Vaccination and Host Marek’s Disease-Resistance Genotype Significantly Reduce Oncogenic Gallid alphaherpesvirus 2 Telomere Integration in Host Birds

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Abstract
Marek’s disease (MD) is an infectious disease characterized by lymphomas and high mortality in susceptible chickens. The causative and ubiquitous alpha-herpesvirus known as MD virus (MDV) integrates into host telomeres during early infection through latency, known to be an important phase for oncogenic transformation. Herein, we sought to determine the influence of vaccination and host genetics on the temporal dynamics of MDV-host genome interactions. We studied integration profiles using 2 MD vaccines that vary in protective efficacy in 2 genetic lines that differ in MD resistance/susceptibility. Virus integration of both oncogenic MDV and vaccine strains was observed in both MD susceptible and resistant birds, however, the lines differed in their dynamic telomere-integration profiles. Notably, the resistant host genotype exhibited a smaller percentage of replicating cells with the virus telomere-integrated only phenotype as compared to the susceptible genotype. Vaccination with Rispens, the most protective MD vaccine, also reduced the establishment of the virus telomere-integrated only phenotype, suggesting a significant role of the phenotype in MD lymphoma development. The effect of Rispens vaccination was most dramatic in the susceptible genotype. These results suggest important connections between vaccinal immunity, MDV telomere integration, virus-induced oncogenesis, and virus-host genome interactions in the context of host genetics and disease susceptibility.

Keywords
Chicken genome · FISH · Genetic resistance · Herpesvirus · Marek’s disease · Rispens vaccine · Telomere integration

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ic repeats [Kaufer et al., 2011; Greco et al., 2014]. In the MDV infection cycle, oncogenic strains transition from a cytopathic replication stage, during which the viral genome takes an episomal and extrachromosomal form, to a latency stage, characterized by evasion of host immune responses [Biggs, 1968; Addlunger and Calnek, 1973; Baigent and Davison, 2004; Osterrieder et al., 2006; Arumugaswami et al., 2009; Gimeno et al., 2011] with the latter having a temporal overlap with MDV-host telomere integration events [Robinson et al., 2010, 2014] and decline in the quantity of extrachromosomal MDV genomes [Kaschka-Dierich et al., 1979; Delecluse and Hammer-schmidt, 1993]. Latently infected host CD4 T lymphocytes are the target cell population for transformation events and subsequent lymphoma development, around 2–3 weeks after infection in susceptible birds [Witter, 1984; Calnek, 2001; Burgess and Davison, 2002; Nair, 2005; Trapp et al., 2006].

Host bird genetics contributes to MD resistance/susceptibility as well as the effectiveness of the MD vaccines [Bacon and Witter, 1994; Sarson et al., 2008; Chang et al., 2010, 2012]. The chicken major histocompatibility complex (MHC) has long been known to have a major influence on MD disease incidence [Briles et al., 1983; Bacon, 1987; Schat et al., 1994; Bacon et al., 2000; Sarson et al., 2008; Miller and Taylor, 2016]. Conveniently, blood group and DNA markers have enabled poultry breeding companies to select birds with superior MD resistance. In addition to the MHC, other genetic factors exert a major influence on MD resistance. For example, Avian Disease and Oncology Laboratory (ADOL) experimental inbred lines 6 and 72 chickens share the same MHC haplotype, yet differ greatly in resistance to MD. As these non-MHC loci are smaller in genetic effect, identifying them has been more challenging though recent efforts based on allele-specific expression in response to MDV challenge have been able to identify a large collection that accounts for over 80% of the genetic variance [Cheng et al., 2015].

