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Chromosomal integration of an avian oncogenic herpesvirus reveals telomeric preferences and evidence for lymphoma clonality

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Abstract

Background: Herpesviruses are a major health concern for numerous organisms, including humans, causing both acute and chronic infections recurrent over an individual's lifespan. Marek's disease virus (MDV) is a highly contagious herpesvirus which causes a neoplastic condition in chicken populations. Several vertebrate-infecting herpesviruses have been shown to exist in an integrated state during latent periods of infection. However the status of MDV during latency has been a topic of debate.

Results: Here we employed high-resolution multi-color fluorescence in situ hybridization (FISH) to show integration of MDV at the telomeres of chicken chromosomes. Cytogenomic mapping of the chromosomal integrations allowed us to examine the clonal relationships among lymphomas within individuals, whereas analysis of tumors from multiple individuals indicated the potential for chromosomal preferences.

Conclusions: Our data highlight that substantive genome-level interactions between the virus and host exist, and merit consideration for their potential impact and role in key aspects of herpesvirus pathobiology including infection, latency, cellular transformation, latency-breaks and viral evolution.

Background

Herpesviruses are a large group of DNA viruses that cause disease in many organisms [1,2]. The ability of herpesviruses to occupy diverse biological niches, the wide range of cell types targeted for primary infection and latency, along with acute and chronic infection symptoms, contribute to the significance and relevance of herpesvirus research. In particular, human herpesviruses are a major public health concern due to the prevalence of such viruses within and among populations worldwide and the associated pathologies. Two of the eight disease-causing human herpesviruses, Epstein-Barr virus (EBV) and Kaposi's sarcoma virus (KSHV), are associated with neoplastic transformation. A closely related avian herpesvirus, Marek's disease virus (MDV; family *Herpesviridae*, genus *Mardivirus*, species *Gallid herpesvirus 2*) also exhibits oncogenic properties inducing lymphoid tumors in chickens.

MDV rapidly infects B cells and subsequently transforms T cells during pathogenesis of the highly contagious neoplastic disease of chickens known as Marek's disease (MD). Since discovery [3], MD has been extensively studied and utilized as a valuable biomedical model for human herpesvirus infection and viral induced malignancies [4,5]. MDV is of great concern in agriculture, causing large economic losses as well as impacting animal well-being. Despite current control through the use of vaccines, viral evolution leads to increasing virulence and vaccine breaks on a cyclical basis [6,7].

A fascinating biological property that MDV shares with all herpesviruses is the ability to persist indefinitely in the host and evade immune response by establishing a latent infection. The molecular mechanisms controlling lytic replication and latent persistence of herpesviruses have been the subject of intense study in recent years, aided largely by advances made in the field of viral genetics. Although new information has emerged from these studies regarding mechanisms of the lytic cycle, factors controlling latency and tumorigenesis

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remain poorly understood. MDV presents a unique model system in which to explore the mechanisms of herpesvirus latency and virus-induced tumor formation.

Another intriguing aspect of herpesvirus infection is that both episomal and linear-integrated forms of the virus are present in the host [8]. One consideration is that latency, and potential reactivation of the virus, could be achieved in part through genomic interactions between the host and virus, evasion through integration and recurring infection through emergence/release. Evidence for herpesvirus integration into human chromosomes was reported for Human Herpesvirus 6 (HHV-6) and EBV [9-14]. Since Delecluse and Hammerschmidt (1993) first reported on the potential for MDV to exist in an integrated state, it has been a subject of debate as to whether the virus integrates stably (and where) into the chicken genome or is merely associated and how such associations might be germane to pathogenesis and oncogenesis. Since that early report, complete genome sequences have been reported for several MDV strains and BAC vectors containing the full-length MDV genome have been cloned [15-20]. The sequencing provided data supportive of the hypothesis that MDV interacts with the chicken genome, as it was discovered that the circular viral genome encodes several orthologous host genes including the telomerase RNA (vTR) gene as well as telomeric repeats (TTAGGG) [17,18,21].

To resolve whether and where MDV integrates in the host chicken genome we employed high-resolution multi-color fluorescence in situ hybridization (FISH) using the tools developed from both the MDV and chicken sequencing efforts. We investigated the details of integration on both intra- and inter-chromosomal levels by mapping the location of MDV integrations. In depth analysis of MDV integration sites allowed us to investigate several specific objectives, including the clonal composition of MDV tumors and integration site preference of the virus. We show here that MDV specifically integrates into the telomeric regions of chicken chromosomes in MDV-induced clonal lymphomas.

Methods

Bird and tumor samples

Two common infection strategies were used to produce tumors. In the first case, progeny from a cross between two genetically susceptible inbred lines (USDA-ADOL 15I₅ X 7₁) [22] were challenged with 1,000 plaque forming units of MDV (GA strain). The resulting 17 tumors from nine birds were processed for chromosome harvest by standard procedures. In the second strategy, peripheral blood lymphocytes harvested from MDV-infected birds were used to infect (rather than purified virus). Nine tumors from four birds were processed for chromosome harvest. Animal experimental procedures, care

and management were approved by the USDA-ARS Avian Disease and Oncology Laboratory Animal Care and Usage Committee.

Fluorescence in situ hybridization (FISH)

The FISH procedures were standard [23,24] except that a telomere oligonucleotide (10 μ M (5' TTAGGG 3')₇) was included in the hybridization to provide competitor DNA to ensure blockage of MDV telomeric repeats [21] from hybridizing to chicken telomeric repeats (see also control experiments described in the results). To allow for multiple hybridizations, slides were cleaned and re-hybridized [25].

