Defective (interfering) viral genomes re-explored: impact on antiviral immunity and virus persistence

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Defective viral genomes (DVGs) are natural products of virus replication that occur in many positive and negative sense RNA viruses, including Ebola, dengue and respiratory syncytial virus. DVGs, which have severe genomic truncations and require a helper virus to replicate, have three well-described functions: interference with standard virus replication, immunostimulation, and establishment of virus persistence. These functions of DVGs were first described almost 50 years ago, yet only recent studies have shown the molecular intersection between their immunostimulatory and pro-persistence activities. Here, we review more than half a century of scientific literature on the immunostimulatory and pro-persistence functions of DVGs. We highlight recent advances in the field and the critical role DVGs have in both the acute and long-term virus–host interactions.

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DVGs, a historical perspective

Defective viral genomes (DVGs) were first described in the early 1940s by Preben Von Magnus when he observed a decrease in infectivity of influenza virus passaged at high titers [1]. Von Magnus posited that particles of ‘incomplete’ virus were produced when viruses were expanded at high titers, and that these particles interfered with viral replication. His hypothesis was supported by the identification in virus stocks of a component with a lower sedimentation rate that associated with interference. This component was named ‘Von Magnus particles’ [2]. Follow-up experiments demonstrated that the production of Von Magnus particles was independent of the number of infective particles used in the infection [3]. Instead, it depended on the ratio of infectious (ID50) to noninfectious particles in the original stock [4]. In the late 1950s, incomplete and interfering particles of Rift Valley fever virus were discovered in viruses passaged in mice at high concentrations [5] and during the 1960s, they were described in several other RNA viruses, including vesicular stomatitis virus (VSV), lymphocytic choriomeningitis virus, and Sendai virus (SeV) [6–11]. However, it was not until the late 1960s that a distinct RNA species shorter than the standard viral genome was reported in incomplete interfering viruses. The amount of this smaller genomic RNA correlated with the ratio of infectious to noninfectious particles [12–14]. It was then speculated that the Von Magnus particles resulted from errors during viral genome replication or during its incorporation into the virion [13,15]. Incomplete viral RNA genomes contained in interfering particles would later be known as defective interfering (DI) particles and, more recently, DVGs [16].

In 1970, Alice Huang and David Baltimore coined the term defective interfering particles, or DIPs, and defined DI particles as follows: “they contain normal viral structural proteins; they contain a part of the viral genome; they can reproduce in the presence of helper virus; they interfere specifically with the intracellular replication of nondefective homologous virus” [17]. Based on infections with VSV and influenza virus, Huang and Baltimore introduced the theory of DI ‘waves’ to explain the asynchronous cycles of DI particles and standard virus observed in persistently infected cultures (details of this theory in the following sections). Importantly, Huang and Baltimore proposed that DI particles played a critical role in determining the course of natural viral infections, including the establishment of virus persistence [17].
During the 1970s and 80s DI particles were described in many other RNA viruses, including poliovirus, rabies, measles, human parainfluenza virus 3, Semliki Forest virus and Sindbis virus [18–27]. It was quickly evident that the presence and amount of DI particles correlated with the establishment of persistently infected cells in vitro [9,19,28–34] and in vivo [35–38]. This was followed by the description of increased survival of infected mice in infections containing DI particles [5,39,40] and by observations that DI particles enhanced the production of interferon (IFN) during infections [26,30,41–43]. By the 1990s, it was clear that most, if not all, RNA viruses could produce DI particles when grown at high multiplicity of infection, including the important human pathogens Ebola, respiratory syncytial virus (RSV) and mumps [44–46].

Despite strong data supporting important biological roles for DI particles, the relevance of these viral products to natural infections was questioned since their discovery. Moreover, DI particles were considered artifacts of in vitro infections and thus irrelevant to natural infections. Authors frequently discussed that despite the interesting properties of DI particles, they are not naturally produced in vivo and are likely caused by highly artificial methods of passaging the virus [5,47]. This line of thought, together with the lack of appropriate technology to identify and distinguish DI particles from the standard virus, largely limited research on DI particles to their use as tools for studying virus replication and as potential antivirals. Recent renewed interest in studying the role of DI particles during natural viral infections and viral persistence was largely motivated by the identification of DVGs in clinical samples [48,49], demonstrating that they indeed occur during natural infections.

