The ATF6 branch of unfolded protein response and apoptosis are activated to promote African swine fever virus infection

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African swine fever virus (ASFV) infection induces apoptosis in the infected cell; however, the consequences of this activation on virus replication have not been defined. In order to identify the role of apoptosis in ASFV infection, we analyzed caspase induction during the infection and the impact of caspase inhibition on viral production. Caspases 3, 9 and 12 were activated from 16 h post-infection, but not caspase 8. Indeed, caspase 3 activation during the early stages of the infection appeared to be crucial for efficient virus exit. In addition, the inhibition of membrane blebbing reduced the release of virus particles from the cell. ASFV uses the endoplasmic reticulum (ER) as a site of replication and this process can trigger ER stress and the unfolded protein response (UPR) of the host cell. In addition to caspase 12 activation, indicators of ER stress include the upregulation of the chaperones calnexin and caioreticulin upon virus infection. Moreover, ASFV induces transcription factor 6 signaling pathway of the UPR, but not the protein kinase-like ER kinase or the inositol-requiring enzyme 1 pathways. Thus, the capacity of ASFV to regulate the UPR may prevent early apoptosis and ensure viral replication.

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African swine fever virus (ASFV) is a double-stranded large DNA virus that induces an acute disease of swine in which apoptosis has a central role in pathogenesis. Virus infection elicits apoptosis in target and immune defense cells.1 The large repertoire of anti-apoptotic proteins encoded by the virus reflects the relevance of apoptosis in limiting viral replication in host cells. Many DNA viruses encode structural and functional homologs of anti-apoptotic Bcl-2 proteins, named viral Bcl2, including the A179L gene of ASFV.2 In common with other viruses, ASFV has several strategies to block apoptotic pathways in order to complete virus replication. ASFV encodes proteins involved in apoptosis inhibition, thereby delaying the final execution step of the apoptotic pathway, which occurs only at late post-infection times. Programmed cell death itself may be relevant for the final release of virus particles within apoptotic bodies, which would provide this virus with a mechanism to spread, thus evading the immune system. The central component of the apoptotic machinery is the caspase proteolytic system. There are two groups of caspases: upstream initiator caspases, such as caspases 8 and 9, which cleave and activate other caspases; and downstream effector caspases, including caspase 3, which cleave a variety of cellular substrates, thereby disassembling cellular structures or inactivating enzymes.3

Caspase 12 is associated with the cytoplasmic face of the endoplasmic reticulum (ER) and it cleaves to an active form in response to ER stress.4 The ER stress response, also called the unfolded protein response (UPR), is mediated by three transmembrane proteins: (i) the protein kinase-like ER resident kinase (PERK); (ii) the activating transcription factor 6 (ATF6); and (iii) the inositol-requiring enzyme 1 (IRE1).5 These three proteins are associated with the ER chaperone Bip/Grp78, which prevents their aggregation and further activation. However, when misfolded proteins accumulate, Bip is released, thus allowing UPR activation. Activated PERK phosphorylates the eukaryotic initiation factor 2α (eIF2α2), thus resulting in translation attenuation to counterbalance enhanced protein accumulation. This counterbalancing effect includes the upregulation of the pro-apoptotic mRNA CHOP (c/EBP homologous protein) and mRNA encoding GADD34 (growth arrest and DNA-damage-inducible protein-34), whose association with the protein phosphatase 1 (PP1) leads to the dephosphorylation of eIF2α.5 Activated IRE1 removes 26 nucleotides from X-box binding protein-1 (XBP1) mRNA, thereby altering the open reading frame and causing the translation of an active transcription factor.6 The spliced form of XBP1 protein (sXBP1) is involved in the transcriptional activation of a number of genes.

