Viral Oncolytic That Targets Raf-1 Signaling Control of Nuclear Transport

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Received 16 July 2009/Accepted 17 November 2009

The central role of Raf protein kinase isoforms in human cancer demands specific anti-Raf therapeutic inhibitors. Parvoviruses are currently used in experimental cancer therapy due to their natural oncotypic and lytic life cycle. In searching for mechanisms underlying parvovirus oncolysis, we found that trimers of the major structural protein (VP) of the parvovirus minute virus of mice (MVM), which have to be imported into the nucleus for capsid assembly, undergo phosphorylation by the Raf-1 kinase. Purified Raf-1 phosphorylated the capsid subunits in vitro to the two-dimensional pattern found in natural MVM infections. VP trimers isolated from mammalian cells translocated into the nucleus of digitonin-permeabilized human cells. In contrast, VP trimers isolated from insect cells, which are devoid of Raf-1, were neither phosphorylated nor imported into the mammalian nucleus. However, the coexpression of a constitutively active Raf-1 kinase in insect cells restored VP trimer phosphorylation and nuclear transport competence. In MVM-infected normal and transformed cells, Raf-1 inhibition resulted in cytoplasmic retention of capsid proteins, preventing their nuclear assembly and progeny virus maturation. The level of Raf-1 activity in cancer cells was consistent with the extent of VP specific phosphorylation and with the permissiveness to MVM infection. Thus, Raf-1 control of nuclear translocation of MVM capsid assembly intermediates provides a novel target for viral oncolysis. MVM may reinforce specific therapies against frequent human cancers with deregulated Raf signaling.

The selectivity of lytic viruses in anticancer therapies depends on cellular targets that favor their growth. Many if not all viral infections interfere with major cellular protein kinase networks, promoting virus entry, repressing antiviral responses, or increasing metabolic activity and virus yield. The effects are, however, virus dependent: viral activation of the MAPK signaling pathway, for example, may induce cell proliferation (oncogenic DNA virus [64]), suppress transcription (hepatitis B virus [83]), reactivate translation (Ebola virus [71]), or enhance nuclear export (influenza virus [52]). The MAPK signaling cascade is essential for cellular proliferation and cancer formation (40, 67), which led to the use of lytic viruses dependent on this pathway (7) in ongoing preclinical and clinical trials (51). Molecular insights into the interactions between viral substrates and components of the MAPK pathway could thus increase specificity and therapeutic efficacy of oncolytic viruses.

The replication of virus members of the Parvoviridae relies on functions provided by proliferative cells (24). Together with diverse factors that are linked to the neoplastic growth, the underlying mechanisms allow parvoviruses to infect and lyse transformed cells preferentially (47), thus interfering with tumor formation in animal models (14, 74); reviewed in references 11 and 54). The complex molecular mechanisms supporting parvovirus intrinsic oncotropism cannot be explained by either suitable receptors responsible for virus uptake or the cytotoxicity, which is triggered by the viral nonstructural proteins (5). Parvovirus productive infection is rather mediated on different levels of the viral life cycle in a cancer-type-specific manner (65, 66). Parvoviruses may provoke death of tumor cells lacking p53 (57, 73) and show increased toxicity to transformed cells with upregulated Ras and other protein kinases (49, 69).
Progeny parovirus production, which is required for the spreading of oncolytic effects, depends on nuclear import of de novo-synthesized structural (VP) proteins, since virion maturation is intranuclear. The parovirus capsid is composed of 60 protein subunits sharing a common C terminus folded in an eight-strand antiparallel β-barrel motif (76). In mammalian cells infected with the parovirus minute virus of mice (MVM), capsid assembly starts by cytoplasmic synthesis of two types of VP1/VP2 trimers in stoichiometric amounts (61). Nuclear translocation of the trimers is driven by an unconventional structured nuclear localization motif (NLM) evolutionary conserved in the parovirus β-barrel (37). This NLM is exposed on the trimer surface but not on the assembled capsid (37, 61). Self-assembly into viruslike particles (VLPs) may occur after expression of the parovirus major capsid protein in heterologous systems (see, for example, references 20 and 82), although little is known of the host factors regulating the nuclear targeting and assembly of parovirus capsid subunits.

