The measles virus V protein binds to p65 (RelA) to suppress NF-kappaB activity

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Abstract

Nuclear factor-κB (NF-κB) transcription factors are involved in controlling numerous cellular processes, including inflammation, innate and adaptive immunity, and cell survival. Here we show that the immunosuppressive measles virus (MV, Morbillivirus genus, Paramyxoviridae) has evolved multiple functions to interfere with canonical NF-κB signaling in epithelial cells. The MV P, V, and C proteins, also involved in preventing host cell interferon responses, were found to individually suppress NF-κB-dependent reporter gene expression in response to activation of TNF receptor, RIG-I-like receptors, or Toll-like receptors. NF-κB activity was most efficiently suppressed in the presence of V, while expression of P or C resulted in moderate inhibition. As indicated by reporter gene assays involving overexpression of the IKK complex, which phosphorylates the inhibitor of κB to liberate NF-κB, V protein targets a downstream step in the signaling cascade. Co-immunoprecipitation experiments revealed that V specifically binds to the Rel-homology domain of the NF-κB subunit p65 but not of p50. Notably, the short C-terminal domain of the V protein, which is also involved in binding STAT2, IRF7, and MDA5, was sufficient for the interaction and for preventing reporter gene activity. As observed by confocal microscopy, the presence of V abolished nuclear translocation of p65 upon TNFα stimulation. Thus, MV V appears to prevent NF-κB-dependent gene expression by retaining p65 in the cytoplasm. These findings reveal NF-κB as a key target of MV and stress the importance of the V protein as the major viral immune-modulatory factor.
Introduction

The innate immune response to viruses involves activation of pattern recognition receptors (PRRs) and transcriptional induction of type I interferons (IFNs) and proinflammatory cytokines. IFN genes are mainly controlled by the activity of interferon regulatory factors (IRF) 3 and 7, but activator protein 1 (AP1) and nuclear factor of the kappa light chain enhancer of B cells (NF-κB) are transcription factors joining the enhanceosome for efficient and regulated transcription of the IFNβ gene (31, 47). NF-κB, in addition, plays an important role in the innate immune system since it controls transcription of a large variety of proinflammatory cytokines upon activation of diverse receptors including not only PRRs like the toll-like receptors (TLRs) and retinoic acid-inducible gene I-like receptors (RLRs); but also members of the tumor necrosis factor receptor (TNFR) family (19). Moreover, NF-κB regulates numerous physiological processes, like immune cell development, proliferation and homoeostasis of the adaptive immune system (24).

The mammalian NF-κB family comprises five members: p65 (RelA), p50 (NF-κB1), p52 (NF-κB2), cRel and RelB. All family members share a structurally conserved N-terminal region of about 300 amino acids, the Rel homology domain (RHD), which is critical for homo- or hetero-dimerization, binding to cognate DNA-sequences, termed κB motifs, and interaction with specific inhibitory proteins. Rel proteins (p65/RelA, cRel, RelB) contain a C-terminal transactivation domain, which is lacking in p50 and p52. Thus, p50 and p52 form heterodimers with a Rel protein for gene activation or homodimers to function as repressors of promoters bearing κB motifs (35). The predominant form of NF-κB is a heterodimer of p65 and p50 subunits.

Most NF-κB dimers are located in the cytoplasm in an inactive form because of their association with inhibitor of κB (IκB) proteins, the most common of which is IκBα (2, 3).
These regulatory proteins mask the NF-κB nuclear localization sequence (NLS) within the NF-κB dimers and thus sequester them in the cytosol. A critical event in the so-called canonical activation of NF-κB is the phosphorylation of IκB proteins by IκB kinases (IKKs).

The IKK complex involved contains two catalytic subunits, IKKα and IKKβ, as well as a regulatory subunit, IKKγ (NEMO (NF-κB essential modulator)). Upon activation, the IKK complex phosphorylates IκBα which is the signal for ubiquitination and proteasomal degradation of the inhibitor. This leads to liberation of the NF-κB dimers, their nuclear translocation and NF-κB-dependent gene transcription. Numerous upstream signaling cascades converge on the IKK complex (27), which is therefore the central mediator of canonical NF-κB activation.

RNA viruses like the *Paramyxoviridae* have developed multiple and powerful strategies to counteract IRF3/7-dependent IFN induction and signal transducer and activators of transcription (STAT)-dependent IFN signaling (21). While numerous recent studies on paramyxovirus innate immune antagonistic activities have brought forth much knowledge on how control of IRF3/7 and STAT is achieved, their potential to interfere with NF-κB is less well studied (25, 26).

Here, we assessed the capability of the immunosuppressive measles virus (MV) to interfere with NF-κB signaling. MV is a non-segmented negative strand RNA virus of the *Paramyxoviridae* family which typically triggers PRRs through interaction of viral RNA with TLR3 or TLR7 in the endosomes or with the RLRs like RIG-I in the cytoplasm (44).