Probes and labeling

Table 1 describes the BACs used to identify specific chicken chromosomes including GGA 6-15, 17-28, W and one linkage group (LG E22/E50C23) [24,26] (<http://bacpac.chori.org>; UCSC, <http://genome.ucsc.edu/>). For GGA 16, an rDNA probe covering the external transcribed sequence (ETS) of the 18S-5.8S-28 S repeat was utilized [27]. Probes for GGA 1-5 and the Z were not needed because these chromosomes are easily identified by their morphological features. The MDV BAC, Md11gDc1.2 was used to query MDV integration in the chicken genome by multi-color FISH (MDV of one color, chromosome probes of another, see below) [18]. A telomere-peptide nucleic acid (PNA) fluorescein probe (Applied Biosystems) was used to identify telomeric sequence repeats. Approximately 300 ng of ETS and 1 μ g of each BAC clone was labeled either with digoxigenin by the DIG Nick Translation Mix (Roche, Indianapolis, IN, USA), or Spectrum Red dUTP, or Spectrum Orange dUTP by the Nick Translation Kit (Abbott Molecular, Inc., Des Plaines, IL, USA), or Cy3 dUTP (Amersham Biosciences) by the Nick Translation System (Invitrogen). Probes labeled with digoxigenin were detected by rhodamine or fluorescein (FITC) anti-DIG antibody incubation (0.70 μ g/slide, Roche Applied Science, Indianapolis, IN, USA).

Chromosome analysis

Chromosome and cell images were analyzed using Applied Imaging CytoVision software (3.92 GENUS). The analysis of mega-telomeres signals was standardized using techniques described previously [24,28]. Forty to 50 cells were examined for each FISH experiment for a total of more than 4, 400 cells analyzed.

Results

Characterization of MDV Integration Sites

The status of MDV integrations was studied by profiling 26 tumors from 13 birds; integrations were observed in every lymphoma examined (Table 2). The overall mode of integration sites was 5 (Figure 1) and the overall

Table 1 Characteristics of probes used in chicken chromosome mapping of MDV integrations

Chromosome [¶]	Clone Identity	Insert Size (Kb)	Features (NCBI Accession)	Marker Position (Mb)*	References
6	CH261-169D14	226 [†]	SCD (AJ297918/X60465)	18.5	[50]
7	CH261-95H15	158 [†]	SP5 (NM_001044684)	19.7	
8	CH261-84K8	235 [†]	ZNF326 (NM_001006533)	15.6	
9	TAM31-29A21	-	TR (AY312571)	21.5	[37]
10	TAM33-42N22	179 [‡]	NEO1 (U07644)	1.30	
11	TAM32-22B17	181 [‡]	ADL210 (G01630)	12.8	
12	TAM32-43M12	118 [‡]	MCW198 (G31980)	12.7	
13	TAM32-5E13	173 [‡]	MCW322 (G32078)	16.3	
14	TAM32-2I7	225 [‡]	ADL200 (G01621)	39.8	
15	TAM32-87J1	-	MCW226 (G31998)	2.4	
16	ETS rDNA	3	NOR	q arm	[26,51]
17	TAM31-22I10	-	GSN (AF042795)	9.1	
18	TAM32-14L7	78 [‡]	SFRS2 (X62446)	4.2	
19	TAM32-21A11	164 [‡]	ACACA (X77240)	8.5	
20	TAM32-13H1	-	BMP7 (AF223970)	11.6	
21	TAM32-12A6	-	TUS0022 (AI979776)	2.5	
22	TAM32-13C2	202 [‡]	CRMP62 (U17277)	0.5	
23	TAM32-27O5	-	ADL299 (G01708)	17.8	
24	TAM32-54P15	219 [‡]	APOA1 (M96012)	5.2	
25	TAM32-49A10	149 [‡]	LEI0345 (AJ240692)	1.7	
26	TAM32-40J16	152 [‡]	MCW209 (G31986)	2.1	
27	TAM32-76B6	169 [‡]	MCW233 (G31915)	1.9	
28	TAM32-4G3	167 [‡]	ADL299 (G01751)	4.3	
W	TAM32-55E18	-	CW01 (D85614)	q arm	[24,52]
LG E50C23	TAM32-2A12	-	LEI0336 (AJ240683)	N/A	
	MDV: Md11gDc1.2	178 [§]	N/A	N/A	[18]
	MDV: P89	43	N/A	N/A	[29]
	MDV: SN16	42	N/A	N/A	[29]

CH = Children's Hospital Oakland Research Institute, CH261 = EcoRI BAC library.

TAM = Texas A&M University, TAM31 = BamHI, TAM32 = EcoRI, TAM33 = HindIII BAC libraries [26,31].

ETS = external transcribed spacer of the 18S-5.8S-28 S rRNA gene repeat (rDNA).

BAC locations and features were obtained from US Poultry Genome Project 'Database of BACs Assigned to Chicken Genes and Markers' (<http://poultry.mph.msu.edu/resources/resources.htm>, May 2006 version) and/or UCSC Genome Browser (<http://genome.ucsc.edu>, Chicken May 2006 assembly).

Features = Genes/Markers and Genbank accession numbers (in parentheses).

[¶] Probes for GGA 1-5 and the Z were not needed because these chromosomes are easily identified by their morphological features

*Position refers to the start position of the BAC or gene/marker on the chromosome in the May 2006 chicken assembly (UCSC Genome Browser). In two cases the chromosomal coordinates are unknown, but the chromosome arm location is indicated.

[†]Obtained from the UCSC Genome Browser.

[‡]Insert size was estimated using the UCSC Genome Browser and Chicken FPC <http://www.bioinformatics.nl/gbrowse/cgi-bin/gbrowse/ChickFPC> as described in O'Hare and Delany 2009.

[§]Based on the Md5 strain (GenBank accession no. AF243438) [17].