The NLM sequence and configuration differs from the best-characterized nuclear localization signals (NLS) causing protein cargo translocation across the nuclear envelope (NE). Nucleocytoplasmic transport of proteins in eukaryotic cells is a saturable and energy-dependent process occurring via the nuclear pore complex (NPC [4, 75]), a supramolecular structure embedded in the NE. The bidirectional exchange of macromolecules across the NE is mostly mediated by proteins of the importin β (karyopherin β) superfamily that comprises importins (including transportins and snurportins) and exportins. Cargo recognition is facilitated by direct or indirect interaction with specialized signals, such as an NLS or a nuclear export signal (NES) (44). In the classical nuclear import pathway, karyopherin β1/importin β binds to the adaptor importin α, which interacts with cargoes carrying NLS consisting of clusters of basic amino acids (22, 28, 62). However, many imported proteins bind karyopherin β1 directly without an adaptor, and there are diverse nonclassical NLS that have structural or linear nonhomologous domains (9, 32). A well-characterized nonclassical import pathway is mediated by karyopherin β2, transportin, which binds different substrates, such as the splicing factor hnRNP A1 (53), carrying the long and structurally disordered NLS termed PY-NLS (29, 31). Independent of the transport receptor, the directionality of the transport is driven by the concentration gradient of the small GTPase Ran in its GTP- and GDP-bound form, which triggers the dissociation of the transport complexes and the release of the cargoes (reviewed in reference 70).

The missing conclusiveness to explain increased parovirus productive infection in cancer cells by receptors or cytotoxicity of the nonstructural proteins, led us search for underlying molecular mechanisms. To identify molecular targets of parovirus oncolysis, we focused on the regulation of MVM assembly in normal and cancer cells. The nuclear transport competence and phosphorylation stage of isolated VP trimers was compared in insect and human cells. We sought the mammalian kinase involved and found that VP proteins phosphorylation by the Raf-1 kinase of the MAPK cascade is essential in the nuclear translocation and capsid formation of MVM assembly intermediates. The implications of these findings for MVM anticancer therapeutic potential are discussed.

MATERIALS AND METHODS

MVMp and mammalian cells. The prototype strain of the autonomous parovirus MVM (MVMp) (13), which infects mouse fibroblasts and human tumor cells (47, 65, 72), was used. Production of viral stocks by transfection with the plasmid (59) or H5 insect cells infected with baculoviruses were harvested at 72 h postinfection (hpi; unless indicated). VLPs of MVMp were purified from AcMNPV-VP2-infected insect cells (47, 65, 72), was used. Production of viral stocks by transfection with the plasmid (59) or H5 insect cells infected with baculoviruses were harvested at 72 h postinfection (hpi; unless indicated). VLPs of MVMp were purified from AcMNPV-VP2-infected insect cells according to previously described protocols (20).

Purification of MVM assembly intermediates. NB324K human cells transduced on a large scale by electroporation (36) using the VP2-K153A expression plasmid (59) or H5 insect cells infected with baculoviruses were harvested in cold HNEM buffer (50 mM HEPES [pH 8.0], 150 mM NaCl, 2 mM MgCl2, 1 mM EDTA) supplemented with the protease inhibitor cocktail set I (Calbiochem), 100 µg of TPCK (tosylphenylalanil chloride) (Roche)/ml, and phosphatase inhibitors (20 mM glycerol phosphate, 5 mM NaF). Cells were disrupted in a cooled water bath sonicator, lysates were cleared by centrifugation at 15,000 × g and 4°C for 30 min in a benchtop centrifuge, and trimers from insect cells were enriched by consecutive 20 and 30% ammonium sulfate preprecipitations. Cellular homogenates were loaded onto 14 ml of a 5 to 20% continuous sucrose gradient in HNEM buffer prepared in advance and centrifuged at 160,000 × g in a SW40 rotor (Beckman) at 5°C and 3 h, with molecular size markers centrifuged in parallel gradients (61). Proteins in the fractions were precipitated with 10% trichloroacetic acid and 10 µg of bovine serum albumin (BSA)/ml, resuspended in Laemmli buffer, and subjected to SDS-PAGE and immunoblot analyses with anti-MVM sera. Fractions containing VP trimers (98) were pooled, dialyzed against transport buffer (see below), concentrated by a Centriplus 50 KDa (Millipore), and stored frozen in aliquots at −70°C until use. The yield was approximately 2 to 5 µg of purified VP-trimer per 107 cells.

