Identification of Multiple Replication Stages and Origins in the Nucleopolyhedrovirus of Anticarsia gemmatalis

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Abstract: To understand the mechanism of replication used by baculoviruses, it is essential to describe all the factors involved, including virus and host proteins and the sequences where DNA synthesis starts. A lot of work on this topic has been done, but there is still confusion in defining what sequence/s act in such functions, and the mechanism of replication is not very well understood. In this work, we performed an AgMNPV replication kinetics into the susceptible UFL-Ag-286 cells to estimate viral genome synthesis rates. We found that the viral DNA exponentially increases in two different phases that are temporally separated by an interval of 5 h, probably suggesting the occurrence of two different mechanisms of replication. Then, we prepared a plasmid library containing virus fragments (0.5–2 kbp), which were transfected and infected with AgMNPV in UFL-Ag-286 cells. We identified 12 virus fragments which acted as origins of replication (ORI). Those fragments are in close proximity to core genes. This association to the core genome would ensure vertical transmission of ORIs. We also predict the presence of common structures on those fragments that probably recruit the replication machinery, a structure also present in previously reported ORIs in baculoviruses.

Keywords: baculovirus; AgMNPV; replication

1. Introduction

Baculoviruses are rod-shaped enveloped viruses with double-stranded-DNA circular genomes from 81,775 (Neodiprion lecontei NPV (NeleNPV)) to 178,733 kbp (Xestia C-nigrum GV (XecnGV)) [1]. Baculovirus infection produces two virion phenotypes: Occlusion derived virions (ODVs) and budded virions (BVs) [1]. The BVs appear during the initial stage of the multiplication cycle and are responsible for systemic infection of the host, while the ODVs are produced at the final stages [2]. This phenotype is so named because it is occluded in a protein matrix mainly based on polyhedrin or granulin (denominated occlusion bodies—OBs) that protects virions from different environmental injuries. ODVs are the agents responsible for primary oral infection in a new host [3]. From a biotechnological point of view, baculoviruses have many interesting features: They can carry large and multiple DNA inserts (at least 38 kbp) [4–6], and they can be easily produced and purified at high titers. These insect viruses (wild type species and recombinant variants) have been used in many applications, such as
biological control agents against insect pests [7–11], protein expression systems [12,13], putative non-human viral vectors for gene delivery [14], vectors for vaccines [15,16], or as models of genome evolution [17,18], among others. The Baculoviridae family is divided into four genera: *Alpha* Baculovirus and *Beta* Baculovirus, comprising lepidopteran-specific viruses; *Gamma* Baculovirus, which includes hymenopteran-specific baculoviruses; and finally, *Deltabaculovirus*, which, to date, comprises only Culex nigripalpus NPV (CuniNPV) and possibly other still undescribed or in the process of description dipteran-specific baculoviruses [1]. A notable feature of these pathogens shared by other viruses with large DNA genomes [19] is the existence of a different gene content among known species. However, they share a set of 38 orthologous sequences known as core genes [20,21], derived from ancestral baculovirus, while the other regions (each genome contain between 89 (NeleNPV) to 181 (XecnGV) open reading frames (ORFs)) are shared by various members or are unique to species [1,22]. This particularity complicates the studies conducted to understand the basic functions encoded in baculovirus genomes.

The core genes provide insights into some of the major functions required of all baculoviruses [22], may be defining what a baculovirus is. Core genes can be considered as ancestral essential genes that diverged in higher or lesser degrees [1]. However, it is not only ORFs that can be conserved among evolution as non-coding DNA sequences associated to regulation processes might also be conserved, hence a relation between core genes and those sequences might exist.

The baculovirus replication process has been extensively studied and similarities with other entities, such as herpesvirus, have been found [23], but many questions remain to be answered. Baculoviruses replicate in cell nuclei, probably using a combination of host and virally encoded proteins [22], but based on the size of the short-lived replication intermediates, the DNA synthesis would proceed by a markedly different mechanism to the host nuclear DNA [24], or the one used by other viruses, such as Polyomaviridae [25]. During replication, DNA molecules larger than the genome length are produced, which could be explained by two mechanisms: Rolling circle [26,27] or recombination-dependent replication [28,29]. In order to understand DNA replication in baculovirus infected cells, short-lived DNA replication intermediates have been characterized, identifying heterogeneous fragments up to 5000 nt, with an average size of 1000 to 2000 nt that could represent Okazaki fragments (Theta mechanism) or could also be fragments synthesized during recombination dependent replication [24].

A Theta or rolling circle mechanism needs an origin of replication (ORI), where a protein can bind to start the process by recruiting the replication machinery or by nicking DNA to allow strand displacement, respectively. In baculovirus, the replication seems to occur from multiple origins [22,23]. To identify this kind of sequence, transfection of plasmids containing viral DNA into virus infected cells has been used, subsequently digesting with DpnI to distinguish newly replicated DNA from input DNA [22]. With this approach, homologous regions (hrs) [30,31], non-hrs sequences [32,33] and early promoters [34] have been identified as causing the infection-dependent replication of plasmids. Wu et al. [35] identified a different non-hr ORI inside a core gene of AcMNPV (p143 or DNA helicase), broadening the spectrum of sequences involved in this function.

To date, different sequences have been described as origins of replication in baculoviruses. The homologous regions (hrs) from AcMNPV are 30 bp imperfect palindromes [36] that were originally postulated as viral origins because of their symmetric location, high AT content, and palindromic structure [37,38]. However, there are some baculovirus that do not have any hr [22], so hrs might not be required by all baculoviruses for replication. Furthermore, Carstens and Wu showed that no single hr is essential for replication in a cell culture [39]. On the other hand, the non-hr ORI found inside the fragment HindIII-K of the same virus [40] has no obvious similarity with any of the hr sequences. This led us to question if different regions can alternatively or simultaneously function as ORI, or if there is a common secondary structure that is recognized by the replication machinery. Besides, this region has been found in defective AcMNPV genomes, generated after extensive serial passage of the virus in an insect cell culture, suggesting an important role in replication [32]. It seems that the AcMNPV DNA replication system does not require hrs for initiation. In fact, virtually any
sequences can be multiplied when plasmids and viral replication-essential genes are co-transfected into susceptible cell lines since the initiation of DNA synthesis could be related to the chromatin structure [34]. However, when given time, after plasmid transfection prior to infection, it has been observed that only the plasmids containing viral promoter or hrs replicate [23,34]. Following this strategy, the previously mentioned non-hr direct repetitions located into the coding region of the p143 gene were functional as ORI in insect and mammalian cells [35].

In this work, an unbiased method on a genome-wide scale was developed to isolate origins of replication in the Anticarsia gemmatalis multiple nucleopolyhedrovirus (AgMNPV), the alphabaculovirus most used worldwide as a bioinsecticide to control crop pests [41]. We combined a genome library with a previously described transient replication assay [34,42]. Exhaustive analyses of recovered sequences obtained in this work and comparison with previous reports are discussed to understand how these entities multiply their genomes. We found three kinds of origins, according to their genome location: Homologous regions (ORI-hrs), intergenic regions (ORI-igrs), and intragenic regions (ORI-grs); particularly, we detected non-hr ORI positioned within or close to core genes.

To find out more about baculovirus replication, viral DNA replication kinetics were monitored post-infection into the susceptible UFL-Ag-286 cells hourly for 30 h. This assay identified four different phases. Two of them were exponential stages (the second and fourth ones) and the other two showed no significant increase in DNA synthesis (the first and the third ones).

Taken together, the results shown here indicate that baculoviruses can use different kinds of sequences to replicate their genomes and they can undertake different replication mechanisms. The common feature that allows the different ORIs to be recognized by the replication machinery remains to be answered. However, this work constitutes a step forward in understanding the viral cycle.

2. Materials and Methods

2.1. Cell and Virus Stocks

The insect cells UFL-Ag-286 [43] were grown at 27 °C in GRACE’s medium (Invitrogen, Carlsbad, CA, USA) containing 10% v/v fetal bovine serum (FBS; GBO, Kremsmünster, Austria) and supplied with antibiotics and antifungal agents (Invitrogen). Initial stocks of AgMNPV-2D [44,45] were multiplied in Anticarsia gemmatalis (third instar larvae) by per os infection using OBs, and by the exposition of monolayers of UFL-Ag-286 cells with BVs in plastic tissue culture flasks. The virus stocks used in all experiments were tittered by plaque assay [46] in UFL-Ag-286 cells and maintained as culture supernatants.