**Interference & immunostimulation by DVGs**

Multiple theories for how DVGs interfere with the replication of standard virus have been tested, including competition for viral receptors, competition for viral components needed for replication, and the induction of IFN [11,30,41,50,51]. These theories are founded on basic knowledge of the structure and properties of DVGs. Though the factors leading to some viruses producing more DVGs than others remain unknown, DVGs form when the viral polymerase loses processivity falling off the template genome and re-attaching elsewhere to complete replication [16]. This alteration during replication leads to truncations of the nascent viral genome resulting in the production of short replication defective genomes. Truncated viral genomes appear in two primary forms: deletion and copyback. Deletion DVGs are formed when the polymerase detaches from the template strand and re-attaches downstream, resulting in the production of DVGs that share their 3′ and 5′ ends with the full-length viral genome. Copyback DVGs are formed when the polymerase detaches from the template and reattaches to the nascent strand, creating a complementary end to the 5′ end of the viral genome. The shorter length of DVGs combined with promoters with increased affinity for viral polymerase in copyback DVGs favor the theory that interference is achieved via competition for viral components, including the viral polymerase (Figure 1A) [11,51–53].

It is well documented that DVGs of several viruses are strong inducers of IFN and they are considered the primary stimuli of antiviral immunity in many infections (Figure 1B) [26,48,54–59]. DVGs activate the intracellular RIG-I-like receptors and turn on the expression of IFNs and proinflammatory cytokines such as IL-1α, TNF, and IL-6. In addition, DVG stimulation optimizes the antigen presentation capacity of specialized antigen presenting cells that initiate adaptive immunity [16,48,57,58,60]. Moreover, accumulating evidence indicates that the immunostimulatory activity of DVGs is maintained in vivo and during natural infections. In mice infected with the respiratory viruses SeV, influenza, or RSV, IFN and proinflammatory cytokines are strongly induced only after DVGs have accumulated to detectable levels [48,56]. Detection of DVGs in respiratory secretions of children infected with RSV correlates with expression of antiviral genes [48], and highly pathogenic influenza virus isolates that fail to induce potent antiviral responses in humans have an impaired ability to generate DVGs [49].

Because of their immunostimulatory activity DVGs have been tested as antivirals and vaccine adjuvants [61,62]. As antivirals, DVGs are expected to interfere with standard virus replication, thereby delaying virus spread and allowing the host time to mount a response [61]. This concept has been extensively proven in mice during infection with Semliki Forest virus, VSV, and influenza virus [39,63,64]. In addition, vaccines complemented with DVGs delivered subcutaneously, intramuscularly or intranasally show improved antibody production and increased protection from virus challenge [65,66]. Moreover, DVGs are present in live attenuated vaccines against polio, measles and influenza viruses [67–70] and are thought to impact their protective activity. Recent work from our laboratory identified a specific secondary structure on SeV DVGs that mediates their strong immunostimulatory activity [71]. Oligonucleotides containing this unique immunostimulatory motif can stimulate protective immune responses during vaccination [66] further illustrating the potential of harnessing DVGs as strong immunostimulants in vivo.
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Review

DVG immunostimulation is also an important factor in modulation of infections in insects. Similar to the pattern recognition receptors RIG-I and MDA5 in mammals, insects have Dicer-2, which senses viral RNA and processes it to produce siRNAs that confer antiviral immunity in insects [72]. Dicer-2 acts preferentially on DVGs relative to standard genome and lead to the control of viral replication and longer survival of infected mosquitos. These observations indicate that the immunostimulatory activity of DVGs is widespread and may have a significant impact in the spread of arboviruses such as chikungunya and Sindbis.

The impact of the immunostimulatory and interfering activities of DVGs for virus spread and clearance during natural infections remains unknown. One possibility is that DVGs act to limit the extent of the infection, thereby delaying the onset of debilitating disease symptoms that would prevent individuals from coming into contact with other susceptible hosts, or in extreme cases kill the host. The strong evidence for DVGs promoting virus persistence argues that keeping the host healthy is not the sole mechanism for viruses to benefit from DVGs. New data provide clues as to how immune stimulation and persistence, two seemingly opposite phenomena, occur in response to DVGs.