Nuclear transport assay in permeabilized cells. Previously described methodologies (18, 18, 56) were followed with minor modifications. HeLa cells were permeabilized with 40 µg of digitonin (Sigma)/ml in DMEM for 5 min at 37°C. Permeabilization was controlled by light microscopy before the cells were washed three times for 10 min at 4°C in transport buffer (2 mM magnesium acetate, 20 mM HEPES [pH 7.3], 110 mM potassium acetate, 2 mM dithiothreitol) containing 5% BSA and 5% goat serum (Gibco). Import reactions were performed for 15 min at 37°C in 30 µl of transport buffer containing 12 mg of rabbit reticulocyte lysate (Promega)/ml, protease inhibitors (as described above), 1 mM ATP, 5 mM creatine phosphate (Sigma), and 20 U of creatine phosphokinase (Calbiochem)/ml. Substrates were 200 ng of gradient-purified MVM capsid or VP trimers. As transport controls, BSA conjugated with the SV-40 Tag NLS (PKKKRKVED) of the SV-40 classical importin route, and the VP2-K153A (Bac-VP2K153A), were previously described (20, 59). Baculoviruses expressing wt Raf-1 and the Raf-22W constitutively active mutant form (79, 80) were obtained from T. M. Roberts (Dana-Farber Cancer Institute, Boston, MA). Baculoviruses were grown in Sf9 cells and titrated as PFU (20) or as fluorescence focus units (36). For recombinant protein expression, growing Sf9 or H5 cells were infected at a multiplicity of infection of 10 PFU/cell for each baculovirus and harvested at 72 h postinfection (hpi; unless indicated). VLPs of MVMp were purified from AcMNPV-VP2-infected insect cells according to previously described protocols (20).

The human HeLa cell line (ATCC CCL2) and the C6 rat (ATCC CCL 107) and U-373MG (ATCC HTB17) tumor glioma cells were purchased from the American Type Culture Collection (ATCC) and maintained as previously described (65). The mouse A9 fibroblasts and the human NB324K simian virus 40-transformed newborn kidney fibroblasts are commonly used cell lines in MVM infections (72, 43). All cells of mammalian origin were maintained under MVM infections (72, 43). All cells of mammalian origin were maintained under MVM infections (72, 43). All cells of mammalian origin were maintained under
were probed with the indicated antisera as described previously (61). Resolved in SDS–8% PAGE and electroblotted onto nitrocellulose membranes. For Western blot analysis, protein samples were subjected to immunoprecipitation with the nuclear import receptors (17), was added at 100 μg/ml in the transport reaction and washing buffer. The transport reaction was terminated by placing the cells on ice, followed by three washes with transport buffer and fixation in 4% paraformaldehyde.

Immunological analyses. Antibodies to components of the MAPK signaling pathway (Santa Cruz Biotechnology) included mouse monoclonal (mo-Raf) and rabbit polyclonal (α-Raf) antibodies recognizing the conserved C terminus of human Raf-1; rabbit polyclonal antibody (α-ERK) recognizing the ERK1/2 proteins, and mouse monoclonal antibody (α-P-ERK) recognizing their phosphorylated forms. A mouse monoclonal antibody (α-NPC) (MAb614; Babco, Richmond, CA) was used as marker of the nuclear envelope. Human casein kinase II was immunoprecipitated with an anti-CKIIα polyclonal antibody (Santa Cruz Biotechnology). Antibodies raised against denatured VP2 protein (α-VPs) for the general localization of VP1 and VP2 capsid protein subunits (to be described elsewhere). Indirect double-label immunofluorescence was performed by using secondary antibodies conjugated to Texas Red or fluorescein isothiocyanate (FITC) (36), and phenotypes were visualized by epifluorescence in a Zeiss Axiovert 200 microscope equipped with a Radiance 2000 confocal laser scanning system (43). Pictures were recorded with lasers giving excitation lines at 488 nm (FITC, filter HQ 500-530), and 543 nm (TXRD, filter HQ 600 LP). For Western blot analysis, protein samples resolved in SDS–6% PAGE and electroblotted onto nitrocellulose membranes were probed with the indicated antisera as described previously (61).