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Abbreviations: ATF6, activating transcription factor 6; ASFV, African swine fever virus; UPR, unfolded protein response; ER, endoplasmic reticulum; PERK, protein kinase-like ER resident kinase; IRE1, inositol-requiring enzyme 1; Bip/Grp78, binding immunoglobulin protein/glucose-regulated protein 78; CHOP/GADD153, c/EBP homologous protein; eIF2α, eukaryotic initiation factor 2α; GADD34, growth arrest and DNA-damage-inducible protein-34; PP1, protein phosphatase 1; XBP1, X-box binding Protein-1; EDEM, ER degradation enhancing α-mannosidase-like protein; HCV, hepatitis C virus; BVDV, bovine viral diarrhea virus; JEV, Japanese encephalitis virus; CMV, cytomegalovirus; SARS-CoV, severe acute respiratory syndrome virus; AEBSF, 4-(2-aminoethyl)benzenesulfonyl fluoride
including the ER mannosidase-like protein EDEM, which participates in protein degradation. Finally, activated ATF6 exits ER compartment to migrate to the Golgi apparatus where it is cleaved by proteases. ATF6 cytosolic fragment is a transcription factor responsible for the transcriptional induction of XBP1 as well as many ER chaperone-encoding genes. However, when cells are unable to recover from ER stress, apoptosis occurs.

The ER is an essential organelle for viral replication and maturation; in the course of productive infection, a large number of viral proteins are synthesized in infected cells, where unfolded or misfolded proteins activate the ER stress response. It is therefore not surprising that viruses have evolved various mechanisms to counteract these responses that limit or inhibit viral replication. For instance, BiP is induced in cells infected with respiratory syncytial virus, hanta viruses, hepatitis C virus (HCV), bovine viral diarrhea virus (BVDV), Japanese encephalitis virus (JEV), dengue virus, and enterovirus 71. The induction of PERK has also been reported in infection with herpes simplex virus, cytomegalovirus (CMV), and BVDV. The IRE1 pathway has been shown to be activated in cells infected with JEV, dengue, and severe acute respiratory syndrome virus, whereas the ATF6 pathway is triggered upon HCV infection and both IRE1 and ATF6 arms are induced during Rotavirus infection. In addition, the activation of all three pathways of the UPR has been reported in infection with West Nile virus and dengue. Furthermore, ASFV encodes the DP71L protein homolog to GADD34, which inhibits the HCV infection and both IRE1 and ATF6 arms are induced genes. However, when cells are unable to recover from ER stress, apoptosis occurs.

To study the impact of apoptosis on ASFV infection, we infected Vero cells treated with an inhibitor of caspase 3 activation (Ac-DEVD-CHO) and with a specific inhibitor of myosin II ATPase activity (Blebbistatin). Myosin II ATPase activity, together with the GTPase ROCK-I, is necessary for cell membrane fragmentation into multiple apoptotic bodies in the execution phase of apoptosis. Also, we analyzed the effect of Y-27632, a ROCK-I inhibitor that blocks membrane blebbing. We then compared final infective virus production at 48 hpi when these drugs were added to the cells, either 2 h before infection, or at 6 hpi. Our results showed that neither blebbing nor caspase 3 inhibition modified total virus production (extracellular plus intracellular virus production; Figure 2a). Caspase 3 activation in early stages of the infection was required for virus exit, whereas the inhibition of membrane blebbing reduced the release of virus particles from the cell and this effect was enhanced when the inhibitor was added 6 hpi (Figure 2b). In order to assess a possible role of caspase 12 in caspase 3 and 9 activation, we analyzed the effect of caspase 12 inhibition in infected cells using Z-ATD-FMK. Addition of this inhibitor before infection did not affect percentages of ASFV-infected cells as detected by flow cytometry (fluorescence-activated cell sorter (FACS); Figure 3a). Also, this inhibitor did not cause a decrease in virus production nor in the exit of virus particles (Figure 3b). Moreover, our results showed that neither caspase 3 nor caspase 9 were inhibited by Z-ATD-FMK and only caspase 12 activation was specifically inhibited (Figure 3c).

**Results**

**Activation of caspases in ASFV-infected cells.** Caspases are key factors in the regulation of apoptotic pathways. These proteins are induced by a variety of apoptotic triggers (stimuli). To identify the apoptotic pathways induced in ASFV-infected cells, we performed caspase activity assays at a range of times after infection (Figure 1). ASFV induced caspases 3 and 9 in Vero (African green monkey kidney) cells from 16 h post-infection (hpi) and increased this activation along infection, with higher levels at 48 hpi and both curves showed a similar pattern (Figures 1a and b). However, caspase 8 activity was not detected in infected Vero cells (Figure 1c). This observation was further confirmed in the infected WSL-R macrophage cell line (Figure 1d); however, caspase 8 activity was found in the specific positive control in both cases.

