INDUCIBLE EXPRESSION OF THE 2-5A SYNTHETASE / RNase L SYSTEM RESULTS IN INHIBITION OF VACCINIA VIRUS REPLICATION

Margarita Díaz-Guerra,¹ Carmen Rivas and Mariano Esteban *

Centro Nacional de Biotecnología, CSIC, Campus Universidad Autónoma, Madrid-28049, Spain

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¹CURRENT ADDRESS: Departamento de Bioquímica (UAM)-Instituto de Investigaciones Biomédicas (CSIC). C/ Arturo Duperier, 4, 28029- Madrid. Spain.

CORRESPONDENCE FOOTNOTE: Mariano Esteban. Centro Nacional de Biotecnología, CSIC, Campus Universidad Autónoma, Madrid-28049, Spain.

Telephone: 341 5854503; FAX: 341 5854506

E-mail: MESTEBAN@SAMBA.CNB.UAM.ES
ABSTRACT

Vaccinia virus (VV) has developed strategies to overcome the action of interferons (IFN), but the mechanisms involved in the differential sensitivity of this virus to IFN have not been elucidated. To investigate whether the IFN-induced enzymes 2-5A-synthetase and 2-5A-dependent RNase L could be responsible for inhibition of VV replication, we have generated recombinant vaccinia viruses containing the corresponding genes (VV-2-5AS and VV-RL, respectively). RNase L produced in cells infected with VV-RL leads to rRNA degradation and inhibition of virus protein synthesis, which correlates with about 92 percent reduction in virus yields by 48h of infection. Combined expression of this enzyme with 2-5A-synthetase further inhibits virus yields. The pattern of rRNA fragments produced by infection with viruses VV-RL and / or VV-2-5AS is the characteristic for activation of the 2-5A pathway by IFN treatment. Combined infection of VV-RL together with vesicular stomatitis virus (VSV) demonstrates inhibition to be specific for VV and not a general effect. Degradation of rRNA is only slightly increased by IFN treatment of the cells previous to VV-RL infection and PKR is not required for induction by RNase L of protein synthesis inhibition. Thus, our findings provide direct evidence for antiviral activity of the 2-5A system on poxviruses and show that it is possible to overcome VV-induced IFN-resistance mechanisms by increasing the levels of enzymes in this pathway in cultured cells, particularly those of RNase L.
Interferons (IFN) are cytokines released from animal cells in response to virus infection or to a variety of other stimuli. By binding to specific cell surface receptors IFN exerts a wide range of functions, including inhibition of virus multiplication (1, 2). The varying effects of IFN are mediated by more than 30 different proteins which are induced after IFN treatment of the cells (1-3) and only the role of a few of them in the antiviral state is beginning to be elucidated. The best characterized IFN-induced proteins are the Mx family of proteins, the two double-stranded (ds) RNA-dependent enzymes, a 68 kDa protein kinase (PKR) and 2-5A-synthetase, and the 2-5A-dependent RNase (RNase L) (3, 4). The proteins 2-5A-synthetase and RNase L constitute, together with 2-5A-phosphodiesterase, what is known as 2-5A pathway or 2-5A system, an IFN regulated RNA degradative pathway (5). Although basal levels of these proteins are found in most, if not all, mammalian cells, treatment with IFN induces at least four different forms of 2-5A-synthetase and a unique RNase L (reviewed in reference 3). Double-stranded RNA activates the 2-5A-synthetases that, in the presence of ATP, synthesize a complex mixture (referred to as 2-5A) of 5'-triphosphorylated oligoadenylic acid, ppp(A2’p5)nA, that binds to and activates endoribonuclease RNase L (2). Activated 2-5A-dependent RNase L cleaves viral and cellular RNAs, with the result of general inhibition of protein synthesis (2).

