

IncRNA UCA1-Mediated Cdc42 Signaling Promotes Oncolytic Vaccinia Virus Cell-to-Cell Spread in Ovarian Cancer

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Oncolytic vaccinia virus (OVV) has demonstrated appropriate safety profiles for clinical development. Although designed to kill cancer cells efficiently, OVV sensitivity varies in individual cancers, and predictive biomarkers of therapeutic responses have not been identified. Here we found that OVV was much more efficient in KFTX paclitaxel-resistant ovarian cancer cells compared to that in KFlow paclitaxel-sensitive cells. Microarray analysis identified long non-coding RNA urothelial carcinoma-associated 1 (UCA1) upregulation, which contributed to both enhanced paclitaxel resistance and OVV spread. In addition, UCA1 expression correlated with efficient OVV spread in other ovarian cell lines and primary cancer cell cultures. When host pathways underlying OVV spread were analyzed, differences were detected in the activation of the Rho GTPase Cdc42, suggesting that filopodia formation enhances OVV cell-to-cell spread and tumor migration. Moreover, we established a clinically relevant mouse model of peritoneal metastasis using KFTX or KFlow cells. Paclitaxel exerted anti-tumor effects on KFlow, but not KFTX, tumors. In mice bearing KFTX cells after paclitaxel failure, OVV treatment induced the regression of residual tumors and improved survival. Our findings demonstrated that UCA1 promotes OVV cell-to-cell spread in ovarian cancer, resulting in enhanced therapeutic outcome.

INTRODUCTION

Oncolytic viruses have emerged as novel candidates for cancer therapy owing to their inherent potential to selectively replicate and destroy tumor cells.¹ With the recent approval of talimogene laherparepvec for advanced melanoma, oncolytic virotherapy has earned its place in clinical settings.² Furthermore, studies have demonstrated that oncolytic vaccinia virus (OVV) is a promising alternative to existing therapies for the treatment of advanced cancer.^{3–5}

In a phase 1 clinical trial of intravenous oncolytic pox virus, a Western Reserve OVV strain, two of 11 patients demonstrated prolonged virus replication in tumor tissues.⁶ Moreover, one

patient showed a mixed response with the resolution of some liver metastases. In a phase 1b clinical trial of intravenous Pexa-Vec, a Wyeth OVV strain, 10 of 15 patients developed radiologically stable disease.⁷ However, there were no complete or partial responses in refractory or heavily pre-treated patients. Therefore, a predictive biomarker to determine patients who are more likely to benefit from treatment or to anticipate therapeutic effects or timing is vital. In addition, studies to elucidate variations in the susceptibility of human malignant cancers to OVV are urgently needed. In this basic and translational research, we address the value of expression of the long non-coding RNA (lncRNA) urothelial carcinoma-associated 1 (UCA1) as a potential predictive biomarker for both OVV and paclitaxel (PTX) chemotherapy in patients with ovarian cancer.

UCA1 was first identified as a sensitive and specific marker for human bladder carcinoma and cancer-upregulated drug-resistant (CUDR) expression.^{8,9} Several groups have reported that UCA1 is highly expressed in ovarian cancer, colorectal cancer, and pancreatic cancer, among others, suggesting that it has the potential to be a biomarker for the diagnoses of these cancers.^{10,11} UCA1 has been reported to function as a decoy for endogenous RNA, specifically by acting as a sponge for microRNAs (miRNAs), resulting in their downregulation, which has been implicated in enhanced drug resistance.^{12,13} Moreover, UCA1 overexpression enhances proliferation and metastasis in several cancer cell lines through ERK, PI3K, Wnt, or Akt signaling.^{14–16} However, UCA1 has not previously been reported to be associated with oncolytic virotherapy outcome.

Here we show the possibility of applying UCA1 expression as a predictive biomarker for both OVV and PTX chemotherapy *in vitro* and

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in vivo for ovarian cancer. Further functional studies revealed the detailed mechanisms underlying the regulatory effect of UCA1 on OVV spread. Importantly, these results could enable the identification of patients more likely to respond to OVV.

RESULTS

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UCA1 Contributes to Enhanced PTX Resistance and Vaccinia Virus-Mediated Oncolysis

PTX-sensitive KFlow cells were isolated from KFTX cells cultured without the selection pressure of PTX. Further, KFlow cells regained resistance by incubating them with PTX, resulting in PTX-resistant KFTXlow cells (Figure 1A). These cell lines were infected with OVV-LG (LucGFP) at an MOI of 0.01. Interestingly, during KFTX infection, OVV-LG induced massive cytopathic effects (CPEs) after robust viral EGFP expression (Figures 1B and 1C). In contrast, weak CPEs and EGFP expression were induced in KFlow cells, whereas intermediate CPEs and EGFP expression were induced in KFTXlow cells. These results suggest that genes that are modulated

Figure 1. Identification of Candidate Genes Involved in Paclitaxel Resistance and Efficacy of Oncolytic Vaccinia Virus Spread

(A) Schema of KFlow, KFTXlow, and KFTX cells. (B) EGFP images of KFlow, KFTXlow, and KFTX cells after infection with OVV-LG (MOI = 0.01) for 72 h. Scale bar, 1,000 μ m. (C) The intensity and area of viral EGFP brightness was measured using a Keyence BZ-X700 fluorescence microscope (n = 3). (D) RNA from KFlow, KFTXlow, and KFTX cells was collected and analyzed by an Agilent Sure Print G3 Human Gene Expression 8 × 60K v.2 Microarray (Takara Bio). The heatmap was constructed using multi-Experimental Viewer (MEV) v.4.9 software. Data with error bars represent mean \pm SEM.

according to PTX resistance are potential host factors that are involved in the oncolytic effects of OVV-LG.

Cellular gene expression profiles among these cell lines were compared by microarray analysis (Figure 1D). Results of 100 significantly dysregulated genes are shown in Figure 1D. Some candidate gene expression patterns among KFTX, KFTXlow, and KFlow cells were correlated with OVV growth efficacy in these cell lines (Table1). Among candidate genes, UCA1 expression was most dysregulated in KFTX (129.2-fold change) and KFTXlow (51.5-fold change) cells, as compared to that in KFlow cells. Moreover, UCA1 expression patterns among KFTX, KFTXlow, and KFlow cells were in complete accordance with OVV growth efficacy, which differed by more than 10-fold between KFlow and KFTXlow cells and 3-fold between KFTXlow and KFTX cells (Figure 1C).