We also tested the activity of caspase 12, an ER membrane-associated cysteine protease that is activated in response to ER stress. Caspase 12 was undetectable in mock-infected cells or at 6 hpi, but at 16 hpi it was induced and increased threefold until 48 hpi (Figure 1e). The subcellular localization of this caspase in ASFV-infected cells was studied by indirect immunofluorescence (Figure 1f). As a control of induced caspase 12, Vero cells were treated with the ER stress-inducing agent tunicamycin, which activates caspase 12 and translocates it into the nucleus. Caspase 12 was localized in the nucleus in infected cells and in cells incubated with tunicamycin, but not in mock-infected ones.

**ASFV infection leads to the induction of ER chaperones.** The induction of caspase 12 after ASFV infection suggests that the virus triggers an ER stress response. This response is an autoregulatory program that upregulates a large number of genes, such as ER chaperones and ER-associated degradation (ERAD) components, which increase the folding capacity of the ER. In order to determine whether ASFV infection induces this response, we used western blot analysis to monitor the expression levels of the ER chaperones BiP, calreticulin, calnexin, ERP57 and PDI (protein disulfide isomerase) over the course of the infection. Calreticulin and calnexin were induced upon ASFV infection at 16 hpi (Figure 4a), while only a modest increase in PDI levels was found at 48 hpi. Interestingly, ASFV did not induce the expression of ERP57 or BiP. Chaperone upregulation was coincident with the expression of viral protein p30. Actin levels remained constant during the course of the infection, indicating that chaperone induction was not caused by a global upregulation of protein translation. We also analyzed the expression of BiP by immunofluorescence. Cells treated with tunicamycin showed a typical ER pattern for BiP, while BiP expression was undetectable in non-treated/non-infected controls and infected cells (Figure 4c). This could be probably due to the low levels of BiP to be detected by the antibody.

Calnexin is an ER resident molecular chaperone that has a key role in the correct folding of membrane proteins. It has
been proposed that calnexin acts as a scaffold for caspase 8-dependent cleavage of the ER transmembrane protein Bap31 and thus for the generation of the pro-apoptotic p20 under ER stress. To study whether overexpression of calnexin during the infection is involved in the cleavage of Bap31, we monitored p20 by western blot. Bap31 remained intact during the infection, thereby indicating the absence of pro-apoptotic signals between the ER and mitochondria (Figure 4a).

ASFV infection does not result in an attenuation of protein translation. We studied the expression of UPR markers in virus-infected Vero cells. The activation of the PERK pathway results in the phosphorylation of the eIF2α subunit, leading to translation attenuation. PERK also activates the expression of ATF4, a transcription factor, thus leading to the upregulation of the pro-apoptotic genes CHOP and GADD34. GADD34 association with the phosphatase PP1 produces the dephosphorylation of eIF2α. Tunicamycin treatment caused an increase in CHOP expression in Vero cells; however, no induction of CHOP was observed in ASFV-infected cells by western blot (Figure 5a). ATF4 expression was detected only in infected cells at 48 hpi. GADD34 protein showed an increase in expression from 16 hpi. Taken together, these results indicate that ASFV inhibits CHOP expression and leads to increased eIF2α dephosphorylation in order to restore protein translation.