Raf-1 phosphorylation of MVM capsid subunits in mammalian cells and in vitro. For [32P]orthophosphate labeling, MVMp-infected cells (5 PFU/cell) were starved for 4 h at 6 h in phosphate-free medium with 10% dialyzed FCS and labeled 10 to 24 hpi by the addition of 1 mCi of carrier-free [32P]orthophosphate (Amersham)/ml. Capsid proteins were precipitated at 4°C by using polyclonal anti-capsid antibodies bound to protein A-Sepharose in 150 mM NaCl–50 mM Tris (pH 8.0)–0.3% SDS–1% NP-40–0.75% 2-mercaptoethanol and protease and phosphatase inhibitors prior to SDS-PAGE analysis. For in vitro phosphorylation, the indicated kinases were immunoprecipitated with antibodies from human NB324A cells at 4°C in 150 mM NaCl–50 mM Tris–HCl (pH 7.5)–0.05% SDS–1% NP-40–0.5% decachloroethane in the presence of protease and phosphatase inhibitors. The Sepharose-bound precipitate was stored at –70°C. The phosphorylation reaction was performed at 37°C for 30 min in 40 μl containing immunocomplex from 106 cells, 8 μg of purified VP2 protein in 10 mM [γ-32P]ATP (10 μCi [Amersham], 3,000 Ci/mmol), 0.1 μM rATP, 25 mM HEPES (pH 7.4), 25 mM glycerophosphate, 1 mM dithiothreitol, and 10 mM MgCl2. The [32P]-labeled VP2 protein resolved in gels was blotted to nitrocellulose membranes (BA85; Schleicher & Schuell) and subjected to two-dimensional tryptic phosphopeptide analysis in 20- by 20-cm thin-layer chromatography plates (Merek, Darmstadt, Germany) according to previously reported procedures (42). The plates were exposed to a radioanalytic imaging system (Fujix BAS 1000; Fuji) or to X-ray films with intensifying screens for 4 days.

Analysis of MVM capsid protein phosphorylation in insect cells. [32P] labeling of VP2 protein and VLPs in S9 and H5 insect cells infected by baculoviruses was attempted for periods of 6 h at either 22, 38, or 48 hpi in TC100 medium without phosphate and 5% dialyzed FCS supplemented with 0.5 to 1 mCi of [32P]orthophosphate carrier-free (Amersham)/ml at 8,500 Ci/mmol. Okadaic acid (Gibco), a phosphatase inhibitor (3), was used at 350 nM during the last hour of labeling to prevent the removal of putative transient phosphate groups. VLPs were purified from the [32P]-labeled cultures by CsCl gradient centrifugation to equilibrium (42). Fractions were tested for [32P] counts and hemagglutination activity (20), and those corresponding to empty MVM particles were pooled and dialyzed. Detection of phospho label in the purified VP2 protein was repeatedly attempted by separating 5 μg of gradient-purified VLPs by SDS-PAGE and exposure to autoradiography with intensifying screens for 7 days. To examine the level of VP2wt and VP2K153A protein phosphorylation in the presence of baculovirus coexpressed Raf-1 proteins, the [32P]-labeled H5 cell monolayers were subjected to immunoprecipitation with the α-VPs and α-Raf-1 antibodies as described above.

RESULTS

Trimers of MVM capsid subunits assembled in insect cells are not competent for nuclear transport. In mammalian cells susceptible to the parvovirus MVM, the capsid proteins (VPs) accumulate intranuclear (12, 37) (see also Fig. 4B below). In contrast, most baculovirus-transduced insect cells, which do not support replication of this mammalian virus, exhibited an exclusive cytoplasmic localization of the VPs (Fig. 1A, upper panels). Particle-specific antibodies characterized by slot blot analyses (see Fig. 1B) showed that, in contrast to mammalian...
cells, VLPs self-assembled inefficiently and primarily localized in the perinuclear cytosolic region (Fig. 1A). This finding, which confirmed previous reports on MVM (20) and canine parvovirus (82) assembly in insect cells, not only showed a cell-type-specific transport but also indicated a different assembly competence.

To avoid the possibility that capsid formation in the cytoplasm may interfere with the nuclear transport competence of VP trimers, the major assembly intermediate (61), we used the VP2-K153A mutant. This mutant forms trimers that are nuclear import competent in mammalian cells but do not assemble into capsids (61). Expression of this mutant in H5 insect cells showed a cytoplasmic phenotype with nuclear exclusion (Fig. 1A, lower panels) as for the wt VP2. Cytoplasmic capsid formation was thus not responsible for the lack of VP2 nuclear import in insect cells. Since nuclear VP2 (or VP2-K153A) localization depends on trimer formation (61), we sought to determine whether insect cells are competent for VP2 trimerization. Both wt and K153A mutant proteins expressed in insect cells sedimented with the velocity of VP2 trimers by analytical sucrose gradient ultracentrifugation (Fig. 1C) and could further be chemically cross-linked as described previously (61; data not shown). Collectively, these findings indicate that the failed VP2 nuclear translocation in insect cells was not determined by impaired trimer formation.