2.2. Quantitative Real Time PCR Assays for AgMNPV and UFL-Ag-286 Cells

A molecular approach to quantify AgMNPV and UFL-Ag-286 cells by two quantitative real time PCRs (qRT-PCR) was developed. Thus, pairs of primers that bind in the virus egt gene (pFw-egt: 5′TAATCGGCAACGCCTCTAC3′; pRev-egt: 5′CGATGCGGCAAACAAACAACS′) and into the cell actin gene (pFw-Ac: 5′TAATCGGCAACGCCTCTAC3′; pRev-Ac: 5′CGATGCGGCAAACAAACAACS′) were synthesized. To design the above oligonucleotides, the AgMNPV genome (NC_008520.2) and our own data (KX139200.1) from the actin gene of Anticarsia gemmatalis were used. First, end-point-PCR assays were performed using standard conditions and Taq DNA pol (PB-L, Bernal, Argentina) to produce the egt and actin amplicons, which were then molecularly cloned in pGEM-T-Easy (Promega, Madison, WI, USA) and Escherichia coli Top 10 [47]. To verify the identity of both constructs (pgT-egt and pgT-act), sequencing reactions with the universal primers, T7 and Sp6, were carried out (Macrogen Services, Seoul, South Korea). After this, both qRT-PCR were optimized in sensitive and dynamic range parameters using serial dilutions of pgT-egt and pgT-act as templates (1 µL), 25 µL of 1× Maxima™ SYBR Green qPCR Master Mix (Fermentas, Waltham, MA, USA), 0.3 µM of every primer, and a Smart Cycler (Cepheid Inc, Sunnyvale, CA, USA) to cycling temperatures and detect fluorescence. Plasmid DNA was previously purified (Qiagen Miniprep Kit; Qiagen, Hilden, Germany), quantified by spectroscopy (Nanodrop 1000; Thermo Fisher, MA, USA), and the copy number was determined using the Thermoscientific copy number calculation tool.
The temperature conditions optimized for both genes were: 10 min at 95 °C (1 cycle), 15 s at 95 °C, 30 s at 60 °C, and 30 s at 72 °C (40 cycles).

2.3. Replication Kinetics

To understand the replication process of AgMNPV multiplying on in vitro cell cultures, a kinetic assay was performed. To this, synchronized monolayers (70% confluence) of UFL-Ag-286 cells growing on 24 multiwell plates were exposed with AgMNPV (multiplicity of infection—moi—10) for 1 h. Then, cells were washed three times with 500 µL of phosphate buffer saline (PBS) and finally incubated with growth medium (500 µL of GRACE’s with 10% v/v SFB) for 1 to 30 h at 27 °C. The whole conditioned medium and cells were recovered from wells every 1 h, collecting three samples of cells and supernatants (three wells) per time. The viral DNA (vDNA) was isolated from supernatants by the miniprep method [46], reserving 50 µL to infection assays. For the remainder of the sample, DNA (including vDNA and cellular DNA—gDNA) was isolated from cells by treatment with Proteinase K (Sigma-Aldrich, St. Louis, MO, USA) in lysis buffer (0.2 M Na₂CO₃, 0.02 M NaCl, 0.2 M EDTA, 0.5% w/v SDS), phenol-chloroform extracted, and subsequently precipitated with ethanol and 0.3 M NaOAc [47]. Finally, to quantify relative levels between vDNA and gDNA, qRT-PCR assays previously optimized were applied. Thus, all DNA samples were analyzed with 1× Maxima™ SYBR Green qPCR Master Mix (Fermentas, Waltham, MA, USA) into a Smart Cycler (Cepheid Inc, Sunnyvale, CA, USA) using as pairs of primers, pFw-Act/pRev-Act and pFw-egt/pRev-egt. To estimate the number of DNA molecules, the recombinant plasmids, pGT-egt and pGT-act, were used as calibrators. For normalization, gDNA was calculated arbitrarily considering one copy of actin gene per cell. The results were plotted as quotients between vDNA and gDNA and the statistical error and ratios were calculated and then propagated. Unpaired T test was done for statistical analysis. The replication rates corresponding to exponential phases were estimated from the slopes of the curves using SigmaPlot software. To verify the presence of BVs in culture supernatants, 50 µL of each conditioned medium sample were used as inoculums on UFL-Ag-286 cells growing on 48 multiwell plates and incubated for 1 week at 27 °C. The presence or absence of infection was determined by microscopy (Nikon Eclipse TS100).

2.4. AgMNPV Genomic Library

A plasmid library of overlapping fragments derived from the AgMNPV genome was produced into the pBlueScript SK (+) vector (pBSK) (Stratagene, San Diego, CA, USA). Briefly, AgMNPV genome (vDNA) was isolated by the full scale method from BVs, which include concentration by ultracentrifugation, proteinase K (Sigma) treatment, phenol-chloroform extraction, and alcohol precipitation steps [47]. The virions were previously produced in controlled conditions in UFL-Ag-286 cells (flask of 175 cm²; 70% cell confluence and state of synchronization [45]; moi 0.1; 5 days of incubation). After verifying the quality by agarose gel electrophoresis and spectroscopy (Nanodrop 1000), vDNA was partially digested with HaeIII (Fermentas, Waltham, MA, USA) to obtain overlapping fragments ranging between 500 and 2000 bp, which were recovered from preparative electrophoresis using a Zymoclean Gel DNA recovery Kit (ZYMO Research, Irvine, CA, USA). Subsequently, vDNA fragments were ligated using T4 DNA ligase (Hoffmann-La Roche, Basel, Switzerland) to pBSK, which was previously treated with EcoRV (Fermentas, Waltham, MA, USA) and dephosphorylated (Shrimp Alkaline Phosphatase, Fermentas, Waltham, MA, USA). Finally, recombinant DNA was transformed by electroporation (BIORAD Gene Pulser II) in Escherichia coli Top 10 (Invitrogen, Carlsbad, CA, USA) and then 500 colonies white colored (in presence of X-Gal; Sigma) and resistant to ampicillin (Sigma-Aldrich, St. Louis, MO, USA) grown on solid Luria-Bertani (LB) plates (Britania, Buenos Aires, Argentina) were selected. The genome coverage of the AgMNPV library was estimated. For this, bacterial clones randomly selected (20%) were used to isolate recombinant plasmids with the goal to determine the insert length by treatment with restriction endonucleases and electrophoresis analyses. All clones were individually maintained at −80 °C in storage medium (LB broth supplied with 15% v/v glycerol).
2.5. Transient Replication Assay

All recombinant plasmids from the AgMNPV library were isolated by the standard alkaline method [47] from 5 mL cultures (LB broth supplied with ampicillin). Then, constructs were mixed in pools of 25 clones (selected without any bias and maintaining the same copy number of each one) and 2 µg of mixed DNA for each group were transfected using CellFectin Reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s instructions on 8 × 10^5 UFL-Ag-286 cells per well grown in standard condition in various six multiwell plates. As a control, some wells were transfected with 2 µg of pBSK without insert. After a period of 6 h, transfected cells were also infected with AgMNPV at moi 1 and subsequently incubated at 27 °C for 48 h, reserving some wells without virus exposure as controls [23,34]. At 48 h, cells from all wells were harvested and plasmid DNA was purified using the alkaline method [47]. The recovered DNAs were overnight treated with 2.5 U of DpnI (Fermentas, Waltham, MA, USA)—reserving some samples without enzyme hydrolysis as controls—to eliminate the DNA template synthesized in bacteria, cleaned up by phenol-chloroform extraction and alcohol precipitation, and transformed in E. coli Top10. Control samples without any plasmid (only UFL-Ag-286), or containing only the empty cloning vector (UFL-Ag-286/pBSK, UFL-Ag-286/pBSK digested with DpnI) were used to adjust the experimental conditions. The insert sequences of transiently replicated constructs were obtained by Sanger sequencing (Macrogen Inc, Seoul, South Korea) with universal primers (T3 and T7), and then compared against the AgMNPV-2D genome (NC_008520.2). The transient replication assay was carried out twice in independent events.

2.6. Bioinformatics Analysis

The sequences obtained from the virus genome library were identified by the BLAST algorithm (www.ncbi.nlm.nih.gov/blast) using the information of AgMNPV-2D (NC_008520.2). The AT content of the whole genome was calculated and profiled using an overlapped sliding window strategy (window = 1000 nucleotides, step size = 100 nucleotides), using a DNA base composition analysis tool developed by Jie Zheng in 2004 (http://molbiol-tools.ca/Jie_Zheng/). The average AT content of the whole genome and library inserts containing AgMNPV’s ORIs were also calculated with the same tool. Besides, the amount of A or T runs (4–8 mers) were dimensioned by a similar approach and profiled using an index, where the number of runs was normalized with the average value of the AgMNPV AT content. To detect putative promoters, the available information regarding typical early and late motifs described in Baculoviridae [48] and the DNA Pattern Find program [49] were applied. Moreover, to predict palindromes, the European Molecular Biology Open Software Suite (EMBOSS) Palindrome tool (http://emboss.bioinformatics.nl/cgi-bin/emboss/palindrome) was utilized [50]. After characterizing the ∆G₀ of stem loops constituted by inverted repeats of different lengths (using RNAstructure Software Package) [51], the following search parameters were selected: Minimum length of palindrome, 5; maximum gap between repeated regions, 18; number of mismatches allowed, 1. DNA folding routine of Mfold web server [52] was used to predict the secondary structures of DNA with the following modified parameters: Folding temperature (27 °C) and maximum distance between the paired bases (40 bp).

3. Results

3.1. AgMNPV Kinetics of Replication

A kinetic analysis of virus DNA synthesis using in vitro cell cultures was performed, to contribute with experimental evidence about the AgMNPV genome’s replication that allow the identification of different phases and rates of synthesis. For this, a qRT-PCR approach and two DNA loci as templates were selected: The egt virus gene (vDNA) and cell actin gene (gDNA). A segment of the actin gene had been obtained using ad hoc designed universal primers in an end-point-PCR assay and subsequently sequenced (GenBank KX139200) since there were no available Anticarsia gemmatalis sequences in the databases. Thus, synchronized UFL-Ag-286 cells were exposed using an excess of AgMNPV to synchronize the
infection, profusely washed to remove the virions that did not enter cells, and finally were incubated in growth conditions. Every 1 h-interval, gDNA and vDNA were measured (Figure 1).