ASFV infection does not induce XBP1 splicing in Vero cells. In response to the accumulation of unfolded proteins in the ER, IRE1 is activated by its oligomerization in the membrane, which causes the splicing of the mRNA encoding XBP1 transcription factor. The XBP1 protein encoded by the spliced mRNA is more stable than the unspliced form and is a potent transcription factor of the basic-leucine zipper (bZIP) family and caspase 12 inhibition flow cytometry (FACS ERAD). PDI eIF2α basic-leucine zipper (bZIP) one of the key
regulators of ER-folding capacity.\textsuperscript{31} Using RT-PCR and specific primers to distinguish between the unspliced (inactive) and spliced (active) forms of the XBP1 transcript, we detected XBP1 splicing in Vero cells treated with the UPR inducer tunicamycin but not in ASFV-infected cells at a range of post-infection times (Figure 5c). Therefore, it appears that the infection did not increase levels of active XBP1 mRNA. XBP1 mRNA is spliced by IRE1 but is induced by ATF6 activation.\textsuperscript{6} The level of endogenous XBP1 mRNA was measured in infected cells using qRT-PCR analysis of RNAs obtained in mock-infected samples. We found an increase in the level of XBP1 transcript in cells treated with tunicamycin or dithiothreitol (DTT) compared with the level of mRNA from ASFV-infected samples. We observed the same increase in the level of XBP1 mRNA in infected cells treated with tunicamycin and dithiothreitol (DTT) compared with the level of mRNA obtained in mock-infected cells. In contrast, ASFV resulted in a gradual decrease in the level of XBP1 mRNA over the course of infection (Figure 5d) indicating the downregulation of this mRNA in ASFV-infected cells.

**ASFV infection activates the ATF6 pathway.** In response to ER stress, ATF6 exits the ER to the Golgi apparatus, where it is processed by proteases to its active form, which in turn translocates to the nucleus. In order to monitor ATF6 activation by fluorescence microscopy, we used an EGFP (enhanced green fluorescent protein)–ATF6 fusion protein (pCMV short-EGFP-ATF6).\textsuperscript{32} This protein was under the control of a shortened CMV promoter, which has a deletion of 430 bp from the 5' side in order to prevent overexpression, which would modify the localization of the fusion protein itself. The short promoter shows considerably less activity than the full promoter and GFP-ATF6 expressed under the short CMV promoter localizes exclusively to the ER and translocates to the nucleus, in a similar way to endogenous ATF6.\textsuperscript{32} Cells transfected with the pCMV short-EGFP-ATF6 plasmid presented a typical ER pattern that relocated to the Golgi apparatus (Figure 6b) or to the nucleus (Figure 6c) in response to tunicamycin treatment. However, this characteristic pattern was modified in infected cells. Cells transfected with this construct and infected with ASFV showed GFP-ATF6 localization either in discrete perinuclear cytoplasmic areas corresponding to viral factories, as indicated by their co-localization with DNA stain (Figures 6d and e), or in the nucleus of infected cells (Figure 6f). This finding indicates that ASFV activates the ATF6 pathway of the UPR. We next studied the effect of a serine protease inhibitor that inhibits ER stress-induced proteolysis of ATF6, AEBSF\textsuperscript{33} on ASFV infection. Addition of this inhibitor before infection affected the percentages of ASFV-infected cells in a dose-dependent manner as detected by FACS (Figure 7a). AEBSF also induced a strong inhibition of viral production resulting in 100% reduction of viral titer at 300 μM concentrations of AEBSF (data not shown). We analyzed caspase activation under AEBSF to find that this agent abrogated caspase 3, 9 and 12 activation in infected cells. Inhibition of caspases was dependent on AEBSF effect on infection, given that this serine protease inhibitor did not prevent activation of caspase 3 induced by staurosporine (Figure 7b).

**Discussion**

Apoptosis is a genetically controlled cell death mechanism involved in the regulation of tissue homeostasis. We analyzed the mechanism by which ASFV induces apoptosis in infected cells and observed that caspases 3 and 9 are activated with similar kinetics as of 16 hpi. However, ASFV-induced apoptosis was independent of caspase 8 activation, a process involved in the extrinsic apoptotic pathways. These results indicated that the virus induces apoptosis through the mitochondrial pathway rather than the death receptor-mediated pathway. ASFV triggers apoptosis in an early stage of the infection process.\textsuperscript{34} Nevertheless, extensive apoptosis of infected cells occurs only at late post-infection times. Programmed cell death itself might be relevant for the release of virus particles into apoptotic bodies, which would provide a mechanism to facilitate virus spread and evasion of the immune system. Our results show that caspase 3 activation in early stages of the infection, was relevant for virus exit and that membrane blebbing contributes to virus dissemination, given the fact that the inhibition of the effector caspase 3 reduced the release of virus particles from the cell and this effect was enhanced when the inhibitor was added after 6 hpi. Thus, ASFV takes advantage of an early induction of apoptosis, through caspase 3 activity, while the final execution of programmed cell death in apoptotic blebs contribute to ASFV invasion of host cells and evasion of the immune surveillance. In addition to its critical role in apoptosis, caspase 3 participates in other cellular processes.\textsuperscript{35} This could also be the case of ASFV infection, where activation of caspase 3 may be required during early stages of infection to ensure subsequent virus exit from the cell. Another example is the influenza virus infection, in which caspase 3 inhibitors prevent the formation of progeny influenza A virus particles.\textsuperscript{36}