Induction of these pathways leads to the activation of both IRF3 and NF-κB and therefore to the transcription of IFNβ and inflammatory cytokines. Measles virus proteins have been shown to inhibit IRF3 and IRF7 activating pathways as well as IFN signaling through different mechanisms (17). Specifically, the three phosphoprotein (P) gene products P, V, and C have been shown to act as the key players of MV mediated immune evasion. A process...
called RNA editing, where an additional G is inserted into the mRNA of the P gene transcript gives rise to the V protein (10). Thus, MV V has a unique, cysteine-rich C-terminal domain (V_{CTD}) and an N-terminal domain which is identical to that of MV P (PV_{NTD}) (Fig. 1A). Notably, the structure of the cysteine-rich and zinc-coordinated V_{CTD} domain is conserved among paramyxovirus family members. Expression of the C protein is achieved through alternative translation initiation (5). In this study we examined the effect of the MV P, V, and C proteins on canonical NF-κB activation. We found that any of the MV P gene products can interfere with NF-κB-dependent gene expression, illustrating that NF-κB is an important target of MV. The V protein displayed the strongest inhibitory effects and was found to specifically bind to the NF-κB subunit p65 and to preclude its nuclear accumulation. Intriguingly, the small V_{CTD}, which is engaged in targeting multiple factors of IFN induction and IFN signaling pathways, was identified as responsible for p65 interaction.
Materials and Methods

Cell culture. Human embryonic kidney HEK-293T cells and HEp2 cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% foetal bovine serum, 1x L-glutamine and penicillin-streptomycin (Gibco®, Invitrogen).

Plasmids and reagents. Generation of expression vectors encoding individual P gene products of the MV Schwarz vaccine strain and their Ig- and flag-tagged versions and fragments were described recently (38). The plasmid encoding rabies virus P (RV P) was described previously (6). The pCR3-Ig vector (where Ig is immunoglobulin) (8) was used to generate a plasmid expressing an N-terminally Ig-tagged NEMO binding domain peptide (Ig-NBD) (34). Oligodeoxynucleotides specifying NBD (forw: 5’ ATA GAA TTC CTA GAC TGG AGC TGG TTA CTC GAG ATA 3’; rev: 5’ TAT CTC GAG T AA CCA GCT CCA GTC TAG GAA TTC TAT 3’) were annealed and cloned into pCR3-Ig using *XhoI/*EcoRI restrictions sites. The NF-κB firefly luciferase reporter plasmid p55A2-luc, comprising three repeats of the PRDII domain of the IFNβ-promoter (50) and the ∆RIG-I plasmid was kindly provided by T. Fujita, Kyoto, (49). Expression vectors for IKKα, IKKβ, and TRIF were kindly provided by K. Ruckdeschel, Munich, the plasmid for IKKγ by F. Randow, Glasgow, and vectors expressing IPS-1 and MyD88 by S. Akira, Osaka. pRL-CMV was purchased from Promega and expression plasmids for p65 (21966) and p50 (21965) from Addgene (4). A plasmid encoding the C-terminal flag tagged p65 was generated by PCR amplification from p65 using the following primers: forw: 5’ ATA AAG CTT GCC ACC ATG GAC GAA CTG TTC CCC 3’ and rev: 5’ ATA CTC GAG CTA TTT ATC GTC ATC G TC TTT GTA GTC GGA GCT GAT CTG ACT CAG 3’ followed by cloning into pCR3 using *HindIII/*XhoI restriction sites. The N-terminal fragment of p65 (1-309) representing the Rel homology domain (RHD) was amplified with forw: 5’ ATA AAG CTT GCC ACC ATG GAC GAA
and cloned into pCR3 using HindIII/XhoI restriction sites. The vector expressing the flag-tagged version of RHD p65 (flag-RHD p65) was generated by PCR amplification using following primers: forw: 5’ ATA AAG CTT GCC ACC ATG GAC GAA CTG TTC CCC 3’ and rev: 5’ ATA CTC GAG TTA TTT ATC GTC ATC GTC TTT GTA GTC GAA GGT CTC ATA TGT 3’ followed by cloning into pCR3 using HindIII/XhoI restriction sites. A plasmid encoding the flag-tagged version of p50 was constructed using the following primers: forw: 5’ ATA AAG CTT GCC ACC ATG GAC TAC AAA GAC GAT GAC GAT AAA GCA GAA GAT GAT CCA 3’ and rev: 5’ ATA CTG GAG TTA AAC TTT CAC AG C GTC CCA 3’ followed by cloning into pCR3 using HindIII/XhoI restriction sites. The flag-tagged RHD of p50 (1-366) was amplified with forw: 5’ ATA AAG CTT GCC ACC ATG GAC TAC AAA GAC GAT GAC GAT AAA GCA GAA GAT GAT CCA 3’ and rev: 5’ ATA CTC CAG TTA TTT CAC AG TTT CCT 3’ and cloned into pCR3 using HindIII/XhoI restriction sites. Recombinant human TNFα was purchased from Biomol.