"-" = unknown

N/A = not applicable

mean for number of viral integration sites was 4.17 (range 0.2 to 8.0). Several control experiments were conducted to aid in MDV signal interpretation including hybridization of the MDV BAC probe to samples from non-infected birds (adults and embryos) and hybridization of two MDV cosmid clones to tumor samples. In the first case, no virus signal was detected in non-infected individuals (Additional Figure 1: Hybridization of the MDV BAC probe to uninfected embryonic chicken cells). In the second case, overlapping MDV

cosmid clones, P89 and SN16, used because they lacked telomeric repeats [21,29], were hybridized against a previously profiled tumor sample (Additional Figure 2: FISH comparison between two probes: the MDV-BAC and MDV cosmid clones). The mode of MDV integrations was found to be the same as when hybridized with the full-genome MDV BAC. Thus, these controls demonstrated that the MDV BAC probe was specific for hybridization to only the MDV DNA versus any elements of the chicken genome.

Table 2 A cytogenomic profile of lymphomas from MDV-infected birds: virus integrations, ploidy, and telomeres.

Bird #	Sample Information		MDV Integration Sites*			Ploidy Levels (metaphase cells) [†]		Mega-Telomere Signals [‡]	
	Sex	Tumor # (Tissue Source)	Mode	Mean	SD	% Tetraploid	% Aneuploid	Mode	W
1	M	1-T1 (heart)	5	4.4	0.9	2	0	7	-
		1-T2 (left gonad)	5	5.4	1.3	0	0	11	-
		1-T3 (lung)	5	4.9	0.7	0	0	9	-
2	M	2-T1 (left gonad)	4	3.8	0.7	4	0	7	-
		2-T2 (right gonad)	4	4.1	1.1	4.5	0	7	-
3	M	3-T1 (lung)	4	4.7	1.6	4	0	6	-
		3-T2 (bursa)	5	4.3	1.3	16	0	7	-
		3-T3 (gonad)	6	5.9	0.8	5	0	6	-
4	F	4-T1 (spleen 1)	3	3.2	1.2	0	0	9	Yes
		4-T2 (spleen 2)	2	1.9	1.2	0	0	7	Yes
		4-T3 (heart)	1	1.4	1.3	0	0	7	Yes
5	M	5-T1 (left gonad)	4	3.8	0.8	0	0	8	-
		5-T2 (right gonad)	4	3.6	0.8	6	0	8	-
6	M	6-T1 (heart)	2	1.6	1.2	0	0	7	-
7	F	7-T1 (spleen)	0	0.2	0.6	0	0	8	Yes
8	F	8-T1 (gonad)	3	2.4	1.4	0	0	10	Yes
9	F	9-T1 (liver)	1	0.5	0.5	0	0	8	Yes
10 [§]	F	10-T1 (gonad)	5	5.0	1.0	25	12.5	9	Yes
11 [§]	M	11-T1 (kidney)	3	3.5	2.2	21	23	5	-
		11-T2 (liver)	6	4.5	2.2	17.5	15	7	-
		11-T3 (thymus)	6	5.4	1.3	20	2.5	8	-
12 [§]	F	12-T1 (gonad)	5	5.6	2.0	27	7	6	Yes
		12-T2 (kidney)	7	5.5	2.2	0	0	6	- [¶]
13 [§]	F	13-T1 (spleen)	9	7.2	3.0	2.5	0	7	Yes
		13-T2 (liver)	9	8.0	1.8	0	10	7	Yes
		13-T3 (kidney)	9	7.7	2.1	10	7	7	Yes

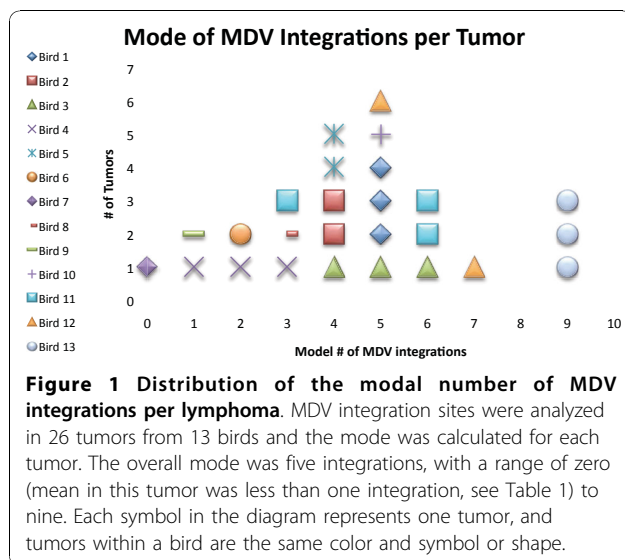
*Mode (most frequent value), mean, and standard deviation (SD) of diploid metaphase cells.

[†]Tetraploid cells were identified as those containing four copies of all macrochromosomes and sex chromosomes, and an estimated number of microchromosomes present in twice the normal amount. Aneuploid cells were identified as those containing an abnormal copy number (less than or greater than two) of one or more macrochromosomes.