The role of DVGs in persistence

A growing number of RNA viruses, previously thought to be acutely infecting, are described to persist in humans, including RSV, human metapneumovirus, Zika, chikungunya, and Ebola [44,73–79]. The mechanisms leading to the establishment and maintenance of persistence of these viruses is poorly understood and contrast with other persistently infecting viruses. For example, retroviruses, such as HIV, integrate within their host's genome, and
DNA viruses, such as herpes, establish episomes in dividing host cells nuclei and produce latent proteins that maintain the persistent infection [80]. RNA viruses are instead thought to remain in the hostile environment of the cytoplasm in the absence of latent viral proteins that help maintain their persistent status. In addition, it is unclear whether persistent RNA viruses produce infectious viral particles either continuously or in response to stimulus as observed in other persistent viruses such as Epstein–Barr virus or HIV [80].

RNA virus persistence is usually pathogenic and associates with the development and exacerbation of chronic diseases. Perhaps the most well studied persistent RNA virus is measles. In persistent measles infections, patients develop subacute sclerosing panencephalitis, a neurologic condition that is often fatal [81]. Other viruses have been associated with chronic diseases, although a causal effect has not been demonstrated. Human metapneumovirus and RSV have been implicated in both asthma and chronic obstructive pulmonary disease [82], and chikungunya virus is associated with chronic arthralgia [78]. In some cases, persistent viruses remain transmissible [36,47,78,81,83,84]. Various RNA viruses persist in immuneprivileged sites including the brain, eyes and testes, and some are capable of being sexually transmitted long after recovery from the disease [76,79,85]. Persistently infected individuals may act as viral reservoirs allowing tourists, emergency healthcare workers, and other travelers to inadvertently carry and spread deadly viruses. For example, there is accumulating data on reactivation and spread of persistent Ebola virus [83,85]. Epidemiological data from the 2014 Ebola outbreak show that a recovered patient transmitted virus 482 days post onset of symptoms infecting 13 different individuals [83]. These epidemiological and biological data highlight the public health hazard of persistent highly pathogenic RNA viral infections.

Although it is well established that DI particles containing DVGs lead to persistently infected cell cultures [9,19,28–34,37], limited data support a role for DVGs in the establishment of persistent infections in vivo. One study reported that mice co-inoculated with Semliki Forest virus and DI particles established a persistent infection through an unknown mechanism [37]. Another study identified measles DVGs in the brain of humans that died due to subacute sclerosing panencephalitis [81]. DVGs may directly lead to persistence, be required for maintenance of persistence, or be produced but not affect persistence. Until recently, the only model of the impact of DI particles on persistence was suggested by Alice Huang in 1970 based on the cycling between high production of standard virus and DI particles during infections in vitro [17]. According to this theory, DI particles arise slowly during virus replication until they reach high concentrations and become predominant. These dominant DI particles interfere with standard virus replication and actively reduce the amount of standard virus. In this process, some cells are infected by standard virus and reinfect the cycle. Other cells are infected only by DI particles, thus yielding no virus. During the next cycle of infection, DI particles are again produced slowly and accumulate repeating the waving pattern of virus production. Asynchronous cycling of DI particles and standard virus occurs in many persistent infections in vitro [86–88] and in vivo [5,88]. The cycling of DI particles and standard virus occur in a predictable pattern and has been mathematically modeled using variations of the predator-prey model [89,90]. Interestingly, in some persistent infections the amount of DI particles appears constant [91]. What drives these cyclic patterns in some viruses but not others, and whether host factors such as the infected cell-type influences the cycling pattern remain unknown.

A molecular mechanism for the establishment of persistence by DVGs was recently reported by our laboratory [92]. Using RNA in situ fluorescent hybridization (RNA FISH) we discovered heterogeneity in the content of viral genome species in epithelial cells infected with SeV in vitro. Some cells are enriched in DVGs, others in standard viral genomes. Cells with a high content of DVGs engage the mitochondria antiviral-signaling protein to induce the production of IFNs and other proinflammatory molecules, including TNF (Figure 2). TNF has two well-characterized opposing functions during infection depending on the signaling pathway it engages. If it signals through TNFR1, it acts as a proinflammatory and proapoptotic factor [93]. In contrast, if it signals through TNFR2, it acts as a pro-survival factor [93]. DVG-high cells express TNFR2 and many of its downstream signaling molecules, including the TNF receptor associated factor 1 (TRAF1). Autocrine TNF thereby extends the survival of DVG-high cells and the infection can persist for months [92]. In contrast, cells enriched in standard viral genome express TNFR1 and TNF promotes their death. A similar phenotype was observed in infections with RSV [92]. This mechanism explains the paradoxical stimulation of both antiviral immunity and establishment of persistence by DVGs and reveals a previously overlooked strategy for host–virus co-existence.

