Oncogenes: The Passport for Viral Oncolysis Through PKR Inhibition

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ABSTRACT: The transforming properties of oncogenes are derived from gain-of-function mutations, shifting cell signaling from highly regulated homeostatic to an uncontrolled oncogenic state, with the contribution of the inactivating mutations in tumor suppressor genes P53 and RB, leading to tumor resistance to conventional and target-directed therapy. On the other hand, this scenario fulfills two requirements for oncolytic virus infection in tumor cells: inactivation of tumor suppressors and presence of oncoproteins, also the requirements to engage malignancy. Several of these oncogenes have a negative impact on the main interferon antiviral defense, the double-stranded RNA-activated protein kinase (PKR), which helps viruses to spontaneously target tumor cells instead of normal cells. This review is focused on the negative impact of overexpression of oncogenes on conventional and targeted therapy and their positive impact on viral oncolysis due to their ability to inhibit PKR-induced translation blockage, allowing virion release and cell death.

KEYWORDS: oncogenes, viral oncolysis, interferon, PKR, clinical trial

Introduction

Oncogenes were first identified in retroviruses and, initially, were regarded as having a retroviral origin, but further studies confirmed that these genes were captured by retroviruses from its mammalian hosts, leading to expression of altered versions of the mammalian genes. The transforming properties of oncogenes are derived from gain-of-function mutations, shifting from highly regulated homeostatic signaling to an uncontrolled oncogenic state. The most well-characterized oncogenes altered in tumors are the receptor tyrosine kinase epidermal growth factor receptor (EGFR), RAS, phosphoinositide 3-kinase (PI3K)/AKT, and MEK/ERK.

Since oncogenes are part of proliferation and survival signaling pathways, their overexpression has been widely related to tumor generation, progression, and resistance to conventional chemotherapy. Accordingly, pharmacological inhibition of these molecules enhances chemotherapy and radiotherapy efficiency, pointing them for targeted therapy. However, the success of target-directed therapy has been challenged by the high mutation rate that alters the target, leading to development of consecutive drug generations for the same target.

Furthermore, the inactivating mutations of the tumor suppressor genes (p53, pRB) and downregulation of proteins involved in death pathways also contribute to tumor resistance. This scenario, however, is extremely suitable for viral oncolysis, the lysis of a tumor cell mediated by viruses that infect and replicate inside them.

Viral oncolysis. Oncolysis may be achieved by the naturally occurring oncolytic viruses, whose viral selectivity toward tumor cells is governed by the absence of factors that impair viral proliferation in the host cell (as INF type I response), absence of functional tumor suppressor proteins (p53 or pRb), and the overexpression of tumor progression factors that lead to survival signaling activation. On the other hand, lysis of normal cells by naturally occurring oncolytic viruses is not successful, since the host defense response, tumor suppressor, and physiological survival signaling are preserved. Additionally, these viruses themselves do not possess proteins that neutralize host defenses of normal cells. Thus, when delivered to the system, they will spontaneously target the tumor instead of normal cells.

Few naturally occurring oncolytic viruses are available for cancer therapy. Viruses that infect human normal cells and cause disease may be modified and become suitable for viral oncolysis. The strategy involves removal of virulence factors and other genes that are not critical for the infection of tumor cells, but are vital for viral replication in normal cells, artificially creating selectivity against tumor.

Due to this selectivity toward tumor cells, oncolytic viruses have the ability to induce cancer regression without...
Viruses are obligatory and/or loss of heterozygosis at 9p21, where INF-κ is a cytosolic sensor of viral dsRNA. In normal cells, this scenario leads to inhibition of viral genome replication and cell lysis mediated by virion release. Furthermore, PKR activation also induces apoptosis through FADD-mediated and mitochondrial pathways (Fig. 1). Most viruses have evolved to overcome this barrier. Some carry proteins in their genomes that inhibit PKR activation through direct interaction, such as E3L and K3L from vaccinia virus, direct binding to PKR without activation of the kinase domain, and inhibition of PKR dimerization by hepatitis C virus. Additionally, cellular proteins can also be recruited by viruses to inhibit PKR function and recruitment of cellular phosphatases acting directly on eIF2 phosphorylation status, bypassing the effects of PKR activation. On the other hand, in tumor cells, the link between INF and PKR can be disrupted by either intrinsic alteration in tumor genetics as deletion and/or loss of heterozygosis at 9p21, where INF-α, INF-β, and other INF-related genes are located, and overexpression of oncogenes such as RAS (Fig. 2).

