The past 2 years have seen several major advances in oncolytic virotherapy. Studies on the interaction between viruses, immune responses and tumor microenvironment have provided important insight, while clinical trials have

shown promise. This review summarizes key findings in this

field over the past 2 years, and provides directions for future success.
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• Suppression of innate immune response enhances efficacy

- Carrier cell strategy avoids immune attack
- Targeting tumor microenvironment enhances viral spread and efficacy
- Oncolytic viruses kill cancer stem cells
- Genetic engineering of oncolytic viruses complements chemo-and molecular-targeted therapies
- Genetic engineering of oncolytic viruses targets cancer signaling pathways
- Novel oncolytic virus species are being explored
- A large number of clinical trials have been carried out

Introduction

Progress

The use of live viruses for the treatment of cancer dates back to a century.¹ Cancer-selective oncolytic viruses replicate preferentially in cancer cells and as a result, destroy those cells at the end of replication cycles; normal cells are spared and hence toxicity is limited. Of note, oncolytic viruses can kill apoptosis-resistant tumor cells, and hence do not have cross-resistance with existing therapies. Engineered oncolytic viruses have been developed over the past 15 years and have various mechanisms-of-actions (MOA; Table 1): (1) inherently tumor-selective virus species (for example, RNA viruses, poxviruses); (2) viral gene-deleted mutants—critical viral gene expendable for growth in tumor cells, but not in

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Gene therapy progress and prospects cancer: oncolytic viruses

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In brief

Prospects

• Logical design of the next generation of oncolytic viruses may take this strategy to the next level

- Ex vivo studies may predict responses
- The demand for target validation is increasing
- New imaging endpoints in clinical trials.

normal cells, were deleted (for example, adenovirus *dl*1520/Onyx-015, herpes simplex virus (HSV) G207); (3) promoter engineered mutants—viral replication was engineered to be dependent on inserted tumor-specific promoters, and as a result, the replication of the virus is restricted to tumor cells that are able to activate the promoters (for example, prostate specific antigen-regulated adenovirus CG7870, telomerase-regulated adenoviruses and HSVs); (4) pseudotyped viruses—normal viral tropism is ablated, and viruses are engineered to attach/bind to specific surface receptors that are expressed exclusively/preferentially on tumor cells (for example, adenovirus Delta-24RGD).

Over the past decade, several oncolytic viruses have been tested in humans, and although the safety results are encouraging, efficacy as single agents was limited.¹ Possible hurdles include attenuation of the virus caused by genetic engineering of the virus that renders cancer selectivity, host immune responses and lack of understanding of tumor microenvironment. However, H101, an oncolytic adenovirus similar to Onyx-015 (E1B-55K/

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Approach to selectivity	Agent(s) example and genetic alterations within virus	Genetic target(s) in tumors	
Inherently tumor-selective species	NDV (none)	IFN resistance	
	Reovirus (none) VSV (none)	Ras pathway IFN resistance	
Deletion of viral gene that is necessary for replication in normal cells, but expendable in tumor cells	G207 (ICP6-/γ34.5-deleted HSV-1)	Proliferation, loss of neurovirulence	
	Onyx-015 (E1B-55K-/E3B-deleted Ad) Delta-24 (E1A-CR2-deleted Ad)	Loss of p53 pathway, late mRNA transport Loss of G1-S checkpoint control; loss of pRB function	
	JX-594 (TK-deleted VV)	Proliferation	
Tumor-/tissue-specific promoter engineering to limit viral gene expression	CG7870 (E1A under rat probasin promoter, E1B under PSA promoter/enhancer Ad)	Prostate cancer	
expression	bM24-TE (Wnt/β-catenin-promoter/ enhancer-driven HSV-1)	Wnt/ β -catenin-over expressing tumors (colorectal, hepatoblastoma and so on)	
Pseudotyped viruses	CAR/integrin-binding deleted Ad, replaced with tumor-targeting ligand	Tumor-specific receptor	

 Table 1
 Cancer-selectivity mechanisms of oncolytic viruses

Abbreviations: CAR, coxsackievirus-adenovirus receptor; IFN, interferon; NDV, Newcastle disease virus; VSV, vesicular stomatitis virus.