Active nuclear import of VP trimers isolated from human cells. Although the nuclear transport process is evolutionarily well conserved, there are some differences between distantly related organisms, since some proteins are imported into the nucleus in a species-dependent manner (10). To exclude that such phenomena cause the lack of import of the VP trimers in insect cells, we purified VP2-K153A trimers from NB324K human and H5 insect cells and tested them with digitonin-permeabilized human HeLa cells. Transport was facilitated by import receptors that are present in rabbit reticulocyte lysate, which supports nuclear entry in various permeabilized mammalian cell lines. A control experiment was performed using native empty capsids purified from MVM-infected cells. Consistent with the localization of the NLM in the internal capsid surface (37), no capsids were detectable within the nucleus of the HeLa cells (Fig. 2A, upper panels). However, human cell line-derived VP2 trimers translocated into the nucleus. In agreement with our previous report (61), this result provided further direct evidence of the important role that the VP trimeric intermediate plays in MVM capsid assembly. Transport was temperature dependent, indicating active nuclear import (Fig. 2A middle panels). Active receptor-mediated transport was confirmed by using the nuclear import inhibitor WGA (Fig. 2A, lower panels). In contrast, H5 insect cell-expressed trimers failed to accumulate in HeLa nuclei even under transport supporting conditions (Fig. 2B).

Site-specific VP2 phosphorylation by Raf-1 in vitro. In searching for the mechanism underlying the different nuclear transport competence of mammalian and insect VP trimers, we investigated the role of phosphorylation, known to be a major regulator of cargo-receptor interactions in nuclear import (60). We observed that wt VLPs and VP2-K153A trimers purified from H5 and S9 insect cells showed no detectable phosphorylation, irrespective of the conditions of metabolic labeling (see Materials and Methods). In contrast, the VP capsid subunits from MVM-infected human and mouse cells are phosphorylated (42, 43). Phosphorylation occurred mainly at Ser-2, -6, and -10 of the VP2 N-terminal domain (2Nt) (42), overlapping with within two contiguous S/TXXXS/T canonical activation loop motifs of the MAPK signaling pathway (40, 78). To explore whether differences in the MAPK signaling pathway between insect and mammalian cells can account for the different VP2 phosphorylation, we subjected insect cell-derived VP2 to in vitro phosphorylation assays with human Ser/Thr protein kinases after precipitation by different antibodies. We focused on the Raf family (A-Raf, B-Raf, and C-Raf [or Raf-1]), which are involved in the development of a high proportion of glioma, melanoma, and other human malignancies (15, 16, 38, 78, 77), by promoting the nuclear translocation of phosphorylated downstream effectors (16, 41). As a control, CKII was used since it affects the nuclear transport capacity of some NLS-containing proteins (60).

Intact VLP could not be phosphorylated, irrespective of the anti-protein kinase antibody used for pulldown. In contrast, heat-disassembled capsid underwent substantial phosphoryla-
tion by Raf-1 and by CKIIα and, to lower extent, by the P-ERK kinase (Fig. 3A). The two-dimensional phosphopeptide pattern obtained with Raf-1 (Fig. 3B) was remarkably identical to the pattern of VP2 with native empty capsids purified from MVM-infected cells (42, 43). In contrast, P-ERK and CKII/H9251 phosphorylations analyzed by two-dimensional chromatography lacked the characteristic prominence of phosphopeptide B (data not shown). These results indicate that Raf-1 can directly phosphorylate VP2 at the sites and to a similar extent found in MVM infections, but on unassembled protein subunits only.

**Raf-1 phosphorylation of capsid assembly intermediates is required for MVM productive infection.** The MVM capsid surface lacked Raf-1 phosphorylation sites but these sites became exposed on heat-treated capsid (Fig. 3A and B). The exposure of Raf-1 phosphorylation sites on MVM capsid disassembly intermediates was further analyzed in vitro by using purified intact VLPs subjected to graded heat-induced conformational rearrangements. MVM native capsid and VLPs undergo conformational changes along a 60 to 90°C temperature range that can be monitored by fluorescence (6) and are associated with the externalization of the coat of the VP2 2Nt that becomes accessible to trypsin digestion (20). As shown in Fig. 3C, the VP2 phosphorylation by Raf-1 gradually increased when the samples had been heated to temperatures that cause partial or total VLP disassembly. It is noteworthy that Raf-1 only phosphorylated residues heat externalized out of the coat in VP2 domains fully accessible to the protease. At the lower temperatures of 60 to 75°C, these phosphorylated residues were localized in the 2Nt domain, since they were removed from the capsid as the VP2 protein was cleaved by trypsin to produce VP3. This experiment confirmed that 2Nt, the major phosphorylated domain of MVM capsid in infected cells (42), is also a major target of Raf-1 phosphorylation in vitro.