**Oncogene-dependent Oncolysis (Natural Tropism and/or Genetic Transformed)**

The convergence of both oncogenes and some molecules encoded by viruses toward PKR is related to the basic condition for tumor proliferation and viral infection: the maintenance or enhancement of protein synthesis rate. Oncolytic viruses with natural tropism find PKR inactivated in tumors due to oncogene overexpression or constitutive activation. Reovirus, a ubiquitous, nonenveloped double-stranded RNA virus, was the first naturally occurring human virus reported to exploit an oncogene signaling in the host cell to induce cell lysis. While the normal tissues, IFN activation leads to inhibition of the viral replication. The antiviral defense system starts to act through viral nucleic acid recognition by intracellular Toll-like receptor (TLR) family. While ssRNA binds TLR-7, dsRNA binds TLR-3. The TLRs are found within the same sites that virus enters the cell. TLRs then induce intracellular signaling that leads to the activation of IFN regulatory factors (IRF)-3, IRF-7, and nuclear factor-kappa beta (NFκB), and the subsequent transcriptional activation of IFNα and IFNβ. Released IFNs bind to its receptors leading, through STAT3, to the transcription of the target genes, which includes PKR, the double-stranded RNA-activated protein kinase. PKR contains a dsRNA-binding domain that binds to duplex regions present in viral RNAs, leading to dimerization, kinase activation, and autophosphorylation of PKR. Activated PKR catalyze the phosphorylation of eIF2α, the translation initiation factor-2 (eIF2), blocking its capacity to recycle Guanosine-5'-triphosphate (GTP). Without recycling, eIF2 becomes unavailable to form the complex with Met-tRNA, impairing initiation of translation. Thus, PKR is a key mediator of the IFN type I-induced antiviral response, acting as a cytosolic sensor of viral dsRNA. In normal cells, this scenario leads to inhibition of viral genome replication and cell lysis mediated by virion release. Furthermore, PKR activation also induces apoptosis through FADD-mediated and mitochondrial pathways (Fig. 1). Most viruses have evolved to overcome this barrier. Some carry proteins in their genomes that inhibit PKR activation through direct interaction, such as E3L and K3L from vaccinia virus, direct binding to PKR without activation of the kinase domain, and inhibition of PKR dimerization by hepatitis C virus. Additionally, cellular proteins can also be recruited by viruses to inhibit PKR function and recruitment of cellular phosphatases acting directly on eIF2 phosphorylation status, bypassing the effects of PKR activation. On the other hand, in tumor cells, the link between INF and PKR can be disrupted by either intrinsic alteration in tumor genetics as deletion and/or loss of heterozygosis at 9p21, where INF-α, INF-β, and other INF-related genes are located, and overexpression of oncogenes such as RAS (Fig. 2). Oncolytic viruses with natural tropism find PKR inactivated in tumors due to oncogene overexpression or constitutive activation. Reovirus, a ubiquitous, nonenveloped double-stranded RNA virus, was the first naturally occurring human virus reported to exploit an oncogene signaling in the host cell to induce cell lysis. While the normal tissues, IFN activation leads to inhibition of the viral replication.