E3B-deleted), was recently approved by the Chinese government to be used in conjunction with radiation therapy for the treatment of head and neck cancers. This is the first oncolytic virus product approved by a governmental agency for human use. To overcome the obstacles toward efficacious virotherapeutics, several major advances have been made to improve the selectivity and efficacy of oncolytic viruses. This review summarizes recent major advances over the past 2 years from over 400 publications and selected unpublished work.

Progress

Suppression of innate immune response enhances efficacy

Virus-immune system interactions have been extensively studied in the context of virotherapy. Innate immune responses to the virus are a major hurdle for long-term gene expression and oncolytic potency. The use of immunomodulatory agents in combination with oncolytic viruses was first reported in the 1970s. Recently, several groups have shown, using different viruses and animal models, that administration of cyclophosphamide, known to inhibit innate immune responses, can significantly enhance viral spread, transgene expression and antitumoral efficacy. These studies demonstrated that cyclophosphamide is able to inhibit neutralizing antibody induction, macrophages, regulatory T cells (Tregs) induction and intratumoral interferon (IFN)-7 production.²⁻⁸ It remains to be determined, however, how much this approach will benefit cancer patients who often have already various degrees of 'pre-existing' immunosuppression due to disease and chemotherapy.

In addition, Haralambieva *et al.*⁹ showed that measles virus-induced gene expression and intratumoral virus spread is inhibited by IFN, which is triggered by virus infection of tumor cells. Interestingly, currently available oncolytic measles viruses are derived from Edmonston tag (Edmtag) strain, which has lost most of the IFNantagonizing activities. Edmtag-based measles virus engineered to express the measles phosphoprotein (P) gene products (P/V/C proteins) from wild-type measles virus, known to antagonize IFN induction and response, exhibited reduced IFN sensitivity and a reduced IFN induction in lymphoma, myeloma and activated peripheral blood mononuclear cells (PBMC).⁹ Measles virus encoding the P gene products also showed significantly enhanced systemic efficacy in a myeloma xenograft model. This study highlights the importance of innate antiviral responses of tumor cells that need to be considered when designing oncolytic viruses.

Importantly, antiviral immunity does not necessarily reduce the efficacy of virus. A recent report by Zhu *et al.*¹⁰ demonstrated that in mice pre-immunized with HSV, subsequent intratumoral administration of oncolytic HSV showed enhanced efficacy compared to HSV-naive mice. The peripheral blood mononuclear cells from the HSV-seropositive mice also exhibited greater *in vitro* cytotoxicity to tumor cells than PBMC from HSV-naive mice, which correlated with an enhanced IFN- γ induction in PBMC from HSV-seropositive mice.¹⁰ This is an important finding that needs to be explored with systemic HSV administration and also with other oncolytic virus species.

Consistent with our previous findings, Diaz *et al.*¹¹ have shown that host CD8 and NK cells are critical for the efficacy of oncolytic virotherapy. Importantly, using an antibody against Treg (PC-61) they showed that depletion of Treg cells inhibited antitumoral efficacy in

the context of virotherapy, as it overcame the suppression of antiviral immune responses. Furthermore, levels of activated T cells can be significantly increased by enhanced expression of tumor antigens. This can be achieved either through adoptive T-cell transfer therapy, or incorporation of tumor antigen into the oncolytic virus.¹¹

Carrier cell strategy avoids immune attack

In addition to blocking the host immune response, one can take advantage of the immune system to boost antitumor responses. Adenovirus and HSV mutants engineered to enhance the expression of class I major histocompatibility complex (MHC) have been shown to enhance enhanced antitumoral immune responses and efficacy in animal models. However, one major challenge for virotherapy is the inefficient uptake of viruses into tumor cells after systemic administration due to systemic antiviral immune response (for example, neutralizing antibodies and complement). Thorne *et al.*¹² described a novel approach to tackle this issue. Cytokine-induced killer (CIK) cells are known to 'home' to and destroy tumors. After isolating the CIK cells from mice, these cells were infected with oncolytic vaccinia viruses and readministered into tumor-bearing animals. The virus replicated in the CIK cells while these traveled to the tumors. As a result, substantially larger amounts of oncolytic viruses were delivered to the tumor, and both the CIK cells and oncolytic viruses were synergistic in tumor killing.12 It remains to be seen, however, whether CIK cells home to tumors in humans within a reasonable time frame. In addition, this approach requires harvesting of cells from individual patients, ex vivo culturing and redelivery to the patients, and therefore requires a substantial amount of laboratory work. Nonetheless, this strategy holds promise.