We have also shown that ASFV infection in Vero cells induces caspase 12, although this induction is not required for...
viral production or for caspases 3 and 9 activation. Caspase 12 is a characteristic ER stress response caspase. Virus assembly begins with the modification of ER membranes, which are subsequently recruited to the viral factories and transformed into viral precursor membranes. Moreover, a large amount of viral proteins are synthesized and accumulated in infected cells, thus overloading the ER-folding capacity. This overload can lead to the activation of the ER stress response of the host cell. Activated caspase 12, a key marker of ER stress-mediated apoptosis, may potentially interact with another pro-apoptotic protein, Bap31, resulting in a feedback mechanism involving the release of cytochrome c elicited through the mitochondrial pathway. However, Bap31 was not activated by the fragmentation of p20 in ASFV-infected cells. This observation indicates the absence of direct pro-apoptotic signals between the ER and mitochondria. Caspase 12 was triggered in ASFV-infected cells but this activation does not necessarily result in apoptosis, because the virus has the capacity to block ER stress at a later step. In general, ER stress starts with the transcriptional induction of ER-localized chaperones as a survival signal. Molecular chaperones are key components of the ER machinery; their main function consists of ensuring the quality of the ER by maintaining newly synthesized proteins in a proper state for folding, and marking misfolded proteins for degradation. Two folding systems have been proposed to contribute to this quality control. One of these is composed of grp78 (BiP), grp94 and PDI. The second system, known as the ‘calnexin–calreticulin cycle’, includes calnexin, calreticulin and ERp57. Chaperone expression analysis of ASFV-infected cells revealed a marked increase in calnexin and calreticulin expression from 16 hpi, and to a lesser extent, of PDI at 48 hpi. However, ASFV did not induce the expression of ERp57, which, together with calnexin and calreticulin, assists disulfide bond formation in glycoprotein folding. Surprisingly, ASFV did not elicit the expression of grp78 (BiP), thereby indicating that the virus contains the ER stress at this level.

To determine whether ER stress is involved in ASFV-induced cellular responses, we analyzed several ER stress-related proteins, including CHOP, GADD34, XBP1, ATF6 and ATF4. Previous results showed that viral infection in Vero cells inhibits the induction of the ATF4-CHOP signaling arm of the UPR. However, we observed expression of ATF4 from 48 hpi when virus is thought to promote programmed cell death in order to spread progeny. The inhibition of this UPR branch results in an attenuation of protein translation. To avoid this, ASFV encodes the DP71L gene, a GADD34 homolog that interacts with the catalytic subunit of PP1, and activates its phosphatase activity and that also promotes the expression of cellular GADD34, as shown here.
Figure 4  ASFV induces the expression of ER chaperones but not the expression of ERp57 or fragmentation of Bap31. (a) Western blot analysis of the ER chaperones in Vero cells infected with ASFV or treated with tunicamycin (Tm). Cell lysates were harvested at the indicated times after infection and then, analyzed. Actin was used as protein load control and viral infection was followed by p30 viral protein expression. (b) Quantification of the bands by densitometry corrected to actin data and normalized to control values. (c) Representative confocal micrographs of Vero cells grown on glass slides were treated with tunicamycin or infected with B54GFP-2 for 16 hpi. Cells were fixed, stained with anti-BiP plus Alexa Fluor 594-conjugated secondary antibody and incubated with Topro-3 for DNA staining and then analyzed by confocal fluorescence microscopy. Bar 10 μM.

