# African Swine Fever Virus CD2v protein promotes β-Interferon expression and apoptosis in swine cells

3 4

5

## Sabal Chaulagain<sup>a</sup>, Gustavo Delhon<sup>b</sup>, Sushil Khatiwada<sup>a</sup>, Daniel L. Rock<sup>a\*</sup>

a. Department of Pathobiology, College of Veterinary Medicine, University of Illinois at UrbanaChampaign, Urbana, IL, United States of America.

8 b. School of Veterinary Medicine and Biomedical Sciences, Nebraska Center for Virology,

9 University of Nebraska-Lincoln, NE, United States of America

10

<sup>\*</sup>Address correspondence to Daniel L. Rock, dlrock@illinois.edu

12

13 ABSTRACT African swine fever (ASF) is a disease of swine characterized by massive 14 lymphocyte depletion in lymphatic tissues due to apoptosis of B and T cells, most likely triggered 15 by proteins or factors secreted by infected adjacent macrophages. Here we describe a role for the 16 ASF virus (ASFV) protein CD2v in apoptosis induction in lymphocytes. CD2v is a viral homolog of host CD2 that has been implicated in viral virulence and immunomodulation in vitro; however, 17 18 its actual function remains unknown. We show that CD2v is secreted into culture medium of CD2v-expressing swine cells; and expression of- or treatment with CD2v led to significant 19 20 induction of IFN-β/ISGs transcription and antiviral state. CD2v expression led to enhanced NFκB-p65 nuclear translocation in these cultures and incubation with a NF-κB inhibitor reduced 21 CD2v-induced NF-κB-p65 nuclear translocation and IFN-β transcription. We show that CD2v 22 23 binds CD58, the natural CD2 ligand, and that CD58 siRNA knock-down results in significant reduction in NF-κB-p65 nuclear translocation and IFN-β transcription. Treatment of swine PBMC 24 with purified CD2v led to enhanced NF-κB-p65 nuclear translocation and induction of IFN-β 25 transcription. Further, induction of caspase-3 and PARP1 cleavage was observed in these swine 26 PBMC at later times, providing a mechanism for CD2v-induced apoptosis of lymphocytes. Finally, 27 IFN-β induction and NF-κB activation was inhibited in swine PBMC treated with purified CD2v 28 pre-incubated with antibodies against CD2v. Overall, our results indicate that ASFV CD2v is an 29 30 immunomodulatory protein that, by promoting lymphocyte apoptosis, may contribute to bystander 31 lymphocyte depletion observed during ASFV infection in pigs.

32

33 **IMPORTANCE** ASF, a severe hemorrhagic disease of domestic swine, represents a significant economic threat to swine industry worldwide. One critical pathological event observed in pigs 34 infected with virulent isolates is an extensive destruction of lymphoid tissue and massive 35 lymphocyte depletion. However, viral factor/s involved in this event are yet to be identified. Here 36 we show that, by inducing NF-kB-dependent IFN signaling, ASFV CD2v is able to promote 37 38 apoptosis in swine PBMC. We propose that CD2v released by ASFV-infected macrophages contributes to the massive depletion of lymphocytes observed in lymphoid tissues of infected pigs. 39 Results here improve our understanding of ASFV pathogenesis and will encourage novel 40 intervention approaches. 41

42 **KEYWORDS** African swine fever virus, CD2v, Interferon-β, NF-κB, CD58, Lymphocyte,

- 43 Apoptosis, Pathogenesis
- 44

### 45 INTRODUCTION

African swine fever (ASF) is a severe hemorrhagic disease of swine caused by ASF virus (ASFV), 46 47 a complex enveloped DNA virus which is currently the sole member of Asfarviridae family and 48 the only known DNA arbovirus. The disease is endemic to sub-Saharan African countries where virus cycles between bushpigs and warthogs, and Ornithodorus sp. ticks (1-4). While infection of 49 wild pigs is asymptomatic, infection of domestic pigs can result in mortality approaching 100%. 50 Clinical forms of ASF disease range from per-acute to chronic depending on the virus strain and 51 host factors (5-7). Lesions are most notably seen in lymphoid organs, and are characterized by 52 massive lymphocyte depletion and lymphatic tissue destruction (6-10). Although ASF virus 53 54 replicates in macrophages (11-14), infection with virulent ASFV causes marked apoptosis in B and T cells, which are not targets for viral replication (3, 15-19). It has been suggested that 55 lymphocyte apoptosis is most likely induced by proteins or factors secreted or released by infected 56 57 macrophages (19-22). Currently, no ASFV protein function has been associated with apoptosis induction in lymphocytes. 58

