: Fiber, predicted structural or primary sequence similarity to bacteriophage tail fibers or coiled-coil proteins; Lectin, predicted structural similarity to galactose-binding domains; SET, sequence similarity to the S-adenosyl methionine-binding pocket of cellular histone-lysine methyltransferases; TRAF, predicted structural similarity to TNF receptor-associated factor 3; UCH, predicted structural similarity to ubiquitin C-terminal hydrolases; IVa2, sequence similarity to adenovirus IVa2 viral genome-packaging ATPases; Dcm, predicted structural similarity to cytosine DNA methyltransferases; TBP, similar to TATA binding proteins; Matrix/Capsid/NC/Pro, similarities to retroviral Gag and retropepsin; S3H, poxvirus D5-like superfamily 3 helicase.

Figure 6: MELD viruses with other replicases. Greek letters indicate genes with sequences similar to Alpha or Beta adintoviruses in BLASTP searches. Sequences similar to adenoviruses are marked with “Ad-.” Yellow flags represent predicted tRNA genes. Abbreviations: dUTPase, similar to poxvirus deoxy-UTP diphosphatases; Fiber, similarity to bacteriophage tail fibers or other coiled-coil proteins;
Tapemeasure, similarity to phage tail tape measure proteins; S1H, RecD/Pif1-like superfamily 1 helicase; YRec, homolog of phage tyrosine recombinases DnaA-like, sequence distantly similar to DnaA and DnaB-like helicases; VI, similar to adenovirus virion core protein six; PolA, DNA polymerase family A (Pfam:00476); S3H superfamily 3 helicase similar to those observed in virophages and megaviruses; S2H, superfamily 2 helicase similar to DEAD-box helicase of Yellowstone Lake virophage 7 (YP_009177696); SLATT, homolog of host SMODS and SLOG-associating 2TM effector domain proteins; VLTF, homolog of mimivirus VLTF3-like transcription factor. See main text for information about other gene names.