Characterization of the biological role of the 2-5A system has advanced by cloning of murine and human RNase L genes (6) which encode proteins of very interesting molecular architecture (7, 8). Human RNase L (83.5 kDa) contains 9 ankyrin-like repeats at the N terminus which are immediately followed by a complete protein kinase domain and next to it, in the C terminus, the residues where the endoribonuclease activity is located. Ankyrin repeats have been involved in mediating intra- and intermolecular protein-protein interactions in many different types of proteins (9). Systematic biochemical studies of IFN-treated virus-infected cells has only revealed the 2-5A system as responsible for virus inhibition in the case of picornaviridae. Direct evidence for antiviral activity of this pathway was shown by constitutive expression of a cDNA encoding human 40 kDa 2-5A-synthetase (10-12). The inhibitory effect is selective for mengovirus and encephalomyocarditis (EMC) virus with little effect on either vesicular stomatitis virus (VSV) or herpes simplex 2 (HSV-2) (10-12). Inhibition of picornaviridae by IFN
also requires functional RNase L as has been demonstrated in a stable cell line expressing high levels of a truncated RNase L which is a dominant negative mutant of this enzyme (7). However, the contribution of the 2-5A system to the antiviral activities of IFN almost certainly goes beyond the picornaviridae family of viruses and is obscured by virus-induced IFN-resistance mechanisms (reviewed in references 2 and 3). There is evidence suggesting that this is probably the case for vaccinia virus (VV), a poxvirus which replicates in the cytoplasm of a wide range of cells, and that has been found to be resistant to IFN in many cultured cell lines (13, 14) while sensitive in others (15) and in experimental animals (16). The reason for the differential sensitivity of VV to IFN in vivo and in vitro might be related to the ability of the virus to synthesize interfering products. Indeed, VV displays an impressive variety of defensive strategies to evade the host response to infection, including counteraction of IFN by strategies at two different levels (reviewed in reference 17): production of soluble type I and II IFN receptors which blocks IFN action at a very early stage (18, 19) and modulation of the activity of the IFN-induced proteins. The product of ORF E3L competitively binds ds RNA and prevents the activation of the IFN-induced ds RNA-activated PKR (20) and probably also 2-5A-synthetase (21). Down regulation of the activation of p68 kinase is also attained by the product of ORF K3L, a protein with sequence similarity to eukaryotic initiation factor 2α (eIF2α), that competitively binds PKR and therefore blocks inactivation by phosphorylation of eIF2α (22). Besides these proteins, there are results suggesting that VV uses additional mechanisms to evade IFN action. The product of ORF A18R, a mediator in virus RNA metabolism (23), and NPH-I, an enzyme involved in virus transcription (24), could be modulating the 2-5A system but if this happens by a direct or indirect mechanism remains to be determined. Finally, there are also results suggesting VV induced ATPase and phosphatase activities being involved in viral interference with the 2-5A system (25, 26).

In this investigation we have examined whether the IFN-induced enzymes 2-5A-synthetase and 2-5A-dependent RNase L could be responsible for inhibition of VV replication in situations of virus sensitivity to IFN. To this aim we have generated VV recombinants able to express individually each one of the IFN-induced enzymes in the 2-5A pathway and analyzed the effect of their expression in VV replication using a cell line where the virus is otherwise mainly
resistant to IFN action. A recombinant virus where expression of RNase L can be regulated was obtained by cloning of the human gene, contained in plasmid ZC-5 (6), under the control of the bacteriophage T7 promoter into VV insertion vector pTM1-E (27) (Fig.1A). We first generated plasmid pPR35-RL by insertion of a Hind III fragment of ZC-5 into the Sma I site of pPR35 (28). Plasmid pRSET-RL for expression in E.coli was then obtained by cloning of a Pst I + Kpn I fragment of pPR35-RL into pRSET-B digested in the same way. It contains the coding sequence for RNase L, lacking only nucleotides coding for the 21 amino acids at the N terminus, fused to a sequence for 38 new amino acids that includes a stretch of 6 histidine residues and the 11 amino acid gene 10 leader peptide. We also generated plasmid pRSET-ΔN by Nco I digestion of pRSET-RL and religation. This plasmid is similar to pRSET-RL but lacking sequences coding for amino acids 228 to 741 of RNase L. Insertion plasmids pTM-RL and pTM-ΔN were then obtained by cloning DNA fragments from plasmids pRSET-RL (Nde I) and pRSET-ΔN (Nde I + Nco I), respectively, into the Sma I site of VV insertion vector pTM1-E (27) and they contain 8 extra amino acids in the N terminus in front of the initiation codon provided by pRSET-B. Recombinant viruses VV-RL and VV-ΔN were generated by homologous recombination of plasmids pTM-RL and pTM-ΔN into thymidine kinase (tk) gene of wild type vaccinia virus (WR) and selection in human TK− 143B cells with 5'-bromodeoxyuridine (25μg/ml) basically as described (29). Two rounds of plaque purification yielded homogeneous virus preparations containing the desired genes, as confirmed by Southern blot DNA hybridization analysis (data not shown). We also prepared a rabbit polyclonal anti-RNase L serum able to recognize proteins produced by these recombinant viruses, using the C-terminus of this protein (amino acids 340 to 741) expressed in E.coli as the antigen. Exponentially growing E.coli BL21 (DE3) cells carrying plasmid pRSET-Δ2S, generated by Sac I digestion of pRSET-RL and religation, were induced with 1 mM isopropyl-β-D-thiogalactopyranoside. The approximately 45-kDa induced protein, containing a polyhistidine metal-binding domain, was purified by using a nickel-charged sepharose resin (Invitrogen) in denaturing conditions. Binding was performed in 6 M guanidine hydrochloride, 0.1 M Na-phosphate, 0.01 M Tris/HCl, pH 8.0, 10 mM imidazole and elution in 6 M urea, 0.5 M NaCl, 20 mM Na-phosphate pH 7.8, 0.5 M imidazole. The protein was precipitated with TCA, the pellet thoroughly washed in acetone and, once dried, suspended in PBS and used as the
antigen for injection into New Zealand white rabbits. Cell extracts obtained from mock-infected monkey BSC-40 cells or from cells infected for 24h with VV-RL or VV-ΔN were therefore analyzed by immunoblotting using this sera (Fig. 1B). Proteins of sizes expected for RNase L (lane 5) and C terminus deleted RNase L (RNase L-ΔN) (lane 6) were only obtained by coinfection with a second recombinant virus that contains a T7 polymerase gene under the control of virus constitutive promoter p7.5 (vT7F7-3, herein vT7) (30). It is noticeable that for RNase L-ΔN, besides a main band of the expected molecular weight for the truncated protein, there are accompanying products of higher and lower molecular weights. Similar results were obtained using T7-tag, a commercial monoclonal antibody directed against the gene 10 leader peptide in the N terminus (data not shown). These bands might reflect complex conformation of RNase L-ΔN due to intramolecular interaction of the ankyrin repeats in the protein. Endogenous RNase L could not be detected in mock-infected BCS-40 cells (Fig. 1B, lane 1), probably because our polyclonal antibodies prepared against the human protein are not able to recognize monkey RNase L.