![Replication kinetics of AgMNPV](image)

**Figure 1.** Replication kinetics of AgMNPV. Synchronized monolayers (70% confluence) of UFL-Ag-286 cells growing on 24 multiwell plates were exposed with AgMNPV (moi 10) for 1 h. Then, cells were incubated with growth medium for 1 to 30 h at 27 °C (hpi: hours post infection). DNA was recovered from the conditioned medium and cells by triplicate for each point. To quantify relative levels between cell and virus DNAs (gDNA and vDNA, respectively), qRT-PCR assays were applied. The results were plotted as quotients between vDNA and gDNA. The statistical error was calculated and then propagated to obtain the error bars. The beginning of replication and budding are indicated with grey arrows. Replication stages: Phase one, from 0 to 4 h; phase two, from 5 to 16 h; phase three, from 17 to 21 h; and phase four, from 22 to at least 30 h. The replication rates corresponding to phase two and four are indicated with a dotted line. An unpaired parametric t-test was done between the vDNA/gDNA values corresponding to contiguous times. The statistically significant differences (* p < 0.0001) that separate the replicative phases are indicated.

Results revealed different stages of AgMNPV replication. The first stage finished at 5 h post virus exposure of infective particles (hpi) when the quantity of vDNA increased significantly with respect to the previous time (p < 0.0001). Then, a second phase between 5 and 16 hpi showed that vDNA progressed in an exponential tendency (μv: 0.37 ± 0.027) without detecting DNA in the conditioned medium. After this time, BV production started because virus genomes were detected outside cells (corroborated by in vitro cell infection assays) and a decrease in the whole production of vDNA was observed (p < 0.0001). This is in accordance with the fact that many genome molecules were packaged and emerged as BVs, and in consequence, they could not have been used as new templates for replication. This particularity occurred for 5 h (from 17 hpi to 21 hpi; p < 0.0001), defining a third replication phase. Finally, the process progressed in another exponential phase of growth (μv: 0.34 ± 0.037) until the end of the assay. Probably, a fifth phase might occur, where the vDNA quantity does not significantly vary over time, but this experiment did not arrive to such a plateau. These results are in accordance with previous results, where DNA replication started 5 to 6 h post infection and reached a maximum rate by 18 h post infection [53].

### 3.2. Discovering ORI Sequences in AgMNPV

As different kinds of sequences have been described as origins of replication in baculoviruses, the genome of AgMNPV was screened using a modified DpnI assay. Briefly, a virus genome library
containing overlapped fragments between 500 and 2000 bp was constructed and then 500 clones, which represent three times the genome coverage, were evaluated in groups of 25 by two independent transient replication assays (Figure S1).

Once the plasmids from infected cells had been recovered after an incubation time of 48 h in growth conditions (period that includes all replication phases previously identified by the kinetic assay), DNA was treated with DpnI to remove the original constructs derived from E. coli. Then, DNA was transformed into bacteria to select the plasmids multiplied by the viral machinery inside the insect cells. From the growing colonies, we randomly chose 16 of them to identify the viral sequences that could function as putative ORIs (DNA inserts of the recovered library constructs) were determined by direct sequencing. Thus, 14 genome regions were found and were classified according to their locations in three main groups: ORI-hr (homologous regions); ORI-igr (intergenic regions located between two annotated ORFs that may also contain small annotated ORFs); and ORI-gr (gene regions corresponding with sequences immersed into some annotated ORFs) (Figure 2; Table 1).

![Figure 2. AgMNPV ORI sequences. The illustration shows the physical maps of AgMNPV loci containing the segments recovered as ORI by the transient replication assay. They are indicated as green boxes and named using the corresponding clone number derived from the AgMNPV library. The annotated ORFs are represented with different grey shaded boxes (dark grey and light grey). Core genes are indicated in violet boxes and named according to the traditional denomination. The homologous regions (hrs) and direct repetitions (drs) are indicated using pink and light green boxes, respectively. Inside the circle that represents the genome, the AT content (gray line) and the profiles of appearance of 4mer (green line), 5mer (violet line), and 6mer (blue line) runs containing only A or T are shown.](image)

As expected by previous reports in other species, two virus regions contained a complete hr (Ori-hr, pBS-AGN39 contains hr9 and pBS-AGN113 contains hr7), supporting the capability of these loci to recruit the replicative machinery.
Table 1. Sequences containing ORIs of AgMNPV.

| ORI-hr | Clone Number | Position (bp) | Identity Sequence | Repetitions |
|--------|--------------|---------------|-------------------|-------------|
|        |              |               | hr9               | 8x          |
|        | 39           | 107,445–106,982 |                  |             |
|        | 113          | 64,519–63,441  | hr7               | 1x          |

| ORI-Igr | Clone Number | Position (bp) | Surrounding ORFs | ORFs Shared by |
|---------|--------------|---------------|------------------|----------------|
|         |              |               |                  |                |
|         | 27           | 45,424–44,663  | ORF60 (he65-like), ORF58 | Most GI, some GII and 3 Beta |
|         | 119/141      | 69,305–69,564  | ORF85 (ac78-like), ORF86 (vlf1) | 5 baculovirus genomes |
|         | 133          | 102,699–101,360 | ORF126, ORF125, ORF124 (lef-6), ORF123 (iap-1) | Gi |
|         |              |               |                  | Gi, most GII and Beta |
|         |              |               |                  | Alpha and Beta |
|         |              |               |                  | Gi, most GII, Beta |
|         | 33/162       | 52,036–52,450  | ORF67 (lef-5), ORF68 (38k) | Baculoviridae |
|         |              |               |                  | Baculoviridae |

| ORI-gr  | Clone Number | Position (bp) | ORF | ORFs Shared by |
|---------|--------------|---------------|-----|----------------|
|         |              |               |     |                |
|         | 26           | 118,470–118,641 | egt | Gi, GII, and most Beta |
|         | 104          | 69,883–70,458  | vlf-1 | Baculoviridae |
|         | 131          | 60,314–60,131  | vlf-1 | Baculoviridae |
|         | 136          | 90,683–91,393  | lef-8 | Baculoviridae |
|         | 144          | 74,559–74,844  | lef-3 | Alpha and Beta |
|         | 153          | 50,359–50,236  | p40  | Baculoviridae |
|         | 168          | 66,087–65,983  | vp91 | Baculoviridae |

The clone number column refers to the AgMNPV library clones selected for its ability to act as ORI ORI-hr (sequence containing homologous regions). ORI-Igr (sequence containing some partial or complete ORFs, including some intergenic region). ORI-gr (sequence immersed into an ORF). “Position” refers to the nucleotide position in the AgMNPV genome (NC_008520.2). Baculovirus classification clades are indicated as follows: GI (group I alphabaculoviruses), GII (group II alphabaculoviruses), alpha (alphabaculoviruses), beta (betabaculoviruses), Baculoviridae (all members sequenced of family Baculoviridae).
For the other recovered sequences, it is important to note that most of the 14 virus segments containing putative ORIs are next to or within ORFs. Interestingly, they are located inside genes shared by all baculoviruses (core genes). Two plasmids (pBS-AGN33 and pBS-AGN162) contain the same insert in the same polarity and four clones (pBS-AGN104, pBS-AGN119, pBS-AGN131, and pBS-AGN141) carry sequences of the same genome locus, adding value to the methodological approach chosen to find putative ORIs.

Regarding the ORI-grs, clones pBS-AGN104/131, pBS-AGN136, pBS-AGN153, and pBS-AGN168 contained virus segments located inside of core genes (vlf-1, lef-8, p40, and vp91, respectively). By contrast, only pBS-AGN26 and pBS-AGN144 clones contained sequences (egl and lef-3) that do not belong to any core genes but correspond to genes that are important to all lepidopteran baculoviruses [20,21]. Plasmids named ORI-igrs contained sequences composed by the 5' or 3' extremities of coding sequences and the non-coding region immersed between ORFs. Plasmid pBS-AGN119/141 contained a fragment of vlf-1 and ORF85 (ac78-like), both being core genes [20]. The clone pBS-AGN162/33 carried fragments of the core genes, ORF67 (lef-5) and ORF68 (ac98-like or 38k) [21]. However, pBS-AGN27, pBS-AGN133, and pBS-AGN150 did not contain fragments of core genes but contained sequences of genes shared by all group I alphabaculoviruses, reflecting that they could be important for AcMNPV and its lineage (Table 1; for more information, see Appendix A).

3.3. Characterization of ORI Sequences from AgMNPV

Since sequences of dsDNA rich in AT content can be more easily unwound and thus act as starting points to replication, we analyzed the AT contents in all the virus segments recovered by the modified transient replication assay previously described. We found that only a few clones had a higher AT percentage value than the average percentage of the AgMNPV genome, so we could not establish any correlation between the AT percentage inside clones and their capability to replicate, but we identified that they all have 4 to 6mer runs of A or T (Figure 2). Particularly the ORI-hrs were not located in genome regions rich in A or T runs, but some ORI-grs (pBS-AGN136, pBS-AGN144) and ORI-igrs (pBS-AGN27, pBS-AGN133) were positioned in the loci where the highest concentration of runs occurred.

Other sequence features were also investigated. Since it was previously reported that early promoters could function as ORI, the existence of typical patterns was researched. The presence of palindromes was predicted, thinking that this kind of sequence may form secondary structures (Table 2).