**Viral oncolysis dependent on PKR inactivation.** The antiviral defense system starts to act through viral nucleic acid recognition by intracellular Toll-like receptor (TLR) family. While ssRNA binds TLR-7, dsRNA binds TLR-3. The TLRs are found within the same sites that virus enters the cell. TLRs then induce intracellular signaling that leads to the activation of IFN regulatory factors (IRF)-3, IRF-7, and nuclear factor-kappa beta (NFκB), and the subsequent transcriptional activation of IFNα and IFNβ. Released IFNs bind to its receptors leading, through STAT3, to the transcription of the target genes, which includes PKR, the double-stranded RNA-activated protein kinase. PKR contains a dsRNA-binding domain that binds to duplex regions present in viral RNAs, leading to dimerization, kinase activation, and autophosphorylation of PKR. Activated PKR catalyze the phosphorylation of eIF2α, the translation initiation factor-2 (eIF2), blocking its capacity to recycle Guanosine-5'-triphosphate (GTP). Without recycling, eIF2 becomes unavailable to form the complex with Met-tRNA, impairing initiation of translation. Thus, PKR is a key mediator of the IFN type I-induced antiviral response, acting as a cytosolic sensor of viral dsRNA. In normal cells, this scenario leads to inhibition of viral genome replication and cell lysis mediated by virion release. Furthermore, PKR activation also induces apoptosis through FADD-mediated and mitochondrial pathways (Fig. 1). Most viruses have evolved to overcome this barrier. Some carry proteins in their genomes that inhibit PKR activation through direct interaction, such as E3L and K3L from vaccinia virus, direct binding to PKR without activation of the kinase domain, and inhibition of PKR dimerization by hepatitis C virus. Additionally, cellular proteins can also be recruited by viruses to inhibit PKR function and recruitment of cellular phosphatases acting directly on eIF2 phosphorylation status, bypassing the effects of PKR activation. On the other hand, in tumor cells, the link between INF and PKR can be disrupted by either intrinsic alteration in tumor genetics as deletion and/or loss of heterozygosis at 9p21, where INF-α, INF-β, and other INF-related genes are located, and overexpression of oncogenes such as RAS (Fig. 2). Oncolytic viruses with natural tropism find PKR inactivated in tumors due to oncogene overexpression or constitutive activation. Reovirus, a ubiquitous, nonenveloped double-stranded RNA virus, was the first naturally occurring human virus reported to exploit an oncogene signaling in the host cell to induce cell lysis. While the normal tissues, IFN activation leads to inhibition of the viral replication. The characterization of the interaction of oncogenes with PKR showed that oncogenes have multiple roles in tumor oncology, and sometimes their role overlaps, with several of them present at the same signaling cascade. Either with inherent tropism or genetically engineered, several oncolytic pathways converge toward the RAS, PI3K, and its downstream effectors, making the oncogenes, such as RAS, RAF, MEK, ERK, and AKT, “The golden tickets” for viral oncolysis to succeed. **RAS.** Physiologically, Kras, a homolog of the Kirsten murine sarcoma virus, and its isoforms couple activated cell surface receptors to their intracellular effector pathways. Upon binding with GTP and Guanosine diphosphate (GDP), RAS alternates between active and inactive forms, respectively. The activating mutations in RAS proteins lead to the maintenance of the GTP-bound state and the receptor-independent constitutive signaling. Upon activation, RAS recruits and activates Raf, that phosphorylates and activates MEK, which the only known substrate are ERK kinases. Active ERK translocates into the nucleus and induces phosphorylation of several targets and transcription of genes involved in cancer progression and inhibition of antiviral response. This modular structure makes RAS the main driver and its downstream effectors may control PKR activity either through activating their effectors downward in the cascade or acting in additional routes to allow viral oncolysis. Ras may regulate PKR activity in several ways, including modulation of INF-induced transcription, which reduces PKR expression. IRF-1 has its expression modulated by MEK-downregulated IFN-inducible genes, which in turn may be negatively regulated by RAS/MEK, allowing viral oncolysis. Ras is important for Reovirus uncoating and infectivity. In modified herpes simplex virus type 1 (HSV-1), Ras...
Impact of overexpression of oncogenes

signaling also dictates host–cell permissiveness. These and other studies are mostly based on pharmacological inhibitors of the Ras signaling pathway and cell transformation with RAS, but there is no evidence of physical interaction between RAS and PKR. Furthermore, there are few direct demonstrations of how the downstream effectors regulate the PKR activity.

**RAF.** Raf, the homolog of v-Raf murine sarcoma viral oncogene, the family of protein kinases (A-RAF, B-RAF, and C-RAF, also known as RAF-1), acts as a central link between Ras and the downstream kinases MEK and ERK. Activating mutations in Raf family members may render independence of Ras activation and resistance to targeted therapy. One of the most studied Raf mutations (B-RAF<sub>V600E</sub>) is also involved in viral oncolysis permissivity.

The crucial involvement of Raf-1 as Ras cascade intermediate allowing the downregulation of JAK/STAT pathway has been shown, since knockdown of Raf-1 inhibits hepatitis C virus replication. Raf-1 is important not only as a part of mitogen-activated protein kinase (MAPK) cascade but also directly allows viral replication success. In parvovirus-mediated oncolysis, VP (viral capsid) proteins phosphorylation by the Raf-1 kinase at Ser-2, -6, and -10 on the VP2 N-terminal.