59 Interferons (IFNs) provide the first line of defense against virus infection. There are three different types of IFNs, types I, II and III. Type I IFNs consist of 13 IFN- $\alpha$  subtypes and IFN- $\beta$ . 60 61 Type II and III IFN consists of IFN- $\gamma$  and 3 subtypes of IFN- $\lambda$ , respectively. Nearly all cell types produce type I IFNs when host pathogen recognition receptors (PRRs) bind different pathogen 62 associated molecular patterns (PAMPs) (23). IFNs exert their effects by signaling through their 63 specific receptors, leading to transcriptional induction of interferon stimulated genes (ISGs). ISGs 64 mediate the anti-viral, anti-proliferative and immunomodulatory functions of IFN (24). 65 Uncontrolled high levels of type I IFN during viral infection might be detrimental to the host as it 66 has been associated with immunopathology, immunosuppression, enhanced pathology and disease 67 progression (25). The importance of interferon for control of virus infections is reflected by the 68 myriad of genes encoded by different types of viruses that target interferon responses (26, 27). 69

IFN-β induction is mediated by two major groups of transcription factors, nuclear factor-kB 70 (NF-kB) and IFN-regulatory factors (IRFs) (28-32). Canonical NF-kB activation involves 71 inhibitor kappa-β kinase (IKK) complex activation that leads to phosphorylation and degradation 72 73 of inhibitor kappa-β (IκB) proteins resulting in NF-κB nuclear translocation and subsequent 74 transcriptional induction of pro-inflammatory genes, including IFN-β (33-38). Lack of NF-κB activity leads to decreased expression of IFN- $\beta$  in cells (39, 40). Because of the central role of NF-75 76  $\kappa B$  in various antiviral responses, viruses target multiple steps on NF- $\kappa B$  activation, from PRR 77 recognition to NF-kB mediated gene transcription (41).

ASFV genome encodes several genes that function to interfere with NF- $\kappa$ B and IFN pathways in macrophages. For example, genes of the multi-gene family 360 and 530 (MGF360/MGF530) suppress IFN induction through yet unidentified mechanisms; DP96 inhibits the cGAS-STING-TBK pathway initiated by viral DNA, leading to inhibition of NF- $\kappa$ B /IRF-3-mediated IFN- $\beta$ induction; A238L inhibits NF- $\kappa$ B activation by binding to the NF- $\kappa$ B subunit RelA, and by inhibiting the acetylation of NF- $\kappa$ B-p65 by p300; and I329L inhibits dsRNA-induced NF- $\kappa$ B and IRF3 activation (42-45). Although virulent ASFV induces very low level of IFNs inconsistently during infection of macrophages *in vitro* (46-51), pigs infected with virulent ASFV show high levels of IFNs in serum and enhanced levels of cytokines (TNF $\alpha$ , IL-1 $\alpha$ , IL-1 $\beta$  and IL-6) in serum and organs (19-22, 52, 53). This suggests a role/s of IFNs of ASFV pathogenesis in pigs.

88 Uncharacterized soluble factors released by ASFV-infected macrophages have been shown to 89 inhibit proliferation of swine lymphocytes in response to lectins (54). CD2v, the ASFV 90 hemagglutinin and homolog of host T cell and NK cell surface antigen CD2, has been shown to 91 inhibit the proliferation of lymphocytes in response to lectins, suggesting that CD2v has immunosuppressive activity in vitro (55). CD2v contains all the domains present in cellular CD2 92 and some of the residues involved in binding to its natural ligand, CD58 (56, 57). CD2-CD58 93 interaction has been shown to activate cellular kinases in lymphocytes; however, the involvement 94 95 of NF-κB signaling downstream CD2-CD58 interaction has not been shown (58-67).

The aim of this study is to investigate immunomodulatory function/s of ASFV CD2v and define a role of CD2v in induction of lymphocyte apoptosis. We show here that expression of- or treatment with- CD2v leads to induction of IFN- $\beta$ , ISGs and the antiviral state in swine cells. We also show that CD2v- induced IFN- $\beta$  expression requires NF- $\kappa$ B activation and CD2v-CD58 interaction. Importantly, CD2v treatment induces apoptosis in swine PBMC.

## 101

## 102 **RESULTS**

ASFV CD2v localizes in perinuclear region, cytoplasm and cell membrane of PK15 cells
 and is secreted into the culture supernatant. CD2v is an ASFV structural transmembrane
 glycoprotein homologous to CD2, a cell adhesion molecule expressed by T- and NK-cells (56, 57).
 CD2v encodes for a protein of 370 amino acids, with a predicted molecular weight of 42 kDa and
 is expressed on the surface of ASFV-infected macrophages. CD2v mediates hemadsorption of
 swine red blood cells (RBCs) (56, 57) and, together with viral C-type lectin, has a role in
 hemadsorption inhibition (HAI) serotype specificity (71).