Subsequently, the 'carrier cell' approach has been tested by several groups.¹³ Using measles viruses, Ong et al.¹⁴ showed that virus-infected T cells can protect from low, but not high, concentrations of antimeasles immune serum. However, even in measles-naive mice, only 1–2% of the virus-infected T cells trafficked to the tumor site after systemic delivery. Power et al.15 tested oncolytic vesicular stomatitis virus (VSV) vectors with different carrier cell types, ranging from leukemia cell lines (which led to systemic delivery of the virus) to cells derived from solid tumors (which accumulated primarily in the lungs). Using dual-enzyme in vivo luminescence imaging, it was shown that whereas the carrier cells were retained in the body for no longer than 1 day, oncolytic VSV continued to replicate and was able to eradicate pre-established tumors.15 Other carrier cells tested include endothelial cells and mesenchymal stem cells.^{16,17} Vile et al. have recently shown that T lymphocytes can be used to harbor oncolytic VSV and release the virus at the tumor site.18 VSV-loaded lymphocytes were also able to purge spleen and lymph nodes of metastatic cells, which in turn primed antitumoral immunity. Furthermore, adoptive transfer of VSV-loaded lymphocytes reduced metastases.18 This strategy might have great potential for many tumor types.

While the 'Trojan Horse' approach avoids the negative impacts of neutralizing antibodies on virotherapy, there are several critical issues that need to be further investigated before this approach can be taken into humans. First of all, the most appropriate carrier cell types need to be carefully determined. Tumor-homing cells (for example, CIK) and systemic disseminating cells (for example, leukemic cells) have both advantages and disadvantages. Importantly, tumorigenicity is a major concern when cancer cells are used as carriers. Finally, although viruses are able to replicate, the newly generated viruses (that is, 'second wave') still face contact with neutralizing antibodies, which will likely restrict significant virus spread. Multiple administrations of viruses in carrier cells may, therefore, be necessary. Finally, whether or not viral antigens will be presented by carrier cells, and the impact on virus delivery and replication, needs to be explored.

Targeting the tumor microenvironment enhances viral spread and efficacy

The efficacy of virotherapy can be limiting when replication-mediated oncolysis is the sole MOA. Indeed, the tumor microenvironment plays an important role in restricting viral spread and promoting tumor growth. To address this issue, several approaches have been taken. The first is to engineer viral vectors with therapeutic transgenes that target the key components of the tumor microenvironment (for example, the tumor vasculature or matrix). Oncolytic adenovirus encoding relaxin, a matrix-degrading protein, was able to enhance viral spread without causing significant toxicity.¹⁹ Oncolytic HSV encoding dominant-negative fibroblast growth factor receptor or antiangiogenic protein platelet factor-4 led to significant reduction in tumor vasculature and as a result, significantly enhanced therapeutic efficacy.20,21 Others have engineered oncolytic virus replication to be activated by tumor matrix metalloproteinases (MMP),²² and shown that MMP-8 gene delivery enhanced the efficacy of oncolytic adenovirus.²³

An alternative approach is to coadminister therapeutic agents with the virus. Coadministration of matrixmodifying agents (bacterial collagenase, MMP-1, 8) has been shown to enhance the spread of oncolytic HSV^{24,25} although concerns about tumor metastases have to be explored in more preclinical models before translation into clinical trials. In addition, infection with wild-type HSV results in reduction in thrombospondin secretion, a protein that has antiangiogenic properties, from extracellular matrix. This leads to increased vascularity in infected tissues. Aghi et al.26 showed that increased vascularity can be counteracted by introduction of certain mutations into oncolytic HSV, or coadministering a thrombospondin-derived peptide 3TSR. A recent study by Kolodkin-Gal et al.27 showed that the difference in amount of extracellular matrix between normal colon and colon cancer tissues determined the infectivity and subsequent cytotoxicity of HSV. This phenomenon needs to be further investigated.