**Reporter gene assay.** HEK-293T cells in 24-well plates were transfected with p55A2-luc (100 ng), pRL-CMV (10 ng), and the indicated amounts of expression vectors using Lipofectamine 2000 (Invitrogen). The total amount of transfected DNA were adjusted by adding empty vector. In case of external stimulation cells were treated with 10 ng/ml TNFα. Cells were harvested in passive lysis buffer (Promega) at the indicated time points and lysates were subjected to reporter gene assay using dual luciferase reporter system (Promega). Luciferase activity was measured with a luminometer (Berthold Lumat LB 960) according to the manufacturer’s instruction. For Western blotting, reporter assay lysates were mixed 1:1 with SDS sample buffer (62.5 mM Tris, 2% SDS, 10% glycerol, 6 M urea, 5% β-mercaptoethanol, 0.01% bromophenol blue, 0.01% phenol red).
Confocal microscopy and antibodies. HEp2 cells grown on coverslips in 24-wells were transfected 24 h post seeding with 500 ng of plasmids for flag-MV V or pCR3 using Lipofectamine 2000, incubated for additional 24 h and then treated with 10 ng/ml TNFα. Cells were fixed after 30 min in 3% paraformaldehyde for 20 min at room temperature, quenched with 50 mmol Ammoniumchlorid for 10 min at room temperature and permeabilized in 0.5% Triton X-100 in phosphate-buffered saline (PBS). After blocking with 2,5% milk in 0,1% Triton X-100/PBS, fixed cells were incubated with primary antibodies, α-p65 (rabbit; sc-109; Santa Cruz) diluted 1:200 and α-flag (mouse; Sigma) diluted 1:200 in 0,1% Triton X-100/PBS for 1 h at 4°C, followed by incubation with fluorescence-labeled secondary antibodies (goat α-rabbit Alexa Fluor 488 and α-mouse tetramethylrhodamine, both from Molecular Probes) at a dilution of 1:200 in 0,1% Triton X-100/PBS for 1 h at 4°C. Nuclear chromatin was stained by adding TO-PRO-3-iodide (Molecular Probes, 1:2000) to the secondary antibodies. Confocal laser scanning microscopy was performed with a Zeiss LSM510 Meta laser system using a Zeiss Axiovert 200 microscope. Excitation of Alexa Fluor 488, tetramethylrhodamine, and TO-PRO-3-iodide occurred at wavelengths of 488 nm, 543 nm, and 633 nm, respectively.

Co-immunoprecipitation. For Co-immunoprecipitation (Co-IP), HEK-293T cells were grown in 6 cm dishes and cotransfected with the indicated plasmids (3 µg each) using polyethylenimine (PEI). Cells were lysed under native conditions 24 h post transfection and IP assays were performed as described recently (7) using Protein A conjugated sepharose beads to pull down Ig-tagged proteins or α-flag M2 affinity gel to immunoprecipitate flag-tagged proteins.

Western blotting and antibodies. Cell lysates were subjected to sodium dodecylsulfate polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to polyvinylidene difluoride (PVDF) membrane (Immobilon-P; Millipore) using a semidry blotter (Peq-Lab). Membranes
were incubated overnight at 4°C with primary antibodies. Protein signals were visualized with horseradish peroxidase-conjugated secondary antibodies and an ECL kit (Perkin Elmer) and detected by applying a film (Amersham Hyperfilm ECL, GE Healthcare; Fig. 1B) or using the FusionFX7 (Vilber Lourmat). α-flag-M2 (Sigma), α-p65 (sc-109; Santa Cruz), α-p65 N-terminus (C22B4; Cell Signaling) and α-p50 (3035; Cell Signaling) were purchased and α-MV P/V (#37069), α-MV C, and α-RV P (P160-5) antibodies were kindly provided by D. Gerlier, Lyon, (11), R. Cattaneo, Rochester, and S. Finke, Greifswald-Insel Riems, respectively. α-MV V\textsubscript{CTD} is a polyclonal rabbit serum raised against a synthetic peptide conjugated to KLH corresponding to the C-terminal domain of V as described in (15).
Results

Suppression of TNFα-mediated NF-κB activation by measles virus P, V, and C proteins.

The canonical NF-κB activation pathway can be initiated by various stimuli, including tumor necrosis factor α (TNFα), a key cytokine regulating immune functions as well as inflammatory responses (18). In order to identify MV proteins influencing NF-κB activity, we performed dual luciferase reporter gene assays in cells treated with TNFα. Increasing amounts of expression vectors for the MV Schwarz proteins, including P, V, and C, were cotransfected in HEK-293T cells together with a plasmid encoding firefly luciferase reporter gene under the control of a trimeric repeat of the NF-κB-binding motif (PRDII) of the IFNβ promoter (50), and a Renilla luciferase expression plasmid. The start codon of the C protein was changed by site directed silent mutagenesis to prevent expression of MV C, in case of all P or V expressing plasmids, as described previously (38). The P protein of rabies virus (RV P), which suppresses activation of IRF3 and STAT1/STAT2 nuclear import, but which has no influence on NF-κB signaling (6, 7, 46), was used as a negative control. Cells were stimulated with TNFα 6 h prior to cell lysis and NF-κB-dependent luciferase activity was determined. Intriguingly, all three P gene products MV P, V, and C exerted a substantial and dose-dependent inhibitory effect on TNFα-mediated NF-κB activation. The V protein showed the strongest suppression (Fig. 1B). The inhibitory effects of P and the C protein were less effective. In contrast to V, P, and C, NF-κB-dependent luciferase activity was not affected in the presence of MV N, F, H, or L (not shown) or of RV P protein. These results indicate that MV P, V, and C proteins can interfere with canonical NF-κB activation, but to a different extent.
NF-κB activation by pattern recognition receptors is suppressed by MV P gene products.