### Table 2. Palindromes and promoters contained in ORIs of AgMNPV.

| ORI | Clone Number | Length (bp) | Early | Late | INR | TATAAA | Perfect | Imperfect |
|-----|--------------|-------------|-------|------|-----|--------|---------|-----------|
| ORI-hr |            |     | 4*  | 3*  | -   | -      | -       | -         |
| 39  | 833         | 10 | 1   | -   | -   | -      | -       | -         |
| 113 | 1080        | 9*  | 11* | -   | 1*  | -      | -       | -         |
| ORI-igr |         |       | 2*  | 3*  | -   | -      | -       | -         |
| 26  | 172         | 2   | 2   | 2   | 2   | -      | -       | -         |
| 104 | 576         | 1   | 4   | -   | 1   | -      | -       | -         |
| 131 | 318         | 1   | 1   | 1   | 1   | -      | -       | -         |
| 136 | 711         | 7   | 10  | 1   | 2   | 1      | -       | -         |
| 144 | 286         | 2   | 5   | -   | -   | 7      | 4       | 3         |
| 153 | 132         | 2   | 2   | -   | -   | 2      | -       | -         |
| 168 | 105         | 2   | -   | -   | -   | -      | 5       | 1         |

The clone number column refers to the AgMNPV library clones selected for its ability to act as ORI. ORI-hr (sequence containing homologous regions). ORI-igr (sequence containing some partial or complete ORFs, including some intergenic region). ORI-gr (sequence immersed into an ORF). Plus (“+” in Promoter Pattern column) or minus (“-“ in Promoter Pattern column) refers to the strand of dsDNA (considering the plus strand the sequence published on NC_008520.2). *Promoters that could be functional by its position close to one ORF. INR: initiator motif.
The analysis of results showed that all segments contained early promoter patterns, but only a few were considered functional by its position relative to adjacent ORFs. As expected, ORI-igrs had a higher proportion of these motifs than the other ones (Table 2; for more information, see Appendix A).

Additionally, all sequences showed the presence of perfect and imperfect palindromes of different lengths. All detected ORIs had perfect palindromes of at least a 5 bp length. ORI-hrs and ORI-igrs revealed a greater number of perfect palindromes than ORI-grs (20.5 ± 13.4, 14.8 ± 8.6, and 5.3 ± 4.3, respectively). The imperfect palindromes found are seven to eight nucleotides in length. Moreover, imperfect palindromes containing more than 10 nucleotides were also detected in the three kinds of ORIs (for more information, see Appendix A).

As the AT content and the distribution of promoter motifs were not a shared property among all detected AgMNPV’s ORIs, the study continued with a focus on the secondary structure since all presented palindromes. Considering that this kind of sequence in double stranded-DNA can form cruciform structures, the virus regions recovered by the transient replication assay were explored to detect shared secondary structures. Notoriously, three hairpins that conform a unit, in some cases, repeated several times, were predicted in all ORIs from AgMNPV (Figure 3, Table 3). This common structure was slightly rich in AT content (50%–60%) and derived from DNA segments containing an average of 65 nucleotides (range of 39 o 111 bp) and could form three stems, where the first and third ones had an average of 4.6 nucleotides (ranges of 2–14 and 2–10, respectively), while the second one had 8.5 nucleotides (range: 5–16). The two linkers located between stems had an average of three nucleotides, ranging between 0 and 6. In all cases, the stem two was the longest one.

![Figure 3](image)

*Figure 3.* ORI sequences of AgMNPV and their putative structure. The illustration shows the putative secondary structures of the sequences recovered as ORI by the transient replication assay. ORIs are classified according their genome location and library clones are identified with their corresponding numbers. The nucleotide genome position of each sequence is indicated at the beginning of the DNA structure.
Considering that a similar structure was reported in 2014 for Suca jujuba NPV's hrs [54], we explored other baculovirus reported ORIs from different species. Thus, the hrs from AcMNPV [55], LdMNPV [56], SeMNPV [57], and non-hr ORIs from AcMNPV [42], SeMNPV [58], OpMNPV [59], and CrleGV [60] were studied. Surprisingly, the structure found in the AgMNPV's ORIs was also detected in the hrs of AcMNPV (hr1, hr1a, hr2, hr3, hr4a, hr4b, and hr5), of LdMNPV (hr1, hr2, hr3a, hr3b, hr3c, hr4, hr5, hr6, hr7a, hr7d, and hr8) and SeMNPV (hr2, hr3, hr5, and hr6), and in the all non-hr repetitive sequences of AcMNPV, SeMNPV, OpMNPV, and CrleGV (Table S1 and Figure S2). In some cases, the linkers are longer than the ones detected in AgMNPV replicative sequences, as in LdMNPV (hr7a and hr7d), but the pattern is conserved. Interestingly, the AcMNPV's hr1a is considered as sufficient for supporting plasmid replication [26] and could adopt a structure similar to stem two.

All analyses performed on the sequences from AgMNPV that acted as ORI were summarized using a Venn diagram (Figure 4). ORI-hrs are rich in AT content and A/T runs, they have palindromes; therefore, ORI-igrs have a similar value of AT content with respect to the genome average but share the other characteristics. Finally, ORI-igrs seem to be rich in A/T runs and palindromes, also enabling the formation of the secondary structure previously described.

Table 3. Putative DNA structures of AgMNPV ORIs.

| Clone Number | Number of Structures | Start (bp) | Length S1 (bp) | Linker 1 (nt) | Length S2 (bp) | Linker 2 (nt) | Length S3 (bp) | AT% S1 | Linker 1 | AT% S2 | Linker 2 | AT% S3 | End (bp) |
|--------------|---------------------|------------|----------------|---------------|----------------|---------------|----------------|--------|----------|--------|----------|--------|----------|
| 39           | 8                   | 107,396    | 3              | 6             | 2              | 4              | 77.7           | 100    | 0        | 53.8   | 107,348  |
| 107,380      | 4                   | 3              | 10             | 4              | 5              | 73.8           | 50              | 0      | 56.2    | 107,394|
| 107,397      | 5                   | 3              | 8              | 3              | 8              | 50            | 100             | 60.5   | 33.3    | 56.5   | 107,227  |
| 107,204      | 14                  | 6              | 15             | 6              | 3              | 50            | 66.7            | 62.5   | 66.7    | 107,010|

113 8 64,384 6 5 9 3 10 40.9 80 36.7 66.7 40 64,295
113 5 10 4 3 81.2 80 72.5 0 76.9 64,187
113 4 2 5 3 2 100 50 57.1 66.7 75 64,130
113 4 4 8 4 84.6 50 96.1 100 63.6 63,923
113 4 2 11 1 6 63.2 100 96.8 100 88.2 63,803
113 3 3 7 5 4 54.5 66.7 65.2 80 76.9 63,790
113 3 3 6 2 2 83.3 100 52.4 0 55.6 63,616
113 3 2 7 5 3 66.7 100 73.5 80 40.0 63,505

27 3 45,429 3 2 10 1 5 33.3 100 0 64.3 45,323
27 3 45,432 3 4 6 2 2 50.0 75.0 10.6 100 45,952
27 3 44,744 3 3 8 3 4 53.8 66.7 65.5 33.3 66.7 44,709

119/141 1 69,393 4 1 9 3 6 92.3 100.0 69.7 66.7 47.4 69,464
119/141 8 102,620 3 3 10 3 8 60.0 96.7 44.1 66.7 33.3 102,547
119/141 6 2 8 2 4 64.7 100.0 43.3 0.0 45.5 102,362
119/141 5 1 7 2 3 86.7 0.0 63.2 50.0 60.0 102,287
119/141 3 3 11 5 4 90.9 100.0 81.3 100.0 82.4 102,017
119/141 3 5 5 5 3 40.0 50.0 45.8 33.3 61.1 101,921
119/141 6 5 6 0 5 56.3 80.0 40.0 0 36.8 101,793
119/141 12 4 11 4 10 51.4 75.0 42.5 75.0 36.7 101,639
119/141 7 5 11 0 4 77.8 60.0 44.4 0 27.3 101,394

150 1 95,126 4 2 5 3 6 50.0 0 70.0 100.0 75.0 95,074
150 2 95,088 5 5 13 2 3 56.0 100 75 50.0 44 95,137
150 2 92,246 8 3 16 8 10 66.7 33.3 71.1 85.3 41.7 92,585

26 1 118,463 4 2 7 4 3 18.2 50.0 28.6 50.0 54.5 118,531

104 3 70,228 4 3 6 0 5 47.1 40 47.4 0 33.3 70,283
104 3 70,264 5 3 8 2 7 33.3 100 64.5 50 45 70,337
104 3 70,279 6 4 13 5 2 65.2 75.0 30 60.0 30.0 70,457

131 2 69,698 3 3 5 3 5 0.6 66.7 39 66.7 66.7 69,882
131 2 69,620 4 4 6 3 5 36.4 50 66.7 66.7 52.9 69,885

136 4 90,742 7 3 11 4 5 70 66.7 51.3 25 46.7 90,822
136 4 90,808 5 2 6 4 2 47.0 50.0 40.9 75 22.2 90,859
136 5 90,972 2 5 9 3 2 40 67 66.7 44 91,025
136 5 91,079 4 6 6 5 3 20 50 37 60.0 71 91,131

144 1 74,567 4 5 14 3 10 83.3 60.0 75.0 66.7 16.1 74,657
144 1 70,464 4 5 14 3 10 83.3 60.0 75.0 66.7 16.1 70,557
144 1 70,379 6 4 13 5 2 65.2 75.0 30 60.0 30.0 70,457

168 1 66,071 4 2 6 4 3 37.5 50.0 36.8 100.0 72.7 66,020

The clone number column refers to the AgMNPV library clones selected for its ability to act as ORI. ORI-hr (sequence containing homologous regions). ORI-igr (sequence containing some partial or complete ORFs, including some intergenic region). ORI-gr (sequence immersed into an ORF). The common structure found in all the AgMNPV ORIs is described in this table, indicating the first and last nucleotide involved (refers as Start and End columns). The length and AT content of each component (S1: stem 1; linker 1; S2: stem 2, linker 2; S3: stem 3). Different rows for each ORI in length and AT% columns contained data from each of the predicted structures.
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Figure 4. AgMNPV ORI sequences and their shared properties. Venn diagram shows shared properties among different ORIs of AgMNPV. The AT percentage has been selected only when values were over the average of the AgMNPV genome. Early promoters were considered functional when they are located close to the first ATG codon. In this graphic, only imperfect palindrome >10 nucleotides were considered.