Tumor hypoxia and its impact on viral replication have also been studied. Previous reports have shown that hypoxia limits group C adenovirus (serotype 5) replication, and new data suggest that the oncolytic activity of other adenovirus serotypes are also affected.²⁸ Importantly, the report showed that the expression level of CD46, a receptor for group B adenoviruses (serotypes 3, 11) as well as measles virus, was not altered in hypoxic conditions. In contrast, we have found that hypoxia enhances the replication of oncolytic HSV (M Aghi *et al.,* unpublished).

Another important issue is to explore how inflammation induced by virus infection impacts on the tumor microenvironment. Breitbach et al.29 showed that administration with VSV and vaccinia viruses resulted in a dramatic transcriptional activation of the proinflammatory neutrophil chemoattractants CXCL1 and CXCL5 and neutrophil attraction. The neutrophils in turn contributed to acute reduction in tumor vasculature. Targeted recruitment of neutrophils to infected tumor beds enhances the killing of malignant cells.²⁹ Recent work by Kurozumi et al.³⁰ also illustrates the importance of targeting the tumor microenvironment to improve the efficacy of oncolytic virotherapy. Using the orthotopic (that is, tumors grown in the organs where it is derived from) immunocompetent rat glioma model, they showed that oncolytic HSV infection increased tumor vascular permeability, host leukocyte infiltration into tumors and intratumoral expression of inflammatory cytokine genes, all of these were part of the inflammatory response after HSV infection. Pretreatment with cyclophosphamide suppressed the inflammation and resulted in reduced tumor vascular permeability.30

Kirn *et al.*³¹ showed that systemically administered vaccinia virus resulted in infection and subsequent destruction of tumor endothelial cells, which led to loss of tumor vascular density.

Oncolytic viruses kill cancer stem cells

In light of recent discoveries in the filed of cancer stem cells, it is becoming clear that those cell populations not only initiate tumorigenesis, but also contribute importantly to resistance to chemo- and radiation therapy. As this cell population is capable of replication and selfrenewal, oncolytic viruses that are designed to target cell cycle-dysregulated tumor cells might also possess the ability to kill cancer stem cells. Indeed, several recent publications have shown that the adenovirus E1A mutant that targets the retinoblastoma-E2F transcriptional factor pathway (Delta-24) is able to kill CD133+ cancer stem cells or CD44(+)/CD22(-/low) cancer initiating cells in vitro, and is also able to eradicate tumors derived from these cancer stem cells.^{32,33} MOA include replication-induced cell lysis (necrosis) and autophagy (degradation of intracellular components in lysosomes).³² It has also been reported that adenovirus serotype 3 was better than serotype 5 in infecting cancer stem cells in vitro. This ability to kill cancer stem cells does not seem to be limited to adenoviruses, as oncolytic HSV can also efficiently kill glioma stem cells (H Wakimoto et al., unpublished). While this is of interest, there are several issues that remain to be solved. As the population of cancer stem cells within a tumor is generally low (often less than 5%), it is a challenge for oncolytic viruses to 'find' and kill these cancer stem cells, especially when viruses are administered systemically. Secondly, there is so far no direct evidence of anticancer stem cells efficacy of this approach *in vivo* in tumors.

Genetic engineering of oncolytic viruses complements chemo- and molecular-targeted therapies

Several novel combination treatments have been tested in combination with oncolytic viruses. Genetic engineering of the viruses allows functional complementation to chemotherapeutic agents and molecular-targeted therapeutics. Aghi *et al.*³⁴ showed that temozolomide-induced DNA repair pathways in glioma cells complemented replication of γ 34.5-deleted oncolytic HSV replication and resulted in enhanced efficacy both *in vitro* and *in vivo*. Stanford *et al.*³⁵ and Lun *et al.*³⁶ showed that treatment of rapamycin, a mammalian target of rapamycin inhibitor which resulted in increased Akt/protein kinase B activation, one was able to enhance myxoma virus replication in tumor, but not normal, cells. We showed that by deleting Us3, oncolytic HSV can synergize with Phosphoinositide-3 kinase (PI3K)/Akt inhibitors *in vitro* and have enhanced efficacy without increasing toxicity *in vivo.*³⁷