Figure 5  ASFV regulates UPR signaling. (a) Western blot analysis of CHOP, GADD34, ATF4, p30 and p72 viral proteins in Vero cells infected with ASFV or treated with tunicamycin, lysated at the indicated times after infection with actin as protein load control. (b) Quantification of the bands corresponding to GADD34 by densitometry was corrected to actin data and normalized to control values. (c) Unspliced (uXBP1) and spliced (sXBP1) bands were RT-PCR-amplified using specific primer pairs. Cells treated with tunicamycin were used as a positive control for the induction of spliced XBP1. (d) XBP1 mRNA levels quantified with real-time RT-PCR. Total RNA was isolated from ASFV-infected Vero cells at a range of times post-infection. The XBP1 message was normalized to the 18S ribosomal message and x-fold changes were calculated as described in Materials and Methods. DTT and Tm-treated cells were used as control. Data are means ± S.D. from three independent experiments.
Figure 6  ASFV infection activates the ATF6 pathway. (a) Representative confocal micrographs of Vero cells transfected with the GFP-ATF6 plasmid. After 24 h, untreated cells showed a typical ER pattern, while in cells treated with tunicamycin, GFP-ATF6 relocated to the Golgi (b) and to the nucleus (c). (d–f) Representative confocal micrographs of ASFV-infected transfected cells. ATF6 translocated to the viral factories and to the nucleus. (d, arrow) A neighboring uninfected cell is shown for comparison. Bar 25 μm. (g) Percentages of GFP-ATF6 subcellular localization events in infected and tunicamycin treated cells in triplicate experiments calculated in a total of 5 × 10^4 cells/well.
IRE1-XBP1 is also critical for the UPR, as it stimulates the expression of proteins involved in ER stress-induced protein degradation. Nevertheless, no splicing of XBP1 was detected during ASFV infection. The absence of splicing may favor viral infection as this process would inhibit viral protein degradation induced by ER stress.

The activation of the ATF6 branch and its transcriptional activation of chaperone-encoding genes might benefit the virus by assisting the folding of accumulated proteins and preventing protein aggregation. We demonstrated ATF6 translocation to the nucleus. Activated ATF6 exits the ER to the Golgi apparatus, where it is processed by proteases to its active form, which in turn translocates to the nucleus.7 We used a short promoter that has considerably lower activity than the full promoter, and GFP-ATF6 expressed under the short CMV promoter localizes exclusively to the ER and translocates to the nucleus in a similar manner to endogenous ATF6.32 The analysis of ATF6 distribution by confocal microscopy in Vero cells infected with ASFV revealed that this protein was translocated into the nucleus. ATF6 was also located in discrete perinuclear cytoplasmic areas that corresponded to viral factories, as indicated by the co-localization of ATF6 with DNA-containing cytoplasmic foci. Furthermore, ASFV viral replication was blocked by inhibition of the ATF6 translocation to the nucleus by AEBSF. To analyze the transcriptional activity of ATF6 in ASFV infection we examined the expression of XBP1 in ASFV-infected cells by qRT-PCR. Surprisingly, ASFV-infected samples showed a decrease in the induction of XBP1 compared with uninfected samples. Activation of ATF6 was necessary for ASFV replication; however, the virus prevented the transcriptional induction of some ATF6-target genes, such as XBP1 and BiP, but not calnexin or calreticulin. These studies have focused interest in ATF6 pathway in viral infection and have paved the way for further characterization of ASFV viral gene regulation of these molecules in the UPR.

In conclusion, apoptosis induction, caspase activation and later modulation of the UPR might be advantageous for ASFV as these processes block the effects that are detrimental to the infection, while maintaining those that are beneficial.