Triggering of PRRs, like RLRs or TLRs can also lead to activation of NF-κB. To test the capacity of MV P gene products to downregulate NF-κB activation mediated by RIG-I signaling, a C-terminal deletion mutant ΔRIG-I (aa 1-284), which was previously shown to constitutively activate NF-κB (49), was co-transfected with increasing amounts of the MV proteins in HEK-293T cells, followed by dual luciferase assays. As a control, increasing amounts of a plasmid encoding RV P were transfected. Expression of ΔRIG-I led to more than 40 fold induction of NF-κB activity. In the presence of MV V at the highest dose, activity was suppressed to nearly basal level (Fig. 2A). MV P protein, which was expressed at much higher levels, as indicated by Western blotting using an antibody recognizing both P and V (Fig. 2A, bottom panel) had a less pronounced inhibitory potency. The C protein had intermediate capacity, while RV P had no significant influence on NF-κB activation.

Signal transduction by the receptors RIG-I and MDA5 is transmitted via IPS-1 (MAVS, VISA, Cardif), while that of TLRs involves the adaptors MyD88 (for TLR1,2,4,5,6,7,8,9) or TRIF (for TLR3) (29). To examine whether MV P, V, and C are able to generally counteract NF-κB activation by RLRs and TLRs, dual luciferase reporter gene assays involving overexpression of IPS-1 (Fig. 2B), MyD88 (Fig. 2C), and TRIF (Fig. 2D) for stimulation of NF-κB activity were performed. In analogy to the previous experiments, all MV P gene products were able to suppress the adaptor-induced NF-κB activation in a dose-dependent way, whereas expression of RV-P had no effect. The reduction of NF-κB activity achieved by the V and P proteins appeared to be similar in these experiments, however, Western Blotting revealed a substantially lower expression of the V protein (Figure 2B-2D, bottom panels), supporting the previous finding that the V protein is the most potent inhibitor of NF-κB activation among the MV P gene products.
MV V inhibits NF-κB signaling downstream of the IKK complex.

Stimulation of TLR, RLR, and TNFR triggers pathways leading to canonical NF-κB activation. These pathways converge on the IKK complex, which is composed of the kinases IKKα, IKKβ, and IKKγ (NEMO). To test whether the inhibition by MV proteins occurs at or downstream of these kinases, NF-κB-dependent luciferase expression was activated by overexpressing IKKα, IKKβ, and IKKγ in HEK-293T cells (Fig. 3A). As a positive control, an Ig-tagged NEMO-binding domain (NBD) was used (Ig-NBD). The NBD peptide binds to IKKγ and thereby inhibits the formation of the IKK complex which is essential for canonical NF-κB activation (34). Expression of Ig-NBD resulted in dose-dependent inhibition of IKK complex-induced NF-κB activity, while the Ig-tag expressed individually did not decrease NF-κB activation. Expression of MV V led to a dose-dependent and effective suppression of NF-κB activity, comparable to that achieved by expression of Ig-NBD (Fig. 3A), indicating a target at the level of, or downstream of the IKK complex. The presence of MV P and C proteins had less pronounced effects on NF-κB activation by the IKK complex.

To further spot the step where MV V inhibits canonical NF-κB activation, we induced NF-κB-dependent luciferase activity by co-expression of the NF-κB subunits p65 and p50 which build the main heterodimer of NF-κB. In the presence of MV V, the NF-κB activity induced by p65/p50 was reduced dose-dependently and significantly, whereas MV P and C showed only minor inhibitory capacities (Fig. 3B). As expected, co-expression of Ig-NBD had no effect on p65/p50 mediated NF-κB activity, as this inhibitor acts upstream of the transcription factor p65/p50. We conclude from these experiments that the V protein of MV can inhibit canonical NF-κB signaling downstream of the IKK complex, while P and C may act upstream in the signal transduction cascade.
MV V binds the NF-κB subunit p65.

In order to clarify the molecular mechanism of MV proteins to suppress NF-κB activation, we analyzed the NF-κB subunits p65 and p50 for potential interactions with viral proteins in co-immunoprecipitation (CoIP) experiments. Extracts from HEK-293T cells coexpressing Ig-tagged MV P, V, or C proteins and p65 from transfected plasmids were purified by protein A-conjugated sepharose beads and precipitates were analyzed by Western blotting using antibodies against p65 or human IgG. Indeed, the NF-κB subunit p65 was specifically co-precipitated with Ig-MV V, whereas no interaction of p65 with the P or C constructs was detectable (Fig. 4A).