**Figure 7: Genome maps for non-animal parvoviruses, bidnaviruses, and proposed bidnaparvoviruses.** Abbreviations: GIIM, similarity to group II intron maturases; tailspike, similarity to bacteriophage short tail fibers.
Figure 1: PolB BLASTP relationships. PolB protein sequences were subjected to all-against-all sequence similarity network analysis with a BLASTP E-value cutoff of 1e-06. Supplementary File S1 presents an interactive version of Fig. 1 that can be viewed using Cytoscape software https://cytoscape.org. Supplementary File S2 contains sequence compilations for PolB and other proteins.
Figure 2: Genome maps of two representative adintoviruses. See main text for descriptions of gene names. Supplementary Table S1 presents accession numbers and full Linnaean designations of animal hosts.
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| Virus name                                                                 | Replicase(s)     |
|---------------------------------------------------------------------------|------------------|
| Astyanax tetra cavefish adintovirus (RNA)                                | Beta PolB        |
| Bos-associated insect adintovirus 1                                      | Alpha PolB       |
| Bos-associated insect adintovirus 2                                      | Alpha PolB       |
| Branchiostoma lancelet adintovirus                                       | Alpha PolB       |
| Caenorhabditis roundworm adintovirus                                     | Alpha PolB       |
| Corynactis coral adintovirus (RNA)                                       | Alpha PolB       |
| Cotetes braconid wasp bracovirus circle 31                                | Alpha PolB       |
| Crocodylus saltwater crocodile endogenized adintovirus                   | Beta PolB        |
| Diachasma braconid wasp adintovirus                                      | Alpha PolB       |
| Eucidaris sea urchin adintovirus                                         | Alpha PolB       |
| Human endogenized adintovirus relic                                      | Beta PolB        |
| Hydra adintovirus                                                        | Alpha PolB       |
| Ladona dragonfly adintovirus                                             | Alpha PolB       |
| Larimichthys croaker adintovirus                                         | Beta PolB        |
| Mayetiola barley midge adintovirus                                       | Alpha PolB       |
| Megastigmus wasp adintovirus (RNA)                                       | Alpha PolB       |
| Mytilus Mediterranea mussel adintovirus                                  | Alpha PolB       |
| Nephila orb-weaver spider adintovirus (RNA)                              | Alpha PolB       |
| Olea-associated insect adintovirus 1                                     | Beta PolB        |
| Olea-associated insect adintovirus 2                                     | Alpha PolB       |
| Oncorhynchus salmon adintovirus                                           | Beta PolB        |
| Orchesella springtail adintovirus                                        | Alpha PolB       |
| Parasteatoda house spider adintovirus                                    | Alpha PolB       |
| Pimephales minnow adintovirus                                             | Beta PolB        |
| Pinctada pearl oyster adintovirus                                        | Beta PolB        |
| Rhodactis coral adintovirus (RNA)                                        | Beta PolB        |
| Spodoptera moth adintovirus 1                                             | Alpha PolB       |
| Spodoptera moth adintovirus 2                                             | Alpha PolB       |
| Steinernema roundworm adintovirus                                        | Beta PolB        |
| Strongylocentrotus sea urchin endogenized adintovirus                    | Beta PolB        |
| Stylophora coral adintovirus                                              | Alpha PolB       |
| Terrapene box turtle adintovirus                                         | Beta PolB        |
| Trichoplusia moth endogenized adintovirus 1                              | Alpha PolB       |
| Trichoplusia moth endogenized adintovirus 2                              | Alpha PolB       |
| Anelosimus tangle-web spider MELD virus                                  | Ad PolB          |
| Blastocystis protist MELD virus                                           | Alpha PolB       |
| Bovine rumen MELD virus                                                  | Alpha PolB       |
| Capsaspora protist MELD virus 1                                          | divergent PolB+S1H|
| Capsaspora protist MELD virus 2                                          | divergent PolB+S3H|
| Coelastrella green algae MELD virus                                      | Beta PolB        |
| Haliotis abalone MELD virus                                               | Alpha PolB       |
| Human feces-associated MELD virus                                        | Alpha PolB       |
| Hydra MELD virus                                                         | Alpha PolB       |
| Kangaroo rat mastadenovirus                                              | Alpha PolB+S1H   |
|                             |
Monoraphidium green algae MELD virus
Monosiga protist MELD virus 1
Monosiga protist MELD virus 2
Physarum slime mold MELD virus
Phytophthora water mold MELD virus
Powellomyces chytrid fungus MELD virus 1
Powellomyces chytrid fungus MELD virus 2
Powellomyces chytrid fungus MELD virus 3
Powellomyces chytrid fungus MELD virus 4
Powellomyces chytrid fungus MELD virus 5
Powellomyces chytrid fungus MELD virus 6
Rhizophas glomeromycete fungus MELD virus
Sheep rumen MELD virus
Synchytrium chytrid fungus MELD element
Trichoplax placozoan protist MELD virus

Abeoforma protist parvovirus
African termite gut bidnaparvovirus
Bombyx silkworm bidnavirus segment 1
Bombyx silkworm bidnavirus segment 2
Dog feces bidnaparvovirus (RNA+DNA)
Drosophila "Vesanto" bidnavirus segment 1
Drosophila "Vesanto" bidnavirus segment 2
False wolf spider bidnaparvovirus (RNA)
Ichthyophonus protist parvovirus
Tasmanian devil feces bidnaparvovirus 1
Tasmanian devil feces bidnaparvovirus 2

Alpha PolB
divergent PolB+S3H
PolA-S3H
S1H+DnaA
D5 S3H
Beta PolB
DEAD S2H
PolA-S3H
divergent PolB+S1H
PolA+D5 S3H
PolA+D5 S3H
Alpha PolB
Alpha PolB
Beta PolB+D5 S3H
RecD S1H
Parvo S3H
Alpha PolB+Parvo S3H
Alpha PolB+Parvo S3H
-
Alpha PolB+Parvo S3H
Alpha PolB
Alpha PolB
Parvo S3H
Alpha PolB+Parvo S3H
Parvo S3H
Alpha PolB+Parvo S3H
Alpha PolB+Parvo S3H
Alpha PolB+Parvo S3H