Late during VV infection of cells in culture, concomitant with synthesis of complementary mRNAs due to symmetrical transcription of late genes, 2-5A is produced at high levels (14, 26) and although is biologically active when assayed in vitro is not able to inhibit VV replication in cultured cells (14). The resistance of VV to the 2-5A system is probably due to the existence of viral products that directly or indirectly counteract the activation of endogenous RNase L. The levels of activatable RNase L and of 2-5A might be critical in the sensitivity / resistance phenomenon of VV to the 2-5A system. To address this hypothesis, we measured RNA stability in BSC-40 cells infected with vT7 and VV-RL at different times of infection. Total RNA was extracted from mock-infected cells or from cells infected either with vT7 (m.o.i. 4) or doubly infected with vT7 + VV-ΔN or vT7 + VV-RL (m.o.i. 2 each virus) for the indicated periods of time (Fig. 2A) and was analyzed by electrophoresis on formaldehyde agarose gels. Interestingly, in cells doubly infected with vT7 + VV-RL for 24h, expression of RNase L induces breakdown of RNA as seen by rRNA cleavage (lane 5). Cleavage of rRNA required the expression of RNase L and was not observed in cells infected with vT7 (lane 2) and only to a very low extent in cells expressing RNase L-ΔN (lane 7). Because recombinant RNase L
produced in insect cells has been shown to dimerize upon binding to 2-5A (31), we cannot exclude that RNase L-ΔN expression might cause minimum activation of endogenous RNase L. Time-course experiments with vT7 + VV-RL showed that rRNA cleavage starts at about 14h after infection (lane 4), coincident with the time when RNase L is first detected by immunoblotting with sera against RNase L (Fig. 2B). Discrete rRNA fragments characteristic of activation of the 2-5A / RNase L pathway (32, 33) are detected up to 24h of infection (Fig. 2A, lanes 4 and 5). Very little rRNA is detected by 48h (Fig. 2A, lane 6) suggesting further degradation of RNA fragments first produced by this or other RNases. Clearly, the results of Fig. 2 demonstrate that RNase L induced from a recombinant VV causes extensive rRNA degradation, in spite of the synthesis of viral interfering products.