For these reasons, we hypothesized that UCA1 might play an important role in vaccinia virus-mediated oncolysis.

To analyze the effects of UCA1 on PTX resistance and OVV-LG oncolysis, KFlow and KFTXlow cells were transfected with pcDNA3.1/UCA1 (UCA1) or pcDNA3.1 (empty) plasmids (Figure 2A). PTX susceptibility was determined by cell viability assays. Based on viability, the resistance of UCA1-KFlow cells was significantly increased, in a dose-dependent manner, as compared to that of KFTX and KFTXlow cells (Figure 2B). Furthermore, the upregulation of UCA1 induced CPEs and EGFP expression in OVV-LG-treated KFlow and KFTXlow cells, which were equivalent to those observed in KFTX cells at MOIs of 0.01 and 0.001 (Figures 2C and 2D). Moreover, viral progeny was quantified by plaque assays. Consistent with the quantification of EGFP levels, UCA1 upregulation in KFlow and KFTXlow cells enhanced viral yields in KFTXlow cells to levels comparable to those observed in KFTX cells (Figure 2E). However, overexpression of candidate genes except

	Gene Symbol	GenBank Accession	Scale Signal KFlow	Scale Signal KFTXlow	Scale Signal KFTX
Upregulated genes	UCA1	NR015379	233	11,996	30,100
	EEF1A2	NM001958	965	31,458	40,666
	S100P	NM005980	2,792	78,119	98,929
	STEAP1	NM012449	103	3,983	3,484
	ABCB1	NM000927	770	40,125	46,599
	SMARCA1	NM003069	100	2,194	2,294
	ITGBL1	NM004791	97	1,669	5,022
	IGF2	NM000612	34	306	797
	LOXL2	NM002318	70	723	704
	PADI1	NM013358	201	2,610	5,596
Downregulated genes	BATF3	NM018664	2,360	88	1,025
	AKR1B15	NM001080538	54,250	16,433	2,456
	NEFL	NM006158	15,321	3,143	561
	FSTL1	NM007085	6,418	60	60
	IRX4	NM016358	32,083	418	346
	CES1	NM001266	14,067	437	365
	GJB6	NM006783	3,192	242	53
	NELL2	NM006159	3,104	154	149
	SMO	NM005631	5,148	256	81
	PHEX	NM0004444	1,041	76	54

UCA1 did not change OVV growth efficacy (data not shown). In addition, the depletion of UCA1 in KFTX cells decreased viral EGFP expression compared to that in control cells (Figure S1). These results suggested that UCA1 regulates viral replication and spread in addition to PTX resistance.

UCA1-Mediated Enhanced Vaccinia Virus-Mediated Oncolysis Is Common in Ovarian Carcinomas

Based on the aforementioned observations, we investigated whether UCA1 expression is involved in the oncolytic effects of OVV-LG in other ovarian cancer cell lines. For this, a comparison of UCA1 expression and oncolytic effects was carried out using SHIN-3, OVCAR3, RMG-1, SKOV3, and ES-2 human ovarian cancer cell lines. We demonstrated that UCA1 expression was obviously inconsistent and that it was overexpressed in SKOV3 and RMG-1 cells compared to expression in SHIN-3, OVCAR3, and ES-2 cells (Figure 3A). These UCA1-high- and UCA1-low-expression cell lines were then infected with OVV-LG at an MOI of 0.001. In SKOV3 and RMG-1 cells, massive CPEs with robust EGFP expression were induced, as compared to those in SHIN-3, OVCAR3, and ES-2 cells (Figures 3B and 3C). In addition, the quantification of EGFP expression and viral progeny yielded consistent results (Figure 3D). Further investigation revealed that UCA1 expression correlated with the oncolytic effects of OVV in primary ovarian cancer cells (Figures 3E-3H). These results suggested that UCA1 regulates the oncolytic effects of OVV and might be considered a promising biomarker to predict oncolytic activity in ovarian cancer.

UCA1 Enhances Vaccinia Virus Oncolysis by Modulating Viral Cell-to-Cell Spread, but Not Binding, Entry, and Replication

To highlight the mechanism associated with enhanced OVV-LGmediated oncolysis, we identified which stage of OVV replication is regulated by UCA1. First, we determined if UCA1 affects OVV-LG binding and entry. It was previously reported that inhibiting OVV entry results in decreased replication.¹⁷ Based on binding assays, the production of viral progeny by cell-attached viruses was not changed with high UCA1 expression (Figure 4A). Regarding entry assays, viral firefly luciferase activity (Fluc) was not affected by UCA1 expression (Figure 4B).

Next, we analyzed the production of early (M1L), intermediate (G8R), and late (A5L) viral transcripts by real-time PCR. In cells infected with OVV-LG, viral genes representing all three phases were not induced by high UCA1 expression (Figures 4C-4E). It was reported that the presence of fewer extracellular enveloped viruses (EEVs), which are required for spread between neighboring cells and distant cells, significantly affects virus production.^{18,19} In the early infectious phase, EEV and cell-associated virus (CAV) titers were not stimulated by high UCA1 expression (Figure 4F). Lastly, because it was shown that the efficacy of intracellular transport and cell-to-cell spread is important for the oncolytic effects of OVV,^{20,21} selective inhibitors of intracellular transport (nocodazole and colchicine) and cell-tocell-spread (cytochalasin D) were used, and plaque assays were performed without the induction of cytotoxicity (Figure S2). In UCA1-KFlow cells, inhibitors of intracellular transport had little



effect on plaque sizes; however, an inhibitor of cell-to-cell spread completely inhibited plaque formation (Figure 4G). These results indicate that UCA1 specifically affects OVV-LG cell-to-cell spread.