Since the early 1970s, MD control has relied heavily on mass vaccination of flocks [Calnek, 2001]. MD vaccination strategies commonly consist of monovalent injection of herpesvirus of turkey (HVT, apathogenic MDV serotype 1) [Rispens et al., 1972; Calnek et al., 1983; Petherbridge et al., 2003; Spatz et al., 2007] or Rispens/CV1988 (attenuated MDV serotype 1) [Rispens et al., 1972; Calnek et al., 1983; Petherbridge et al., 2003; Spatz et al., 2007]. The MD vaccines protect birds against disease progression and lymphoma development induced with oncogenic strains [Chang et al., 2010; Haq et al., 2010] through non-sterilizing immunity [Nair, 2005] in both susceptible and resistant host birds. The genomes of these strains are highly similar to the oncogenic strains, but crucial differences exist [McPherson and Delany, 2016]. The MD vaccine viruses replicate in host cells and transmit to other birds [Baigent et al., 2005; Islam et al., 2007; Tan et al., 2007; Gimeno et al., 2011; Islam et al., 2013; McPherson and Delany, 2016; McPherson et al., 2016], but do not induce disease or lymphomas [Calnek and Witter, 1997]. Despite improved understanding of vaccine-related immunity [Witter, 1984; Osterrieder et al., 2006], an explanation for the anti-tumor effect of vaccination has not yet been established. Vaccine protective efficacy can vary with the vaccine serotype, host MHC haplotype, and other aspects of the host genetics [Bacon, 1987; Bacon and Witter, 1992, 1993, 1994; Chang et al., 2010, 2012, 2014]. Rispens provides the most effective protection against the most virulent MDV strains in susceptible birds [Haq et al., 2012; Chang et al., 2014].

The MD vaccine strains have been detected in a latent form in vivo [Calnek and Witter, 1997], although infrequently and highly delayed as compared to oncogenic MDV, and are capable of host telomere integration [McPherson et al., 2016], partially resembling oncogenic MDV. Nevertheless, the vaccines do not result in a transformation-related phenotype, known as MDV telomere-integrated only, in dividing host cells [Robinson et al., 2010, 2014; McPherson et al., 2016]. Noteworthy, a similar pattern of host chromosome association and integration [Robinson et al., 2014] is observed with the Meq-deleted (ΔMeq) MDV, a recombinant MDV that lacks the viral oncogene and is a candidate MD vaccine [Silva et al., 2010].

Host genetic background (MHC and non-MHC) contributes to the effectiveness and behavior of the MD vaccine strains, which has been investigated for lines 63 (resistant) and 72 (susceptible) [Bacon and Witter, 1994; Chang et al., 2010, 2012]. In general, the serotype 1 vaccines, particularly Rispens, are more effective compared to serotype 2 vaccines. The bivalent serotype 2 + 3 vaccine is also more protective than serotype 3 HVT alone [Bacon and Witter, 1994]. The impact of host genotype on MD vaccine strain replication and shedding, and other aspects of the viral cycle within host lymphoid cells are poorly understood.

Our central hypothesis for this research was that oncogenic MDV and MD vaccine strain interactions with the host genome are influenced by the host genotype and contribute to the outcome of MD pathogenesis. Furthermore, we hypothesize that vaccination with Rispens MD...
vaccine influences oncogenic MDV interactions with the host genome in challenged birds in either an MD-resistant or -susceptible genetic background. Here, we tested these hypotheses by investigating chromosomal association and integration profiles of oncogenic MDV and MD vaccine viruses in inbred chicken disease genotypes 6
(resistant) versus 7
(susceptible) via previously defined cytogenomic methods. Our results contribute to a better understanding of the impact of host genotype and vaccination on MDV interactions with the host genome in dividing cells during early infection. Our findings also enlighten how the MD vaccines may work to reduce the oncogenic effect of virulent MDV strains.