Degradation of RNA observed in cells that express RNase L might have as a consequence the inhibition of virus protein synthesis. Thus, we performed a time-course experiment where, due to shut-off of cellular protein synthesis as a consequence of infection, we measured de novo synthesis of viral proteins (Fig. 2C). Monolayers of BSC-40 cells were either mock-infected (lane 1) or infected with vT7 + VV-RL (lanes 2-5) or vT7 + VV-ΔN (lanes 6-9) using the same multiplicity for the indicated times. Then, cells were pulse-labeled with 35S-methionine+cysteine (100 μCi/ml) in methionine and cysteine-free media for 15 min. After washing with PBS, cells were collected in lysis buffer, protein extracts were fractionated in SDS-polyacrylamide gels that once dried gels were autoradiographed. Inhibition of virus protein synthesis by RNase L is clearly observed starting at about 14h after infection (compare lanes 3 and 7) when de novo protein synthesis is maximum (compare lanes 6-9). After 24h of infection, low protein synthesis activity is found in control infections (lanes 8 and 9) but inhibition by RNase L can still be observed (lanes 4 and 5). A very abundant protein of size expected for RNase L-ΔN is produced in cells infected with vT7 + VV-ΔN, while levels of full-length RNase L synthesized are probably very low (compare lanes 7 and 3). These findings provide direct evidence that degradation of RNA induced throughout expression of RNase L by a recombinant VV results in inhibition of virus protein synthesis.
We next examined if by increasing the levels of 2-5A through expression of 2-5A-synthetase in combination with RNase L, further enhances the inhibition of VV protein synthesis and what is the net effect in viral replication. To this aim we generated VV recombinant VV-2-5AS where expression of human 40-kDa 2-5A-synthetase is driven by the virus constitutive early-late promoter p7.5. The recombinant was obtained by homologous recombination of plasmid pSC-2-5AS into wild type virus (WR) thymidine kinase (tk) gene basically as described (29). The DNA was obtained by cloning of a 1.2 kb fragment from plasmid pTL4-2-5AS (Nco I + Xho I digested and ends filled with Klenow fragment) into the Sma I site of VV insertion vector pSC 11 (34). It contains the complete human 40-kDa 2-5A-synthetase coding sequence lacking methionine residue in the N terminus and the \textit{E.coli} \( \beta \)-galactosidase gene (lac \( z \)) under the control of a late viral promoter (p11). Recombinant viruses were selected in human TK-143B cells using X gal and 5'-bromodeoxyuridine (25\( \mu \)g/ml) (29) and two rounds of plaque purification yielded homogeneous virus preparations. Monkey BSC-40 cells were then infected with different combinations of vT7, VV-RL, VV-\( \Delta \)N or VV-2-5AS using 5 pfu/cell of each virus but maintaining all along the experiment a total multiplicity of 15 pfu/cell by using an auxiliary recombinant virus VV-LUC (35), as other recombinants used in this work a tk(-) virus. We have compared the stability of rRNA at two different times of infection (24 and 48h) with levels of expression of the recombinant proteins (RNase L and 2-5A-synthetase), with \textit{de novo} synthesis of virus proteins and with virus yields. As shown, cleavage of rRNA is observed at 24h of infection with vT7 + VV-RL (Fig. 3A, lane 5) but degradation increases upon addition of VV-2-5AS (compare lanes 7 to 5). Control experiments using VV-\( \Delta \)N instead of VV-RL demonstrate that lack of RNA stability is due to expression of nearly full-length RNase L (compare lanes 6 and 8 to 5 and 7, respectively). At 48h, in cells infected with vT7 + VV-RL or vT7 + VV-RL + VV-2-5AS, rRNA degradation progresses (Fig. 3A, compare lanes 5 and 7 in upper and lower panels) and for triple infections most rRNA is degraded by this time of infection. Interestingly, expression of 2-5A-synthetase by itself induces some rRNA cleavage at 48h of infection (lane 2). This is probably due to low levels of activation of endogenous RNase L by the high levels of 2-5A oligos produced by 2-5A-synthetase expressed by the recombinant virus. Activation of endogenous RNase L might be also the cause of low levels of rRNA degradation observed in triple infection vT7 + VV-\( \Delta \)N + VV-2-5AS at 48h of infection (lane 8). Noteworthy, the pattern
of rRNA fragments produced by infection with recombinant viruses expressing RNase L and / or 2-5A-synthetase is similar to the one obtained in cells infected for 24h with wild type virus (m.o.i. 15) in the presence of isatin-β-thiosemicarbazone (IBT) (15 µM) added to the culture after viral adsorption and maintained for the duration of the experiment (23) (Fig. 3A, upper panel, lane VV-(IBT)). This is an anti-poxviral drug which causes a significant increase in the amount of ds RNA synthesized during VV infection, therefore inducing cleavage of rRNA with a pattern that is the characteristic one for activation of the 2-5A pathway (36). The levels of recombinant proteins RNase L and 2-5A-synthetase were analyzed by using specific rabbit polyclonal sera and the results obtained at 48h of infection are presented in Fig. 3B. Cleavage of rRNA correlates in all cases with expression of RNase L and / or 2-5A-synthetase (compare lower panel in Fig. 3A and Fig. 3B). Samples presenting the highest levels of rRNA degradation also show clear signs of protein synthesis inhibition with amounts of RNase L and 2-5A-synthetase being, respectively, moderate and severely inhibited. Analysis of de novo virus protein synthesis at 24h of infection (Fig. 3C) showed very low levels of protein synthesis activity in cells infected with vT7 + VV-RL that are further reduced by co-expression of 2-5A-synthetase (compare lanes 5 and 7).