The assembly-dependent exposure of Raf-1 phosphorylation sites in vitro suggested that VP phosphorylation in cells must occur prior to capsid assembly, which follows VP nuclear import (61). To confirm the latter hypothesis in vivo, the effect of Raf-1 activity on VP transport and assembly was investigated. We used Radicicol, which selectively depletes Raf kinase (68). Inhibitor treatment instead of a genetic silencing approach was chosen since permanent Raf-1 inhibition affects cell proliferation, which is mandatory for parvovirus replication (24). The inhibitor was added on synchronous cultures of three MVMp permissive cells with different origins and tumorigenic potentials, at postinfection times and in a concentration that allowed both effective Raf-1 depletion and accumulation of capsid proteins to high levels (Fig. 4A). The Radicicol treatment caused a significant cytosolic retention of the VP proteins that was partial in human NB324K and mouse A9 fibroblasts (Fig. 4B and C) and absolute in U373 glioma cells (Fig. 4D). The different degrees of VP transport inhibition among the cell lines occurred consistently, for reasons that were not further investigated. However, a severely reduced capsid formation was consistently observed in all three cell cultures (Fig. 4B to C).
D, α-capsid/+Rad panels), corresponding to an inhibition in infectious virus yields by ~1,000-fold within a single replication cycle (data not shown). Collectively, these results indicated that inhibition of Raf-1 signaling cascade interferes with the nuclear accumulation of VP proteins, which is a prerequisite for capsid assembly (61). Consistently, a severe impairment of MVM capsid formation and progeny virus production was observed.

**Raf-1 phosphorylation confers nuclear transport competence to the VP trimer.** To directly relate Raf-1 phosphorylation to the nuclear transport of MVM capsid proteins, we used insect cell-expressed VP2 trimers coexpressed with the wt or with a constitutively active form of the Raf-1 kinase (Raf-22W). Expression of Raf-22W was performed since—in contrast to mammalian cells—insect cells require the coexpression of p21ras, pp60src, or JAK2 oncoproteins to obtain full wt Raf-1 activation (79, 81). Phosphorylation of the VP2 protein in the insect cells was evaluated by in vivo labeling with [32P]orthophosphate. VP2 phosphorylation was almost undetectable in baculovirus infections even under a very high level of accumulation of phosphorylated wt Raf-1 protein (Fig. 5A). However, wt and K153A mutant VP2 proteins were phosphorylated to similarly high levels in insect cells coexpressing the constitutively active Raf-22W protein (Fig. 5A, middle and lower panels). We next purified such phosphorylated VP2 trimer and its nonphosphorylated equivalent and analyzed their transport competence. In agreement with the results shown in Fig. 2, the nonphosphorylated VP2 trimer did not enter the nuclei of digitonin-permeabilized HeLa cells (Fig. 5B, upper panels). In contrast, VP2 trimers isolated from Raf-22W-expressing insect cells were efficiently and actively imported (Fig. 5B, lower panels). We thus concluded that Raf-1 phosphorylation confers nuclear transport competence to the VP2 trimer in mammalian cells.

**Raf-1 activity in mouse and human transformed cells increases permissiveness to MVM infection.** We finally investigated the relationship between Raf-1 activity and the susceptibility to MVM infection and oncolysis. The simian virus 40-transformed human NB324K and the mouse A9 fibroblasts are two common MVM cell hosts (72). Virus production was significantly higher in the NB324K cells (43, 72), which corresponded to an increased degree of VP2-specific phosphorylation (43). We thus analyzed whether Raf-1 activation correlated with these parameters. Figure 6A demonstrates that the amount of Raf-1 was similar in both fibroblast cell lines. Activation of the MAPK signaling pathway was, however, higher in NB324K