UCA1 Enhances the Cell-to-Cell Spread of OVV via Cdc42 Activation

We hypothesized that UCA1 regulates actin polymerization, thereby promoting cell-to-cell spread based on two observations. First, gene ontology analysis based on a microarray comparing KFlow and UCA1-KFlow cells indicated alterations in cell adhesion pathways; however, functional assays were then required (Figure 5A). Second, high UCA1 expression mediated dramatic increases in the numbers and lengths of filopodia, which are actin-rich structures involved in

Figure 2. Both Paclitaxel Resistance and Oncolytic Vaccinia Virus Spread Are Recovered by the Overexpression of UCA1

(A) Enhanced UCA1 expression was confirmed by qRT-PCR (n = 3). (B) KF cell lines were treated with PTX (0, 2, 5, 20, 50, 100, 200, and 500 nM). Cell viability was determined 48 h after PTX treatment (n = 3). (C) EGFP (left) and bright-field (right) images of KF cell lines are shown after infection with OVV-LG (MOI = 0.01 and 0.001) for 72 h. Scale bar, 1,000 μ m. (D) The intensity and area of viral EGFP brightness were measured using a Keyence BZ-X700 fluorescence microscope (n = 3). (E) Plaqueforming units (PFUs) of OVV were determined based on titration using RK-13 cells (n = 3). Data with error bars represent mean ± SEM.

numerous cellular processes like cell migration and cell-cell interactions (Figure S3). In support of this hypothesis, wound-healing assays demonstrated that UCA1 enhances cell migration and adhesion upon serum starvation (Figure 5B).

Since previous studies suggested that cell adhesion and migration are regulated by Rho GTPases, which have an important role in the assembly and disassembly of the actin cytoskeleton,^{22,23} we examined whether the suppression of Rho, Rac1, and Cdc42, members of the Rho family of small GTPases, could inhibit plaque growth. The addition of ML141, a Cdc42 inhibitor, resulted in a significant decrease in plaque size, as compared to that with other inhibitors against Rac1 and Rho (Figure 5C; Figure S2). We also examined the effects of Rho, Rac1, and Cdc42 suppression on OVV-LG plaque size in UCA1-high- and UCA1-low-expression cell lines, namely SKOV3 and ES-2 cells, respectively (Figure 5D). As expected, the addition of a Cdc42 inhibitor resulted in inefficient cellto-cell spread in SKOV3, but not in ES-2, cells. Moreover, Cdc42 expression was not changed

between KFlow (scale signal: 41,996) and UCA1 KFlow cells (scale signal: 37,793) based on microarray analysis. Nevertheless, when the expression of active Cdc42 was tested, we observed that UCA1 expression increased Cdc42 activation (Figure 5E); we also noted a correlation between UCA1 and activated Cdc42 expression in primary ovarian cancer cells (Figure 5F). These results suggest that UCA1-mediated Cdc42 regulation enhances OVV-LG cell-to-cell spread through filopodia formation.

OVV Exerts Increased Oncolytic Effects on PTX-Resistant KFTX Cells in a Mouse Model of Peritoneal Metastasis

We previously found that transgene insertion into the hemagglutinin (HA) gene of OVV-LG does not affect native vaccinia virus



replication.²⁴ Further, OVV-LG is suitable to elucidate correlations between UCA1 expression and OVV spread. However, OVV-LG maintains viral pathogenicity.²⁴ We thus utilized OVV-VGF Δ / O1L Δ for therapeutic experiments. Since two viral proteins, specifically VGF and O1, contribute to viral spread and replication through activation of the epidermal growth factor receptor (EGFR)-dependent mitogen-activated protein kinase (MAPK)/ERK pathway, deletion of both genes inhibits pathogenic viral replication in normal cells, whereas therapeutic replication is retained in tumor cells with constitutive ERK1/2 activation.²⁵

To assess the oncolytic effects of OVV-VGF Δ /O1L Δ , we examined its cytotoxicity using KFlow, UCA1-KFlow, Empty-KFlow, and KFTX

Figure 3. Correlation between UCA1 and Oncolytic Vaccinia Virus Spread as Observed in Other Ovarian Cancer Cell Lines

(A and E) UCA1 expression was detected by qRT-PCR in SHIN-3, OVCAR3, RMG-1, SKOV3, and ES-2 cells (A) and primary ovarian cancer cells (E) (n = 3). (B and F) EGFP (left) and bright-field (right) images of SHIN-3, OVCAR3, RMG-1, SKOV3, and ES-2 cells (B) and primary ovarian cancer cells (F) after infection with OVV-LG (MOI = 0.001 or 0.01) for 48 h. Scale bar, 1,000 μ m. (C and G) Intensity and area of viral EGFP brightness in SHIN-3, OVCAR3, RMG-1, SKOV3, and ES-2 cells (C) and primary ovarian cancer cells (G) were measured using a Keyence BZ-X700 fluorescence microscope (n = 3). (D and H) Plaque-forming units (PFUs) of OVV in SHIN-3, OVCAR3, RMG-1, SKOV3, and ES-2 cells (D) and primary ovarian cancer cells (H) were determined by titration using RK-13 cells (n = 3). Data with error bars represent mean \pm SEM.

cells (Figure 6A). OVV-VGFΔ/O1LΔ mediated enhanced oncolysis in PTX-resistant KFTX and UCA1-KFlow cells compared to that in PTXsensitive KFlow and Empty-KFlow cells. Therefore, we applied a peritoneal dissemination model using PTX-resistant KFTX or PTX-sensitive KFlow cells expressing Renilla luciferase. Cells were injected into BALB/cAJcl-nu/nu mice, and, after confirming tumor growth based on Renilla luciferase activity, mice were intraperitoneally administered OVV-VGFA/O1LA or control PBS (Figure 6B). On day 1 after viral injection, mice bearing KFTX cells showed tumor-specific high virus-associated signals, whereas mice bearing KFlow cells exhibited little viral replication (Figures 6C and 6D). On day 10 after viral injection, viral signals in mice bearing KFTX cells disappeared, which was accompanied by a reduction in tumor signals. The treatment of mice harboring KFTX cells with OVV-VGF Δ /O1L Δ resulted in the significant inhibition of tumor growth, by more than two log orders, compared to that in con-

trol PBS-treated animals (Figure 6D). In terms of animal survival, treating KFTX-harboring mice with OVV-VGF Δ /O1L Δ led to a significant improvement, but the same treatment in KFlow-bearing mice had no effect (Figure 6E). These data demonstrated that OVV-VGF Δ /O1L Δ is an effective therapy for PTX-resistant ovarian cancer formed from KFTX cells.