In further experiments, binding of p50 (NF-κB1) to flag-tagged MV proteins was assessed. Flag-p65 was included as positive control. Cell extracts were analyzed by immunoblotting using antibodies specific for p50 and flag. While flag-p65 efficiently co-precipitated p50, no interactions of the p50 NF-κB subunit with any of the flag-MV proteins could be demonstrated (Fig. 4B). Taken together, these experiments revealed that the V protein of MV specifically interacts with the NF-κB subunit p65, but not with p50.

To determine if MV V binds to the N-terminal RHD of p65, which is responsible for dimerization, DNA-binding, and nuclear import, we constructed the fragment of p65 spanning from aa 1-309 (RHD p65) and performed Co-IP experiments with flag-MV P, V, C. As revealed by Western blotting with an antibody specific for the N-terminus of p65, the p65 RHD was efficiently pulled down by flag-MV V (Fig. 4C). Notably, RHD p65 also showed some affinity to MV P, though considerably weaker than to MV V. A minor affinity of the MV P protein to bind p65 was observed occasionally in pull down experiments with flag tagged p65 and authentic, untagged MV proteins (data not shown). Taken together, we observed that MV V binds with a strong affinity to the RHD of p65, while a weak interaction of MV P with RHD p65 was suggested.
To further verify the specificity of MV V to the RHD of p65 we tested binding of flag-tagged p65, RHD p65, p50 and RHD p50 to V within one experiment. Therefore, we constructed the RHD of p50 spanning from aa 1-366 and the other constructs with a flag tag. CoIP experiments of the flag tagged proteins with authentic, untagged MV V revealed specific binding of V to flag-p65 and flag-RHD p65, while no interaction of V and flag-p50 or flag-RHD p50 could be detected (Fig. 4D). These findings confirm the specificity of MV V to the RHD of p65.

V prevents nuclear translocation of p65.

Since MV V is a cytoplasmic protein, binding of V to p65 might interfere with trafficking of this NF-κB subunit. In order to address this hypothesis, HEp2 cells were transfected with a flag-MV V encoding plasmid, or empty vector, and stimulated with TNFα for 30 min, followed by immunostaining of p65 and flag-tagged MV V. Whereas in unstimulated cells transfected with empty vector p65 is located predominantly in the cytoplasm (Fig. 5, upper panel), TNFα treatment resulted in almost complete translocation of p65 from the cytoplasm to the nucleus (Fig. 5, middle panel). In contrast, accumulation of p65 in the nucleus upon TNFα stimulation in cells expressing flag-MV V was severely impaired (Fig. 5; lower panel; closed arrows), while cells failing to express detectable flag-MV V showed a normal nuclear accumulation of p65 (Fig. 5B; lower panel; open arrows). Thus, binding of the MV V protein to p65 is sufficient to prevent the nuclear accumulation of NF-κB and thereby to preclude its transcriptional activity.
The CTD of MV V is required and sufficient for p65 binding and suppression of p65/p50-mediated NF-κB activity.

Since MV P and V have identical amino-terminal domains (PV_{NTD}), but distinct carboxy-terminal domains (P_{CTD}; V_{CTD}) (see Fig. 1A), we reasoned that p65 binding is mediated via the V-specific CTD. To verify this, HEK-293T cells were transfected with expression plasmids encoding the individual protein domains fused to an Ig-tag (Ig-MV PV_{NTD}, P_{CTD}, V_{CTD}), or the full length proteins, together with p65. Indeed, Ig-MV V_{CTD} was sufficient for precipitation of p65 with a binding affinity apparently similar to that of full length Ig-MV V, while the other constructs did not reveal interaction with the NF-κB subunit (Fig. 6A). To clarify whether binding of the small V_{CTD} is also sufficient for inhibition of NF-κB transcriptional activity, p65 and p50 were overexpressed in HEK-293T cells along with Ig-MV V_{CTD} or Ig-MV V. A similar and dose-dependent reduction of NF-κB-dependent luciferase expression confirmed that binding of V_{CTD} is sufficient for full inhibition (Fig. 6B).