The effect of expression of enzymes in the 2-5A pathway on virus yields is shown in Table 1. Production of infectious virus at 24 and 48h was quantitated by plaque assay and values referred to results obtained for cell infected with VV-RL in the absence of vT7. Expression of 2-5A-synthetase by itself causes inhibition of virus replication by 60 and 77.5 % at 24h and 48h of infection, respectively. Expression of RNase L inhibits virus yields by 75 and 92.1 % at 24 and 48h of infection, respectively, while combined expression of RNase L and 2-5A-synthetase enhances the inhibition of virus growth to 96.6 % at 48h. Reduction in virus yields due to RNase L expression is a late event and it is only apparent after 12h of infection (data not shown). The limited reduction in viral yields observed in these experiments is probably due to the kinetics of RNase L activation. Synthesis of this protein is detected for the first time 14 h after infection (Fig. 2A, lane 4) and inhibition of protein synthesis is clearly observed starting at about the same time (Fig. 2C, lane 3), a relatively late time point in VV infection when most viral protein
synthesis has already occurred. Therefore, the one log reduction observed in our expression system probably is an underestimate of the potency of the 2-5A system in VV sensitivity to IFN.

In order to establish that inhibition of VV replication is specific, we have also analyzed the effect of RNase L expression in the replication of vesicular stomatitis virus (VSV) which is very sensitive to IFN treatment, although has been previously described as resistant to the 2-5A system (12). Monolayers of BSC-40 cells were infected with VSV (m.o.i. 10), alone or in combination with either wild type VV (m.o.i. 10) or recombinant viruses: vT7 (m.o.i. 5) + VV-RL (m.o.i. 5) or vT7 (m.o.i. 5) + VV-ΔN (m.o.i. 5) and VSV viral yields were established at different times of infection. Production of active RNase L, demonstrated by protein analysis (data not shown), has no effect on VSV replication as indicated in Table 2.

We have also investigated if RNase L is the only protein responsible for the effects observed in cells infected with vT7 + VV-RL or some other IFN-induced enzyme could be also playing some role. Therefore, we measured RNA stability in BSC-40 cells treated for 18h with IFN (1000 international units [U] per ml of human lymphoblastoid IFN-α, 1 x 10^8 U/μg, Wellcome Res.) and then infected with vT7 (m.o.i. 4) or doubly infected with vT7 + VV-ΔN or vT7 + VV-RL (m.o.i. 2 each virus) for 24h (Fig. 4A). It should be noted that replication of wild type VV is nearly resistant to IFN treatment of BSC-40 cells as described above with only a 35% reduction in viral yields observed at 24h of infection of IFN-treated compared to untreated cells. In agreement with that, in cells pretreated with IFN, cleavage of rRNA could not be observed in vT7 or vT7 + VV-ΔN infections (lanes 2 and 4, lower panel) but only when recombinant RNase L was produced (lane 3, lower panel). In addition, the extent of rRNA degradation induced by vT7 + VV-RL infection of BSC-40 cells was only slightly increased by IFN treatment (compare lane 3, upper and lower panels). Therefore, induction by IFN of endogenous enzymes in the 2-5A pathway together with PKR and additional proteins has no significant contribution to effects produced by recombinant RNase L. However, due to synthesis by VV of proteins E3L and K3L that modulate the activation of PKR, the possibility still remains that the inhibition of protein synthesis observed at late times of infection with vT7 + VV-RL is due to activation of PKR in response to low levels of those viral interference products.
To exclude this possibility, we have infected mouse 3T3-like cell lines derived from homozygous PKR knockout mice (Pkr \(^{0/0}\)) or wild type animals (Pkr \(^{+/+}\)) (37) (Fig. 4B). Absence of murine 65 kDa protein kinase in the cell line derived from Pkr \(^{0/0}\) mice is shown by immunoblot of protein extracts using a rabbit polyclonal antiserum specific for mouse PKR (left panel, Fig. 4B). De novo protein synthesis activity was then measured in mock-infected cells or in cells infected for the indicated times with vT7 + VV-RL or vT7 + VV-\(\Delta\)N (m.o.i. 10 each virus) by pulse-labeling with \(^{35}\)S-methionine + cysteine as described (Fig. 4B, right panels). Levels of inhibition of virus protein synthesis by RNase L are comparable in cells devoid of PKR (lower panel) or control cells (upper panel). The result of inhibition of protein synthesis is a reduction in vT7 + VV-RL yields produced after 24h of infection of both Pkr \(^{0/0}\) (63%) and Pkr \(^{+/+}\) cells (84%). Therefore, our results exclude activation of p68 kinase as playing any fundamental role in our expression system.