Materials and Methods
Cell culture, viruses and plasmids. Vero cell lines were obtained from the American Type Culture Collection (ATCC, Richmond, VA, USA) and cultured at 37 °C in Dulbecco’s modified Eagle’s medium supplemented with 5% fetal bovine serum (FBS), 100 IU/ml penicillin, 100 μg/ml streptomycin and 2 mM L-glutamine. WSL-R line 379, a macrophage cell line of wild swine origin that was kindly provided by Günther Kiel (Friedrich-Loeffler Institut, Greifswald, Germany), was grown in Iscove’s medium (Gibco-BRL, Grand Island, NY, USA) plus F-12 nutrient mixture (Gibco) plus 10% FBS. Where indicated, cells were treated with 1 μg/ml Staurosporine (Sigma-Aldrich, Steinheim, Germany), or with 10 μg/ml tunicamycin (Sigma-Aldrich) or 2.5 mM DTT (Sigma-Aldrich) as a positive control of ER stress. The Vero-adapted ASFV isolate BA71V was used. For flow cytometry (FACS) analyses we used an infectious recombinant ASFV, B54GFP-2, which expresses and incorporates into the virus particle a chimera of the p54 envelope protein fused to the EGFP.41 Preparation of viral stocks, titrations and infection experiments were carried out in Vero cells as previously described.42

The plasmid pCMVshort-EGFP-ATF6 was kindly provided by Dr. Mori (Kyoto University, Japan) and was transfected into Vero cells using FuGene 6 (Roche Biochemicals, Mannheim, Germany), following the manufacturer’s protocols.
Western blotting. Cells seeded in 24-well plates were harvested in Laemml sample buffer, boiled for 5 min at 95 °C, resolved by SDS-PAGE and transferred to a nitrocellulose membrane (Bio-Rad, Hercules, CA, USA). The non-specific antibody binding sites were blocked with skimmed milk in phosphate buffered saline (PBS) and then incubated with the specific primary and the HRP (Horseradish peroxidase)-conjugated secondary antibody.

Bands obtained after development with ECL reagent (GE Healthcare, Picataway, NJ, USA) were detected using a Chemidoc XRS imaging system (Bio-Rad).

Antibodies used for western blotting included the following: rabbit polyclonal antibody to calreticulin (ab29077 abcam, Cambridge, UK) at a dilution of 1: 3000; rabbit polyclonal antibody to calnexin (StressGen, San Diego, CA, USA) 1: 3000; rabbit polyclonal to Bap31 (ab37120 abcam) 1: 3000; rabbit polyclonal antibody to ERp57 (ab10287 abcam) 1: 1000; goat polyclonal to GADD34 (ab9869 abcam) 1: 1000; rabbit polyclonal antibody GADD153(CHOP) (F-168) (Santa Cruz Biotechnology, Santa Cruz, CA, USA) 1: 200; rabbit polyclonal antibody to GRP78 (Bp) (Santa Cruz Biotechnology) 1: 200; rabbit polyclonal antibody to ATF4 (ab85049) 1: 500 mouse monoclonal antibody to PDI (ab2792 abcam) 1: 1000; anti-p-actin (Sigma, St. Louis, MI, USA) 1: 1500; anti-ASFV p30 monoclonal antibody 1: 1000; and anti-p72 monoclonal antibody clone 18BG3 (Ingenasa, Madrid, Spain) 1: 5000. The following secondary antibodies were used: goat anti-rabbit IgG HPR conjugate (Bio-Rad) and sheep anti-mouse IgG HPR conjugate (GE Healthcare), observing the manufacturer’s instructions. Band densitometry was performed with TINA image analysis software and data were normalized to control values.

Caspace activity assays. Using fluorometric caspase 3, 8 and 9 assay kits (Sigma-Aldrich), the activity of caspases 3, 8 and 9, respectively, were measured at a range of times post-infection in Vero cells and in WS-L-R cells infected with ASFV at a multiplicity of 1 pfu/cell. Caspase 12 activity was determined using a fluorometric caspase 12 assay kit (Bovision, Mountain View, CA, USA), following the manufacturer’s instructions.

Effect of inhibitors on viral production. Monolayers of Vero cells were infected with ASFV at a multiplicity of infection of 1 pfu/cell and treated with Ac-DEVD-CHO (N-acetyl-L-aspartyl-L-glutamyl-N-(2-carboxyl-1-formylethyl)-L-valinamide) (Sigma-Aldrich), Blebbistatin (Sigma-Aldrich), Y-27632 (Calbiochem, Merck, Dormstadt, Germany), Z-ATAD-FMK (benzyoxycarbonyl-Ala-Thr-Asp-fluoromethylketone) (Bovision) or AEB55 (4-(2-Aminoethyl)-Benzenesulfonylfluoride) (Sigma-Aldrich) at final concentrations of 25, 30, 10, 2 and 300 μM respectively, for 2 h before the infection or 6 hpi. Infection was allowed to proceed for 48 h and then virus titers in cell lysates (intracellular virus) and culture medium (extracellular virus) were titrated by plaque assay in triplicate samples on cells.