UCA1 Expression Potentially Predicts the Therapeutic Strategy for Ovarian Cancer Based on Conventional PTX or Novel Oncolytic Virotherapy

For the development of a clinically relevant mouse tumor model, nude mice bearing tumors derived from intraperitoneally injected KFTX or KFlow cells were injected with PTX. After 3 weeks of



Figure 4. UCA1 Expression Specifically Targets Oncolytic Vaccinia Virus Cell-to-Cell Spread and Increases the Oncolytic Effect of OVV-LG (A and B) KFlow, UCA1-KFlow, and Empty-KFlow cells were infected with OVV-LG (MOI = 10) for 30 min. After ample washes, the number of plaque-forming units (PFUs) of OVV was determined by titration using RK-13 cells (A), and viral luciferase activity was determined (B) (n = 3). (C–E) Gene expression in KFlow, UCA1-KFlow, and Empty-KFlow cells after either mock infection or infection with OVV-LG (MOI = 5). RNA was extracted at 1, 2, 4, 6, 8, 10, and 12 h postinfection (hpi). gRT-PCR was performed with

administration, PTX efficiently exerted anti-tumor effects on KFlow tumors, whereas KFTX tumors showed only marginal regression for 1 week after treatment and then gradually regrew (Figures 7A and 7B). Importantly, PTX-treated mice bearing KFTX tumors were injected with OVV-VGF Δ /O1L Δ or control PBS, resulting in the dramatic regression of residual tumors. On day 9 after OVV-VGF Δ /O1L Δ injection, a reduction in tumor signals was observed in mice injected with virus (Figures 7C and 7D). In terms of animal survival, treating KFTX-bearing mice with OVV-VGFA/ O1L Δ led to a significant improvement in survival, compared to that in control PBS-treated animals. After all treatments, the survival of mice bearing KFTX tumors and treated with virus was prolonged compared to that in animals treated with PBS (Figure 7E). Further, UCA1 expression was quantified in excised tumor tissue before and after PTX administration. Consistent with in vitro data, high UCA1 expression was maintained in KFTX tumors (Figure 7F). Taken together, these results indicate that tumors expressing high levels of UCA1 should be treated with OVV rather than PTX, whereas those with low UCA1 expression should receive the opposite therapeutic regimen.

DISCUSSION

Various viruses such as adenovirus, herpes simplex virus, Newcastle disease virus, and vaccinia virus have been engineered and clinically investigated as oncolytic viruses.^{2,5,26,27} OVV has many advantages for oncolytic virotherapy.²⁸ First, the range of tropism for OVV in mammalian cells is broad, and, thus, it could be applied to many types of cancers. Second, the whole life cycle of OVV occurs in the cytoplasm, indicating that it is not incorporated into the host genome, which thereby remains intact. Third, OVV spreads via the blood-stream; it is thus expected that OVV will exert oncolytic effects not only in primary tumors but also in metastatic cancer via remote infection. Finally, the replication cycle of OVV is fast, which should facilitate rapid viral replication and spread, maximizing therapeutic effects. Accordingly, OVV is a promising cancer therapy, and the efficiency of viral replication and spread will ultimately play a crucial role in therapeutic outcome.²⁹

Promising clinical data based on various strains of OVV have indicated both therapeutic efficacy and safety.^{6,7,30} However, targeting appropriate cancer types and grades using OVV remains difficult, and combinations with other cancer therapies is likely to be primarily attempted. In this study, xenograft mouse models of PTX-resistant and PTX-sensitive ovarian cancer were found to be efficiently treated with PTX and OVV, respectively. Although high UCA1 expression was maintained in resected tumors after PTX treatment, UCA1 expression was not detected in normal tissues, such as ovary and urothelial tissues (data not shown). Therefore, this therapeutic strategy could accurately reflect clinical situations with the use of UCA1 as a correlative marker (Figure 7).

First, the expression patterns of some genes, including UCA1 and ITGBL1, were found to be correlated with OVV growth in KFTX, KFTXlow, and KFlow cells (Table 1). However, the overexpression of candidate genes other than UCA1 did not change OVV growth efficacy. For this reason, we selected UCA1. Importantly, multiple studies have shown that it plays an important biological role in tumor growth, metastasis, and malignancy and that it might act as a potential biomarker and therapeutic target for various cancers.^{9,31,32} In clinical specimens, high UCA1 expression was observed in cancer tissue compared to normal tissue.³¹ Furthermore, the function of UCA1 as an oncogene has gradually become elucidated. For example, elevated expression induces cisplatin resistance via SRPK1 activation in ovarian cancer.³³ Further, high UCA1 expression induces PTX resistance through the miR-129/ABCB1 pathway in ovarian cancer.³⁴ Considering that UCA1 regulates chemotherapy sensitivity through multiple pathways, it might modulate PTX resistance through a mechanism other than that required for viral spread.

UCA1 is highly expressed in several cancers and promotes metastasis and progression by inducing ERK signaling.^{15,35} Consistent with previous reports, ERK phosphorylation was consistently high and UCA1 was found to further activate ERK in serum-stimulated KFlow and UCA1-KFlow cells (Figure S4A). Furthermore, ERK activation is known to play an important role in the early stages of OVV infection, and it is also crucial for the generation of virus progeny.³⁶ Thus, we examined whether ERK activation plays a key role in promoting OVV spread. The ERK pathway was found to be activated by epidermal growth factor (EGF) stimulation, and the activation of ERK was measured (Figure S4A). Further, KFlow and UCA1-KFlow cells were infected with OVV and treated with EGF stimulation. OVV-mediated EGFP expression in KFlow cells was not enhanced by ERK activation following EGF stimulation (Figure S4B). In addition, we found that OVV replication and progeny were not promoted by high UCA1 expression (Figures 4C-4F). These results suggested that further activation of ERK by EGF stimulation had no effect on OVV replication and progeny in serum-stimulated tumor cells but that another pathway regulated by UCA1 is crucial for OVV spread.

OVV cell-to-cell spread is an important process and is supported by some viral proteins through actin-based motility. During the process of fusion between the extracellular membrane of intracellular enveloped virus and the cell membrane, B5 is exposed at the vaccinia virus surface of cell-associated enveloped virus (CEV).³⁷ The detailed function of B5 is unknown with respect to actin-based motility. However, deletion of B5 dramatically changes vaccinia

primers specific for M1L (C), G8R (D), A5L (E), and GAPDH (C–E, as an internal control) (n = 3). (F) One-step growth curve. KFlow, UCA1-KFlow, and Empty-KFlow cells were infected with OVV-LG (MOI = 5). Cells (CAV) and supernatant (extracellular enveloped virus [EEV]) were collected, and the number of OVV PFUs was determined by titration using RK-13 cells (n = 3). (G) KFlow, UCA1-KFlow, and Empty-KFlow cells were treated with nocodazole (25 nM), colchicine (2.5 nM), and cytochalasin D (KFlow and Empty-KFlow, 25 nM; UCA1-KFlow, 50 nM) and infected with OV-LG (MOI = 0.001). Representative plaques formed under semisolid medium were photographed and calculated with the analysis application measurement module (Keyence) (n = 10). Data with error bars represent mean ± SEM.