**Figure 1. Viral oncolysis dependent on RNA-activated protein kinase (PKR) inactivation.** In normal cells (A), the presence of oncolytic viruses activates interferon type I pathway, leading to PKR expression and activation. Activated PKR phosphorylates eIF2α, impairing GTP recycling and blocking cell translation. The presence of PKR may induce apoptosis of the infected cell. In tumor cells (B) where interferon type I response is disrupted, oncolytic viruses does not induces PKR expression, allowing GTP recycling in eIF2α and viral protein translation.
Figure 2. The role of oncogenes in translation blockage-induced by PKR. Under overexpression of oncogenes as RAS and its downstream effectors, the interferon type I response is impaired in several points. The point mutations that make these targets resistant to therapy, maintains their activity, which affects transcription of PKR and its activation. Along with PKR inhibition, the overexpression of PI3K/Akt/mTor lead to repression of translation repressor 4E-BP1, releasing eIF4E to form the Translation initiation complex, promoting viral protein translation.
domain is essential for the nuclear translocation and capsid formation of MVM assembly intermediates.69

**MEK/ERK.** MEK is a key regulatory kinase activated by RAF kinases. The activating mutations in upstream members of the cascade lead to constitutive activation of MEK and its substrates, ERK1 and ERK2, in a large percentage of tumors, pointing MEK and ERK1/2 as pharmacological targets in cancer therapy.60

The resistance to MEK inhibitor is related mainly to activating mutation MEK6061, making it independent from RAS activation,61 and the activation of alternative activation of PI3K/AKT, leading to combination strategies, by inhibiting both MEK and AKT.62 When ERK is phosphorylated by MEK, genes involved in IFN production as IRF-1 and STAT2 are downregulated.20,50 Thus, tumors that resist to inhibitors of MEK will maintain the downregulation of PKR, allowing viral oncolysis.

Additionally, with its role in the inactivation of PKR through its downstream effectors, the importance of MEK itself for tumor oncolysis was also shown. Smith et al63 showed that activated MEK suppresses activation of PKR in mutant HSV, and they concluded that MEK could directly or indirectly inhibit PKR activity. This suggestion was reinforced by studies in vascular intimal hyperplasia using HSV-1.64

**MAPKAPKs.** The activation of MAPK signaling module has additional downstream players that also are able to inactivate PKR response. Among the targets of MAPKs are the MAPK-activated protein kinases (MAPKAPKs) that include MK2 and MK3. The activation of the three main cascades of the MAPK module, ERK, p38, and JNK, converge to MK2 and MK3 with ERK and p38 as their main activators.65 MK2 and MK3 possess important roles in inflammation,66 cell cycle regulation,67 and cell migration.68 In tumor cells where its upstream regulators (such as RAS and RAF) of the cascade are overexpressed, active MK2 and MK3 may interact with a complex containing repressor of the inhibitor of PKR (p88IPK), PKR inhibitor p58IPK, and PKR itself, leading to suppression of the PKR activity and the consequent preservation of the eIF2 function.69 Interestingly, overexpression of PKR inhibitor p58IPK may induce malignant transformation.70

Additional findings of this study showed that MK2 and MK3 kinases are phosphorylated and activated in lung cancer cell line under infection of influenza virus; that influenza A virus propagation is strongly reduced in mouse fibroblasts deficient in either MK2 or MK3; and that the activation of MKs reduces the activity of PKR.69

**PI3K/Akt/mTOR.** Akt is a serine/threonine kinase, is one of the downstream targets activated by phosphoinositide 3-kinase (PI3K), and, as other oncogenes, AKT was named after its viral oncogene homolog (v-Akt) present in the murine leukemia AKT8 retrovirus.71 Akt is a key regulator of important cellular functions, including cell survival, proliferation, glucose metabolism, and protein synthesis.72 In the majority of human cancer cells, the Akt pathway is either mutated or constitutively activated, leading to cancer progression through uncontrolled cell proliferation and apoptosis inhibition.73

The importance of AKT for viral oncolysis has been demonstrated for myxoma virus replication in human lung cancer cells, where kinase activity of Akt is required,74 while cells with no phosphorylated Akt were not infected. The same was observed in metastatic ovarian cancer that is also affected directly by AKT.75 The influence on protein synthesis by mTOR relies on regulation of translation initiation factor 4E-binding proteins (4E-BPs), promoting translation initiation.76 A pathway may be cooperatively modulated by both RAS and AKT downstream effectors.77 Sarinella et al78 found that oncolysis of pancreatic tumor by HSV-1 does not rely on RAS activation, but on PI3K in a suggested mechanism that involves modulation of eIF2B through inhibition of GSK-3, leading to independence of PKR status.