The results presented in this investigation provide direct evidence for an anti-VV role of ds RNA-dependent 2-5A-synthetase and 2-5A-dependent RNase L, key enzymes in the molecular mechanisms of IFN action. By increasing the levels of these enzymes we have been able to overcome the factors that VV uses to evade the IFN action. Activation of the 2-5A-pathway induced by infection with VV-2-5AS or VV-RL alone most probably requires the participation of endogenous enzymes: RNase L that becomes active upon increased levels of 2-5A are produced by infection with VV-2-5AS, or 2-5A synthetase causing accumulation of 2-5A (14, 26) that activate the increased levels of RNase L achieved by VV-RL. Basal levels of enzymes in the 2-5A system are present in variable amounts in most if not all mammalian cells. RNase L is a low abundance protein which is increased from 2 to 10 fold in some cells after treatment with IFN (38). On the contrary, the amount of 2-5A-synthetase usually increases substantially in response to IFN (10 to 10,000-fold) although the high levels of 2-5A-synthetase produced in cells transfected with the 40-kDa 2-5A-synthetase human gene are not sufficient by themselves to protect against picornaviruses, and minimum levels of RNase L are also required (39). These data can be interpreted to mean that the antiviral state in the cell is compatible with amounts of 2-5A-synthetase in a very broad range while small changes in levels of activated RNase L are really critical. Our results, obtained in cells infected with the recombinant viruses,
provide evidence for this hypothesis and showed that expression of RNase L by itself causes greater inhibition of VV replication than expression of 2-5A-synthetase (Table 1). Because there is endogenous RNase L in the cells used in these experiments, we have tried to separate the contribution of endogenous and recombinant enzymes by using stable cell line SVT2/ZB1.4 expressing a dominant negative mutant of RNase L (7). However, infection of these cells with vT7 + VV-RL produced degradation of rRNA similar to control cells SVT2/pSVL. A possible explanation is that levels of mutant RNase L, which has been described as insufficient to counteract the anti-EMCV effect of IFN (7), are also below the amounts of RNase L produced by vT7+VV-RL infection.

The VV cell system described here for expression of enzymes in the 2-5A pathway has two main advantages: first, it makes possible to increase the levels of these enzymes in a controlled fashion, and second, virus infection provides at the same time a very powerful activator of the system, i.e. ds RNA, a byproduct of virus replication. We believe that RNase L activation requires first of all production of the protein and then synthesis of ds RNA, the activator of endogenous 2-5A-synthetase that produces 2-5A oligonucleotides at late times of infection (14, 25, 26). We have previously described that 2-5A levels in untreated BSC-40 cells are 1.4 nM by 8h of infection with VV and increase to 3.0 nM by 24h of infection (26). Low concentrations of 2-5A (≤ 1nM) are normally required to activate the 2-5A-dependent RNase in vitro (14) and 2-5A extracted from VV infected cells is biologically active when assayed in vitro (14). Because degradation of RNA occurs during infection with VV-RL, our results strongly suggest that 2-5A accumulated in infected cells is also biologically active in vivo. Then, how can we explain that 2-5A produced during wild type VV infection of BSC-40 cells does not induce rRNA breakdown?. We propose the existence of a mechanism directly responsible of interference with 2-5A-dependent RNase L. Therefore, during infection of cells with the recombinant virus expressing RNase L, activation of this enzyme would be fulfilled only when enough levels of RNase L and its specific activator dsRNA accumulate as to out-number viral-induced resistance mechanisms specific for this enzyme. Indeed, induced expression of PKR with a recombinant VV similar to VV-RL has been shown to counteract viral mechanisms of interference with this pathway and cause inhibition of its own replication (40). We envisage
resistance or sensitivity of VV to IFN as a tightly regulated equilibrium between levels of the IFN-induceable proteins (basal, IFN-induced or recombinant virus-induced enzymes in our expression system), their activators, such as ds RNA and 2-5A provided upon viral infection, and virus-encoded resistance products. Subtle changes in the levels of some of the components in the 2-5A system can direct the cell toward an IFN sensitive or resistant phenotype that might also explain differences in IFN sensitivity found for VV infecting different cell lines (13-15). Although in our expression system several positive or negative regulatory factors are involved, maintaining the multiplicity of infection equal in each experiment with auxiliary recombinant tk(-) viruses warranties the same contribution of all these factors. Therefore, effects observed can be specifically attributed to expression of the recombinant proteins.