Histone deacetylase (HDAC) inhibitors are currently being investigated in combination with various oncolvtic virus species. Trichostatin A (TSA) has been shown to upregulate cellular coxsackievirus-adenovirus receptor (CAR) expression and hence, infectibility of tumor cells to adenoviruses. We showed that in addition to CAR upregulation, TSA possesses antitumoral and antiangiogenic activities, and shows synergistic tumor-killing effect with oncolytic HSVs in vitro and enhanced efficacy in vivo (Liu et al.38). In contrast, valproic acid, another HDAC inhibitor used for epilepsy disorders, showed antagonistic effect with oncolytic adenoviruses in vitro, presumably due to enhanced apoptosis that limits viral replication.³⁹ It will be interesting to see what the effect other types of HDAC inhibitors will have on different oncolytic virus species.

Genetic engineering of oncolytic viruses targets cancer signaling pathways

Through genetic engineering, viruses can be designed to target cancer cells through certain activated signaling pathways. Apart from the Akt pathway targeted by myxoma virus described above, HSV y34.5-deleted mutants showed enhanced replication in cells with activated mitogen-activated protein (MAP) kinase or extracellular signal-regulated (ERK) kinase, which in turn inhibited protein kinase R activity, thus circumventing the negative impact of the IFN signaling pathway.^{40,41} Similarly, viruses such as VSV showed preferential replication in cells with an activated Ras-ERK pathway and defective IFN pathways.42 A vaccinia virus mutant with a deletion in B18R, whose gene product neutralizes type I IFNs, showed IFN-dependent cancer selectivity and efficacy.³¹ It has also been shown that adenovirusinduced ERK activation is critical to viral replication.⁴³ Oncolytic viruses can also be 'programmed' to replicate in cells through certain cellular signaling activities, such as β -catenin,⁴⁴ to carry therapeutic transgene that targets tumorigenic pathways,²⁰ or retargeted to cellular receptors that are essential for signaling (for example, epidermal growth factor receptor (EGFR)).45

Novel oncolytic virus species are being explored

As most oncolytic viruses have produced less than optimal efficacy in clinical trials as single agents, there is great interest in exploring novel viral species. These studies assess oncolytic activity and/or investigate tumor selectivity. For example, the porcine Seneca Valley virus has recently been discovered to possess antitumoral activity against certain cancers of neuroendocrine origin.⁴⁶ It has been speculated that the tumor selectivity is based on differential receptor binding in cancer and normal cells, but more work needs to be done to verify this and to study the impact of the immune system has on the virus. Myxoma virus, a rabbit virus, has also been assessed as an oncolytic agent. New studies reveal that the tumor selectivity of myxoma virus is based on overexpression of Akt in human cancer cells, which facilitates replication and oncolysis.⁴⁷ While these are important findings, the safety profiles of these viruses need to be cautiously examined, especially when the natural host of the virus is not human.

Intratumoral administration of UV-inactivated, replication-deficient Sendai virus induced a robust antitumoral immune response (including cytotoxic T lymphocyte (CTL) induction, dendritic cell maturation and antagonism of Tregs) and resulted in significant efficacy in CT26 syngeneic murine colorectal cancer model.⁴⁸ It will be interesting to study how much of this vaccine effect contributes to antitumoral efficacy in oncolytic Sendai virus studies. A 'vaccine' effect was also seen with parvovirus H-1 in a rat lung tumor metastases model.⁴⁹

A large number of clinical trials have been carried out Virotherapy has several features that are distinct from other therapeutics. Its multiple novel MOAs include replication-mediated oncolysis, antitumoral immunity induction, antiangiogenesis, apoptosis and autophage induction. There is no cross resistance with other therapeutics, and synergistic interaction is seen with other treatment modalities. Safety in human has been demonstrated in more than 800 patients. In addition, current biotechnology allows us to rapidly address issues encountered in clinics at the bench. There are several reports on virotherapy clinical trials over the past 2 vears. Readers are referred to other articles for a more comprehensive review.¹ A list of oncolytic virus agents that have completed, or are currently in late phase trials, are listed in Table 2. In a recent study examining the OV 001 (HUJ) strain of Newcastle disease virus (NDV), OV 001 was administered intravenously to 11 patients with glioblastoma with no dose-limiting toxicity.⁵⁰ Following biweekly maintenance therapy, one complete remission with a duration of 3 months was described. Virus was recovered from blood, urine and saliva samples. Infectious NDV was recovered from a tumor biopsy. In a separate study testing intravenous delivery of a different NDV strain PV701, using 'two-step' desensitization (dosing with significantly smaller doses prior to full dose) has proven to significantly reduce acute adverse events.⁵¹