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| Hexon | Sampled organism                  | Source Contig   | Bioproject/sample |
|-------|----------------------------------|----------------|-------------------|
| Beta  | Astyanax mexicanus               | GFIE01023086    | PRJNA285201       |
| Alpha | Bos taurus                       | NKLS02000104    | SAMN03145444      |
| Alpha | Bos taurus                       | NKLS02001728    | SAMN03145444      |
| Beta  | Branchiostoma belcheri           | AYSR01015205    | PRJNA214454       |
| Alpha | Caenorhabditis elegans           | AL110478        | -                 |
| Alpha | Corynactis australis             | GELM01072747    | PRJNA313487       |
|       | Cotesia congregata               | -               | -                 |
| Beta  | Crocodylus porosus               | JRXG01022388    | PRJNA163131       |
| Alpha | Diachasma alloeum                | LDKA01017305    | SAMN03701895      |
| Beta  | Eucidaris tribuloides            | JZLH010489059   | SAMN00717266      |
| Beta  | Homo sapiens                     | AC083860        | -                 |
| Alpha | Hydra vulgaris                   | ABRM01008631    | SAMN00000081      |
| Alpha | Lagenaria siceraria              | FGKS01003641    | -                 |
| Alpha | Larimichthys crocea              | JRP01001925     | -                 |
| Alpha | Mayetiola destructor              | AEGA01002012    | PRJNA45867        |
| Alpha | Megastigmus spermotrophus        | GCPB01002833    | PRJNA274192       |
| Beta  | Mytilus galloprovincialis        | LNJA010099434   | SAMN03084337      |
| Beta  | Nephila clavipes                 | GFKT01464732    | PRJNA356433       |
| Beta  | Olea europaea                    | FKYM01057659    | PRJEB4992         |
| Alpha | Olea europaea                    | FKYM01057927    | PRJEB4992         |
| Beta  | Oncorhynchus tshawytscha         | PIPH01094194    | SAMN07614989      |
| Beta  | Orchesella cincta                | LJJ01003246     | PRJNA294050       |
| Beta  | Parasteatoda tepidariorum        | AOMJ02256338    | PRJNA167405       |
| Beta  | Pimphales promelas               | JNCE01310190    | SAMN02418978      |
| Beta  | Podagrica attenuata              | NIJO01055010    | -                 |
| Beta  | Rhodactis indosinensis           | GELO01003383    | PRJNA313487       |
| Alpha | Spodoptera frugiperda            | KY042018        | PRJNA344686       |
| Alpha | Spodoptera frugiperda            | KY042019        | PRJNA344686       |
| Beta  | Steinernema feltiae              | AYZB01015367    | -                 |
| Beta  | Strongylcentrotus purpuratus     | AAGJ05080195    | SAMN0829422       |
| Alpha | Styphophora pistillata           | LSMT01002030    | SAMN03491624      |
| Beta  | Terrapene mexicana triunguis     | PRFB02010501    | PRFBO-SAMN08285364|
| Alpha | Trichoplusia ni                  | WHVK01000405    | SAMN06765378      |
| Alpha | Trichoplusia ni                  | NKQN01000414    | SAMN07304761      |
| MELD  | Anelosimus studiosus             | VSFD01006507    | SAMN12325188      |
|       | Human fecal metagenome           | OJKC01000757    | ODFM-SAMEA4350162 |
| MELD  | Bovine rumen metagenome          | -               | -                 |
|       | Capsaspora owczarzaki            | ACF028000430    | SAMD00141126      |
|       | Capsaspora owczarzaki            | ACF028000498    | SAMD00141126      |
| MELD  | Coelastrella sp. UTEX B 3026     | NSMO01000305    | SAMN07595962      |
| MELD  | Haliotis diversicolor            | LC027333        | PRJNA374926       |
|       | Human fecal metagenome           | OOPJ01000021    | -                 |
| MELD  | Hydra oligactis                  | PJUT01375957    | SAMN08153368      |
| Adeno | Dipodomys ordii                  | ABRO02087964    | SAMN02900551      |