**Tumor suppressor protein status and PKR.** The inactivating mutations found in tumor suppressors Rb and p53, associated with overexpression of RAS, may lead to tumorigenesis,79 tumor progression, and drug resistance. Among the functions performed by p53 are the cell cycle control and transcriptional activation of proapoptotic proteins, also, p53 interacts with PKR promoter, leading to its expression.80 The central role of p53 in cell death induction led several viral species to develop strategies to inhibit p53 and Rb activity in normal cells.81 In tumor cells, viral tropism is enhanced where these proteins are not functional.19

Protein Rb (retinoblastoma susceptibility protein I) inhibits cell cycle progression from G1 to the S-phase. Under phosphorylation by cyclin D, cyclin-dependent kinase 4 (CDK4), and CDK6, Rb became inactive allowing proliferation. Overactivation of their upstream regulators, mutation, or deletion of Rb favors tumor proliferation.82

Recently, Kline et al83 showed that activation of transcription factor 4 (ATF4) in response to a small molecule required PKR activity with consequent eIF2α phosphorylation. This pathway also induced dephosphorylation of the retinoblastoma (Rb) protein, leading to inhibition of cell cycle progression. This is one example of how, under pharmacological activation, PKR and Rb contribute to tumor cell death.

Dysfunctional or deleted p53, ATM, and Rb conferred enhanced susceptibility to reovirus and myxoma viral infectivity, replication, and cytolysis in tumor cells when compared to the cells where these molecular activities are preserved.17 Modifications in adenovirus to target Rb-defective cells have been performed.84 The absence of p53 activity and the presence of activated RAS strongly reduce PKR expression and activity, along with inactivation of Rb, which sets the favorable environment for replication of oncolytic viruses.

**Clinical Trials**

The targeting of several oncolytic viruses under clinical trials may be spontaneous since these viruses recognize tumor cells that overexpress surface molecules such as sialic acid, CD46,
and nections, among others, which defines the tropism toward these cells.\textsuperscript{85} Viruses may also be engineered to recognize cell surface receptors of specific tumor types, such as adenovirus Ad5/3-A24, which was modified to bind to integrins of ovarian cancer cell surface.\textsuperscript{86} The interplay among oncogene overexpression, tumor suppressor gene status, and IFN-1 response has been explored in clinical trials using viruses that take advantage of tumor molecular alterations.

The failure to engage type I IFN signaling provides a replicative advantage for spontaneous oncolytic viruses such as Newcastle disease virus (NDV), a paramyxovirus that is low pathogenic for humans, but highly infectious in avian species. NDV is highly sensitive to type I IFN response that this virus elicits in normal cells, what confers its cancer cell specificity.\textsuperscript{87} Attenuated NDV was tested in patients with melanoma, glioblastoma, head and neck, advanced colorectal cancer, and other malignancies.\textsuperscript{88} It was used as live virus\textsuperscript{89} or NDV oncolysates.\textsuperscript{90} Other viruses of this group whose oncolytic potential were already tested clinically are Sendai virus,\textsuperscript{91} measles virus,\textsuperscript{92} and mumps virus.\textsuperscript{93} The patients generally developed flu-like symptoms and there was no death associated with the viral infection.

The production of viral RNAs by reovirus, one of the most well-studied spontaneous oncolytic virus, leads to activation of PKR in normal cells. On the other hand, overexpression of RAS impairs PKR pathway, leading to preferential infection of tumor cells by reovirus.\textsuperscript{44} Reolysin\textsuperscript{®}—Reovirus Serotype-3-dearing Strain, Oncolytics Biotech Inc., a purified live replication competent reovirus, induces cytolysis in tumor cells with an activated ras pathway due to inhibition of the PKR. Reolysin administration was safe and well tolerated, showed partial response and/or stable disease in patients with breast cancer, gliomas, melanoma, and ovarian cancer.\textsuperscript{94} colorectal cancer, metastatic pancreatic adenocarcinoma, and lung cancer,\textsuperscript{95} melanoma, and head and neck sarcoma.\textsuperscript{96} On the other hand, Reolysin showed no objective responses in relapsed and refractory multiple myeloma patients.\textsuperscript{97}

The importance of the tumor suppressor proteins for viral oncolysis was also explored in clinical trials. E1B is a protein that interacts with p53, allowing infection of normal cells by wild-type adenovirus.\textsuperscript{98} ONYX-015/H101 (Oncorine\textsuperscript{R}) is an E1B-55 kDa gene-deleted adenovirus engineered to selectively replicate in and lyse p53-deficient cancer cells.\textsuperscript{99} ONYX-015 was tested for the first time in humans in Phase I and II studies with intratumoral and peritumoral injection in 2000,\textsuperscript{100,101} leading to promising results, overall in p53-deficient cancer cells, as initially intended.