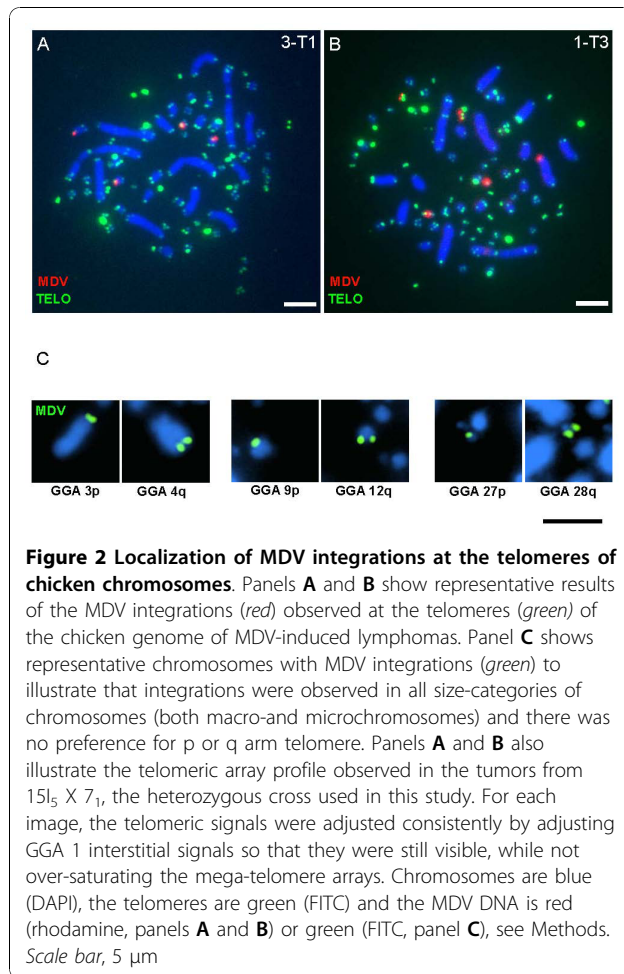
[‡]Mega-telomere signals were counted in tumor metaphase cells using the telomere-PNA probe, and the mode was calculated for each tumor.

[§]Birds from infection strategy 2 (see Methods section for details).

[¶]Tumor 12-T2 lacked a W chromosome (as verified by the W BAC probe, see Table 1).



In all tumors examined, the MDV integrations were localized at the telomeres (Figure 2) and interstitial chromosome integration was not observed on the macrochromosomes or the microchromosomes, to the extent allowed by cytogenetic resolution. When sister chromatids were visibly separated [22], two MDV signals were observed, one on each sister chromatid. Only one integration site per chromosome was observed, i.e., MDV was only found at either the p or the q arm of the involved chromosome, but not at both arms. Chromosomal p versus q arm telomere integration was easily determined in the macrochromosomes (GGA 1-5) based on cytogenetic observations (e.g., GGA 3p and 4q, Figure 2C). The p versus q arm telomere integrations into intermediate (GGA 6-10) and microchromosomes (GGA 11-28) were established using cytogenetic observations and the sequenced genome assembly location for the



chromosome-specific BAC probes used in the mapping experiments. For example, see GGA 9p, 12q, 27p and 28q in Figure 2C. For a few of the smallest microchromosomes, the p versus q arm position of MDV integration could not be established because BAC assembly information and sequence coordinates were not available.

Identification of Involved Chromosomes by FISH Mapping

Chromosome-specific BACs were utilized (Table 1) to identify the specific chicken chromosomes harboring MDV integrations. Our approach was to hybridize more than one chromosomal BAC (~ three at a time) to one of the tumor samples for a bird (e.g., 13-T1, Figure 3A, B, and 3C) and any dual-signal positive chromosomes were recorded (e.g., GGA 9, Figure 3A). The entire suite of chromosomal BACs was tested on the same tumor until all of the MDV integrations had been identified or until all available BACs were tested. Next, the positive chromosomal BACs were hybridized to the other tumor samples from that bird (e.g., 13-T2 and 13-T3, Figure 3D,E,a). If the tumors showed different integration

profiles then each tumor was tested separately with all available BACs. Integrations in fifteen tumors from six birds were mapped using this iterative approach and the results are found in Table 3. Some viral integration positions remain “unknown” due to the lack of sequence assembly information and BAC probes for microchromosomes (GGA 29-38).

In terms of the uniformity of integration positions among tumors from a single individual, four different categories of results were observed. First, three individuals (Birds 2, 5, and 13) each had multiple tumors that showed complete uniformity among tumors. Figure 3 is a composite of FISH images from three tumors from one individual (Bird 13) each showing nine identical integration sites and provides a representative of the iterative process of mapping the involved chromosomes. The second category of results observed was heterogeneity within a tumor, e.g., the Bird 1 profile. Two of three tumors from Bird 1 (1-T2 and 1-T3) included sub-populations of cells with integration sites differing from the main population of cells (which were shared). The sub-populations of each tumor shared the same profile with eight total integration sites, different from the main profile, which had five integration sites (Table 3). The third results category was observed in Bird 3, having in-common sites in each of its three tumors (3-T1, 3-T2, and 3-T3) but also unique integration sites in each tumor. Tumors 3-T1 and 3-T2 had four in-common locations at GGA 10, 11, 12, and 18 and tumor 3-T2 contained an additional integration site at GGA 13. Tumor 3-T3 had two in-common sites with both 3-T1 and 3-T2 (GGA 12 and 18); however, it also contained four unique sites. And finally, the fourth results category was found in Bird 12, which was the only individual with two tumors (12-T1 and 12-T2) that had no in-common chromosomal integration sites. Tumor 12-T1 displayed a uniform integration profile, which differed from the main profile of 12-T2. In addition, 12-T2 had a subpopulation of cells which had two integration sites which could not be mapped (lack of probes and assembly information) (Table 3).

Karyotype Abnormalities and Mega-telomere Loci

Several karyotype anomalies were found in the tumors, a phenomenon well established in human cancers. Variations included ploidy changes, commonly tetraploid and aneuploid (Figure 4). The tetraploid cells typically contained double the number of MDV integrations of the diploid cells suggesting the ploidy change occurred after integration.