Figure 5. Cdc42 Activation Is Critical for the Cell-to-Cell Spread of Oncolytic Vaccinia Virus-LG Induced by UCA1 Expression

(A) Results of gene ontology analysis after microarray analysis. Gene lists correspond to 2-fold changes in KFlow and UCA1-KFlow cells. (B) A pipette tip was used to introduce a scratch in a monolayer of KFlow, UCA1-KFlow, and Empty-KFlow cells in the absence of serum. The percentage of wound closure was estimated using the analysis application measurement module (Keyence) (n = 6). (C and D) KFlow, UCA1-KFlow and Empty-KFlow (C) and SKOV3 and ES-2 cells (D) were treated with Rhosin



morphology. A36 localizes to the inner cell membrane after CEV is exposed on the cell surface. Phosphorylated A36 has a binding site for Nck, which mediates the recruitment of neural Wiskott-Aldrich-syndrome protein (N-WASP), resulting in Arp2/3 complex binding to existing actin filaments.³⁸ The activated Arp2/3 complex helps to propel the virus into neighboring cells. However, based on our results, viral gene expression was not changed by UCA1 expres-

Figure 6. Oncolytic Vaccinia Virus-VGF∆/O1L∆ Efficiently Exerts Oncolytic Effects against Paclitaxel-Resistant KFTX Cells *In Vitro* and *In Vivo*

(A) KFlow, UCA1-KFlow, Empty-KFlow, and KFTX cells were infected with OVV-VGF Δ /O1L Δ (MOI = 0.5). After 72 or 96 h of infection, cell viabilities were determined (n = 3). (B and C) BALB/cAJcl-nu/nu mice were intraperitoneally injected with KFTX or KFlow cells stably expressing *Renilla* luciferase (n = 6). After confirming tumor growth, mice were administered an intraperitoneal injection of OVV-VGF Δ /O1L Δ (1 × 10⁶ PFUs) or PBS. Tumor imaging (B) and viral imaging (C) are shown. (D) Bioluminescence signals (in photons/s) were calculated from the imaging data of tumor (Rluc) and viral (Fluc) signals. (E) Survival curves of mice are shown after virus injection. A log-rank (Mantel-Cox) test was used to analyze significance. **p < 0.01, ***p < 0.001. Data with error bars represent mean ± SEM. Two-way ANOVA was used for (D).

sion, and host cell-signaling pathways related to actin filament branching had a critical role in viral virulence. The small guanosine triphosphate (GTP)-binding protein Cdc42 has a major role in inducing filopodia formation by regulating actin polymerization.³⁹ A recent report demonstrated that infection by the IHD-J vaccinia virus strain induces Cdc42 activation and filopodia formation.⁴⁰ ITSN1 is induced by viral infection-activated Cdc42 and associated N-WASP-actin polymerization.^{38,41,42} These results indicate that Cdc42 has an important role in viral spread by inducing filopodia formation. Consistent with this, we found that Cdc42 inhibitors blocked effective viral spread mediated by high UCA1 expression (Figures 5C and 5D).

This research is the first to indicate that host factor-regulated small G proteins mediate efficient OVV cell-to-cell spread; specifically, vaccinia virus proteins directly regulate G

protein activation. Recent studies have shown that vaccinia virus encodes F11, a viral protein with a conserved Rho-binding domain that inhibits its downstream signaling.^{43,44} The loss of RhoA signaling results in the modulation of cortical actin filaments beneath the plasma membrane and contributes to increased virus release. In this study, inhibition of the Rho GTPases RhoA and Rac1 had no effect on viral cell-to-cell spread at non-cytotoxic

⁽targeting Rho; 50 nM), ML-141 (targeting Cdc42; KFlow, UCA1-KFlow, and Empty-KFlow, 20 nM; SKOV3, 18 nM; ES-2, 15 nM), and NSC23766 (targeting Rac1; KFlow, UCA1-KFlow, Empty-KFlow, and SKOV3, 30 nM; ES-2, 15 nM) inhibitors and infected with OV-LG (MOI = 0.001). Representative plaques formed under semisolid medium were photographed and calculated with the analysis application measurement module (Keyence) (n = 10). (E and F) Activation of Cdc42 in KFlow, UCA1-KFlow and Empty-KFlow (E) and primary ovarian cancer cells (F) was analyzed by performing a colorimetric G-LISA assay (n = 3). Data with error bars represent mean ± SEM. One-way ANOVA was used for (C) and (D).*p < 0.05, **p < 0.01, ***p < 0.001.



concentrations (Figures 5C and 5D). Further, activation of RhoA and Rac1 were consistently low before and after infection (data not shown). These results suggest that actin filaments are not altered and that cell-to-cell spread is triggered by filopodia formation induced by Cdc42 activation in ovarian cancer. In primary ovarian cancer cell lines, a correlation between UCA1 expression and Cdc42 activation was observed. However, low UCA1 expression was observed in the A2780 ovarian cancer cell line, despite highly activated Cdc42 signaling, resulting in efficient cell-to-cell OVV spread (data not shown). Taken together, these results suggest that UCA1 regulates the therapeutic effect of OVV by activating Cdc42 in most ovarian cancer cells and that other pathways might be involved in the activation of Cdc42 in some ovarian cancers and other cancer types. Like UCA1, Cdc42 activation is

Figure 7. UCA1 Expression Predicts the Therapeutic Effect of Paclitaxel and Oncolytic Vaccinia Virus-VGF Δ /O1L Δ

BALB/cAJcl-nu/nu mice were intraperitoneally injected with KFTX or KFlow cells stably expressing Renilla luciferase. After confirming tumor growth, PTX (1 mL, 10 mg/kg) in saline was intraperitoneally injected twice per week. Then 3 weeks later, mice were administered a single intraperitoneal injection of OVV-VGFA/O1LA $(1 \times 10^6 \text{ PFUs})$ or PBS. (A) Tumor imaging is shown before and after PTX treatment (n = 14). (B) Bioluminescence signals (in photons/s) were calculated from the tumor imaging data. (C) Tumor imaging is shown after OVV-VGF Δ /O1L Δ injection (n = 7). (D) Bioluminescence signals (in photons/s) were calculated from the tumor imaging data. (E) Survival curves for the mice are shown after virus injection. A log-rank (Mantel-Cox) test was used to analyze significance. (F) At 3 weeks before and after PTX treatment, for the extraction of KFTX and KFlow tumors. mice were sacrificed, and tumors were collected and frozen quickly using liquid nitrogen. Collected tumors were disrupted in lysis buffer and UCA1 expression was detected by qRT-PCR (n = 9–15). **p<0.01, ***p<0.001. Data with error bars represent mean ± SEM. Two-way ANOVA was used for (B) and (D).

a potential biomarker to predict the therapeutic effect of OVV in other cancer types.