Materials and Methods

**Chickens, MDV and MD Vaccines, and Procedures**

Experimental White Leghorn birds were from highly inbred lines 6
(resistant to MD-lymphoma development, also known as "resistant") or 7
(susceptible to MD-lymphoma development and related mortality, also known as "susceptible") from the USDA ADOL [Sharma and Stone, 1972; Stone, 1975], cared for under approved animal care protocols. All institutional and national guidelines for the appropriate care and use of laboratory animals were followed. For the single-treatment group, i.e., viral challenge or vaccination only, day-of-hatch chicks from both lines were intra-abdominally injected with 2,000 pfu Rispens/CVI988 (hereafter referred to as "Rispens") only, 2,000 pfu HVT only, 2,000 pfu Md5 strain MDV only, or received a mock inoculation. For the second group, i.e., dual treatment of vaccination and viral challenge, day-of-hatch chicks from both genotypes were intra-abdominally vaccinated with 2,000 pfu of Rispens or mock vaccine (saline injection), then all birds were challenged with 2,000 pfu of Md5 virulent MDV at 4 days post hatch (dph). Chicks were hatched and maintained in separate Horsfall-Bauer isolation chambers at ADOL.

**Tissue Collection and Processing**

Mitotic chromosome preparations were harvested from 4 splenic-derived samples (1 sample per bird) collected at 1, 4, 7, 14, and 21 days after viral challenge or vaccination (dpc, dpv) for single-treatment birds and at 1, 4, 7, 14, and 21 dpc for dual-treatment birds; thus, birds in the latter groups were older at the time of tissue sampling (i.e., 5, 8, 11, 18, and 25 dph). At least 3 tissues were processed per treatment as outlined previously [McPherson et al., 2016].

**MDV-Specific FISH**

Slide hybridization was carried out as previously described [Delay et al., 2007], with adaptations for labeling MDV as detailed in McPherson et al. [2016]. Image capture occurred within 24 h of MDV FISH and DAPI counterstaining.

**Cytogenetic Analysis**

Metaphase and interphase cell images were collected using an Olympus Bx41 epifluorescence microscope equipped with an automatic filter wheel (Chroma Technology 82000, DAPI/FITC/TRITC filter set), X-Cite 120 Series metal-halide fiber optic lamp, and Applied Imaging software (CytoVision 7.4 GENUS, Leica Biosystems). DAPI (host chicken DNA) and FITC (MDV DNA) exposure times were consistent across slide cell image captures. From 20 to 80 metaphase (dividing) cell images (average of 50) were captured and analyzed per individual sample or bird in each FISH experiment. Negative control (no treatment) samples were incorporated in all MDV FISH experiments to ensure that the MDV genome-containing BAC probe was hybridizing specifically to the MDV genome, as indicated by the absence of FITC signals from all terminal and interstitial telomeres, as described [Robinson et al., 2010]. All captured mitotic metaphase cells were categorized as 1 of 4 cytogenomic host-virus phenotypes as previously described [Robinson et al., 2014; McPherson and Delany, 2016] and are depicted in Figure 1: null (no signals detected); chromosome-associated (dispersed signals over and around the chromosomes); chromosome-associated/telomere-integrated (associated signals along with distinct, punctate and bright signal(s) at the telomeres); or telomere-integrated only (only the distinct telomeric signals detected).

For the Rispens-vaccinated/Md5-challenged (dual treatment) birds in the second set of experiments, mitotic cells with the telomere-integrated only virus-host phenotype represent solely the oncogenic MDV (Md5) infection, due to previous results and data here within demonstrating that the MD vaccines alone do not, or very rarely, exhibit the telomere-integrated only phenotype in dividing host splenic cell populations [McPherson et al., 2016]. However, the chromosome-associated phenotype or the chromosome-associated/telomere-integrated phenotype represent host mitotic cells containing the Rispens vaccine strain, the oncogenic Md5 strain, or both [McPherson et al., 2016].