In contrast, expression of the C-terminal portion of the P protein had no considerable effect on p65/p50 mediated luciferase activity. In summary, the C-terminal domain of the V Protein is sufficient for mediating p65 binding and inhibition of p65/p50 mediated NF-κB activity.
Discussion

NF-κB is a key mediator of antiviral host response and inflammation, as well as of immune cell development, survival, and function (24), and therefore a prime candidate for viral interference (26). In particular, hematotropic viruses like the immune-suppressive and immune-modulatory measles virus (22) should have means to interfere with NF-κB signaling. In fact, recent work showed upregulation of the ubiquitin modifying enzyme A20 in monocytes, but not in epithelial cells, infected with MV, or expressing the MV P protein (48), indicating that MV has at least indirect means to affect NF-κB signaling. Our present data revealed that all of the MV Schwarz P gene products, including the essential P protein, and the “accessory” proteins V and C, which are established MV virulence factors, are able to interfere with NF-κB activation. The V protein, in particular, revealed a potent inhibitory capacity. This is correlated with specific binding of V to the central NF-κB subunit, p65, and therefore a lack of p65 nuclear accumulation. P and V of a MV wild type isolate (genotype D5) bear some point mutations in their common PV_{NTD} domain, while the unique P_{CTD} have two conservative aa exchanges compared to the Schwarz strain. V_{CTD}, however, which was shown to be responsible for p65 binding, is completely conserved in the wild type isolate and the Schwarz strain. Consistently with this fact, binding of wild type V to p65 and similar inhibition of reporter gene activity was also observed (data not shown). The C protein of the D5 strain shows only some aa exchanges, that seem to have no effect on suppression of NF-κB activity, since all wild type P gene products showed the same suppression pattern like MV Schwarz proteins in the NF-κB-dependent luciferase assays (data not shown). Thus, the capacity of MV P gene products seems not to be affected in the vaccine strain compared to wild type.

Expression of the individual MV proteins P, V, and C, was sufficient to suppress NF-κB-mediated reporter gene transcription triggered through different signaling cascades, including
TNFR, RLR, and TLR. Importantly, upregulation of the NF-κB inhibitor A20 was not observed in HEK-293T cells with any of the MV proteins (not shown), excluding a contribution of this recently described mechanism (48). In any of the pathways investigated, the specific inhibitory capacity of V was greater than that of P or C, while it appeared to be particularly pronounced in cells stimulated by overexpression of ∆RIG-I. Since RIG-I is thought to be the main sensor of paramyxovirus infection (28, 39), interference with RLR signaling cascades is a promising mechanism to evade host immune responses. MV V is known to strongly bind to the helicase domain of MDA5, but not RIG-I (1, 12, 13). In addition, genetic knock-out of MDA5 was previously shown to reduce RIG-I-mediated IFN induction in transgenic mice (20), indicating that MDA5 is synergistic to RIG-I signaling. Although ∆RIG-I consists only of the RIG-I CARD domains, a potential ∆RIG-I/MDA5 interplay, which is disturbed by V binding, cannot be formally excluded and might contribute to the potent inhibition of ∆RIG-I-mediated NF-κB activity by V. However, we suggest that binding of MV V to the downstream transcription factor p65 is the major mechanism for suppression of RLR-mediated NF-κB activation.

Signaling cascades initiated by TNFR, TLR, and RLR, or their respective adaptor proteins MyD88, TRIF, and IPS-1, converge on the IKK complex which controls the phosphorylation-dependent proteasomal degradation of the inhibitor IκB and therefore is the central regulator of the canonical NF-κB pathway (23, 27). The finding that MV V is able to suppress not only IKK-mediated NF-κB activity but also the activity of overexpressed NF-κB (p65/p50) revealed a universal, downstream inhibitory mechanism of the V protein (Fig. 3B).

Inspired by the fact that MV V and P are cytoplasmic proteins and interfere with import of STAT1 and STAT2 to the nucleus (9, 16, 36), we performed co-immunoprecipitation assays revealing specific binding of V to p65 (Fig. 4) and immunofluorescence assays indicating V-mediated retention of p65 in the cytoplasm (Fig. 5). We found that binding of MV V to p65 is
mediated through the Rel homology domain (RHD) of this NF-κB subunit, while no interaction of with the RHD of p50 is detected. The N-terminal RHD is characteristic for all NF-κB subunits (19) and is responsible for homo- and heterodimerization of the NF-κB proteins, the binding to IκBs as well as sequence specific DNA binding. The RHD also contains the nuclear localization signal (NLS) which is masked by the IκBs in nonstimulated cells. Upon stimulation of the NF-κB signaling cascade IκB is degraded and the NLS is liberated which results in the nuclear translocation of NF-κB. We propose that binding of V to the RHD domain of p65 shields the NLS of the NF-κB subunit such that translocation of NF-κB into the nucleus is impaired, therefore suggesting an IκB-like function for the MV V protein.

We recently described binding of MV V to IKKα (38), which is involved in the activation of IRF7 through TLR7/8/9, but is also a subunit of the canonical IKK complex. We could show in vitro kinase assays, that the IKKα-dependent phosphorylation of IRF7 was diminished in the presence of MV V, whereas the phosphorylation of the NF-κB inhibitor IκBα by IKKα was not altered (38). Furthermore, activation of NF-κB by overexpression of either IKKα or IKKβ was inhibited equally by MV V (not shown). Therefore, we suggest that binding of MV V to IKKα suppresses IRF7 activation but does not affect activation of NF-κB, whereas binding of V to p65 interferes with NF-κB signaling. This is also emphasized by the fact that IKKα plays only a minor role in the canonical NF-κB pathway, since IKKα may support canonical NF-κB activation, but is dispensable, whereas IKKβ is the essential kinase (27).