4. Discussion

To successfully carry out the replication of dsDNA genomes, multi-protein complexes are needed to distinguish the starting points of DNA synthesis (ORIs), promote replisome formation, and then progress along the whole replicon. This whole process has been extensively studied for prokaryote and eukaryote organisms. However, the study of viruses becomes more complicated since it involves viral and host proteins. Two models had been proposed for baculovirus initiation: (a) Replication is coordinated with packaging, initiated in multiple ORIs to greatly amplify the viral DNA that can be produced in a given time; (b) replication is independent of packaging and undergoes extensive recombination [22]. However, data published until date does not permit discernment between those two possibilities. Furthermore, these models do not consider the two stages in the baculovirus life cycle: BVs and ODVs production. It is known that in eukaryotes, multiple origins of replication are activated during the cell cycle in a program precisely coordinated in space and time [61].

The results obtained on AgMNPV in this work support the existence of different phases during baculovirus infection. In agreement with previous publications [62], we detected an early phase (phase one) in which BVs require about 5 h to enter the cell, migrate across the cytoplasm, enter into the nuclei, and hijack host transcription machinery to finally start replication. Then, a first exponential phase of replication occurs until the beginning of budding (phase two), an event that initiates a decrease in DNA productivity (phase three). Later, this stage is followed by phase four, characterized by another increase in genome synthesis. This behavior could be due to a change of the mechanism involved in DNA replication probably associated with the production of late virus proteins. DNA synthesis rates (μV) were similar in the two exponential phases. Since part of the DNA that could be used for replication is constantly removed to be packed into BVs and ODVs during phase four, it is possible to say that a different process of replication is being carried out.
The different loci identified here could recruit the replication machinery at some point during the infection cycle and allow the different replication stages by using different kinds of origins. It has been discussed before that despite the mechanism of replication (Theta or Sigma), baculoviruses may utilize multiple origins with different levels of initiation efficiency, starting primarily at the hr origin [63].

Analyzing the results from the transient assay, we found that two of the plasmids able to replicate contained hrs sequences. Most baculoviruses have several hrs dispersed over the genome [22,33], varying from 3 in CrleGV [64] to 17 in SpltNPV-G2 [65] but cannot be considered core sequences. All group I alphabaculoviruses have hrs but not all group II species, such as ChchNPV [66] and TnSNPV [67]. Although hrs are not conserved in all baculoviruses, they have been detected as ORI as demonstrated by transfection replication assays in different species, including AcMNPV [39,68], SeMNPV [57], CfMNPV [69], and LdMNPV [56]. In AcMNPV, the study showed that plasmids containing hrs replicate 5- to 19-fold higher than those containing other genomic DNA fragments, and the replication rates were associated with the number of palindromes or their orientation [68]. However, recombinant viruses, where hr1, hr1a, hr2, hr3, hr4, or hr4b were knocked-out, revealed that none of hrs would be essential for DNA replication [39]. In our work, the selection of fragments was done without bias, thus the appearance of hr adds new evidence of its function as ORIs.

Furthermore, our data opens a new panorama regarding the kind of sequences that can be used as origins of replication in AgMNPV. Apart from hrs, other ORIs containing tandemly repeated sequences have been characterized in AcMNPV [39,42], SpliMNPV [70], SeMNPV [58,71,72], OpMNPV [33,59], and CrleGV [60]. Also, in AcMNPV, two other kinds of non-hr ORIs have been described: Early promoters [34], and a segment of the DNA helicase ORF [35]. In our work, 12 different sequences of AgMNPV classified as intergenic and intragenic regions were detected and named ORI-igrs and ORI-grs, according to the presence of putative functional early promoters or the presence of fragments of ORFs, respectively. In this work, one of the fragments (pBS-AGN150, classified as ORI-igr) contains a direct repetition (dr12, 95,016-95,057 nt). Why are the sequences found in AgMNPV able to initiate replication? Possible answers should include the role of IE-1 protein because both early promoters and hrs are sequences able to bind to that polypeptide [73,74]. For example, AcMNPV IE-1 binds to the hr palindromic sequences [75] and this interaction would be essential to function as transcriptional enhancers [76]. Besides, it has been proposed that the palindromic core of AcMNPV hr (28 bp) acts as ORI, whereas the flanking sequences are required for enhancer activity [68]. Thus, IE-1 could be an ORI binding protein or be necessary to transactivate all early genes involved in DNA replication [20]. As mentioned above, AdorGV, AgseGV, CpGV, CrleGV, PrGV, and SpliGV have no hrs, although they have ie-1 orthologs [77–79]. Therefore, the recovery of hrs (pBS-AGN39, pBS-AGN113) and regions containing early promoters (pBS-AGN27, pBS-AGN119/141, pBS-AGN133, pBS-AGN141, pBS-AGN150, and pBS-AGN33/162) makes sense considering that IE-1 would be an ORI binding protein.

The reason why intragenic regions are able to initiate replication processes still remains to be answered. Probably, a common secondary structure might be the need factor recognized by the replication machinery to initiate DNA synthesis. In fact, there are reports in eukaryotic genomes about the presence of loops on mitochondrial ORI [80] as well as in circovirus [81] and herpesvirus ORIs [82]. It is possible that particular palindromic regions form a curve or cruciform structure that are recognized by the proteins responsible for the initiation of DNA synthesis [29,55]. As it is known, both transcription and replication require the local unwinding of double stranded-DNA. In these processes, the generated single stranded-DNA is used as a template for the synthesis of RNA transcripts or to produce daughter strands. Theoretically, the hrs palindromes may form cruciform structures although some evidences [55] indicate that this situation only occurs when the imperfect palindromes are modified to be perfect. However, the formation of a cruciform structure under certain conditions during viral replication or the stabilization by protein attachment or by particular neighboring sequences cannot be discarded. Again, IE-1 could play an important role in replication by recruiting a replication complex, since it has been shown that it can bind to hrs in their cruciform or extended shape [55]. In AcMNPV, a single palindrome repeat from an hr is
a minimal requirement for plasmid dependent DNA replication [26], and a single palindrome could adopt a hairpin structure.

The presence of palindromes was detected in all the sequences found here. Thus, the DNA secondary structure could be the key feature recognized by the replication machinery in baculoviruses to initiate DNA synthesis and be positively selected in preference of a primary sequence. Many origins of replication contain inverted repeats, since the formation of a cruciform structure is energetically more favorable than simple DNA melting and can absorb part of the strand generated folding it into hairpins [83].

An ORI was also reported inside the p143 gene of AcMNPV. This sequence contains 13 imperfect palindromic sequences of 12 to 18 bp in size [35]. Furthermore, the intergenic and intragenic fragments identified as ORI in our work are located inside or close to core genes, the common ancestral set of sequences conserved by all baculoviruses. This proximity could ensure their transfer to progeny and thus be sustained through the evolution, allowing them to conserve at least one mechanism of replication, which is still functional in all species. The presence of many ORIs around the genome could provide a redundancy required to ensure DNA replication. In vivo evidence supports the hypothesis that AcMNPV replication involves multiple ORIs that are activated with different efficiency during the viral infection cycle [42]. According to our results, the same could occur in AgMNPV.

The fact of having found and predicted a shared structure in all the sequences that act as ORI in baculoviruses, both those found by this work in AgMNPV and those found by other authors in other species, is in itself very relevant because such types of structures are usually key factors in replicative processes in other biological systems [83]. For example, the phage G4 (an ssDNA virus from Escherichia coli) carries three hairpins with stems of 5 to 19 bp and loops of 4 to 8 bases in the ORI of replication, which seems to direct the binding of single-stranded binding proteins [84]. The dsDNA bacterial plasmid R6K replicates by the theta mechanism and a DNA hairpin named M13-A is the core of the priming initiation [85]. In the plasmid RSF1010, which replicates by strand displacement, RepC protein binds to iterons and unwinds the dsDNA in a region that folds into hairpins [86]. In turn, the plasmid pT181 replicates by the rolling circle and contains a hairpin with the Rep nicking site, which initiates replication [87]. Thus, the structures that we have predicted in baculoviral ORIs are frequent in nature and are associated with replication functions under various mechanisms compatible with those that could occur in baculoviruses. Consequently, they would constitute the essential characteristic that requires a baculoviral sequence to be the initiator of the DNA synthesis process.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/1999-4915/11/7/648/s1, Figure S1: Workflow to discover ORI sequences in AgMNPV, Figure S2: Baculovirus ORI sequences and their putative structure, Table S1: Putative DNA structures of Baculovirus ORIs.