**Statistical Analysis**

Raw cell counts for each of the 4 cytogenomic viral phenotypes were collected by analyses of mitotically dividing cell populations from 3 individual bird samples within each genotype (6
and 7
), treatment group (Md5 oncogenic MDV challenge, Rispens or HVT MD vaccination), and timepoint (1, 4, 7, 14, 21 dpc/dpv). The raw count data were transformed using a logarithm of the odds (logit) transformation [logit (k/n) = ln[(k + 1/n) - k + 1]], in which k out of n total analyzed cells were observed for any given viral phenotype within a bird, treatment group, and timepoint, as described [Robinson et al., 2014]. The logit transformation is applied to fit categorical data in the form of proportions to a linear model. The transformed data were evaluated by two-factor ANOVA and Tukey’s multiple comparisons of means post-hoc test, with a 95% family-wise confidence interval, in R (version 3.3.1) [R Development Core Team, 2016] to establish statistically significant differences within the genotypes. The count of metaphase cells identified within each viral phenotype was the dependent variable in ANOVA analyses. The Tukey’s method of multiple comparison analysis tests all pairwise differences between groups and reduces the probability of type I error. The effects of MDV strain (treatment group and dpc/dpv) on viral phenotype within a given chicken genotype were calculated by this method. Furthermore, statistically significant differences in the normalized phenotype data between genotypes (for each treatment and timepoint) were determined by multiple one-way ANOVA analyses in R.

The data from the dual-treatment bird groups were collected, logit-transformed, and evaluated by two-factor ANOVA and
Fig. 1. Representative cytogenomic phenotypes of oncogenic Marek’s disease (MD) virus (Md5 strain) or MD vaccine (Rispens and HVT) interactions with the host genome at 21 days after challenge or vaccination. The FISH images signify the status of the herpesvirus with regard to the chicken host genome in mitotic splenic-derived cells of disease-susceptible birds. The null phenotype lacks viral FISH signals (FITC, green) around the chromosomes (DAPI, blue). The viral chromosome-associated phenotype is defined by diffuse viral FISH signals surrounding the host chromosomes, while the chromosome-associated/telomere-integrated phenotype consists of both the diffuse associated signals and bright, punctate integrated-virus signals at the telomeres. The telomere-integrated only viral phenotype is exclusively composed of distinct, punctate FISH MD virus signals at the telomeres.
Tukey’s post-hoc test, as described above, to establish statistically significant differences within the genotypes (i.e., between the unvaccinated + MDV-challenged and the vaccinated + MDV-challenged treatment groups). The effects of vaccination (treatment) and dpc (timepoint) on viral phenotype within each chicken genotype were calculated by this method. Statistically significant differences in the normalized viral phenotype data between the genotypes (for each timepoint) were, similarly, determined by multiple one-way ANOVA analyses, as described above.

**Results**

All cytogenomic data are represented in the figures and tables as average values from 3 individual splenic-derived samples (3 different birds) per treatment type, timepoint, and genotype. The oncogenic and vaccine virus-host cell population phenotypes are shown in Figure 1. The dynamic nature of the phenotypes over time (temporal) for the 2 genotypes and single or dual treatments over 1–21 days post treatments are found in on-line supplementary Figures 1 and 2 (for all online suppl. material, see www.karger.com/doi/10.1159/000495174), with all data comprehensively reported in on-line supplementary Tables 1 and 2. Here, we emphasize the key results for genotype and vaccine treatment virus/host interactions, as outlined in Figures 1–3.

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**Fig. 2.** Virus-host cytogenomic interactions following oncogenic Marek’s disease (MD) virus (Md5 strain) or MD vaccine strain (Rispens or HVT) inoculation of disease resistant (63) or susceptible (72) genotypes. Each stacked bar graph represents the virus-host cytogenomic phenotypes (see Fig. 1) observed for oncogenic MDV, the Rispens vaccine, or the HVT vaccine strain, respectively, at 1 day after challenge/vaccination (1-day-old birds) or 21 days after challenge/vaccination (3-week-old birds). The colored bars within each plot represent the mean percentages of mitotically dividing host cells with a given phenotype for 3 birds per treatment group. A Virus phenotypes for the 63 (MD-resistant) birds. B Virus phenotypes for the 72 (MD-susceptible) birds.