Intriguingly, the 68 aa short C-terminal domain of the V protein (V_{CTD}) was found responsible and sufficient for specific binding of V to the p65 RHD and inhibition of canonical NF-κB activation (Fig. 6). The structure of the cysteine rich and zinc-coordinated V_{CTD} is the conserved part of Paramyxovirinae V proteins and responsible for most of the host antagonistic functions described for V so far (21). Therefore, it is not surprising that V_{CTD} of
different *Paramyxovirinae* family members have common functions and binding partners, such as STAT2 (42, 43, 45) and MDA5 (1, 12, 37). However, it appears that the V<sub>CTD</sub> of *Paramyxovirinae* can further adapt to different targets, according to their requirements and niches. While V<sub>CTD</sub> of the respiratory pathogen parainfluenza virus type 5 (PIV5) and of related rubulaviruses was found to bind IKKε, and thereby prevents IRF3 activation and RLR-dependent IFNβ induction (33), we could recently show that the MV V<sub>CTD</sub> instead binds IKKα and IRF7 and thereby prevents TLR7/8/9-mediated IFNα induction, which is instrumental in hematopoietic cells like pDC (38). Similarly, a general inhibition of canonical NF-κB activation due to binding to p65 as observed here for MV seems not to be a common feature of *Paramyxovirinae* V proteins. Though the V protein of PIV5 was reported to suppress NF-κB activation upon triggering with synthetic dsRNA or due to viral infection, inhibition of LPS- or TNFα-dependent NF-κB activity was not observed (32, 40). This is in accordance with our own NF-κB reporter gene experiments where expression of the PIV5 V was ineffective in preventing TNFα-mediated NF-κB activity (data not shown). Further experiments should reveal whether NF-κB p65 targeting by V is specific for the human MV or also for the related animal morbilliviruses.

As indicated in reporter gene experiments, in addition to MV V, both MV P and C interfered with canonical NF-κB signaling, though mostly less prominent. However, neither the C nor the P protein of MV did reveal a pronounced interaction with p65. We therefore presume that the P and C proteins contribute to MV mediated NF-κB escape by targeting other steps of the canonical activation pathways which remain to be elucidated. In case of P, the PV<sub>NTD</sub> appears to be required for counteracting NF-κB signaling in epithelial cells, since the P<sub>CTD</sub> displayed no inhibition of TNFα-mediated NF-κB reporter gene activation (data not shown). Notably, the PV<sub>NTD</sub> of P and V is also involved in association with STAT1 and thereby contributes to inhibition of IFN signaling (9, 42). In MV-infected monocytes, P may in addition lead to
upregulation of the NF-κB inhibitor A20 (48). As far as C is concerned, an explanation is not close at hand. C proteins of other Paramyxovirinae family members show also NF-κB inhibitory capacity, as illustrated by the Sendai virus C proteins which suppress dsRNA- and Newcastle disease virus-mediated NF-κB activation, however, the mechanism is also elusive (30).

In summary, we demonstrated that measles virus applies multiple mechanisms to manipulate NF-κB signaling pathways. The major activity could be attributed to the V protein, and more specifically to the V\textsubscript{CTD}, which interferes with canonical NF-κB activation by binding to the RHD of the NF-κB subunit p65 and therefore prevents nuclear accumulation of the transcriptionally active NF-κB subunit. The V protein is a well established virulence factor and the V\textsubscript{CTD} turns out to be a hub for specific binding of numerous cellular proteins. This includes not only targets of innate immunity but also proteins related to proliferation and cell death like the p53 family member p73, which down regulates expression of the proapoptotic target gene PUMA and might therefore function as a viral antiapoptotic factor (14). Revealing the exact binding sites for specific proteins as recently shown for MDA5 and STAT2 (41, 42) and generation of recombinant MV deficient only in single functions of V should help in elucidating the contributions of individual V functions to measles virus cell biology and immune modulation.

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References


Figure legends

Figure 1: Suppression of TNFα-mediated NF-κB activation by measles virus P, V, and C proteins.

(A) The P gene of MV encodes the P protein and the nonstructural proteins V and C. The V mRNA is generated by insertion of an additional guanosine between nucleotides 751 and 752 of the mRNA by RNA editing. Therefore, the MV P and V proteins share a common amino-terminal domain (PV_{NTD}) stretching from aa 1 to 231, but have distinct carboxy-terminal domains (P_{CTD}; V_{CTD}). The C protein is produced by translation of an alternative ORF initiated 19 nucleotides downstream of the P/V start codon.

(B) Increasing amounts (200 ng, 400 ng, 600 ng) of expression plasmids encoding for measles virus (MV) proteins (P, V, C), rabies virus (RV) P protein or empty vector (EV) were cotransfected into HEK-293T cells with the NF-κB-dependent reporter plasmid p55A2-luc and pRL-CMV for normalization. After 18 h cells were stimulated with 10 ng/ml recombinant human TNFα and incubated for additional 6 h followed by cell lysis. NF-κB-driven luciferase activity was determined by dual luciferase assay. Values given are average and standard deviation of two independent experiments. Depicted is a representative experiment out of four repeats. Lower panel: Cell lysates were subjected to SDS-PAGE and separated proteins were probed with α-MV P/V, α-MV C or α-RV P antibodies by Western blotting to determine the expression levels.
Figure 2: NF-κB activation by pattern recognition receptors is suppressed by MV P gene products.