In addition to this modification, the ONYX-015 virus is “armed” with genes that enhance the immunological response toward the tumor, such as granulocyte-macrophage colony-stimulating factor (GM-CSF).\textsuperscript{22} This construction was also used in combination with mitomycin C, doxorubicin, and cisplatin adjuvant chemotherapy.\textsuperscript{102} The most common side effects were flu-like symptoms. No deaths were associated with viral infection, and no limiting dose was determined.\textsuperscript{88}

The capacity of HSV-1 to induce disease in humans depends on proteins that are able to impact the INF-1 signaling, PKR activity, and tumor suppressor proteins. Therefore, important modifications were performed in order to make this virus safe for clinical use: (a) deletion of thymidine kinase (TK), a protein involved in DNA synthesis and repair,\textsuperscript{103} where expression of its cellular homolog is correlated with malignancy; (b) deletion of ICP 34.5 gene, which codes for a phosphatase that dephosphorylates PKR, allowing protein synthesis; and (c) deletion of ICP47, which blocks the antigen presentation in infected cells by inhibiting TAP1 and TAP2 transporters. These alterations were combined in HSV strain JS1 (JS1/34.5/-47),\textsuperscript{105} and its clinical use has been evaluated to treat head and neck tumors and showed high rates of complete response, the absence of recurrence, and the prolonged progression-free survival seen in two-thirds of the patients strongly supporting further clinical studies.\textsuperscript{106} Equally impressive results were also achieved with unrespectable metastatic melanoma.\textsuperscript{107}

JX-594 (Pexa-Vect) is an oncolytic vaccinia engineered from the Wyeth vaccine strain. It has both a disruption of the TK gene and expression of human GM-CSF, which induces tumor-specific immunity.\textsuperscript{108} JX-594 was tested in hepatocellular carcinoma using systemic delivery, and the results showed a strong impact on the reduction of tumor mass and patient survival.\textsuperscript{109} JX-594 was also tested in patients with neuroblastoma and Ewing sarcoma with limited benefits.\textsuperscript{110}

Taken together, the clinical trials with both attenuated and genetically modified viruses showed mostly that virotherapy success is dependent on several factors, such as administration route, with intratumoral injection showing better results over infusion due to the development of neutralizing antibody against several strains,\textsuperscript{111,112} if the virus is used as monotherapy or in combination with other drugs, or immunomodulatory strategies.\textsuperscript{108} The benefits include variable degrees of tumor remission, enhancement of free survival rate, and patient quality of life.\textsuperscript{113} Virotherapy alone was well tolerated and did not induce patient death, and the most frequent side effects were flu-like symptoms, such as fatigue and fever.\textsuperscript{100}

**Resistance to Viral Oncolysis**

Even with its high efficiency in eliminating tumors where chemotherapy has failed, viral oncolysis also faces resistance. If several of the oncogenes that make viral oncolysis possible are resistance factors in both conventional and target-directed therapy, how resistance takes place in viral oncolysis? Maintenance of INF type I responses contributes to resistance to viral oncolysis.\textsuperscript{114} Despite the importance of RAS constitutive signaling to allow viral oncolysis, persistent infection with the reovirus led to oncolysis resistance in fibrosarcoma cells, with PKR constitutively activated, even in the presence of high RAS activity.\textsuperscript{115}

Overexpression of CUG2 oncogene upregulates STAT1, leading to tumor resistance to vesicular stomatitis virus through
maintenance of PKR activity. In addition to the PKR activity through INF type I response persistence, oncolysis resistance may occur as a result of the systemic virus clearance by the host immune system; the reduction of viral receptors on the target cells became an important barrier for viral oncolysis, once surface receptor is the main feature used by viruses to infect host cells. Survivin stabilization may lead to resistance of sub-populations of cancer cells to NDV, and resistance to viral oncolysis may be acquired during malignant progression.