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**MD-Resistant Chicken Genotype 63: Comparisons of Oncogenic MDV and MD Vaccines at the Level of the Host Genome**

Notable features of the oncogenic MDV-host genomic interactions within MD-resistant birds were a significant increase in the viral telomere-integrated only phenotype concomitant with a decline in chromosome-associated phenotype by 21 dpc (Fig. 2A). The nonlinear changes of the oncogenic MDV viral phenotype across timepoints indicated a variable or sporadic nature of oncogenic MDV genomic interactions during early infection in the MD-resistant birds (online suppl. Fig. 1).

The cytogenomic interactions of the MD vaccines with the host genome in the MD-resistant genotype significantly contrasted with those of oncogenic MDV at multiple points (online suppl. Table 1). The vaccine treatment groups almost completely lacked the telomere-integrated only phenotype across almost all experimental timepoints; as a result, this viral phenotype appeared in a significantly lower percentage of dividing cells as compared to the oncogenic MDV group from 7 dpc/dpv onwards. The Rispens-vaccinated birds had a higher mean percentage of null phenotype (no virus) cells in the splenic-derived samples as compared to the MDV-challenged birds at all timepoints of early infection except at 4 dpc/dpv. The HVT-vaccinated group showed a significant phenotypic pattern...
fluctuation at 7 dpv with a sharp rise in the associated phenotype and dramatic decline in the “transitional” chromosome-associated/telomere-integrated phenotype (online suppl. Fig. 1). The MDV-challenged group showed a lower mean percentage of cells with the chromosome-associated viral phenotype than both MD vaccinated groups at 4 and 14 dpc/dpv and a significantly lower percentage as compared to the vaccine groups at 21 dpc/dpv.

MD-Susceptible Chicken Genotype 7₂: Comparisons of Oncogenic MDV and the MD Vaccines at the Level of the Host Genome

In the MD lymphoma-susceptible genotype 7₂ birds, there was a significant increase in the viral telomere-integrated only phenotype and a decrease in chromosome-associated phenotype between 1 and 21 dpc for the oncogenic MDV-challenged group (Fig. 2B), similar to the observation within genotype 6₃. These phenotypic changes over the course of early infection were near linear, indicating a more predictable pattern of oncogenic MDV-host genomic interactions in MD-susceptible birds (online suppl. Fig. 1). Additionally, the mean percentage of cells with the MDV null phenotype showed a sharp decline at 21 dpc (around the timing of lymphoma development), which was significantly less compared to all previous timepoints.

The patterns of chromosome association and integration for the MD vaccine-viruses within the MD-susceptible birds sharply contrasted with that of oncogenic MDV (Fig. 2B). The MD vaccine treatment groups contained almost no telomere-integrated only viral phenotype mitotic cells. The mean percentage for the integrated only phenotype was significantly less in the vaccinated groups versus the oncogenic MDV challenge group from 7 to 21 dpc/dpv (online suppl. Table 2). Notably, the chromosome-associated viral phenotype was nearly absent in the oncogenic MDV-challenged birds at 14 and 21 dpc but remained in a high (>50%) percentage of splenic cells in the vaccinated birds across both timepoints. At 21 dpc/dpv, the challenged group lacked a population of MDV-free cells, while the null (virus-free) phenotype persisted at significantly higher levels in the vaccinated birds (Fig. 2B). Unlike in the MD-resistant genotype, the HVT and Rispens vaccine strains did not reveal significant viral-phenotype pattern fluctuations between timepoints.