In conclusion, in this report we provide direct evidence with cultured cells that by increasing the levels of enzymes in the 2-5A pathway we generate an antiviral state in the cell that is effective upon VV replication but does not affect VSV, which has been previously shown to be insensitive to this pathway. Since replication of VV is severely inhibited in IFN-treated animals, the biological significance of our findings is to point out that in vivo the 2-5A system is likely to be a mediator of the anti-poxvirus action of IFN and that modulation of the 2-5A pathway might have therapeutic value. The unique system described here provides the means to search for virus interference factors throughout their interaction with RNase L. In addition, by expressing mutant or truncated forms of enzymes in the 2-5A system, it should be possible to define in vivo functional domains in these important proteins.

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LEGENDS TO FIGURES

**Figure 1. Inducible expression of nearly full-length and truncated RNase L by recombinant vaccinia viruses.** (A) Diagram showing the structure of the transfected genes pTM-RL and pTM-ΔN. pT7, bacteriophage T7 polymerase promoter; EMC, encephalomyocarditis virus sequences for cap-independent translation; solid boxes, pRSET-B sequences comprising an ATG, 6 histidine residues and the gene 10 leader peptide; open circles, ankyrin-like repeats in 2-5A-dependent RNase L; gray oval, protein kinase homology region; gray rectangle, region required for RNase activity; Ter, T7 transcriptional terminator. (B) Expression of the recombinant proteins. Monolayers of BSC-40 cells were either mock-infected (lane 1), single infected with VV-RL, VV-ΔN or vT7 (m.o.i. 4) (lanes 2-4) or doubly infected with vT7 + VV-RL (lane 5) or vT7 + VV-ΔN (lane 6) (m.o.i. 2 each virus) for 24 h. Cell extracts (50 µg) were fractionated in SDS-10% polyacrylamide gels, transferred to nitrocellulose, and analyzed by immunoperoxidase staining after reactivity with rabbit polyclonal serum against RNase L.

**Figure 2. Induced RNase L cleaves rRNA and inhibits virus protein synthesis in BSC-40 cells.** (A) Time-course of RNA degradation. Monolayers of BSC-40 cells were either mock-infected (lane 1), single infected with vT7 (m.o.i. 4) (lane 2), or doubly infected with vT7 + VV-RL (lanes 3-6) or vT7 + VV-ΔN (lane 7) (m.o.i. 2 each virus). Control infections proceeded for 24 h (lanes 2 and 7) or for the indicated times in the case of vT7 + VV-RL (lanes 3-6). Total RNA was purified using Ultraspec-II RNA Resin Purification System (Biotecx), and the amount of RNA corresponding to 3 x 10⁵ cells was fractionated in 1% agarose-formaldehyde gels and stained with ethidium bromide. Abundant RNAs, 28 S and 18 S rRNA, are indicated. (B) Time-course of RNase L synthesis. Cells were either mock infected or infected as before. Cell extracts (50 µg) were fractionated in SDS-10% polyacrylamide gels, transferred to nitrocellulose, and immunoblotted using rabbit polyclonal antiserum against RNase L as described. (C) Time-course effect of RNase L on de novo protein synthesis. Monolayers were mock-infected or infected with vT7 + VV-RL or vT7 + VV-ΔN (m.o.i. 5 each virus) for the indicated times. Cells
were labeled for 15 min with $^{35}$S-labeled methionine+cysteine (100 $\mu$Ci/ml) and protein extracts (50 $\mu$g) were fractionated in SDS-polyacrylamide gels and autoradiographed. Lane 1 is over-exposed reflecting lack of shut-off in protein synthesis in mock-infected cells compared to infected cells (lanes 2-9).