The HSV-1 mutant NV1020 (R7020) virus was originally developed as a vaccine. The virus has a deletion in one of the two copies of the γ -34.5 gene. A phase I trial of NV1020 administered by hepatic arterial infusion (HAI) was performed in HSV-seropositive patients with colorectal liver metastases.52 No significant toxicity was noted in patients receiving doses up to 1×10^8 pfu per infusion. No replication data were reported. An ongoing phase I/II trial is evaluating repeat HAI of NV1020 followed by second-line chemotherapy in seropositive patients with colorectal cancer. The oncolytic HSV vector OncoVEX^{GM-CSF} has been generated with deletions in γ -34.5 (to reduce pathogenicity) and ICP47 (to restore MHC I presentation). In addition, OncoVEX^{GM-CSF} has a granulocyte monocyte colony-stimulating factor (GM-CSF) transgene insertion. An early passage clinical isolate was used to generate OncoVEX^{GM-CSF} because it had enhanced potency relative to available laboratory strains. A phase I trial of intratumoral injection of OncoVEX^{GM-CSF} into cutaneous metastases from solid tumors and melanomas was carried out.53 Treatment did not result in significant systemic toxicities; injection-site inflammation was dose limiting. Viral genomes were detected shedding from the skin over ulcerated tumors. Injected tumor histology showed inflammation and necrosis. No distant efficacy was reported. Phase II trials of intratumoral injection using OncoVEX^{GM-CSF} are underway in patients with melanoma and other tumor types. A Phase III trial of HSV-1 mutant 1716 in brain tumor has also been announced.

In addition, an oncolytic vaccinia virus with deletion in thymidine kinase (TK) and expressing GM-CSF, JX-594, has been tested in patients with liver tumors. Unlike other virus species described above, systemic exposure and evidence of replication *in vivo* has been

 Table 2
 Oncolytic viral agents in completed or ongoing late phase trials

Product	Species	Genetic modification	Target tumor type	Phase
Reolysin	Reovirus	None	Bone/soft-tissue sarcoma	II
NDÝ (MTH-68H)	Newcastle disease virus	None	Metastatic solid tumors	II
JX-594	Vaccinia	Thymidine kinase deletion;	1. Hepatocellular	II
		GM-CSF expression	2. Melanoma	II
H101	Adenovirus	E1B-55 K, E3 deletion	Head and neck (+ chemotherapy)	III
Ad5-yCD/ mutTKSR39rep-ADP	Adenovirus	E1B-55 K deletion; CD/TK fusion gene expression; ADP overexpression	Prostate (+ radiotherapy)	Π
OncoVex ^{GM-CSF}	HSV 1	γ34.5 and ICP47 deletion; GM-CSF expression	Melanoma	II
1716	HSV 1	γ 34.5 deletion	Brain	III

Abbreviations: ADP, adenovirus death protein; GM-CSF, granulocyte monocyte colony-stimulating factor; HSV, herpes simplex virus; TK, thymidine kinase; VSV, vesicular stomatitis virus.

demonstrated in this study. Tumor response was observed in the majority of treated tumors, and distant tumor responses were demonstrated in several patients with target tumor responses. Replication and biological activity *in vivo* have also been shown.⁵⁴ With the advance in our knowledge of cancer and oncolytic virus biology, we expect that carefully designed clinical studies will not only show proof-of-concept for this novel treatment, but also result in evidence of clinical benefit in patients.