Genotypes 6₃ versus 7₂: Interactions between Oncogenic MDV Strain and the Host Genome

The oncogenic MDV-challenged birds from both genotypes showed a decline in the chromosome-associated viral phenotype and a rise in the telomere-integrated only phenotype within dividing cells during early infection (Fig. 2).
From 7 dpc to the later timepoints, the MD-resistant genotype had higher mean percentages of chromosome-associated phenotype cells (\( p \leq 0.01 \) at 7 and 14 dpc and \( p \leq 0.05 \) at 21 dpc) and lower mean percentages of telomere-integrated only phenotype cells (\( p \leq 0.1 \) at 7 dpc and \( p \leq 0.05 \) at 14 dpc) (online suppl. Fig. 1). The mean percentage of MDV-free (null phenotype) cells as compared to the susceptible chicken genotype at 2 weeks after vaccination (\( p \leq 0.01 \)) (online suppl. Fig. 1). Furthermore, the telomere-integrated only phenotype was not detected in splenic-derived cells of the Rispens-vaccinated MD-resistant and -susceptible birds, with a few exceptional incidences (online suppl. Tables 1, 2).

The HVT-vaccinated treatment groups had inversely fluctuating mean percentages of viral chromosome-associated phenotype and viral associated/integrated phenotype cells in both genotypes (online suppl. Fig. 1). The difference in the mean values for these viral phenotypes between the genotypes was significant at 4 and 7 dpc. These fluctuations indicate a highly variable nature of HVT-host genome interactions in the dividing cell population. The difference in the associated/integrated HVT phenotype between the genotypes was statistically significant at 21 dpc. The high percentage of cells with the associated/integrated viral phenotype in the resistant birds and the large decline of this phenotype in the susceptible birds at 21 dpc was accompanied by a respective decrease and increase in the percentage of HVT-free (null phenotype) dividing cells (Fig. 2). Similar to the Rispens-vaccinated group, the HVT-vaccinated group did not develop a population of telomere-integrated only splenic-derived cells at any timepoints, with a few cells as rare exceptions (online suppl. Tables 1, 2).

**Genotypes 6\(_3\) versus 7\(_2\): Comparisons of Unvaccinated and Vaccinated Birds after Challenge with Oncogenic MDV**

The MDV-host genomic interactions within MD-resistant birds were altered by day-of-hatch vaccination with Rispens, determined by comparison to unvaccinated and MDV-challenged resistant birds (online suppl. Fig. 2). At later timepoints (14 and 21 dpc), the population of splenic-derived cells with the viral telomere-integrated only phenotype was higher in unvaccinated birds. These cells represent solely an MDV infection (i.e., no vaccine present). At 1 dpc, the vaccinated resistant birds, versus unvaccinated, showed a significantly higher population of splenic-derived dividing cells free of virus (null phenotype) concomitant with a lower population of viral chromosome-associated/telomere-integrated phenotype cells (Fig. 3A). At 21 days after MDV challenge, the vaccinated birds indicated significantly more viral chromosome-associated phenotype in splenic-derived cells. Aside from these instances, the percentages of the host mitotic cell populations with each virus-host phenotype were not significantly different between the vaccinated and unvaccinated birds (online suppl. Fig. 2).

**MD-Susceptible Genotype 7\(_2\): Comparisons of Unvaccinated and Vaccinated Birds after Challenge with Oncogenic MDV**

The cytogenomic interactions of MDV with the host genome in the MD-susceptible genotype greatly contrasted between the Rispens-vaccinated and unvaccinated groups (Fig. 3B). The vaccinated and challenged treatment group almost completely lacked the MDV telomere-integrated only phenotype across all of the experimental timepoints (online suppl. Table 3). As a result, this viral phenotype appeared in a significantly lower percentage of cells as compared to the unvaccinated group from 4 dpc onwards (online suppl. Fig. 2). Importantly, this phenotype represents solely MDV infection of the dividing host cell. The Rispens-vaccinated/MDV-challenged group had a higher mean percentage of viral null phenotype (no virus) and chromosome-associated (replicating virus) phenotype cells in the spleen as compared to the unvaccinated group. This higher percentage was significant for the null phenotype from 7 through 21 days post MDV challenge and for the associated phenotype at 21 dpc. The percentage of host cells with the transitional, chromosome-associated/telomere-integrated phenotype was generally similar between the unvaccinated and vaccinated susceptible birds, except for at 7 days post MDV challenge.