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| MELD       | Monoraphidium neglectum | AYTC01002326 | - |
| MELD       | Monosiga brevicollis ATCC 50154 | ABFJ01000467 | SAMEA81495418 |
| MELD       | Monosiga brevicollis ATCC 50154 | - | SAMEA81495418 |
| -          | Physarum polycephalum | ATCM03027720 | SAMN02839457 |
| -          | Phytophthora cambivora | AUVH01007938 | SAMN02178787 |
| MELD       | Powellomyces hirtus | QEAQ01000220 TPX53SAMEA104555614 |
| MELD       | Powellomyces hirtus | - | SAMEA104555614 |
| MELD       | Powellomyces hirtus | - | SAMEA104555614 |
| MELD       | Powellomyces hirtus | - | SAMEA104555614 |
| MELD       | Powellomyces hirtus | - | SAMEA104555614 |
| Gag        | Rhizophagus irregularis | AC163889 | SAMD00053945 |
| MELD       | Sheep rumen metagenome | ODAJ014727734 | SAMN02144231 |
| -          | Synchytrium microbalum | QEAO01000105 | SAMEA104555616 |
| Alpha      | Trichoplax sp. H2 | ABGP01001206 | SAMEA5175452 |
| Parvo      | Abeoforma whisleri | PDIO01005734 | SAMN06200030 |
| Parvo      | Cubitermes ugandensis | - | SAMN06268476 |
| Parvo      | Bombyx mori | - | - |
| Parvo      | Bombyx mori | - | - |
| Parvo      | Canis lupus familiaris fecal virome | - | PRJNA473018 |
| Parvo      | Drosophila melanogaster | - | - |
| Parvo      | Drosophila melanogaster | - | - |
| Parvo      | Tengella perfuga | GGOF01037889 | PRJNA471192 |
| Parvo      | Ichthyophonus hoferi | NAOQ01000873 | PRJNA290639 |
| Parvo      | Sarcophilus harrisii | - | SAMN10229747 |
| Parvo      | Sarcophilus harrisii | - | SAMN10229747 |
| SRA          | GenBank   |
|--------------|-----------|
| SRR2045404-14. M BK010998 |          |
| SRR5753414 SRR57BK010889 |          |
| SRR5753414 SRR57BK010999 |          |
| SRR1174915. SRR1: BK010888 |          |
| Can't find SRR |          |
| SRR3198576, SRR31BK011000 | NC_006658|
| parent reads unavailable |          |
| SRR2043726 - "synl" matching RNASeq: |          |
| SRR870190. Origir BK012049 |          |
| SRR870190. SRR87: BK0111001 |          |
| SRR1258893. Also r BK0111002 |          |
| Used Spades to rec BK0111003 |          |
| SRR1805073 SRR18BK011004 |          |
| SRR1598987 SRR15BK011005 |          |
| SRR5139361. Also r BK011006 |          |
| MagicBlast finds ali |          |
| MagicBlast finds ali |          |
| SRR6312215, SRR6: |          |
| SRR2222657(Illumin BK011007 |          |
| SRR891578 83 85 h BK011108 |          |
| SRR1301972. SRR1: BK011109 |          |
| ion torrent metage |          |
| SRR3201278 SRR32BK011101 |          |
| SRR4308584. Match BK0111025 |          |
| SRR4308584. Match BK0111026 |          |
| Can't find SRR |          |
| SRR7211988. Conti BK0111011 |          |
| Megahit reassemble BK012063 |          |
| SRR6435272 BK010890 |          |
| ORFs appear degra |          |
| ORFs appear degra |          |
| SRR9900418 BK012043 |          |
| ERR1543996 SRR06 BK012044 |          |
| SRR094424 BK012045 |          |
| DRR183067 BK012046 |          |
| DRR183067 BK012047 |          |
| parent reads unavailable BK012048 |          |
| SRR5986943 but co |          |
| MG-RAST mgm462 |          |
| SRR6364634 BK012051 |          |
| SRR1646412-B |          |
TPA_inf (parent re: BK012050
ERR1824636   BK012085
ERR1824636   BK012086
SRX000029 (old 45+ BK012052
SRR870494. SRR871/BK012053
ERR2451317. Didn't BK012054
ERR2451317. Didn't BK012055
ERR2451317. Didn't BK012056
ERR2451317. Didn't BK012057
ERR2451317. Didn't BK012058
ERR2451317. Didn't BK012059
SRR1046100. DRR0 BK012060
SRR1222431   BK012061
ERR2451318   BK012062
ERR2992734   BK012064
SRR5195172 SRR51BK010894
SRR5247064   BK012042
-   NC_020928
-   NC_020927
SRR7287177   BK012084
-   KX648533
-   KX648534
SRR7186263. Only: BK011024
SRR2134855. ReassBK011022
SRR8048117   BK010893
SRR8048117   BK011023