![FIG. 5](http://jvi.asm.org/) Raf-1 phosphorylation suffices VP2 trimer for nuclear import. (A) VP2 phosphorylation by Raf-1 in insect cells. H5 cells infected with the indicated baculoviruses were labeled at 22 to 28 hpi with [32P]orthophosphate. The upper section blot shows VP2 and Raf-1 expression in the [32P]orthophosphate-labeled cultures. For the middle panel, autoradiography of the same filter was performed. The lower panel shows phosphorylation of VP2wt and VP2-K153A proteins in Raf-22W-expressing insect cells demonstrated by immunoprecipitation with the α-VPs antibody. Approximately 10^7 labeled cells were subjected to each analysis. (B) Confocal laser scanning microscopy of the nuclear transport of phosphorylated VP2 trimer in mammalian cells. Gradient-purified VP2 trimers from insect cells infected by the Bac-VP2wt (VP trimer) or by the Bac-VP2wt and the Bac-Raf1-22W baculoviruses (VP trimer-P) were subjected to transport analysis in permeabilized HeLa cells. The lower panels show the results of a control experiment with permeabilized cells kept on ice showing no significant nuclear localization caused by diffusion.

![FIG. 6](http://jvi.asm.org/) Activation of the MAPK signaling cascade in MVM permissive cells. (A) Levels of Raf-1, and of phosphorylated ERK protein isoforms, accumulated in the indicated cell lines, as determined by SDS-PAGE and immunoblotting. Cells were starved 24 h with low serum and stimulated 6 h with 10% FCS before harvest. Equivalent graded amounts of total protein extracts were loaded per slot. (B) Pattern of VP2 phosphorylation in glioma cells. The figure shows two-dimensional phosphopeptide maps of VP2, immunoprecipitated from the indicated MVM-infected glioma cells labeled with [32P]orthophosphate. Scale bar, 2 cm.
cells. To exclude a cell-type-specific activation that is independent to permissiveness, we further analyzed the human U373 and the rat C6 glioma cell lines, which are both highly permissive to MVM cytotoxicity and productive infection (65). Both glioma cells exhibited MAPK signaling pathway activation (Fig. 6A) that corresponded to the characteristic VP2 phosphorylation pattern found in MVM infections (Fig. 6B). Of note, phosphopeptide B, representing the VP2 2Nt (42), was particularly prominent in the human U3T3 glioma cells. Conclusively, the levels of Raf-1 activation corresponded with increased virus production and VP2-specific phosphorylation in MVM permissive transformed fibroblasts and glioma cells.

To gain direct evidence that an activated Raf-1 does increase cell permissiveness to MVM, mouse A9 cells were stably transfected with the constitutively active RafCAAX mutant (33). Two clones transfected by the RafCAAX expression vector harboring an increased Raf-1 activity with respect to the parental A9 cells were selected (Fig. 7A). Upon MVM infection, these clones showed a moderately enhanced VP phosphorylation (Fig. 7B). Moreover, the two-dimensional phosphopeptide map of the expressed VP2 (Fig. 7C) remained identical to that found after infection of the parental A9 cells (43).

Raf-1 activation thus enhanced VP2 specific phosphorylation in vivo. To correlate the extent of Raf-1 phosphorylation with the level of productive infection, we analyzed MVM plaque-forming capacity in the parental and Raf-1-transfected A9 cells. As shown in Fig. 7D, the higher extent of VP2 phosphorylation in RafCAAX-expressing cells correlated well with a three- to fourfold increase in the number of MVMwt plaques formed in the transfected clones. For further confirmation, we infected these cells with an MVMp mutant harboring the three serine residues targeted by Raf-1 at the 2Nt domain mutated to glycine (42). This mutant (S/G) lacking 2Nt phosphorylation grows poorly but can be purified from NB324K cells (42), although it can form clear plaques in A9 fibroblasts (43). In contrast to the wt, this mutant showed no significant alteration in its plaque phenotype (regarding both number and size of the plaques) comparing A9 and A9-RafCAAX cells (Fig. 7D, right). Thus, a constitutively active Raf-1 does not enhance the plaque-forming capacity of a MVM mutant lacking the target Ser residues at 2Nt.

DISCUSSION

Lack of MVM capsid protein phosphorylation in insect cells. Most vertebrate proteins expressed by baculoviruses in insect cells are correctly phosphorylated. However, quantitatively altered and nonphysiological phosphorylation sites were described for a number of mammalian virus proteins (21, 34). The lack of MVM capsid protein phosphorylation in insect
cells described in the present study is thus uncommon. It was unexpected given the overall MAPK signaling pathway conservation in evolution, including the Raf homolog (Draf) found in Drosophila melanogaster (46). Although Draf contains all three conserved Raf domains, its sequence similarity to Raf-1 is lower across their N-terminal regulatory region (35), suggesting that insect Raf proteins may be not homologous enough to interact with and phosphorylate VP2. Our results are consistent with the requirement of coexpressed oncoproteins of this pathway in insect cells for mammalian Raf-1 activation (79–81).