**Figure 3.** Expression of 2-5A-synthetase together with RNase L increases effects on RNA stability and protein synthesis. (A) Analysis of RNA degradation. Monolayers were either mock-infected (lane 1) or infected with indicated combinations of viruses vT7, VV-RL, VV-ΔN and VV-2-5AS (m.o.i. 5 each virus) (lanes 2-8) for 24 h (upper panel) or 48 h (lower panel). Total multiplicity was maintained 15 at all times using VV-LUC when required. Cells were also treated with IBT (15 $\mu$M) and infected with wild type virus (VV) for 24 h (m.o.i. 15) (lane VV-IBT). Total RNA was purified as before, and the amount of RNA corresponding to 3 x 10$^5$ cells was fractionated in 1% agarose-formaldehyde gels and stained with ethidium bromide. Abundant RNAs, 28 S and 18 S rRNA, are indicated. (B) Synthesis of recombinant proteins. Monolayers were infected for 48h as before, cell extracts (50 $\mu$g) were fractionated in SDS-10% polyacrylamide gels, transferred to nitrocellulose, and immunoblotted using rabbit polyclonal antiserum specific for RNase L (upper panel) or 2-5A-synthetase (lower panel). (C) De novo protein synthesis. Monolayers were infected for 24h as before and labeled with $^{35}$S-labeled methionine+cysteine (100 $\mu$Ci/ml) for 15 min as described. Protein extracts (50 $\mu$g) were fractionated and dried gels were autoradiographed. Lane 1 is over-exposed reflecting lack of shut-off in protein synthesis in mock-infected cells compared to infected cells (lanes 2-8).

**Figure 4.** Effects produced by VV-RL are not dependent on other IFN-induced enzymes. (A) Analysis of RNA degradation in IFN-treated cells. Monolayers of BSC-40 cells were treated for 18h with 1000 international units [U] per ml of human lymphoblastoid IFN-α (lower panel) or left untreated (upper panel). Cells were either mock-infected (lane 1), single infected with vT7 (m.o.i. 4) (lane 2), or doubly infected with vT7 + VV-RL (lane 3) or vT7 + VV-ΔN (lane 4) (m.o.i. 2 each virus) for 24h. Total RNA was purified as before, and the amount of RNA corresponding to 3 x 10$^5$ cells was fractionated in 1% agarose-formaldehyde gels and stained with ethidium bromide. Abundant RNAs, 28 S and 18 S rRNA, are indicated. (B) Effect of
RNase L on de novo protein synthesis in cells devoid of PKR. Left panel. Immunoblot of protein extracts prepared from 3T3-like cells derived from homozygous PKR knockout mice (Pkr^{0/0}) or wild type animals (Pkr^{+/+}) using a rabbit polyclonal antiserum specific for mouse PKR. Right panels. Monolayers of Pkr^{++} (upper panel) or Pkr^{0/0} cells (lower panel) were mock-infected (lane 1) or infected with vT7 + VV-RL (lanes 2-5) or vT7 + VV-ΔN (lanes 6-9) (m.o.i.10 each virus) for the indicated times. Cells were labeled as before and protein extracts (50 μg) were fractionated in SDS-polyacrylamide gels and autoradiographed.

**Table 1. Viral yields in cells infected with recombinant viruses expressing RNase L and/or 2-5A-synthetase.** Monolayers of BSC-40 cells grown in 24-well plates were infected with viruses vT7, VV-RL and VV-2-5AS (m.o.i. 5 each virus). Total multiplicity was maintained 15 at all times, using VV-LUC when required. Cells were collected in media at 24 and 48 h of infection, freeze-thaw three times, sonicated and virus yields titrated on BSC-40 cells. The results are mean values of three independent experiments and standard deviations are given in parenthesis.

**Table 2. Effect of RNase L expression on VSV replication.** Monolayers of BSC-40 cells were either mock-infected or infected with wild type VV (m.o.i. 10) or recombinants: vT7 (m.o.i. 5) + VV-RL (m.o.i. 5) or vT7 (m.o.i. 5) + VV-ΔN (m.o.i. 5). After 30 minutes of adsorption, VSV (m.o.i. 10) was also added. Following another 30 minutes the virus inoculum was removed, cells washed and medium containing 2% NCS was added. Infections were also performed with VSV or vaccinia viruses alone as controls. Cell supernatants were collected at the indicated times of infection and VSV virus yields titrated on BSC-40 cells for 14h. The results are presented in pfu/ ml.