Intriguingly, the spread of other oncolytic viruses such as adenovirus and measles virus was not found to be induced by UCA1 (data not shown). The life cycle of these oncolytic viruses is independent of actin assembly.^{45–47} The OVV LC16mO strain was used for this study. Furthermore, many widely used vaccinia virus strains were shown to exhibit enhanced cellto-cell spread by inducing Cdc42-dependent actin polymerization.^{38,40} Moreover, there are different strains of OVV, such as WR, Copenhagen, LC16mO, and GLV-1h68. LC16mO was

found to be highly similar to GLV-1h68 based on phylogenetic analysis, and it shares orthologous open reading frames (ORFs) with WR and Copenhagen based on sequence comparisons.⁴⁸ These results also suggest that UCA1 does not interact with other pathogenicity factors and that it could predict the therapeutic effects of other vaccinia virus strains like WR, Copenhagen, GLV-1h68, and Wyeth; further efforts are needed to establish UCA1 as a predictive biomarker.

In summary, UCA1 triggers a variety of intracellular mechanisms and independently regulates PTX resistance and Cdc42 activation in ovarian cancer. Cdc42-induced filopodia formation promotes tumor invasion and migration, which is necessary for cell-to-cell spread, and dramatically enhances the therapeutic effect of OVV. Since these mechanisms were proven in primary ovarian cancer cells and *in vivo*, UCA1 expression has the potential to be a biomarker that could predict responses to both PTX chemotherapy and OVV.

MATERIALS AND METHODS

Cell Culture

SHIN3,⁴⁹ SKOV3, RMG-1,⁵⁰ OVCAR3,⁵¹ KFTX,⁵² KFlow, and KFTXlow cells were cultured in RPMI-1640 with 10% fetal bovine serum (FBS). A2780 cells⁵³ were cultured in DMEM with 10% FBS. ES-2 cells⁵⁴ were cultured in DMEM/Ham's F-12 plus 10% FBS. RK13 cells were cultured in E-MEM plus 5% FBS. A549 cells were cultured in Ham's F-12K plus 10% FBS. SKOV3, RK13, and A549 cells were purchased from the American Type Culture Collection (ATCC). All clinical P1, P2, and P3 samples were taken from recurrent ascites after chemotherapy, including platinum-based drugs and performed at the Saitama Medical University International Medical Center. Ascite cells were cultured in RPMI-1640 with 20% FBS. After reaching confluence, cells were trypsinized and cultured. All cultures were supplemented with 10% FBS and 1% penicillinstreptomycin (Gibco, Grand Island, NY, USA). Approval for this project was obtained from the Ethics Committee of Saitama Medical University International Medical Center (12-096) and Tottori University (2543). All cells were maintained at 37°C in a humidified atmosphere containing 5% CO₂

Microarray

Total RNA from each cell line was extracted using the Nucleo-SpinRNA kit according to the manufacturer's protocol (Takara Bio, Shiga, Japan). Using total RNA, microarray experiments were outsourced for Agilent Array expression analysis (Takara Bio). The RNA expression array was performed using the Agilent Sure Print G3 Human Gene Expression 8 \times 60K version 2 (v.2) Microarray. These data were deposited (https://www.ncbi.nlm.nih.gov/geo/; GEO: GSE122358).

Hierarchical Clustering Analysis

After extracting the expression values from the gene expression profile, a bidirectional hierarchical clustering heatmap was constructed using multiExperimental Viewer (MEV) v.4.9 software.

Gene Ontology Analysis

Microarray data comparing KFlow and UCA1-KFlow cells were subjected to gene ontology analysis. Biologically relevant features were constructed using the Database for Annotation, Visualization, and Integrated Discovery (DAVID) tools (https://david.ncifcrf.gov/). Gene lists corresponding to 2-fold changes between KFlow and UCA1-KFlow cells were uploaded to DAVID for gene ontology analysis.

Cell Viability Assays

PTX resistance was determined using the CellTiter-Glo Luminescent Cell Viability Assay (Promega, Madison, WI, USA). Each cell line was plated at 6×10^3 cells/well in 96-well plates. After 12 h, PTX was added to the media at pre-determined concentrations. After another 48 h, cell viability was measured.

UCA1 Stable Transfectants

The UCA1 gene was amplified using Phusion High-Fidelity Polymerase (New England Biolabs, Ipswich, MA, USA) with primers (5'-CTG GAT CCT GAC ATT CTT CTG GAC AAT GAG-3' and 5'-CTG CGG CCG CAT ATT AGC TTT AAT GTA GGT GGC-3'), using *KFTX* cDNA as the template. The PCR product was digested with *BamH*I and *Not*I and cloned into the same restriction site of pcDNA3.1. KFlow and KFTXlow cells were transfected with pcDNA3.1-UCA1 or pcDNA3.1 using FuGENE HD Transfection Reagent (Roche, Indianapolis, IN, USA), according to the manufacturer's instructions. Then, transfected cells were selected with G418 (500 µg/mL) for 3 weeks, resulting in UCA1-KFlow or Empty-KFlow cells, respectively.

OVV Preparation

Recombinant viruses were constructed as described previously.²⁵ Briefly, RK13 cells were infected with LC16mO²⁴ at an MOI of 0.02, and then they were transfected with pVNC110-SP-LucGFP or pUC19-VGF-SP-LucGFP plasmid, resulting in the generation of OVV-LG or OVV-VGF Δ viruses, respectively. After harvesting progeny viruses 2–5 days later, each virus was selected based on LucGFP expression, through three serial plaque purifications. Finally, the insertion of LucGFP was verified by sequencing the modified region. Similarly, OVV-VGF Δ O1L Δ was constructed by infecting RK13 cells with OVV-VGF Δ viruses, as described, and then transfecting them with the pUC19-O1-P-DsRed plasmid. All viruses were propagated in RK13 or A549 cells and stored at -80° C. For *in vivo* experiments, all viruses were purified using OptiPrep (Axis-Shield, Oslo, Norway), according to the manufacturer's protocol.