Raf-1 phosphorylation regulates the MVM life cycle. The Raf-1 kinase plays an essential role in MVM cycle, since Raf-1 activity increased MVM infection (Fig. 6 and 7). The site-specific in vitro phosphorylation of fully and partly disassembled VLP protein subunits by Raf-1 (Fig. 3) strongly suggests that newly synthesized VP proteins in MVM-infected mammalian cells are phosphorylated by the normally cytoplasmic Raf-1 kinase (33). Raf-1 phosphorylation was required (Fig. 2) and sufficient (Fig. 5) to confer competence for nuclear translocation to the VP subunits. Phosphorylation was thus needed for the interaction of VP trimers with nuclear transport factors. The mechanism by which Raf-1 phosphorylation enhances NE translocation of MVM assembly intermediates is in agreement with the largely recognized association of parvovirus infections with mitogenic and proliferative cell processes (2, 24).

In insect cells devoid of Raf-1 activity, VP2 proteins lacking any phoshp label (Fig. 3A and 5A and data not shown) self-assembled into cytoplasmic VLPs but rather inefficiently (Fig. 1A), as was previously shown for the canine parvovirus (82). We further show here that the VP2 proteins accumulate as trimers and large aggregates in the insect cytoplasm (Fig. 1C). Therefore, VP phosphorylation may be also important in the orchestrated parvovirus morphogenesis, in particular at the stage of assembly of the trimeric intermediate into the T=1 capsid. Interestingly, the capsid subunits must indeed undergo a conformational change along this stage of the assembly process, since trimers expressed in cells (61) (Fig. 1A) or isolated as protein complex in vitro (Fig. 1B) failed to configure the epitope recognized by the B7 monoclonal antibody at the threefold axis of the capsid (23). VP phosphorylation may facilitate this conformational change or the recognition of the trimers by a morphogenetic cellular factor. However, VLPs were shown to exhibit the same three-dimensional crystal structure as native phosphorylated MVM capsid (20, 27, 42), indicating that phosphorylation does not shape the final configuration of the amino acid residues in the formed capsid.

Raf-1 targeting in MVM anticancer potential. The Raf kinase isofoms are frequently overexpressed, activated, or mutated in glioma, melanoma, and pancreatic carcinomas (15, 16, 38, 77, 78). The findings that Raf-1 phosphorylation (i) plays a key role in the nuclear entry of MVM capsid assembly intermediates, (ii) is required for virus maturation, and (iii) supports propagation of the lytic infection in glioma and other human transformed cells is thus supporting current searches for inhibitors of the Raf signaling cascade for cancer treatment (19, 45, 63). Disruption of the Raf-1–Rb interaction by RRD-251 suppresses tumor growth (26), and Sorafenib or Nexavar (Bayer 43-9006), designed to counter Raf kinases, is currently in phase III clinical trials for melanoma treatment and is approved for metastatic renal cancer (30). However, concerns about the specificity and efficacy of these inhibitors in the clinic were raised, since therapeutic benefit may be caused by side inhibition of V-EGFR needed for angiogenesis (reviewed in reference 39) and Raf-1 may be activated as consequence of B-Raf inhibition (77). Parvoviruses, which show antiproliferative activity in animal models (11, 54), may thus become alternative and more specific agents in combined therapies against certain cancer types. Our results support MVM, a mouse parvovirus that is apathogenic for humans, as a cytotoxic and replicative inhibitor targeting common cancers with a deregulated Raf signaling cascade.

ACKNOWLEDGMENTS

We gratefully acknowledge C. J. Marshall (CRC, Institute of Cancer Research, London, United Kingdom) for the plasmid expressing the RafCAAX active form of Raf-1 and T. M. Roberts (Dana-Farber Cancer Institute, Boston, MA) for the Bac-Raf-1 and Bac-Raf22F constructs.

This study was supported by grants from the Spanish Ministerio de Ciencia e Innovación (SAF2008-03238) and Comunidad de Madrid (S-SAL/0185/2006) to the laboratory of J.M.A. and by an institutional grant from Fundación Ramón Areces to the Centro de Biología Molecular “Severo Ochoa.” L.R. was supported in part by a short-term EMBO fellowship.

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