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Appendix A

Genes Contained in the Different Clones (Table 1)

pBS-AGN133 contained fragments of different ORFs (ORF126, ORF125, ORF124, ORF123). ORF126 (ac30-like) has homologs in the genomes of all group I Alphabaculoviruses, ORF125 (ac29-like) is found in all group I and most group II NPVs and GV genomes, ORF124 (ac28-like or lef-6) has homologs in all lepidopteran NPVs and GVs, and ORF123 (ac27-like or iap-1) is found only in group I alphabaculoviruses [22]. Albeit, most of those genes are not essential, a deletion of lef-6 on BmNPV severely compromised the virus [88]. The clone pBS-AGN150 had fragments of ORF115 (sequence with homologs in all group I alphabaculoviruses and in SeMNPV) and ORF114 (odv-e66), a sequence shared by all group I alphabaculoviruses and betabaculoviruses and most group II alphabaculoviruses.

Early Promoters (Table 2)

pBS-AGN27 contained a non-coding region flanked by the first 569 bp of he65 (ORF60) and the first 8 bp of pnk/pnl (ORF57), which presented three early promoters in the orientation of the first gene and four in the other one. In the cases of pBS-AGN119 and pBS-AGN141, segments found should contain the promoter region of the vlf-1 gene because this region is located 161 bp upstream of the initial codon and three early and one late promoter were detected in the proper orientation. PBS-AGN133 contained the last 371 bp of ORF126, a small intergenic region (41 bp), complete ORF125 (210 bp), another intergenic region (32 bp), ORF124 (432 bp), and the last 248 bp of ORF123. Therefore, the first 415 bp and the last 248 bp could be regulatory regions in opposite polarities. Particularly, 11 early promoters were detected in the plus strand, but only one was considered functional as it was 155 bp upstream of ORF125’s ATG. Another 11 early promoters were detected in the other polarity. In this case, two motifs could be functional as they were 80 and 118 bp upstream of ORF124’s ATG, and one late promoter was located 166 bp upstream of the ORF. PBS-AGN150 contained the last 127 bp of ORF115, an intergenic region (50 bp) and the first 226 bp of ORF114. In this zone 5 early promoters were detected, but only 2 were in the intergenic region. Also, an initiation motif (INR) of the viral RNA polymerase was found in the intergenic region 16 bp upstream of the ATG, and a late promoter was detected 91 bp upstream of the ORF. PBS-AGN162/33 contained the first 197 bp of lef-5, an intergenic region, and the first 112 bp of ORF68 in the opposite polarity. An INR was detected in the intergenic region. Also, one early promoter was detected 88 bp upstream of ORF68. In the other polarity, three early promoters were located at 97, 137, and 185 bp away from the ATG of lef-5. By contrast, the late promoters, initiator motifs (INR), and TATA boxes were not very represented in this segment.

Palindromes (Table 3)

As mentioned before, all sequences showed the presence of perfect and imperfect palindromes. All detected ORIs had perfect palindromes of at least 5 bp length, with a minimum of 2 for the ORI-grs contained in pBS-AGN26/pBS-AGN153/pBS-AGN-168 and a maximum of 30 for the ORI-hr pBS-AGN113. Anyway, ORI-hrs and ORI-igrs revealed a greater number of perfect palindromes than ORI-grs (20.5 ± 13.4, 14.8 ± 8.6, and 5.3 ± 4.3, respectively). Even this observation applies for longer perfect palindromes or for imperfect palindromes of any length. Moreover, all fragments presented at least 1 imperfect palindrome of 7 or 8 nucleotides in length and imperfect palindromes containing more than 10 nucleotides were also detected in the three kinds of ORIs.

References
1. Miele, S.A.B.; Garavaglia, M.J.; Belaich, M.N.; Ghiringhelli, P.D. Baculovirus: Molecular Insights on Their Diversity and Conservation. *Int. J. Evol. Biol.* 2011, 2011, 1–15. [CrossRef]
2. Blissard, G.W.; Rohrmann, G.F. Baculovirus Diversity and Molecular Biology. *Annu. Rev. Entomol.* 1990, 35, 127–155. [CrossRef]
3. Braconi, C.T.; Ardisson-Araujo, D.M.P.; Leme, A.F.P.; Oliveira, J.V.; Pauletti, B.A.; Garcia-Maruniak, A.; Ribeiro, B.M.; Maruniak, J.E.; Zanotto, P.M. Proteomic analyses of baculovirus Anticarsia gemmatalis multiple nucleopolyhedrovirus budded and occluded virus. *J. Gen. Virol.* 2014, 95, 980–989. [CrossRef]

4. Cheshenko, N.; Krougliak, N.; Eisensmith, R.C.; Krougliak, V.A. A novel system for the production of fully deleted adenovirus vectors that does not require helper adenovirus. *Gene Ther.* 2001, 8, 846–854. [CrossRef]

5. Bieniossek, C.; Imasaki, T.; Takagi, Y.; Berger, I. MultiBac: Expanding the research toolbox for multiprotein complexes. *Trends Biochem. Sci.* 2012, 37, 49–57. [CrossRef]

6. Gupta, K.; Tölzer, C.; Sari-Ak, D.; Fitzgerald, D.J.; Schaffitzel, C.; Berger, I. MultiBac: Baculovirus-Mediated Multigene DNA Cargo Delivery in Insect and Mammalian Cells. *Viruses* 2019, 11, 198. [CrossRef]

7. Inceoglu, A.B.; Kamita, S.G.; Hinton, A.C.; Huang, Q.; Severson, T.F.; Kang, K.; Hammock, B.D. Recombinant baculoviruses for insect control. *Pest Manag. Sci.* 2001, 57, 981–987. [CrossRef]

8. Inceoglu, A.B.; Kamita, S.G.; Hammock, B.D. Genetically Modified Baculoviruses: A Historical Overview and Future Outlook. *Adv. Virus Res.* 2006, 68, 323–360.

9. Szewczyk, B.; Hoyos-Carvajal, L.; Paluszek, M.; Skrzecz, I.; Lobo de Souza, M. Baculoviruses — re-emerging biostatistics. *Biotechnol. Adv.* 2006, 24, 143–160. [CrossRef]

10. Popham, H.J.R.; Nusawardani, T.; Bonning, B.C. Introduction to the Use of Baculoviruses as Biological Insecticides. *Methods Mol. Biol.* 2016, 1350, 383–392.

11. Van der Merwe, M.; Jukes, M.D.; Rabalski, L.; Knox, C.; Opoku-Debrah, J.K.; Moore, S.D.; Krejmer-Rabalska, M.; Szewczyk, B.; Hill, M.P. Genome Analysis and Genetic Stability of the Cryptophlebia leucotreta Granulovirus (CrleGV-SA) after 15 Years of Commercial Use as a Biopesticide. *Int. J. Mol. Sci.* 2017, 18, 2327. [CrossRef]

12. Kost, T.A.; Kemp, C.W. Fundamentals of Baculovirus Expression and Applications. *Adv. Exp. Med. Biol.* 2016, 896, 187–197.

13. Irons, S.L.; Chambers, A.C.; Lissina, O.; King, L.A.; Possee, R.D. Protein Production Using the Baculovirus Expression System. In *Current Protocols in Protein Science*; John Wiley & Sons, Inc.: Hoboken, NJ, USA, 2018; pp. 5.5.1–5.5.22.

14. Ono, C.; Okamoto, T.; Abe, T.; Matsuura, Y. Baculovirus as a Tool for Gene Delivery and Gene Therapy. *Viruses* 2018, 10, 510. [CrossRef]

15. Felberbaum, R.S. The baculovirus expression vector system: A commercial manufacturing platform for viral vaccines and gene therapy vectors. *Biotechnol. J.* 2015, 10, 702–714. [CrossRef]

16. Mansouri, M.; Berger, P. Baculovirus for gene delivery to mammalian cells: Past, present and future. *Plasmid* 2018, 98, 1–7. [CrossRef]

17. Wennmann, J.T.; Keilwagen, J.; Jehle, J.A. Baculovirus Kimura two-parameter species demarcation criterion is confirmed by the distances of 38 core gene nucleotide sequences. *J. Gen. Virol.* 2018, 99, 1307–1320. [CrossRef]

18. Thézé, J.; Lopez-Vaamonde, C.; Cory, J.S.; Herniou, E.A. Biodiversity, Evolution and Ecological Specialization of Baculoviruses: A Treasure Trove for Future Applied Research. *Viruses* 2018, 10, 366. [CrossRef]

19. Gao, L.; Qi, J. Whole genome molecular phylogeny of large dsDNA viruses using composition vector method. *BMC Evol. Biol.* 2007, 7, 41. [CrossRef]

20. Javed, M.A.; Biswas, S.; Willis, L.G.; Harris, S.; Pritchard, C.; van Oers, M.M.; Donly, B.C.; Erlandson, M.A.; Hagedus, D.D.; Theilmann, D.A. Autographa californica Multiple Nucleopolyhedrovirus AC83 is a Per Os Infectivity Factor (PIF) Protein Required for Occlusion-Derived Virus (ODV) and Budded Virus Nucleocapsid Assembly as well as Assembly of the PIF Complex in ODV Envelopes. *J. Virol.* 2017, 91. [CrossRef]

21. Garavaglia, M.J.; Miele, S.A.B.; Iserte, J.A.; Belaich, M.N.; Ghiringhelli, P.D. The ac53, ac78, ac101, and ac103 Genes Are Newly Discovered Core Genes in the Family Baculoviridae. *J. Virol.* 2012, 86, 12069–12079. [CrossRef]

22. Rohrmann, G.F. *Baculovirus Molecular Biology*, 3rd ed.; National Center for Biotechnology Information (US): Bethesda, MD, USA, 2013.