OVV Infection

Cells $(6-10 \times 10^3/\text{well})$ were seeded in 96-well plates 36 h before OVV infection. Each cell line was infected with OVV at an MOI of 0.01 or 0.001. After 48–72 h, 96-well plates were photographed under phase-contrast or fluorescence microscopy using a BZ-X700 microscope (Keyence, Osaka, Japan). EGFP fluorescence was quantified based on intensity and area using Hybrid Cell Count (Keyence, Osaka, Japan).

OVV-VGF Δ /O1L Δ Cytotoxicity Assays

Cells (6 \times 10³/well) were seeded in 96-well plates 36 h before OVV infection. Each cell line was infected with OVV-VGF Δ /O1L Δ at an MOI of 0.5. At 72 and 96 h post-infection, cell viabilities were determined using a CellTiter 96 AQueous One Solution Cell Proliferation Assay (Promega), according to the manufacturer's instructions.

CAV and EEV Titration

Cells were plated at 3×10^4 /well in 24-well plates. After 36 h, cells were infected at an MOI of 5 plaque-forming units (PFUs)/cell for 1 h at 37°C and then washed with medium and incubated for the indicated times. Supernatants, as EEV, were collected and centrifuged at 700 \times g for 10 min to separate the cells as CAV. Titers of viruses present in the supernatant and cellular fractions were then determined using RK13 cells.

Quantification of UCA1 Expression

Cells were plated at 2×10^5 /well in 6-well plates. After 36 h, total RNA from each cell line was extracted using the NucleoSpinRNA kit according to the manufacturer's protocol (Takara Bio). Each total RNA sample was subjected to reverse transcription using the High-Capacity cDNA Reverse Transcription Kit, according to the manufacturer's protocol (Life Technologies, Carlsbad, CA, USA). Then, expressions of *UCA1* and the endogenous control *GAPDH* (glyceral-dehyde-3-phosphate dehydrogenase) were quantified by real-time PCR using TaqMan probes (Applied Biosystems, Carlsbad, CA, USA). Real-time PCR conditions were 95° C for 10 min, followed by 40 cycles of 95° C for 15 s and 60° C for 60 s. The relative expression of UCA1 was calculated by the comparative threshold cycle method.

Gene Expression of Vaccinia Virus Genome

Cells were plated at 2×10^5 /well in 6-well plates. After 36 h, cells were infected with OVV-LG at an MOI of 5. Total RNA was extracted at 1, 2, 4, 6, 8, 10, and 12 h post-infection and converted to cDNA. Realtime PCR was performed using SYBR Green Master Mix with the following conditions: 95°C for 10 min, followed by 40 cycles of 95°C for 15 s, 55°C for 30 s, and 72°C for 30 s. Transcripts were detected with the following primer pairs: M1L forward (5'-AAC GGA CCA CAT CCT TCT TC-3') and M1L reverse (5'-ATC CAA ACG CGT GTGATA AA-3'); G8R forward (5'-GCG GAT CTG TAA ACA TTT GG-3') and G8R reverse (5'-CCT TGGACA CAG GAA GAT TAA A-3'); and A5L forward (5'-TTT CCA TCC GAT TGT TGT GT-3') and A5L reverse (5'-AGT TCA CTC CTT CCA GCG TT-3'). GAPDH was used as the internal loading control, and standardization among samples was performed using the primer pair GAPDH forward (5'-CCT GTT CGA CAG TCA GCC G-3') and GAPDH reverse (5'-GA CCA AAT CCG TTG ACT CC-3').

Plaque Formation Assays

Cells were plated at 2×10^5 /well in 6-well plates. After 36 h, cells were infected with OVV-LG at an MOI of 0.001. After a 1-h incubation, E-MEM containing 0.8% methyl cellulose supplemented with 5% FBS and reagents was added to each well, and plaque sizes were calculated with a BZ-X700 microscope. Reagents used for plaque assays were as follows: nocodazole (Wako, Osaka, Japan), colchicine (Wako), cytochalasin D (Wako), Rhosin (Merck, Boston, MA, USA), ML-141 (Sigma, St. Louis, MO, USA), and NSC23766 (Abcam, Cambridge, UK).

Wound-Healing Assays

Cells were plated at 6×10^5 /well in 6-well plates. After 24 h, confluent cells were scratched with a pipette tip in serum-free media. Cells were fixed at 0 and 12 h and processed for microscopic analysis. Phase-contrast images were captured using a BZ-X700 microscope.

Virion-Binding Assays

To analyze viral binding affinity, cells were plated at 1×10^5 /well in 12-well plates. After 36 h, cells were infected with OVV-LG at an MOI of 10 at 4°C for 30 min. After ample washes, cell-attached viruses were collected, and virus titers were determined using RK13 cells.

To measure early viral gene expression, cells were plated at 1×10^4 /well in 96-well plates. After 36 h, cells were infected with OVV-LG at an MOI of 10 at 4°C for 30 min. After ample washes, cells were incubated for 1 h and the ONE-Glo Luciferase Assay System was used.

GTPase Activation Assays

Cdc42 activation was measured using Cdc42-specific G-LISA activation kits (Cytoskeleton, Denver, CO, USA), according to the manufacturer's instructions. Briefly, KFlow, UCA1-KFlow, and Empty-KFlow cells, as well as primary ovarian cancer cell cultures, were plated at 1×10^5 cells/well in 6-well plates. After 36 h, G-LISA kits comprising 96-well plates coated with the binding domain of the corresponding GTPase effector protein and GTPase-GTP were detected using specific antibodies, followed by absorbance measurements at 490 nm.

In Vivo Experiments

Protocols for the following animal experiments were approved by the Animal Experiment Committee of the Institute of Medical Science, University of Tottori, Japan.