Prospects

Logical design of the next generation of oncolytic viruses may take this strategy to the next level

Design of the next generation of oncolytic viruses should be based on current knowledge of virology, immunology and cancer biology. More importantly, findings from clinical trials should be incorporated/addressed. Two recent publications illustrate this concept.31,55 The first article describes rational species and strain selection and genetic engineering based on updated knowledge. As described before, poxvirus was selected for this study based on human data showing that systemic delivery of poxvirus is safe and can induce significant tumor responses.⁵⁶ A highly potent vaccinia virus strain that also trafficked efficiently to human tumors after intravenous administration was first identified. This strain was then engineered to target cancer cells with activated transcription factor E2F and the EGFR pathway, and further engineered to express human GM-CSF for induction of tumor-specific CTL. The new vaccinia construct, JX-963, demonstrated significant cancer selectivity in human tumor cell lines, tumor-bearing rabbits and primary human surgical samples. Intravenous administration led to systemic efficacy against both primary carcinomas and widespread organ-based metastases in immunocompetent animals.55 The second study describes the use of a vaccinia virus background that selectively targets IFN pathway resistance in tumor cells. Further engineering with TK deletion and IFN-β insertion results in a multimechanistic oncolytic vaccinia virus.31

Ex vivo studies may predict responses

Genetic markers are being developed for chemotherapy and molecular-targeted therapeutics to predict responses and/or idiosyncratic reactions, and the results have been implemented into practice. Given the complexity of MOA of virotherapeutics, there is therefore a long way to go before such markers/predictors can be developed for virotherapy. However, testing patients' samples ex vivo prior to treatment may provide important information and complement our current studies. In addition to the difference between results obtained from immortalized cell lines and primary cancer cells, explant samples contain extracellular matrix and a three-dimensional structure that more closely mimics clinical situation. Importantly, when adjacent normal tissues are included, a therapeutic index can be obtained, which is critical for local administration protocol. Several recent publications described the use of this approach.27,55,57,58 The results of oncolytic viruses on ex vivo tissues and its clinical outcome correlation have yet to be established, but researchers are encouraged to include explants

studies whenever possible to maximize clinical relevance. Future clinical developments might include pre-/post-treatment *ex vivo* assessment of infectivity, cytopathic effects and viral replication.

The demand for target validation is increasing

The demand for target validation in small moleculebased kinase inhibitors is increasing. For virotherapy to be successful, it has to pass similar hurdles. The most important biological endpoint that needs to be demonstrated with all species of oncolytic virus is tumor-selective virus replication, therapeutic transgene expression and biological function (if applicable). This has been effectively achieved with several virus species, but still required for others. Depending on the strategy demonstration of targeting cancer-specific features pathways may also be necessary. For instance, if viruses are designed to target the Ras/Raf/MAPK pathway, a reduction in pathway activity should be proven. High TK activity in cancer cells are needed for TK-deleted viruses, while viruses replicating exclusively in IFNresistant cells (cancer cells) need to demonstrate local IFN induction in vivo. These are difficult tasks but warrant further exploration. The effect of oncolytic viruses on tumor microenvironment, as shown in various preclinical studies, will need to be validated in clinical trials. For instance, future clinical studies should include tumor vascularity assessment to see if virotherapy reduces tumor vasculature.

New imaging endpoints in clinical trials?

Tumor size measurement has been a gold standard for defining responses in clinical practice. Tumor progression is defined as increased in tumor size above certain degrees. However, recent studies on molecular therapeutics have shown that some agents induce tumor necrosis without affecting the sizes of the tumors. To address this issue, new criteria (for example, Choi criteria⁵⁹) have been incorporated in tumor response assessment by imaging. Since most of the virotherapy clinical trials were done by local or locoregional administration, it is likely that the effect of the viruses is localized within the tumors. Thus, we need to consider whether adopting these new criteria is feasible. Correlation between tumor size, tumor density and survival will be necessary in late phase trials.

Abbreviations

CAR, coxsackievirus-adenovirus receptor; CIK, cytokineinduced killer; CTL, cytotoxic T lymphocyte; EGFR, epidermal growth factor receptor; HDAC, histone deacetylase; HSV, herpes simplex virus; IFN, interferon; MMP, matrix metalloproteinase; MOA, mechanism-ofaction; PBMC, peripheral blood mononuclear cells; TK, thymidine kinase; Treg, regulatory T cells; TSA, trichostatin A; VSV, vesicular stomatitis virus

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