We investigated the telomere profile of tumor cells to consider whether there was an observable preference for integration into the extremely large (aka mega-) telomeres of the chicken genome, see Figure 2A,B and

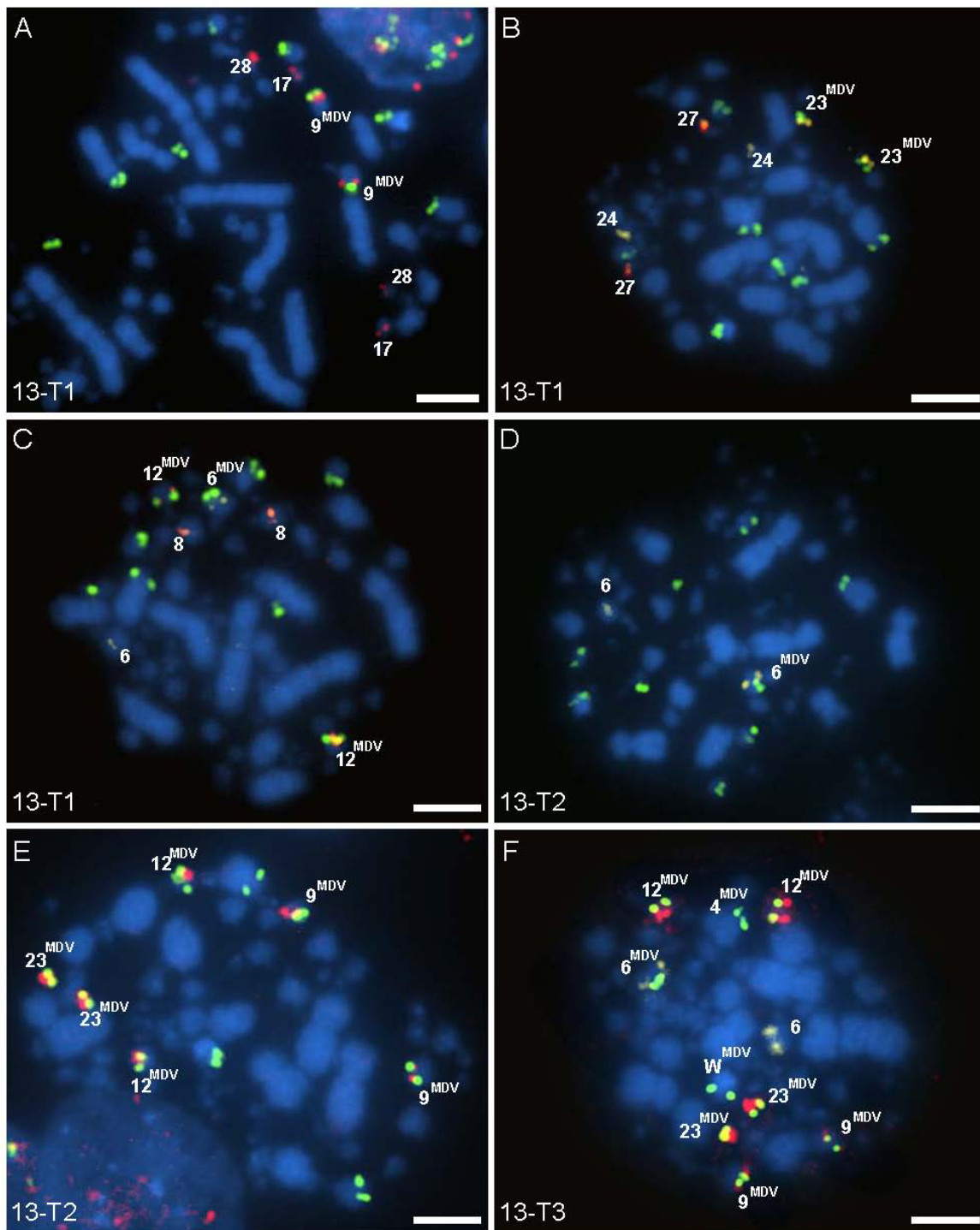


Figure 3 Mapping of MDV integrations in chicken chromosomes. An iterative process of mapping by multicolor FISH was used to identify the involved chromosomes. Images shown are from Bird 13, which had nine integration sites (4p, 6p, 9q, 9q, 12, 12, 23, 23, and Wp) for all three tumors. Cells shown in **A**, **B**, and **C** are from tumor 13-T1, which was used to begin mapping. Once the positive sites were identified for 13-T1, they were subsequently tested on 13-T2 (**D** and **E**) and 13-T3 (**F**). Multiple chromosomal BAC probes were tested for possible integration in panels **A-C**. The MDV positive chromosomes are indicated by superscript. Chromosomal BAC probes are labeled either red (Spectrum Red) or yellow (Spectrum Orange) and in all panels MDV DNA is labeled green (FITC), see Methods. Scale bar, 5 μm

Table 3 Chromosome locations of MDV integrations in lymphomas: Telomeres of autosomes and sex chromosomes.

Tumor Identity*	% of cells	Chromosomes with Integrations	# of Unmapped Integration Sites [†]
1-T1	100	3p, 23, 24p	2
1-T2	68	3p, 23, 24p	2
"	32	1q, 4q, 8p, 9p, 9p, 20, 28q, Z	0
1-T3	87	3p, 23, 24p	2
"	13	1q, 4q, 8p, 9p, 9p, 20, 28q, Z	0
2-T1	100	15q, 17q, 20q, 20q	0
2-T2	100	15q, 17q, 20q, 20q	0
3-T1	100	10, 11, 12q, 18	0
3-T2	100	10, 11, 12q, 13, 18	0
3-T3	100	4q, 6p, 12q, 18, 19p	1
5-T1	100	7q, 12q, 20q, 27p	0
5-T2	100	7q, 12q, 20q, 27p	0
12-T1	100	12, 26	3
12-T2	74	3p, 6p, 24, 25	3
"	26	none	2
13-T1	100	4p, 6p, 9q, 9q, 12, 12, 23, 23, Wp	0
13-T2	100	4p, 6p, 9q, 9q, 12, 12, 23, 23, Wp	0
13-T3	100	4p, 6p, 9q, 9q, 12, 12, 23, 23, Wp	0

*See Table 2 for additional tumor sample information (Bird #, sex, and tissue source).