For the first *in vivo* experiments to assess the oncolytic effect of OVV-VGF Δ /O1L Δ , 5–10 × 10⁶ cells KFlow and KFTX ovarian cancer cells stably expressing *Renilla* luciferase in 200 µL PBS were intraperitoneally injected into 5-week-old female BALB/cAJcl-nu/nu mice (Clea, Tokyo, Japan). To confirm tumor growth non-invasively, 150 µL ViviRen *In Vivo* Renilla Luciferase Substrate (0.925 mg/kg, Promega) was intraperitoneally injected before virus injection. Then, mice were administrated a single intraperitoneal injection of OVV-VGF Δ / O1L Δ in 200 µL PBS or control PBS. After virus injection, 200 µL VivoGlo Luciferin, *In Vivo* Grade (150 mg/kg, Promega) was injected subcutaneously on days 1, 2, 3, and 10 to monitor viral replication and spread. On day 11, tumor size was measured based on *Renilla* luciferase activity. Each luminescence image was obtained using the NightSHADE LB985 (Berthold, Tokyo, Japan) imaging system, and, during imaging, mice were anesthetized with isoflurane.

For the second *in vivo* experiments to develop a clinically relevant mouse tumor model, $5-10 \times 10^6$ KFlow and KFTX cells in 200 µL PBS were intraperitoneally injected into 5-week-old female BALB/ cAJcl-nu/nu mice (Clea). After confirming tumor growth, PTX (10 mg/kg, Mylan, PA, USA) in 1 mL saline was intraperitoneally injected twice per week. Then 3 weeks later, tumor size was measured based on *Renilla* luciferase activity. On the following day, mice were administrated a single intraperitoneal injection of OVV-VGF Δ /O1L Δ in 200 µL PBS or control PBS. On day 11 after injection, tumor sizes were measured as stated.

For the isolation of KFTX and KFlow tumors, $5-10 \times 10^6$ cells in 200 µL PBS were intraperitoneally injected into 5-week-old female BALB/cAJcl-nu/nu mice. After confirming tumor growth, PTX (10 mg/kg, Mylan) in 1 mL saline was intraperitoneally administered twice per week. Then 3 weeks later, mice were sacrificed, and tumors

were collected and frozen quickly using liquid nitrogen. Tumors were disrupted using lysis buffer (Nucleo SpinRNA) and a Multi-Beads Shocker (Yasui Kikai, Osaka, Japan) and centrifuged at 2,000 \times g for 30 s.

Statistical Analysis

Differences in cytolytic activity and *in vivo* viral replication between treatment groups were analyzed for statistical significance by performing one-way or two-way ANOVA, and a Bonferroni test when ANOVA showed overall significance. p values < 0.05 were considered statistically significant. Survival curves were constructed using the Kaplan-Meier method. Survival times were statistically analyzed by performing a log-rank test. Data were analyzed using GraphPad Prism v.5.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10. 1016/j.omto.2019.03.003.

AUTHOR CONTRIBUTIONS

T.N. supervised the entire project. T.N. and K. Horita designed the study and interpreted all results and concluded that UCA1 promotes OVV cell-to-cell spread in ovarian cancer, resulting in enhanced therapeutic outcome. T.N. and K. Horita also wrote the manuscript. K. Horita performed the majority of the experiments. H. Kurosaki, M.N., and N.K. constructed recombinant vaccinia viruses. H.I., T.O., and T.H. isolated and maintained paclitaxel-resistant ovarian cancer cell lines. K. Hasegawa and S.S. established primary cancer cell cultures. H. Kono and M.I. purified and titrated recombinant vaccinia viruses.

CONFLICTS OF INTEREST

The authors declare no competing interests.

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Supplemental Information

IncRNA UCA1-Mediated Cdc42 Signaling

Promotes Oncolytic Vaccinia Virus

Cell-to-Cell Spread in Ovarian Cancer

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Fig. S1. UCA1 depletion impairs oncolytic vaccinia virus (OVV)-LG spread. (A) si-UCA1 and si-NC construction. (B) si-UCA1 or si-NC was transfected into KFTX cells with LipofectamineTM RNAiMAX Transfection Reagent according to the manufacturer's protocol. After 36 hours, RNA was collected and UCA1 expression was confirmed by qRT-PCR; n = 3 (C) Cells were plated at 6×10^3 /well in a 96-well plate and transfected with si-RNA. EGFP images are shown after infection with OVV-LG (MOI = 0.01) for 48 hours. Scale bar: 1000 µm. (D) The intensity and area of viral EGFP brightness was measured using a Keyence BZ-X700 fluorescence microscope; n = 3. Data with error bars represent mean ± s.e.m. An un-paired t-test was used for (B,C). **p < 0.01, ***p < 0.001.



Fig. S2. Cytotoxicity of the inhibitor against intracellular transport, cell-to-cell spread and the Rho family of small GTPases. KFlow, UCA1-KFlow, and Empty-KFlow cells were plated at 6×10^3 /well in 96-well plates. Cell viability was determined 48 hours after treatment with an inhibitor using the CellTiter-Glo® Luminescent Cell Viability Assay (Promega, Madison, WI).; n = 3. Data with error bars represent mean ± s.e.m.



Fig. S3. UCA1 expression enhances filopodia formation. Cells were plated at 1.3×10^4 /well in 8-well chambers and cultured with 10% FBS. After 24 hours, cells were infected with oncolytic vaccinia virus (OVV)-LG (MOI=1) for 12 hours. (A) Mock and (B) infected cells were fixed and stained. Images were captured under an immunofluorescence microscope with DAPI in blue, β -actin (13E5, Cell Signaling Technology, MA, USA) in red, viral EGFP in green, and merged pictures. Scale bar: 20 µm.



Fig. S4. Activation of ERK with high UCA1 expression had no effect on oncolytic vaccinia virus (OVV) spread. (A) Cells were plated at 6×10^3 /well in 96-well plates and cultured with 10% FBS. After 36 hours, cells were treated with epidermal growth factor (EGF) for 30 minutes and tERK and pERK signals were detected using an In-Cell ELISA Kit according to the manufacturer's protocol (Thermo Fisher Scientific, Rochester, NY); n = 3. (B) After 30 minutes of EGF stimulation, KFlow and UCA1-KFlow cells were infected with OVV-LG (MOI = 0.01) for 72 hours. Intensity and area of viral EGFP brightness were measured using a Keyence BZ-X700 fluorescence microscope; n = 3. Data with error bars represent mean \pm s.e.m. Two-way ANOVA was used for (A,B). ***p < 0.001.