Though DVGs lead to persistent infections, evidence for other mechanisms of RNA virus persistence exists. For example, during parainfluenza virus infection high expression of neuraminidase in infected cells leads to the depletion of viral attachment factors preventing cell to cell fusion and cell death, thereby promoting persistence independently of DVGs [94]. Thus, it is unlikely that DVGs are the only mechanism for the establishment of RNA virus persistence. Host factors, such as cell and tissue types or immunological status of the patient also have a large
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Defective viral genome (DVGs) are sensed by the RIG-I-like receptors RIG-I and MDA5 and activate the downstream signaling component MAVS (1). MAVS signaling leads to the production of TNFR2, TRAF1, IFN, and TNF (2). IFN and TNF are secreted to act in an autocrine or paracrine manner (3). In DVG-high cells expressing TNFR2 and TRAF1, a prosurvival pathway is activated and virus persistence is established. In cells high in standard virus genome but with few or no DVGs the TNFR1 pathway is activated leading to cell death (4).

IFN: Interferon; MAVS: Mitochondria antiviral-signaling protein.

impact on the establishment of persistent infections. How host and viral factors, specifically DVGs, interact is the subject of active investigation.

It remains unclear how the enhanced survival of DVG-high cells leading to persistence fits into the DVG and standard virus cycling observed in many infections. Two nonmutually exclusive models may be envisioned: cycling may occur at the intracellular level, where each infected cell goes through cycles of standard viral genome or DVG enrichment driven by competition and interference with the viral replication machinery (Figure 3A). In this model, a single cell infected by a standard virus accumulates DVGs that will eventually take over the replication machinery thereby reducing standard virus replication to the verge of elimination. With limited viral polymerase available and failure to produce more due to interference, production of DVGs ceases allowing the standard virus to resurge and take over. This model makes the assumptions that cells enriched in standard virus can transition to a DVG-high status before they are killed, and that DVGs are eliminated once polymerase levels are reduced; cycling may occur at the population level, where individual infected cells will determine the composition of the pool of standard virus or DI particles available for infection of new cells (Figure 3B). In this model, cells infected with a standard virus may either die or transition to a DVG-high content. Cells transitioning into a DVG-high state shed both standard virus and DVGs until reaching a high DVG status where viral polymerase complexes are no longer available and
Figure 3. Models for defective viral genome cycling. (A) Intracellular cycling begins with a cell infected with standard virus. Upon virus replication DVGs slowly accumulate and are shed in Defective interfering (DI) particles. Cells that accumulate DVGs may escape cell death and instead transition to a DVG-high status where virus is no longer produced and viral proteins are not longer made due to interference. As a result, viral RNA decreases and the cycle is reinitiated given that DVGs are somehow eliminated. (B) In a population model, cycling also begins with a standard virus infection. As the infection propagates, DVGs form and DI particles are shed. Cells failing to produce DVGs die while cells able to replicate DVGs survive accumulating DVGs until reaching a high content accompanied by low standard virus. As DVGs interfere with viral protein production within cells, DVG and virus production diminishes. Cells that have died from infection with standard virus are eventually replaced by new cells allowing the cycle to begin again as standard virus infects those new uninfected cells. DVG: Defective viral genome.

no more virus is produced. As most standard virus-high cells die and DVG-high cells begin shedding less virus, dead cells are replaced allowing for infection to take hold. As these new cells are naïve, standard virus must infect them in order to establish a productive infection and reinitiate the cycle. This model does not specifically consider other forms of virus spread, such as cell-to-cell fusion, cell division, or virus transfer through cytoplasmic channels, although similar dynamics are expected to apply in these conditions.
Conclusion

Despite decades of work on DVGs and their functions, much remains unknown. As there are different types of DVGs, it is conceivable that immunostimulation, interference and persistence exist as separate functions for at least some. In addition, in light of recent developments, it is important to consider how new models of DVG-standard virus dynamics align with older models of DVG-induced persistence. It is possible that persistently infected cells are not stable but behave dynamically and switch between DVG-high and low states (Figure 3B). This switch would be dependent on the degradation of viral RNAs yet it is unclear how this would occur and whether DVGs are degraded at different rates than standard viral genome. Data for this dynamic intracellular behavior remain limited yet DVG cycling as described in the literature refers to population dynamics as opposed to cellular dynamics. It is thus possible that population cycling is independent of cellular cycling as long as persistently infected cells continue producing viruses. Further work must be done to elucidate cellular dynamics from population dynamics.