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| Tissue     | SRR            | Mbases | Paired Ad into Reads | Reads/Gb |
|------------|----------------|--------|----------------------|----------|
| Ovary      | SRR2045404     | 11,670 | 2460                 | 211      |
| Brain      | SRR2045405     | 6,631  | 2604                 | 393      |
| Gills      | SRR2045406     | 5,039  | 2236                 | 444      |
| Heart      | SRR2045407     | 5,779  | 1052                 | 182      |
| Muscle     | SRR2045408     | 5,358  | 388                  | 72       |
| testis     | SRR2045409     | 13,255 | 2200                 | 166      |
| Head kidney| SRR2045410     | 5,855  | 3996                 | 682      |
| Bones      | SRR2045411     | 5,647  | 1024                 | 181      |
| Intestine  | SRR2045412     | 8,625  | 5012                 | 581      |
| Testis     | SRR2045413     | 5,660  | 2190                 | 387      |
| Embryos    | SRR2045414     | 8,273  | 104                  | 13       |
The identity of this gene was assigned based on sequence similarity to other adintovirus Cupiennin genes (which in turn generate hits for cupiennin in HHpred searches, see below)
Note: this result is for the single GasderminX sequence of the example virus. Multiple alignments of related GasderminX genes give higher probability hits in HHpred searches.
1. PF37517.13 (DNA_pol_B-2): DNA polymerase type B, organellar and viral

Probability: 100%, E-value: 2e-47, Score: 470.37, Aligned cols: 426, Identities: 29%, Similarity: 3.322,

```
| Q   | P   | AA1 | AA2 |
|-----|-----|-----|-----|
| KOG3288 | DNA_pol_B-2 | DNA |
| 38Y_A | 2PY5_B |
| 460U_A | SW5X_B |
| 6DX1_A | -IR |
| 460U_B | KOG2482 |
| 3PUH_A | 6E94_B |
| 3HAN_A | - | 3TAY_A |
```

Template alignment | CDD

```
| Q   | P   | AA1 | AA2 |
|-----|-----|-----|-----|
| KOG3288 | DNA_pol_B-2 | DNA |
| 38Y_A | 2PY5_B |
| 460U_A | SW5X_B |
| 6DX1_A | -IR |
| 460U_B | KOG2482 |
| 3PUH_A | 6E94_B |
| 3HAN_A | - | 3TAY_A |
```
The ORF designated “oncoid” is named based on synteny with other adintovirus cysteine-rich ORFs that encode short linear motifs of known oncogenes. The oncoid ORF in this example does not itself show clear hits for the linear motifs detailed in the text.
1. **3NTT** A Capsid protein; Adeno-associated virus 5, gene therapy; 3.45A (Adeno-associated virus - S)