Solid tumors are less organized in its structure and vascularity, due to defects on tumor edges and aberrant angiogenesis. Furthermore, extracellular matrix affect oncolysis in solid tumors, leading to the reduced virus spread. These challenges led to therapies that combine oncolytic virotherapy through intratumoral injection with antitumor conventional or targeted drugs, with promising results.

**Modulation of immune response.** Modulation of immune response is also an important factor in resistance to oncolysis. Preexisting or virus-induced antibodies against the oncolytic strain may be present by previous vaccination, cross-reaction, and response to oncolysis, impairing the systemic spread of the virus. An interesting example is the cross-reaction between clinically tested oncolytic Sendai virus (rodent pathogen) with human parainfluenza virus type 1 (hPIV-1). Thus, the vaccination history of the patient has an important impact on the therapy outcome. Additionally, in several studies where infusion was used as the administration route, the efficiency of the oncolysis was reduced overall in metastatic tumors when compared with intratumoral injection.

One important feature of viral oncolysis is that, by killing infected tumor cells, viruses provide tumoral antigens that allow the immune system to recognize and kill more tumor cells, an approach that is particularly important for the treatment of metastatic lesions. However, the tumor microenvironment may impair the effectiveness of immune response, leading to another important source of resistance to viral oncolysis: the engagement of the immune checkpoint, a consequence of activation of a particular set of pathways that downregulate immunological efficiency against tumors.

The programmed cell death protein 1 (PD1) and its ligand (PD-L1) may be expressed by both tumor cells and macrophages in the tumor microenvironment, turning off T-cells. The expression of PD-L1 may also occur as a consequence of translocations, chromosome aberrations, and EGFR activation, leading to tumor evasion. Cytotoxic T-lymphocyte antigen 4 (CTLA-4) suppresses T-cells within the lymphoid compartment by limiting T-cell proliferation, preventing expansion of antitumor T-cell responses in cancer patients. Immune checkpoint blockade involves the direct use of antibodies to neutralize immune checkpoint pathways (anti-PD-L1 and anti-CTLA-4), also, oncolytic attenuated viruses (such as measles virus) may be used as vectors to encode antibodies against CTLA-4 and PD-L1, thus enhancing the effectiveness of viral oncolysis.

**Biomarkers for viral oncolysis resistance.** There are quite a few proposed markers for resistance to viral oncolysis, one of these is YAP-1 (yes-associated protein) in head and neck cancers. Its expression was correlated with resistance to reovirus, whereas low YAP-1 expression was correlated with sensitivity to reovirus infection. YAP-1 is a candidate human oncogene, found overexpressed in lung, colon, ovarian, and breast tumors. This protein is a nuclear effector of the Hippo pathway, which is a key regulator of organ size. However, this study is in its infancy, and further investigation is needed.

**Concluding Remarks**

The problem of conventional chemotherapy impairment by oncogene overexpression was thought to be solved when targeted therapy was introduced. Again, with targeted therapy, the target mutated, leading to consecutive generations of drugs to overcome resistance. Even though cancer virotherapy has been discussed and tested for the past 60 years, we are finally moving to what may be the golden age of virotherapy, with modified viruses with natural tropism toward tumors currently undergoing clinical trials as cancer therapeutic.

The exploitation of molecular profile of tumors to achieve success in viral oncolysis highlights the importance of multidisciplinary approach that made this therapeutic alternative available. The lesson learned with in vitro and clinical experiences regarding virotherapy is that none of the previously established approaches to clinically treat cancers are dispensable. The multifactorial nature of resistance needs to be counterbalanced by multiple strategies, instead of monotherapies.

The use of multiple strategies with zero mortality involving the use of oncolytic viruses in clinical trials makes this approach efficient and safe. Although not curative yet, it certainly increases the quality and expectancy of patient’s life. Different from other therapeutic strategies to treat refractory tumors, in viral oncolysis, the overexpression of oncogenes mostly drive the solution, rather than the problem, partially due to INF-1 response and PKR inhibition.

**Author Contributions**

Conceived and designed the experiments: JF. Analyzed the data: JF. Wrote the first draft of the manuscript: JF. Contributed to the writing of the manuscript: JF. Agree with manuscript results and conclusions: JF. Jointly developed the structure and arguments for the paper: JF. Made critical revisions and approved final version: JF. Author reviewed and approved of the final manuscript.

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