23. Carstens, E.B. AcMNPV as a model for baculovirus DNA replication. *Virol. Sin.* 2009, 24, 243–267. [CrossRef]

24. Mikhailov, V.S.; Rohrmann, G.F. Characterization of short-lived intermediates produced during replication of baculovirus DNA. *Virus Res.* 2009, 145, 106–111. [CrossRef]

25. Anderson, S.; DePamphilis, M.L. Metabolism of Okazaki fragments during simian virus 40 DNA replication. *J. Biol. Chem.* 1979, 254, 11495–11504.
26. Leisy, D.J.; Rohrmann, G.F. Characterization of the replication of plasmids containing hr sequences in baculovirus-infected Spodoptera frugiperda cells. *Virology* 1993, 196, 722–730. [CrossRef]
27. Oppenheimer, D.I.; Volkman, L.E. Evidence for rolling circle replication of Autographa californica M nucleopolyhedrovirus genomic DNA. *Arch. Virol.* 1997, 142, 2107–2113. [CrossRef]
28. Okano, K.; Vanarsdall, A.L.; Rohrmann, G.F. A baculovirus alkaline nuclease knockout construct produces fragmented DNA and aberrant capsids. *Virology* 2007, 359, 46–54. [CrossRef]
29. Wu, Y.; Liu, G.; Carstens, E.B. Replication, integration, and packaging of plasmid DNA following cotransfection with baculovirus viral DNA. *J. Virol.* 1999, 73, 5473–5480.
30. Pearson, M.; Bjornson, R.; Pearson, G.; Rohrmann, G. The Autographa californica baculovirus genome: Evidence for multiple replication origins. *Science* 1992, 209, 1392–1396. [CrossRef]
31. Kool, M.; van den Berg, P.M.M.M.; Tramper, J.; Goldbach, R.W.; Vlak, J.M. Location of Two Putative Origins of DNA Replication of Autographa californica Nucleopolyhedrosis Virus. *Virology* 1993, 192, 94–101. [CrossRef]
32. Lee, H.Y.; Krell, P.J. Generation and analysis of defective genomes of Autographa californica nuclear polyhedrosis virus. *J. Virol.* 1992, 66, 4339–4347.
33. Pearson, M.N.; Bjornson, R.M.; Ahrens, C.; Rohrmann, G.F. Identification and characterization of a putative origin of DNA replication in the genome of a baculovirus pathogenic for Orgyia pseudotsugata. *Virology* 1993, 197, 715–725. [CrossRef]
34. Wu, Y.; Carstens, E.B. Initiation of baculovirus DNA replication: Early promoter regions can function as infection-dependent replicating sequences in a plasmid-based replication assay. *J. Virol.* 1996, 70, 6967–6972.
35. Wu, Y.-L.; Wu, C.-P.; Huang, Y.-H.; Huang, S.-P.; Lo, H.-R.; Chang, H.-S.; Lin, P.-H.; Wu, M.-C.; Chang, C.-J.; Chao, Y.-C. Identification of a high-efficiency baculovirus DNA replication origin that functions in insect and mammalian cells. *J. Virol.* 2014, 88, 13073–13085. [CrossRef]
36. Guarino, L.A.; Summers, M.D. Interspersed Homologous DNA of Autographa californica Nuclear Polyhedrosis Virus Enhances Delayed-Early Gene Expression. *J. Virol.* 1986, 60, 215–223.
37. Cochran, M.A.; Faulkner, P. Location of Homologous DNA Sequences Interspersed at Five Regions in the Baculovirus AcMNPV Genome. *J. Virol.* 1983, 45, 961–970.
38. Kool, M.; Voeten, J.T.; Goldbach, R.W.; Tramper, J.; Vlak, J.M. Identification of seven putative origins of Autographa californica multiple nucleocapsid nuclear polyhedrosis virus DNA replication. *J. Gen. Virol.* 1993, 74 (Pt 12), 2661–2668. [CrossRef]
39. Carstens, E.B.; Wu, Y. No single homologous repeat region is essential for DNA replication of the baculovirus Autographa californica multiple nucleocapsid nuclear polyhedrosis virus DNA replication. *J. Gen. Virol.* 1997, 88, 114–122. [CrossRef]
40. Friesen, P.D.; Miller, L.K. Divergent transcription of early 35- and 94-kilodalton protein genes encoded by the HindIII K genome fragment of the baculovirus *Autographa californica* multiple nucleopolyhedrovirus. *J. Virol.* 1987, 61, 2264–2272.
41. Oliveira, J.V.; Wolfl, J.L.C.; Garcia-Maruniak, A.; Ribeiro, B.M.; de Castro, M.E.B.; de Souza, M.L.; Moscardi, F.; Maruniak, J.E.; Zanotto, P.M. Genome of the most widely used viral biopesticide: Anticarsia gemmatalis multiple nucleopolyhedrovirus. *J. Gen. Virol.* 2006, 87, 3323–3325. [CrossRef]
42. Kool, M.; Goldbach, R.W.; Vlak, J.M. A putative non-hr origin of DNA replication in the HindIII-K fragment of Autographa californica multiple nucleocapsid nuclear polyhedrosis virus. *J. Gen. Virol.* 1994, 75 (Pt 12), 3345–3352. [CrossRef]
43. Sieburth, P.J.; Maruniak, J.E. Growth characteristics of a continuous cell line from the velvetbean caterpillar, *Anticarsia gemmatalis* Hübner (Lepidoptera: Noctuidae). *Vitr. Cell. Dev. Biol.* 1988, 24, 195–198. [CrossRef]
44. Zanotto, P.M.; Sampaio, M.J.; Johnson, D.W.; Rocha, T.L.; Maruniak, J.E. The Anticarsia gemmatalis nuclear polyhedrosis virus polyhedrin gene region: Sequence analysis, gene product and structural comparisons. *J. Gen. Virol.* 1992, 73 (Pt 5), 1049–1056. [CrossRef]
45. Menguál Gómez, D.I.; Balach, M.N.; Rodriguez, V.A.; Ghiringhelli, P.D. Effects of fetal bovine serum deprivation in cell cultures on the production of *Anticarsia gemmatalis* multinucleopolyhedrovirus. *BMC Biotechnol.* 2010, 10, 68. [CrossRef]
46. Kost, T.A.; Condraey, J.P. Recombinant baculoviruses as expression vectors for insect and mammalian cells. *Curr. Opin. Biotechnol.* 1999, 10, 428–433. [CrossRef]
47. Green Michel R, S.J. *Molecular Cloning: A Laboratory Manual*, 4th ed.; Cold Spring Harbor Laboratory Press: New York, NY, USA, 2012; ISBN 978-1-936113-42-2.
48. Chen, Y.-R.; Zhong, S.; Fei, Z.; Hashimoto, Y.; Xiang, J.Z.; Zhang, S.; Blissard, G.W. The transcriptome of the baculovirus Autographa californica multiple nucleopolyhedrovirus in Trichoplusia ni cells. *J. Virol.* **2013**, *87*, 6391–6405. [CrossRef]

49. Stothard, P. The Sequence Manipulation Suite: JavaScript Programs for Analyzing and Formatting Protein and DNA Sequences. *Biotechniques* **2000**, *28*, 1102–1104. [CrossRef]

50. Carver, T.; Bleasby, A. The design of Jemboss: A graphical user interface to EMBOSS. *Bioinformatics* **2003**, *19*, 1837–1843. [CrossRef]

51. Mathews, D.H. Using the RNAstructure Software Package to Predict Conserved RNA Structures. In *Current Protocols in Bioinformatics*; John Wiley & Sons, Inc.: Hoboken, NJ, USA, 2014; Volume 46, pp. 12.4.1–12.4.22.

52. Zuker, M. Mfold web server for nucleic acid folding and hybridization prediction. *Nucleic Acids Res.* **2003**, *31*, 3406–3415. [CrossRef]

53. Tjia, S.T.; Carstens, E.B.; Doerfler, W. Infection of *Spodoptera frugiperda* cells with *Autographa californica* nuclear polyhedrosis virus II. The viral DNA and the kinetics of its replication. *Virology* **1979**, *99*, 399–409. [CrossRef]

54. Liu, X.; Yin, F.; Zhu, Z.; Hou, D.; Wang, J.; Zhang, L.; Wang, M.; Wang, H.; Hu, Z.; Deng, F. Genomic sequencing and analysis of *Sucra jujuba* nucleopolyhedrovirus. *PLoS ONE* **2014**, *9*, e110023. [CrossRef]

55. Rasmussen, C.; Leisy, D.J.; Ho, P.S.; Rohrmann, G.F. Structure-function analysis of the *Autographa californica* multinucleocapsid nucleopolyhedrosis virus homologous region palindromes. *J. Gen. Virol.* **1996**, *78* (Pt 6), 1497–1506. [CrossRef]

56. Pearson, M.N.; Rohrmann, G.F. Lymantria dispar nuclear polyhedrosis virus homologous regions: Characterization of their ability to function as replication origins. *J. Virol.* **1995**, *69*, 213–221.