   - **Q ss_pred**
   - **Q_MayetiolaPLA**
   - **Q Consensus**
   - **T Consensus**

   - **3NTT_A**: 404786
   - **cd04786**: 127 (315)
   - **Phospholipase_P2**

2. **PP50829.12**: Adeno PX; Adenovirus late L2 mu core protein (Protein X)

   - **Q ss_pred**
   - **Q_MayetiolaPLA**
   - **Q Consensus**
   - **T Consensus**

   - **pp50829.12**: 127 (315)
   - **128**: 155 (315)

---

**Astyanax Cupiennin**

1. **PF17563.2**: Cu; Cupiennin

   - **Q ss_pred**
   - **Q_AstyanaxCup**
   - **Q Consensus**
   - **T Consensus**

   - **PF17563.2**: 74 (159)
   - **2K30.0**: 49 (159)
   - **2K10.0**: 49 (159)
   - **2G6L.0**: 49 (159)
   - **Antimicrobial_7**: 49 (159)
   - **ODF4449**: 49 (159)

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Branchiostoma

Bcl2

Mytilus

IAP
unidentified

Parasteatoda house spider

Megasporagia wasp (RNA)

Mytilus mussel

Nephila orb-weaver spider (RNA)

Olea-associated insect

Orchesella springtail

Parastatoida house spider

Spodoptera moth 1

Spodoptera moth 2

Stylophora coral

Asmatman tetra cavityfish (RNA)

Branchiostoma lancelet

Eucidrilus sea urchin

Larimichthys croaker fish

Olea-associated insect

Oncorhynchus salmon

Adinococ7 Adinococ5

Pimephales minnow

Pinctada oyster

Rhodactis coral (RNA)