FACS analyses. Vero cells were pretreated with inhibitors at the indicated concentrations in growth medium for 1 h at 37 °C, followed by infection with ASFV B34GFP-2 recombinant virus at a moi of 1 pfu/ml. After 24 hpi cells were washed with PBS, harvested by trypsinization, and then washed with FACS buffer (PBS, 0.01% sodium azide, and 0.1% bovine serum albumin). In order to determine the percentage of infected cells per condition, 10,000 cells/time point were scored and analyzed in a FACS Calibur flow cytometer (BD Sciences, Franklin Lakes, NJ, USA). Untreated control infected cultures yielded 25–30% of infected cells from the total cells examined. Infected cell percentages obtained after drug treatments were normalized to values found in control plates.

Immunofluorescence. Cells seeded on glass coverslips in a 24-well plate were infected with ASFV. At several time points after virus infection, the cells were fixed with 4% paraformaldehyde in PBS for 15 min at room temperature and permeabilized in 0.1% Triton X 100 in PBS for 15 min at room temperature. Cells were consecutively stained with primary and secondary antibodies and then incubated with Topro-3 (Invitrogen, Eugene, OR, USA) 1:1000 in PBS/10X for DNA staining. After washing, coverslips were finally mounted on glass plates and cells were observed under a Leica TCS SPE confocal microscope (Leica-Microsystems, Wetztair, Germany). The primary antibodies used were: rabbit polyclonal antibody to GRP78 (Bp) (Santa Cruz Biotechnology) 1: 50, rabbit polyclonal antibody to caspase 12 (Biovision) diluted 1: 25 and anti-ASFV p30 monoclonal antibody 1: 100. Anti-rabbit IgG Alexa 488 and anti-mouse IgG Alexa 594-conjugated antibodies (Invitrogen) diluted to 1: 200 were used as secondary antibodies respectively.

RNA preparation and RT-PCR analysis for XBP1. Total RNA from cultured cells was isolated with TRI-Reagent (Sigma-Aldrich) following the manufacturer’s instructions. Reverse transcription reactions (RT-PCR) were carried out using the SuperScript One Step RT-PCR (Invitrogen), following the manufacturer’s instructions. The primers used were: forward primer 5’-CCCTGTAQTTGGACAG CAGG-3’ and reverse primer 5’-GGGCTTGTATATGTTGG-3’. Amplification conditions included an initial step of 45 °C for 30 s, followed by a denaturation step of 94 °C for 2 min, 30 cycles of 94 °C for 30 s, 55 °C for 30 s and 70 °C for 30 s. PCR products were resolved by electrophoresis in a 2% agarose gel and stained with SYBR Green.

For quantitative PCR, cDNA was synthesized using the QuantiTect Reverse Transcription kit (Qiagen, Hilden, Germany) following the manufacturer’s instructions. After heat inactivation at 95 °C, cDNA levels were quantified by using quantitative PCR (qPCR). Amplification was performed in a Rotor-Gene 3000 (Corbett Life Sciences, Sydney, Australia) using a QuantiTect SYBR Green PCR kit (Qiagen) following the manufacturer’s recommendations. Primer sets used for 18s and XBP1 are available from Qiagen (QuantiTect Primer Assay). Fold change was calculated using the ∆∆CT method of relative quantification with 18s RNA as the endogenous control for normalization. Melting curves were used to verify the specificities of products.

Data analysis. One-way analysis of variance was performed using INSTAT 3 for Windows. For multiple comparisons, Dunnet’s correction was applied. Data are presented as means ± S.D. Differences were considered statistically significant at a P value of <0.05.

Conflict of Interest
The authors declare no conflict of interest.

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