[†]Unidentified chromosomes with MDV integration are microchromosomes (GGA 29-38) that lacked assembled sequence information or BAC probes at the time of this study.

Table 2. The number of mega-telomeres ranged from 5 to 11 among individuals and intra-individual diversity was observed. For example, Bird 1 indicated different mega-telomere profiles in each of its tumors with modes of 7 (1-T1), 9 (1-T3) and 11 (1-T2). As reported in prior work (36), the females exhibited an extremely large (mega-) telomere on the W sex chromosome. However, there was no observable preference for MDV integration into the mega-telomere loci.

Discussion

Delecluse and colleagues provided initial evidence for viral integration of MDV into chicken chromosomes [30,31]. At the time, this work was necessarily limited in scope and resolution by the lack of sequence information and probes for both the MDV and the chicken chromosomes. Having the advantage of genomics tools and high resolution cytogenetics, we are able to now show that the MDV genome is present in an integrated state at the telomeres of specific chicken chromosomes of tumors resulting from MDV-infection. MDV was observed on both sister chromatids indicating the viral DNA replicates in a semi-conservative fashion along with the host genomic DNA. We also noted that signal intensity varied among integration locations, suggestive of different numbers of integrated copies. The

sequencing of the chicken genome coupled with the availability of chromosome specific BACs to identify chicken microchromosomes [24,25,32] created the opportunity for mapping of integrations and allowed us to address several crucial questions related to integration site preference and lineages of transformation events.

Preferences of Viral Integration: Subchromosomal and Chromosomal

MDV integrates at the chicken telomeres

Our initial working hypothesis was that MDV preferentially integrates into the chicken mega-telomeres, which are ultra-long telomeric arrays ranging in size from 100 s of Kb to several Mb [24,28,33-35]. Although our data did not support this hypothesis, the MDV genome was found exclusively at the telomeric regions of the host chromosomes. Viral integration into the interstitial telomeric sequences was not observed. We hypothesize that the terminal ends of chromosomes may provide an environment more conducive to integration, perhaps through a specialized chromatin confirmation (e.g., the t-loop of the telomere as it is replicated, and/or the actual telomerase extension apparatus). Interestingly, HHV-6 was found integrated into the subtelomeric region of human chromosomes and, similar to MDV,

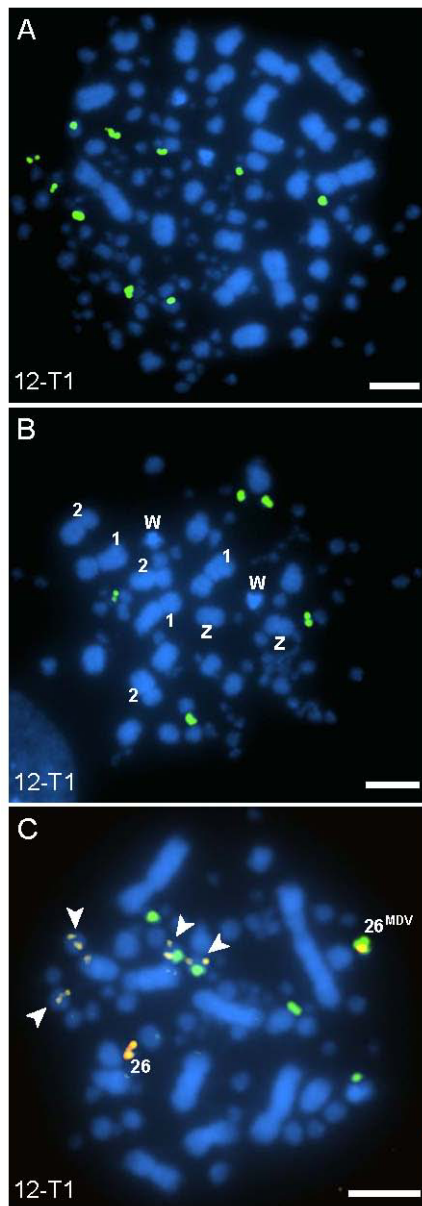


Figure 4 MDV-induced lymphomas exhibit a variety of ploidy-level abnormalities. Karyotype abnormalities were common in the tumors and the tetraploid cells typically had double the number of MDV integration sites (green) as compared to diploid cells. As shown here, a tetraploid cell (A) from tumor 12-T1 of Bird 12 had ten MDV integration sites (the majority of diploid cells for 12-T1 had five integration sites). Aneuploid cells (B and C) were also observed in the tumors, in the cell shown in panel B there are three copies of GGA 1 and 2, and two copies of both GGA W and Z. The cell shown in panel C from 12-T1 has several karyotype abnormalities, including four copies of GGA 12 (arrowheads, yellow signals), two of which have MDV on them. The mode of integrations for diploid cells in 12-T1 was five, where as the cell in panel C displays six MDV integration sites with the addition of the tetrasomic GGA 12, indicating non-disjunction occurred after viral integration. GGA 26 (red) is also shown here as having one homolog positive for MDV integration (indicated by superscript text). Scale bar, 5 μm

possesses telomere repeat arrays within its genome [9,36]. The telomeric array repeats were hypothesized to assist the HHV-6 integration mechanism through homologous recombination [9].

In exploring our initial hypothesis regarding MDV preference for integration into mega-telomeres we found additional evidence for extensive genotypic variation in telomere array sizes within and among chicken genetic lines and cell lines [28,35]. Although variation for the number of mega-telomere loci observed among the birds in this study was to be expected from an F1 generation (see methods), the variation among tumors within birds was not expected. Inter-tumor variation (within an individual) could be a manifestation of gain or loss of telomeric repeat sequence due to genomic instability. Another consideration is the unknown effect that integration of MDV DNA may have upon the telomere sequence itself.