Steinerna roundworm

Stronglyocentrotus sea urchin

Turrataene box turtle
Supplementary Fig. S3: Sequence similarity network analysis for Hexon and Penton proteins. Hexon and Penton protein sequences from curated adintovirus and MELD virus genomes were subjected to all-against-all BLASTP analysis with an E-value cutoff of 1e-5. Asterisks mark the *Mayetiola* (triangle) and *Terrapene* (square) exemplars. The figure shows that some genomes with Alpha PolB genes have Hexon and Penton proteins that cluster with Beta-type Hexon and Penton proteins. The figure also shows that non-animal MELD virus Hexon and Penton protein sequences generally cluster with one another and not with animal adintovirus sequences (see Results section “Viruses with adinto-like genes in non-animal eukaryote datasets”).
LOCUS AC083860    7129 bp    DNA    linear    SYN 16-Nov-2019
DEFINITION .
FEATURES Location/Qualifiers
  source 1..7129
    /mol_type="genomic DNA"
    /organism="Homo sapiens"
    /chromosome="7"
    /clone="RP11-3P18"
    /clone_lib="RPCI human BAC library 11"
    /db_xref="taxon:9606"
    /segment="37604428..37597300"
    /sex="male"
    /tissue_type="blood"
  CDS join(1..246,249..362,365..499,501..614,618..767,769..894,
          1093..1233)
    /product="Hexon"
  repeat_region 1..75
    /note="MLT1H"
  repeat_region 1138..1312
    /note="MER58A"
  CDS 1230..1514
    /product="Adenain?"
  CDS complement(join(1504..1578,1581..1646,1648..1926,
                     1963..2061,2096..2299))
    /product="Adintoc2?"
  CDS complement(join(2278..2328,2331..2450))
    /product="GasderminX"
  repeat_region 2666..3121
    /note="MLT1C"
  CDS complement(join(2848..2886,2888..2902,2905..2985,
                     3021..3122,3156..3311,3337..3477))
    /product="FtsK?"
  CDS complement(join(3441..3557,3559..3687,3689..3877,
                     3909..4316,4322..4405,4408..4467,4469..4561,4564..4956,
                     4964..5101,5301..5369,5372..5518,5599..5700,5791..6009,
                     6036..6242))
    /product="PolB"
  CDS complement(join(6245..6367,6370..6621,6630..6704,
                     6706..6975,6977..7129))
    /product="Oncoid?"
BASE COUNT 2069 A 1429 C 1416 G 2215 T 0 OTHER
ORIGIN
  1 ATGACAAATA ATCAATGTG TTTAAAGCCA CTAAGTTTTA GGAAGCTTGT
   TACGCAGTAA
  61 AGTCTAACTG ATACACAAGA AGGTAATTT TACAAAGATG CTGCCAGTAA
   GCAGAAAATAC
2881 TATGGATTAT TCTATTATTA GGAAGATTAG GATTAGATTA TGCTGCCACA AACCAGGAA
2941 TGCTCCTGGGC AACAAAAACG CGGAAGAGTC AAGGAAGGTT
CTCCCTTGAAG GTCTACGGAA
3001 GTCCTCTGGG CACTTTGATT TCAGACTAGG GACCTCCAGA AATGTAAGAG
AATCAATT
3061 TGCTCTTTGG AGGCATCTAC TTTATGTTAA TTTACTATGG CAGCCCCAGG
AAATGACAC
3121 AATCGACACA GGTATCCTTG GAGACTATGG TAGCATCATG CAAGGTACAC
GAAAAAAAGA
3181 GTGCCTGTTTT CATGGACATC TGTGGGGCCA GTGTACCTG TAATTAGTCC
CAAGCATTT
3241 GAAAAAGAAC CATGACTTTT TCTATTGTTG GCACAGTTTC CCTATCCTG
CAAAATAT
3301 TTGGAAACCAT GTTAAAGAGG CTGAGATTCA GAGATGTACAT ACTTGTTGGG
ATTTTTTAAA
3361 AAATTATTTG TTGCCATGGG CCGCTTGTGCT CTGAAACTTA CACTGCACAC
ACTCAACAC
3421 GCAGGCTTCAT CCTTCCAGAG TTAGAGGGCT TGTTCCGTTTT TTTGAAAAGC
GTTGTGTCA
3481 CAGACTACTG AAATCTCAGC AGTAAGAAGG TGCTTTAAGG CCAACAAACT
ATGTGTAGA
3541 CAGATAACTG ATGTTTTTTC ACATGTGTCTG TCTCAATTGG TCAAAACATG
GCTGCTCTTT
3601 TAAAATGTC TTCAATTATTA CCTGCTTTTT TGAGGAGATA AACAAGCATG
AGTTTCCTAA
3661 GACATACAAA TTGACTTTGG TACAGTTCTA GCTTTCTGTG TAATGTCTCT
GAGATGGGCC
3721 AGACCATTAAC CCGCTGATGGA TGAGGTGCCC CAGAAGAAGT CTCTGCAATA
TACACCTGAAG
3781 GAACAAATCC CCTACTCAAT TCAGCCAGCT TACAGCTCC AGGATGGTTC
CAGTTCTGT
3841 TCTGTGCTGC TAATAAAAAAG GAATTTCCTTG TTACTGATTA AACatatTTAA
TTAAGATCAT
3901 CAAGTAGGATT ATACTCCTCT GTGAATGGAAT GCTAAATATA AGTAGGCAAT
GACAGTATTTA
3961 GCCCCTCCTCT CTATTAGGCA GGTGATTCTA AAATCACCTT TCCAACAAAT
CATAAGGATTA
4021 TAGTCATTAT TGGATTTGGC CCGACACAT GCATAAGAAA
GAGACACAG GAAATGGTCA
4081 AGCTCTACAT GTATCCTCAC TAGGCAATG TGGAAGAGG ATTAACCTTGTG
GGCAACCTTTA
4141 TAATCCCCGTG AAATAAAAAA TGAGTTAGCA ATCTCTCTGC TGGGGCTTC
TCTTACAGC
4201 GCTGGGGATT CAAGCAAACT CCTACAGGG TGATAATCCT CGAGACATT
GTTTTTCTTG