**Genotypes 6\(_3\) versus 7\(_2\): Genotype Effects on Virus-Host Genomic Interactions in Rispens-Vaccinated and Oncogenic MDV-Challenged Birds**

The unvaccinated disease-resistant versus -susceptible genotype birds greatly differed in their virus-host cytoge-
nomic interactions after challenge with oncogenic MDV at 4 dph (online suppl. Fig. 2), as was observed for the birds challenged with MDV at hatch (online suppl. Fig. 1). However, day-of-hatch vaccination for MD, prior to challenge, altered the virus-host interactions to a similar outcome in disease-susceptible and -resistant birds (Fig. 3). Rispens-vaccinated/MDV-challenged (i.e., dual treatment) resistant (6\textsubscript{r}) and susceptible (7\textsubscript{s}) birds indicated analogous patterns of viral chromosome association and integration phenotypes (online suppl. Fig. 2), and statistically significant differences between the genotypes were scarce (online suppl. Table 4). The resistant birds indicated a significantly lower percentage of viral chromosome-associated phenotype dividing cells at 1 and 4 dpc as compared to the susceptible birds ($p \leq 0.05$). The MD-resistant birds also indicated significantly more virus-free dividing cells at 1 and 4 dpc. Notably, the telomere-integrated only phenotype was rarely detected in the spleen mitotic cell populations of vaccinated/challenged birds from both genotypes (Fig. 3).

**Discussion**

MD in commercial chicken populations is largely controlled by selective breeding for viability and disease resistance coupled with widespread vaccination. Our understanding of the biological and molecular mechanisms behind these preventative measures and the interplay between MDV and the host genome, especially with regard to bird genotype, is limited. The inbred chicken genotypes 6\textsubscript{r} (MD resistant) and 7\textsubscript{s} (MD susceptible) share the same MHC haplotype, yet have unique genetic backgrounds and immune responses that contribute to their dissimilarity in MD susceptibility [Palladino et al., 1977; Lee et al., 1981; Vallejo et al., 1997; Baigent et al., 1998; Kaiser et al., 2003; Zhang et al., 2006]. Our cytogenomic results provide new insights with regard to host genetic background effects on pathogenic MDV and MD vaccine-virus behavior, encompassing viral replication, integration, and latency, with single-cell resolution. Our data also outline the effect of Rispens vaccination on oncogenic MDV interactions with the host genome in resistant and susceptible birds, as compared to the interactions observed in unvaccinated birds. The viral infection stages represented by the virus-host phenotypes in splenic-derived, mitotically dividing cells were previously described [Robinson et al., 2014; McPherson and Delany, 2016; McPherson et al., 2016], and this work confirms that MD vaccines can integrate into the host genome, but do not generate a robust telomere-integrated only cell population as seen with oncogenic viruses [McPherson et al., 2016]. The exceptionally rare instances of the Rispens integrated-only viral phenotype may represent transformation-susceptible host cells induced by MD vaccine virus. Furthermore, the cytogenomic profiles for all 3 MDV strains indicated MDV-positive dividing host cells at surprisingly high levels during early timepoints after challenge or vaccination (1 and 4 dpv/dpc). This finding indicates that oncogenic MDV and MD vaccines may preferentially infect replicating host lymphocytes and/or induce cellular proliferation upon initial infection, thus resulting in elevated representation within mitotic cell-specific datasets.