Future perspective

Recent data on DVG biology and dynamics during infection raise a number of new questions. Out of particular interest is whether all RNA viruses can persist. SeV is a widely used model virus for the Mononegavirales order of viruses, which includes paramyxoviruses, such as parainfluenza virus, pneumoviruses, such as RSV, filoviruses, such as Ebola and rhabdoviruses, such as VSV. Interfering DVGs have been characterized for all these viruses and many have been observed to persist [7,22,44,45,73,91]. Whether the same mechanism for establishment of persistence applies to similar viruses of this order requires further research and the relevance of this mechanism during natural infections needs to be determined. In addition, many positive sense RNA viruses have been shown to produce DVGs and persist; yet it remains unclear whether they use a similar strategy for persistence. With evidence of DVGs functioning as immunostimulators in insects, it is also important to consider how arboviral DVGs impact infections of mammals versus insects and what is their role during virus spread and transmission between species.

Many additional questions on the contributions of DVGs to antiviral immunity and persistence remain unanswered: what controls DVG generation? What regulates their heterogeneous distribution in an infected population? Do surviving cells remain DVG-high or is there a mechanism for switching to a virus-producing cell? Do persistent viruses adopt unique localizations in order to escape the cell's antiviral sensors? How does DVG production impact viral protein production and what are the impacts on infection? What are the determinants for DVGs stimulating the TNFR2 prosurvival pathway as opposed to the TNFR1 pathway? Though a clear correlation between DVGs and persistence exists, this remains an understudied field. The ability for RNA viruses to persist raises public health concerns and additional questions about the biology of persistence: how long does a persistently infected individual shed virus? Is the individual constantly shedding virus or do events such as stress and subsequent homologous or heterologous infections lead to reactivation of virus? What are the cellular reservoirs? With renewed interest in the study of DVGs, we hope to see many of these questions answered in the near future. In addition, identifying factors that influence establishment and maintenance of persistence is critical to manage the onset and impact of persistent infections. For example, it is important to know if persistent virus may reactivate during immunosuppression or whether some patients are more susceptible to viral persistence, thus being at a higher risk to develop chronic pathologies. As defective virus detection methods become more sensitive and more persistent RNA viruses are identified, we will be better equipped to understand how DVGs contribute to the persistence of important human pathogens.

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Executive summary

Historical studies on defective viral genomes
- Discovery of defective viral genomes (DVGs) more than half a century ago originated from the observation of decreased infectivity from virus passaged at high titers.
- DVGs were shown to occur in most RNA viruses.
- Critical functions of DVGs have been described including correlation with persistent infections, interferon induction and decreased severity of infection in mice.
- Lack of evidence for DVGs occurring in natural infections delayed progress in the field until a recent resurgence of interest accompanied by better technologies to distinguish standard and defective viral genomes.

Interference & immunostimulation
- DVGs can interfere with full length virus replication by competing for the viral polymerase.
- For many viruses DVGs have been described as the primary stimulus for induction of antiviral immune responses. This observation highlights the potential for DVGs to be used as vaccine adjuvants or antivirals.
- Immunostimulatory functions of DVGs are ubiquitous with data suggesting similar function in model mammals, insects and humans.

DVGs & persistence
- Unlike other viruses known to persist, RNA viruses do not form episomes or integrate into host genome.
- RNA virus persistence associates with many chronic diseases and may lead to re-emergence of viruses from persistent reservoirs.
- In vivo data of persistence are scarce yet data do support DVGs drivers of persistent infections in animal models.
- Asynchronous cycling between high production of DVGs and full length virus is often observed during persistence.
- A mechanism for the establishment of persistence by DVGs has recently been proposed. In this model, DVG enriched cells trigger a prosurvival response to TNF while cells high in standard full length virus follow the TNF apoptotic pathway. It remains unclear how ubiquitous this mechanism is and what other mechanisms may exist for the establishment of persistence.

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