57. Broer, R.; Heldens, J.G.; van Strien, E.A.; Zuidema, D.; Vlak, J.M. Specificity of multiple homologous genomic regions in *Spodoptera exigua* multinucleocapsid nucleopolyhedrosis virus replication. *J. Gen. Virol.* **1998**, *79* (Pt 6), 1563–1572. [CrossRef]

58. Heldens, J.G.; Broer, R.; Zuidema, D.; Goldbach, R.W.; Vlak, J.M. Identification and functional analysis of a non-hr origin of DNA replication in the genome of *Spodoptera exigua* multicapsid nucleopolyhedrovirus. *J. Gen. Virol.* **2002**, *83*, 2025–2034. [CrossRef]

59. Ahrens, C.H.; Pearson, M.N.; Rohrmann, G.F. Identification and Characterization of a Second Putative Origin of DNA Replication in a Baculovirus of *Orgyia pseudotsugata*. *Virology* **1995**, *207*, 572–576. [CrossRef]

60. Jehle, J.A. The expansion of a hypervariable, non-hr ori-like region in the genome of *Cryptophlebia leucotreta* granulovirus provides in vivo evidence for the utilization of baculovirus non-hr oris during replication. *J. Gen. Virol.* **2002**, *83*, 2025–2034. [CrossRef]

61. Ciardo, D.; Goldar, A.; Marheineke, K. On the Interplay of the DNA Replication Program and the Intra-S Phase Checkpoint Pathway. *Genes* **2019**, *10*, 94. [CrossRef]

62. Oliveira, J.V.; de Brito, A.F.; Braconi, C.T.; de Melo Freire, C.C.; Iamarino, A.; de Andrade Zanotto, P.M. Modularity and evolutionary constraints in a baculovirus gene regulatory network. *BMC Syst. Biol.* **2013**, *7*, 87. [CrossRef]

63. Habib, S.; Hasnain, S.E. Differential activity of two non-hr origins during replication of the baculovirus Autographa californica nuclear polyhedrosis virus genome. *J. Virol.* **2000**, *74*, 5182–5189. [CrossRef]

64. Lange, M.; Jehle, J.A. The genome of the *Cryptophlebia leucotreta* granulovirus. *Virology* **2003**, *317*, 220–236. [CrossRef]

65. Pang, Y.; Yu, J.; Wang, L.; Hu, X.; Bao, W.; Li, G.; Chen, C.; Han, H.; Hu, S.; Yang, H. Sequence Analysis of the *Spodoptera litura* Multicapsid Nucleopolyhedrovirus Genome. *Virology* **2001**, *287*, 391–404. [CrossRef]

66. Van Oers, M.M. Genome sequence of *Chrysodeixis chalcites* nucleopolyhedrovirus, a baculovirus with two DNA photolyase genes. *J. Gen. Virol.* **2005**, *86*, 2069–2080. [CrossRef]

67. Willis, L.G.; Siepp, R.; Stewart, T.M.; Erlandson, M.A.; Theilmann, D.A. Sequence analysis of the complete genome of *Trichoplusia ni* single nucleopolyhedrovirus and the identification of a baculoviral photolyase gene. *Virology* **2005**, *338*, 209–226. [CrossRef]

68. Habib, S.; Pandey, S.; Chatterji, U.; Burma, S.; Ahmad, R.; Jain, A.; Hasnain, S.E. Bifunctionality of the AcMNPV homologous region sequence (hr1): Enhancer and ori functions have different sequence requirements. *DNA Cell Biol.* **1996**, *15*, 737–747. [CrossRef]
69. Xie, W.D.; Arif, B.; Dobos, P.; Krell, P.J. Identification and analysis of a putative origin of DNA replication in the Choristoneura fumiferana multinucleocapsid nuclear polyhedrosis virus genome. *Virology* **1995**, *209*, 409–419. [CrossRef]

70. Levin, D.B.; Huang, J. Identification and functional analysis of a putative non-hr origin of DNA replication from the Spodoptera littoralis type B multinucleocapsid nucleopolyhedrovirus. *J. Gen. Virol.* **1999**, *80*, 2263–2274.

71. Pijlman, G.P.; Dortmans, J.C.F.M.; Vermeesch, A.M.G.; Yang, K.; Martens, D.E.; Goldbach, R.W.; Vlak, J.M. Pivotal role of the non-hr origin of DNA replication in the genesis of defective interfering baculoviruses. *J. Virol.* **2002**, *76*, 5605–5611. [CrossRef]

72. Pijlman, G.P.; Vermeesch, A.M.G.; Vlak, J.M. Cell line-specific accumulation of the baculovirus non-hr origin of DNA replication in infected insect cells. *J. Invertebr. Pathol.* **2003**, *84*, 214–219. [CrossRef]

73. Guarino, L.A.; Gonzalez, M.A.; Summers, M.D. Complete Sequence and Enhancer Function of the *Adoxophyes orana* granulovirus genome. *Virology* **2003**, *311*, 350–365. [CrossRef]

74. Guarino, L.A.; Gonzalez, M.A.; Summers, M.D. Complete Sequence and Enhancer Function of the Homologous DNA Regions of Autographa californica Nuclear Polyhedrosis Virus. *J. Virol.* **1986**, *60*, 224–229.

75. Rodems, S.M.; Friesen, P.D. Transcriptional enhancer activity of hr5 requires dual-palindrome half sites that mediate binding of a dimeric form of the baculovirus transregulator IE1. *J. Virol.* **1995**, *69*, 5368–5375.

76. Leisy, D.J.; Rasmussen, C.; Kim, H.T.; Rohrmann, G.F. The Autographa californica nuclear polyhedrosis virus homologous region 1a: Identical sequences are essential for DNA replication activity and transcriptional enhancer function. *Virology* **1995**, *208*, 742–752. [CrossRef]

77. Wormleaton, S.; Kuzio, J.; Winstanley, D. The complete sequence of the *Adoxophyes orana* granulovirus genome. *Virology* **2003**, *311*, 350–365. [CrossRef]

78. Crook, N.; Luque, T.; O’Reilly, D.R.; Winstanley, D.; Finch, R. The complete sequence of the *Cydia pomonella* granulovirus genome. *J. Gen. Virol.* **2001**, *82*, 2531–2547.

79. Wang, Y.; Choi, J.Y.; Roh, J.Y.; Liu, Q.; Tao, X.Y.; Park, J.B.; Kim, J.S.; Je, Y.H. Genomic Sequence Analysis of Granulovirus Isolated from the Tobacco Cutworm, *Spodoptera litura*. *PLoS ONE* **2011**, *6*, e28163. [CrossRef]

80. Ki, J.-S.; Hop, H.; Kim, S.-J.; Kim, I.-C.; Park, H.G.; Lee, J.-S. Complete mitochondrial genome sequence of the Arctic gammarid, Onisimus nanseni (Crustacea; Amphipoda): Novel gene structures and unusual control region features. *Comp. Biochem. Physiol. Part D Genom. Proteom.* **2010**, *5*, 105–115. [CrossRef]

81. Cheung, A.K. Porcine circovirus: Transcription and DNA replication. *Virus Res.* **2012**, *164*, 46–53. [CrossRef]

82. Macao, B.; Olsson, M.; Elias, P. Functional Properties of the Herpes Simplex Virus Type I Origin-binding Protein Are Controlled by Precise Interactions with the Activated Form of the Origin of DNA Replication. *J. Biol. Chem.* **2004**, *279*, 29211–29217. [CrossRef]

83. Bikard, D.; Loot, C.; Baharoglu, Z.; Mazel, D. Folded DNA in action: Hairpin formation and biological functions in prokaryotes. *Microbiol. Mol. Biol. Rev.* **2010**, *74*, 570–588. [CrossRef]

84. Sun, W.; Godson, G.N. Structure of the *Escherichia coli* primase/single-strand DNA-binding protein/phage G4oric complex required for primer RNA synthesis. *J. Mol. Biol.* **1998**, *276*, 689–703. [CrossRef]

85. Masai, H.; Nomura, N.; Arai, K. The ABC-primosome. A novel priming system employing dnaA, dnaB, dnaC, and primase on a hairpin containing a dnaA box sequence. *J. Biol. Chem.* **1990**, *265*, 15134–15144.

86. Miao, D.M.; Honda, Y.; Tanaka, K.; Higashi, A.; Nakamura, T.; Taguchi, Y.; Sakai, H.; Komano, T.; Bagdasarian, M. A base-paired hairpin structure essential for the functional priming signal for DNA replication of the broad host range plasmid RSF1010. *Nucleic Acids Res.* **1993**, *21*, 4900–4903. [CrossRef]

87. Noirot, P.; Bargonetti, J.; Novick, R.P. Initiation of rolling-circle replication in pT181 plasmid: Initiator protein enhances cruciform extrusion at the origin. *Proc. Natl. Acad. Sci. USA* **1990**, *87*, 8560–8564. [CrossRef]

88. Lin Guangyun, B.G.W. Analysis of an Autographa californica Nucleopolyhedrovirus lef-11 Knockout: LEF-11 Is Essential for Viral DNA Replication. *J. Virol.* **2002**, *76*, 2770–2779.

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