MDV integration: random versus targeted?

Unlike HHV-6, which typically integrates into one or two chromosomes [10,14,37] MDV integrates into more chromosomes, typically five (and as many as nine). We found integrations across all size categories of chromosomes, i.e., both macro- and microchromosomes (Figure 2C). A few chromosomes stand out as interesting candidates for future studies, as highlighted by Figure 5. In the two instances that MDV was found integrated on GGA 9, both homologs were involved. GGA 9 encodes the single copy chicken telomerase RNA (cTR) component of the telomerase enzyme [36]. Sequencing of the MDV genome showed that it also encodes two copies of TR (viral (v)TR) with high sequence identity (85-88%) to cTR [16,17,38,39]. In comparison to cTR, vTR was more efficient at restoring telomerase activity in telomerase negative cell systems [39]. Furthermore, when vTR was deleted from the MDV genome, the vTR-deleted virus showed reduced capabilities in causing lymphomas, suggesting that vTR is a key element in promoting T-cell lymphomagenesis [40]. Whether MDV is specifically utilizing vTR to aid in integration by taking advantage of telomerase activity at the telomeres requires further study, and would be a novel role for TR (chicken or viral).

Several other chromosomes frequently exhibited MDV integrations, for example GGA 4, 6, 12, and 20, were found in common among tumors between three or more birds (Figure 5). Interestingly, GGA 4 encodes the chemokine interleukin-8 (IL-8) gene sequence [41] and MDV sequencing showed the virus also encodes an IL-8 gene [16,42]. The presence of host sequences in the virus genome is noteworthy because a plausible mechanism for MDV to obtain such chicken-derived genes (vTR and vIL-8) and sequences (e.g., TTAGGG repeats) is by integration into the chicken chromosomes, and then subsequent emergence with exogenous host

sequence. Since telomeric integration appears to be the preferred location, we propose that viral integration near host genes is a rare event. However, if a “captured” host gene sequence provides a selective advantage to the virus, e.g., vTR’s ability to promote oncogenesis or perhaps facilitate integration, it would persist in future generations of virus.

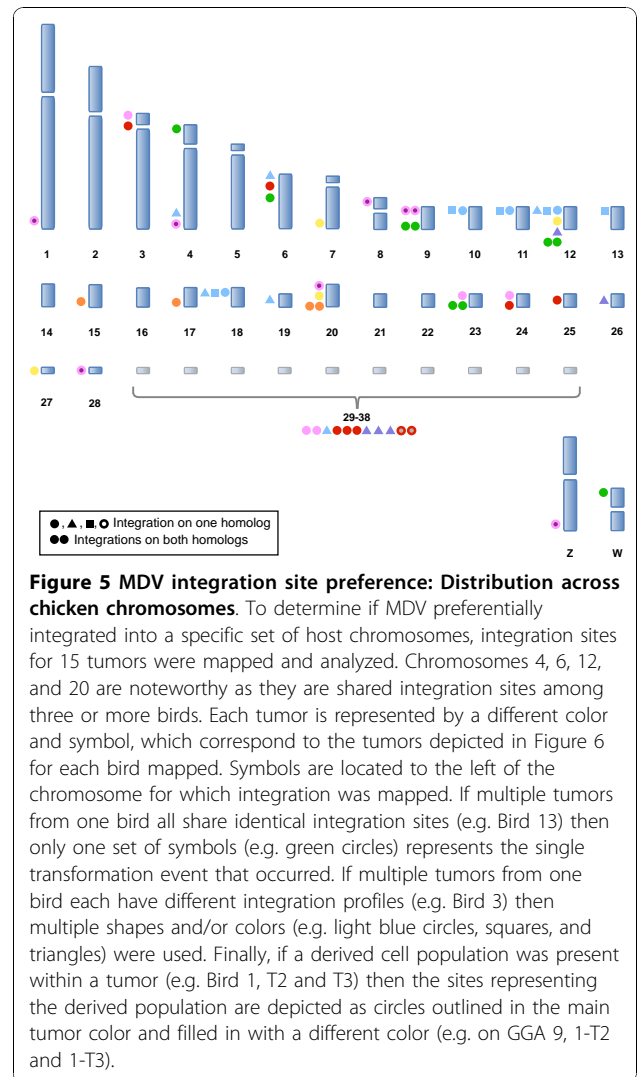
Clonality of Late-Stage Lymphomas: Lineage relationships among tumors within birds

Another objective of this study was to improve our understanding regarding clonality of MDV-induced lymphomas. That is, are the multiple tumors of a bird derived from a single transformation event (monoclonal origin) or do the tumors arise from independent events (polyclonal origin)? Clonality of tumors has been addressed by immunoglobulin analysis, T-cell receptor gene rearrangements, X-chromosome methylation patterns, various molecular markers for somatic mutations, and analysis of viral integration [reviewed in [43-48]]. Viral integration sites and viral sequencing have proved to be useful markers when assessing clonality of tumors [49]. Here we utilized the mapped MDV integrations to gain insight into tumor lineage(s).