Increasing amounts (200 ng, 400 ng, 600 ng) of vectors encoding for the indicated measles virus (MV) proteins, rabies virus (RV) P protein or empty vector (EV) were cotransfected into HEK-293T cells with either 200 ng of expression plasmids for ΔRIG-I (A), MyD88 (B), IPS-1(C) or TRIF (D) and the NF-κB-dependent reporter system (100 ng p55A2/ 10 ng pRL-CMV). After 24 h cells were lysed and NF-κB activity was determined by dual luciferase assay. Values given are average and standard deviation of two independent experiments. Depicted are representative experiments out of three repeats. Lower panels: Expression of the viral proteins was assessed by Western blotting.

Figure 3: MV V inhibits NF-κB signaling downstream of the IKK complex.

(A) HEK-293T cells were cotransfected with expression plasmids encoding for IKKα, IKKβ, and IKKγ (100 ng each) together with increasing amounts (200 ng, 400 ng, 600 ng) of vectors for the indicated proteins or empty vector (EV) and the NF-κB-dependent reporter system (100 ng p55A2/ 10 ng pRL-CMV). After 12 h the cells were lysed and the luciferase activity was measured by dual luciferase assay. Values given are average and standard deviation of two independent experiments. A representative experiment out of three repeats is shown. Lower panel: Expression levels were determined by Western blotting.

(B) Plasmids encoding for p65 and p50 (150 ng each) and increasing amounts (300 ng, 600 ng) of vectors for the indicated proteins or empty vector (EV) were cotransfected into HEK-293T cells together with dual luciferase reporter system. Cells were lysed 12 h after transfection followed by determination of normalized NF-κB-dependent luciferase activity. The given values are average and standard deviation of two independent experiments. Shown
is a representative experiment out of three repeats. Lower panel: Expression levels of viral proteins were assessed by Western blotting.

**Figure 4: MV V binds the NF-κB subunit p65.**

(A) HEK-293T cells were used to express p65 in combination with the indicated Ig-tagged proteins or the Ig-tag (Ig) itself (3 µg each). After 24 h cells were lysed under native conditions and Ig-tagged proteins were pulled down using protein A conjugated sepharose beads. Binding of p65 to measles proteins was visualized by Western blotting. Depicted is a representative experiment out of four repeats.

(B) p50 was coexpressed in HEK-293T cells with the indicated flag proteins or empty vector (EV) (3 µg each). Cells were lysed 24 h post transfection and flag-tagged measles proteins were immunoprecipitated using anti-flag M2 affinity gel. Interaction of p50 and flag-proteins were analyzed by Western blotting. A representative experiment out of three is shown.

(C) HEK-293T cells were cotransfected with vectors encoding for the indicated flag-tagged constructs or empty vector (EV) and the RHD of p65 (aa 1-309). CoIP assay was performed as described above and RHD-p65 was stained using α-p65 (Cell Signaling; 3035). A representative experiment out of three is shown.

(D) MV V was coexpressed with the indicated flag tagged proteins or empty vector (EV) (3µg each) in HEK-293T cells. CoIP experiments were performed as described above and MV V was stained using α-MV VCTD. Depicted is a representative experiment out of three repeats.
Figure 5: V prevents nuclear translocation of p65.

HEp2 cells were transfected with a vector for flag-MV V or empty vector. 24 h post transfection cells were either treated with 10 ng/ml TNFα for 30 min (+) or left untreated (-). Subsequently, cells were fixed and stained with the indicated antibodies. Green: p65 (stained with specific antibody); red: flag-MV V (stained with α-FLAG® M2); blue: ToPro3 nuclear staining; open arrow: not flag-MV V expressing cell; closed arrow: flag-MV V expressing cell. Depicted is a representative experiment out of three repeats.

Figure 6: The CTD of MV V is required and sufficient for p65 binding and suppression of p65/p50-mediated NF-κB activity.

(A) HEK-293T cells were cotransfected with vectors encoding for the indicated Ig-tagged constructs or the Ig tag itself (Ig) and p65. CoIP assay was performed as described above. A representative experiment out of four is shown.

(B) Increasing amounts (300 ng, 600 ng) of the indicated Ig-tagged constructs were coexpressed in HEK-293T cells together with p65, p50 (150 ng each) and the NF-κB reporter system (100 ng p55A2/ 10 ng pRL-CMV). 12 h post transfection cells were lysed and NF-κB activity was determined by dual luciferase assay. Values given are average and standard deviation of two independent experiments. Depicted is a representative experiment out of three repeats. Lower panel: Expression levels were determined by Western blotting.