For both chicken genotypes in this study, oncogenic MDV showed a decline in viral replication and significant emergence of latent virus, as represented by the telomere-integrated phenotypes, around the timing of MD tumor development (21 dpc). The MD vaccines, which lack oncogenic transformation potential, conserved the viral replication stage later into the infection (through 21 dpv). These results indicate that vulnerability to development of MD lymphomas, or the oncogenic potential of an MDV infection, is correlated with an elevated dividing cell population with telomere-integrated oncogenic MDV and a loss of the virus-free cells in the lymphoid organs. The reemergence of Rispens and HVT vaccines viral replication may stem from the absence of the “classic” switch of pathogenic MDV to the latency and oncogenic transformation herpesvirus stages. In contrast, some studies have indicated a positive correlation between MDV virulence and replication levels in host cells [Yunis et al., 2004; Dunn et al., 2014]. The contrast in our findings could be explained by a less productive vaccine-virus replication (lower viral genome production detected by quantitative PCR methods) occurring in a large population of dividing splenic lymphocytes within the vaccinated host [McPherson et al., 2016].

The improved clearing of highly oncogenic MDV in the MD-resistant 6\textsubscript{r} birds seems unsurprising, as the reduced presence of pathogenic viral genomes in host cells would decrease the probability of disease progression and tumor development. The greater presence of the telomere-integrated only phenotype in MD-susceptible hosts supports a strong correlation between this MDV phenotype and MD-induced oncogenic transformation and tumor cell populations, as seen in prior studies [Robinson et al., 2010, 2014; McPherson et al., 2016]. This correlation may be underlined by increased proliferation of transformed cells with integrated virus, thereby increas-
ing the proportion of MDV-integrated phenotype cells in lymphoid tissue(s). Integrated-only phenotype mitotic cells were also detected (at significantly lower levels) in challenged MD-resistant hosts, suggesting that the profusion of this viral phenotype, rather than its presence alone, in the lymphoid tissue is the strongest indicator of MD tumor development susceptibility. As a whole, these cytogenomic findings indicated unique early-infection patterns of virus-host phenotypes for pathogenic MDV and the apathogenic HVT and Rispens MD vaccines.

Our investigations of vaccination and subsequent MDV challenge in the same host bird genotypes (i.e., dual-treatment group) provided insights on the impact of vaccination on oncogenic MDV behavior. Analyses of immune cell populations in vaccinated/challenged host birds, in the absence of differential labeling between vaccine and oncogenic MDV strains, were possible due to the scarcity of the telomere-integrated only phenotype in Rispens-vaccinated only birds. Vaccination with Rispens significantly reduced the percentage of splenic-derived dividing cells with the telomere-integrated only phenotype and resulted in more virus-free cells after MDV challenge in disease-susceptible hosts. The disease-resistant genotype demonstrated less contrast in vaccinated versus unvaccinated outcomes, likely because of the genotype-based resistance to MD tumor development in these birds. The larger population of chromosome-associated phenotype cells in the vaccinated birds aligns with our knowledge that vaccination for MD does not induce sterilizing immunity and, thus, host immune cells may contain replicating vaccine and/or oncogenic MDV without inducing disease symptoms. The low presence of the telomere-integrated only phenotype in the vaccinated (i.e., protected) birds may be a hallmark of effective vaccination against MD and, further, supports the hypothesis that this viral integration phenotype represents a cell primed for virus-induced transformation. Through use of single-cell cytogenomic approaches for evaluation of viral dynamics, as described here, the MDV telomere-integrated-only phenotype abundance in lymphoid tissues may prove to be a strong candidate for evaluating MD vaccine protective efficacy. Furthermore, the signature virus-host dynamics involved in vaccinal protection, disease progression, and MD-induced oncogenesis, established by this study, provide analytical tools in the effort to continuously protect the world’s poultry populations from serious MD outbreaks.

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Statement of Ethics

The authors declare that all experiments performed in this study comply with the current laws of the United States of America. All animal experiments were carried out in accordance with the National Institutes of Health guide for the care and use of Laboratory animals (NIH Publications No. 8023, revised 1978) and the guidelines set forth by the USDA, ARS, ADOL Institutional Animal Care and Use Committee (IACUC).

Disclosure Statement

The authors have no conflicts of interest to declare.

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