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5641 CTCACTTTCT CTAAACAGG TGTTGATGAT ACATAGAATT TGGGTATTTA
TCTCAAGTTTCC
5701 TTACTTTACG CTGAAATCAG ATACACACTCA ATCCTGGAAG AGTTTAGAGT
TTCCAAAAACC
5761 ATAGCCCCAG AAAATAGATG TTATCAATT TAAAGCAGA CTTTTTAGGG
ACACCATATA
5821 AAAAGGCTTTC ACTTTACTTT TCCTGGGGAG GGAATCCCT GAAGCATCTC
CCAAACAATA
5881 TTTTTTTATG CTCAATTTCG CGCCAGCAAGC TTCTCATTTC TGCCATTTTA
AGCAAAATTT
5941 GGAGCAGCAG ATGGCCTCACT AGTTAGGAGTTA TCCCATATGGG TTCTCCACCC
TTCTCCCAAA
6001 AACACTGTATT TATACACATC CTTGGGTTTG TGTCAGGAGAC CACTGTAACA
CTGATCAGCG
6061 ATGGGAATCAT CACCTGAGA TCCTACATGC ACGACATGAC ACAAACAGGG
TTGCTCTGAC
6121 AAAAAGCTGT TTGCATCACT ACAAGCTGGG GACTGAGGTC AACAACAGGG
TTGCTCTGAC
6181 CTATGCCCTGT TATAAGATAT GATAATAGTA AAGGTAGGTT AGCCCTTACA
CACAGACTGC
6241 ACTCAGGTGG ATAATTTTCT AAATGAAAAA TCTCAATGTC TTCTGAAGCC
TTCTGGTTG
6301 GGCAATCAGT TTGACTGCACT TCTACCTTTA AAAAGGCTTG CTGTGAACC
ACTGAAATCC
6361 AGATAAATCAA TACTGGATAT TAGTGATCCC AATAAATGTA ACTACTGAAT
TCCAGGCTGA
6421 TCCAGGTGTC AAGCAGGACCG TGTTTCTACAG GAGGAAAACT TCTGCACCTCA
CTCCCTGCAA
6481 AGTTCTTACT GTTGTTACTG TTAACATAAAA CAACACTGCA CAGATAAAACT
CTGTATTGTT
6541 CTGTACCACA TTAGCAAAAT TAGTTAGAAA TTCCAAGCA TTTAGCTCGC
CAGTCCTGGA
6601 TTGAAGATAGG LTGAAGTGGG TTTAAAACTTA AAATTCATATA ACCTACTAAA
ATATTTCTA
6661 TGAATACTCAT ATATGAATCT CTGCAAGTTTC CAGTTAATCA CTICCAGTTTC
TTCTGAAAT
6721 GATGGATCTG TGGATAAGTT TATTAAAAA CTCATATTTT ATTGTAATTAT
TCAACATTTT
6781 GGAAACCTCT ATTATCCATT GATGTGTTTT CAGGCCCTAG CTTCCTAGTG
TCTGTGTGCA
6841 TTTTAATCAC CCTTTTCTTC TCTCTAGTGAC TCCAGAACCCT CTTTTTTTAT
TTCTGAGGA
6901 TTCTGAGGGA CTGTGTGTTG GATTTGTGTG GAGTGAGTCC AAGGGATGCC
AGGCTGACATC
6961 GTAGCTCTGG AATTTACTTG CAACCACAGG CTCTCCCAAC CCTGTATGG
GGCTTACAGAG
7021 TTTGGAGGTG GTAAGAGGT GGCCAAAAGC TCTTCAACCC CACAGGTGCT GCACATCCAA
7081 TGTATATGAG GCTGACCACA TCTTGGAGAA GGCTTGATAT GCTGACCTC

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