Mapping lineage analysis indicates that four (of six) birds had multiple tumors of monoclonal origin, inclusive of one individual (Bird 1) showing a derived cell population (Figure 6), which is not unexpected given lack of genome stability in cancers and/or the possibility that the virus could be mobile. The remaining two birds (Bird 3 and 12) displayed tumors consist with a mixed or polyclonal origin (Figure 6). The most conservative analysis of the three tumors from Bird 3 places them in a polyclonal category, that is, three separate transformation events occurred to form each tumor. Alternatively, the tumors from Bird 3 could be related, which raises the possibility that the virus is capable of reemerging from its integrated state and moving to new integrated locations. Alternatively, tumors 3-T1 and 3T-2 could be of monoclonal origin given the loss of one of the integration sites (GGA 13). However, the third tumor (3-T3) showed only two MDV integration sites (GGA 12 and 18) in common with the other two tumors, suggestive of a separate transformation event. The tumors from Bird 12 may also represent a polyclonal set of tumors, indicating tumors 12-T1 and 12-T2 have different founder cells altogether (Figure 6). However, it must be noted that 12-T1 and 12-T2 both have three unidentified MDV integration sites and the possibility remains that those three sites could be in common.

Conclusions

The indications for predominately, but not exclusively, monoclonal tumor origins, as well as derived cell



populations within the MDV lymphomas, illustrates the complexity of tumor biology and the potentially diverse strategies employed by the herpesvirus. Understanding the timeframe during pathogenesis when MDV integrates into the chicken genome will be essential to assessing both the utility of integration profiles as a marker for clonality and the significance of integration on the latency cycle. In addition, further investigation of the effect of viral integration on cellular transformation is necessary. Integration may have a profound effect on transformation, or it may simply serve as a mechanism of maintaining the latent genome. We are currently investigating the stages at which MDV integrates, from early infection to late oncogenesis, and have preliminary indications that MDV integrates prior to transformation and tumorigenesis. Further research on the molecular mechanism for telomere-targeting of MDV integration should provide new insight regarding the role of

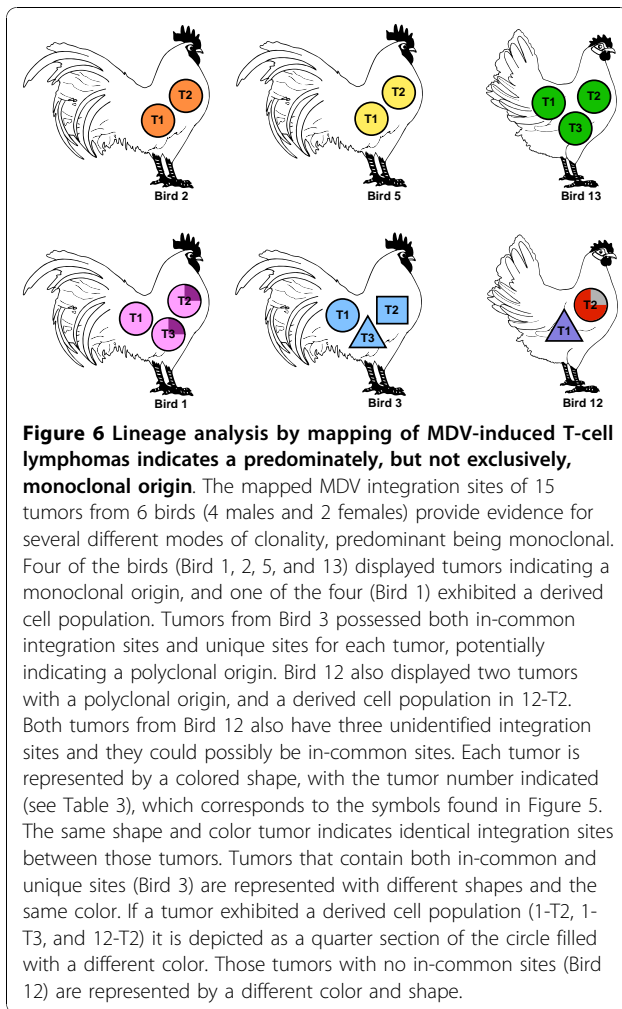


Figure 6 Lineage analysis by mapping of MDV-induced T-cell lymphomas indicates a predominately, but not exclusively, monoclonal origin. The mapped MDV integration sites of 15 tumors from 6 birds (4 males and 2 females) provide evidence for several different modes of clonality, predominant being monoclonal. Four of the birds (Bird 1, 2, 5, and 13) displayed tumors indicating a monoclonal origin, and one of the four (Bird 1) exhibited a derived cell population. Tumors from Bird 3 possessed both in-common integration sites and unique sites for each tumor, potentially indicating a polyclonal origin. Bird 12 also displayed two tumors with a polyclonal origin, and a derived cell population in 12-T2. Both tumors from Bird 12 also have three unidentified integration sites and they could possibly be in-common sites. Each tumor is represented by a colored shape, with the tumor number indicated (see Table 3), which corresponds to the symbols found in Figure 5. The same shape and color tumor indicates identical integration sites between those tumors. Tumors that contain both in-common and unique sites (Bird 3) are represented with different shapes and the same color. If a tumor exhibited a derived cell population (1-T2, 1-T3, and 12-T2) it is depicted as a quarter section of the circle filled with a different color. Those tumors with no in-common sites (Bird 12) are represented by a different color and shape.

telomeres in aspects of host genome stability and also viral evolution.

Additional material

Additional file 1: Hybridization of the MDV BAC probe to uninfected embryonic chicken cells. The MDV BAC was hybridized to chromosomes from uninfected chicken embryo fibroblasts. As shown, the MDV BAC probe did not hybridize to uninfected chicken cells, i.e., there are no FISH signals observed. Scale bar, 5 μ m

Additional file 2: FISH comparison between two probes: the MDV-BAC and MDV cosmid clones. Panel A displays tumor 10-T1 labeled with the MDV cosmid clones (green) with a total of five integration sites indicated by arrowheads. Panel B displays tumor 10-T1 labeled with the full-genome MDV BAC probe (red) also with a total of five integration sites. Scale bar, 5 μ m

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Authors' contributions

CMR, HDH, HHC, and MED conceived of and designed the research project. CMR conducted the experiments. CMR and MED wrote the manuscript. All authors have read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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