To examine the subcellular localization and expression kinetics of CD2v, PK15 cells were 110 mock transfected or transfected with pCMV plasmid expressing C-terminally HA-tagged CD2v 111 (CD2v-HA) and examined at various times post-transfection (pt) by confocal microscopy. CD2v 112 was observed adjacent to the nucleus at 2 h pt, and in the cell membrane, perinuclear area and 113 cytoplasmic vesicles at later times pt (Fig. 1A). To confirm that CD2v expressed by PK15 cells is 114 capable of hemadsorption, PK15 cells were transfected with pEmpty-HA (control plasmid) or 115 pCD2v-HA and incubated with swine red blood cells (RBCs) 24 h later. PK15 transfected with 116 pCD2v-HA but not with control plasmid hemadsorbed swine RBCs as evidenced by rosette 117 formation (Fig. 1B, arrowheads). 118

119 The expression kinetics of CD2v was assessed by Western blot after transfection of PK15 cells 120 with CD2v-HA. Two major protein species of approximately 100 kDa and 25 kDa, and a less-121 abundant 15 kDa species were detected at 6 h pt, with increasing protein levels observed at later 122 time points (Fig. 2A). A similar expression pattern was observed in 293T cells and Vero cells 123 (Fig. S1 in supplemental material). The observed molecular weight of the full length protein was approximately 58 kDa higher than predicted. A single 42 kDa band was detected when CD2v was 124 125 expressed in presence of tunicamycin, an inhibitor of N-linked glycosylation, confirming that the 126 protein is heavily modified through N-linked glycosylation (Fig. S2 in supplemental material) (72). Because the 25 kDa species is absent in presence of tunicamycin, we speculate that the 25 kDa 127

protein product might result from processing of the full length protein in the endoplasmic reticulum. A study by Goatley and Dixon (2011) has shown that CD2v in virus-infected cells is cleaved in the endoplasmic reticulum or Golgi compartments. A faint 100 kDa and a predominant 25 kDa CD2v band were detected in the culture supernatant of PK15 cells 24 h post transfection with pCD2v-HA, indicating that CD2v is secreted into the culture supernatant (Fig. 2B).

Overall, our results are in agreement with previous studies on CD2v expression in ASFVinfected cells (56, 72, 73). In addition, this is the first study showing that CD2v is secreted into the culture medium, as previously speculated by others (72, 74).

Expression of ASFV CD2v in PK15 cells induces IFN- $\beta$  and ISGs transcription, and the 136 antiviral state. Preliminary RNA-Seq experiments were conducted to examine the effect of 137 secreted CD2v on cellular gene transcription. PK15 cells were incubated with CD2v-containing 138 supernatant (1:2 dilution) or supernatant from cells transfected with pEmpty-HA (control plasmid), 139 140 and total RNA was collected at 1 h, 2 h and 3 h post-treatment. RNA-Seq analysis showed upregulation of several interferon-stimulated genes (ISGs), including MX1, OAS1, and IRF9 at 2 141 h post treatment with further increase at 3 h, suggesting a potential role of CD2v in IFN-ß signaling 142 (data not shown). To assess the effect of CD2v in IFN-β and ISG transcription, PK15 cells were 143 transfected with pCD2v-HA or control plasmids pEmpty-HA and pORFV120-Flag, the latter 144 encoding for Orf virus protein ORFV120 and IFN-β transcription was assessed by RT-PCR. 145 Compared to controls, cells transfected with pCD2v-HA plasmid showed significant upregulation 146 147 of IFN- $\beta$  (5.6-fold) as early as 6 h pt and similar upregulation was observed at all other subsequent time points (Fig. 3A). Consistent with upregulation of IFN- $\beta$ , significant increase of ISGs MX1 148 (17.7-fold) and OAS1 (12.8-fold) transcription was observed at 30 h pt with pCD2v-HA plasmid 149 compared to controls (Fig. 3B and C). 150

To investigate the functional significance of IFN-β and ISGs induction by CD2v, the antiviral 151 state of cells was examined using an IFN bioassay. PK15 cells were transfected with pCD2v-HA 152 or control plasmids (Empty-HA vector or plasmids expressing Orf virus proteins ORFV120 and 153 ORFV113), and then infected with reporter VSV<sup>GFP</sup> (50 PFU/well) at various times pt. PK15 cells 154 transfected with pCD2v-HA but not with control plasmids showed inhibition of VSV<sup>GFP</sup> 155 replication as determined by both fluorescent microscopy and flow cytometry at 12 h. 24 h and 30 156 h pt (Fig. 4A and B). Inhibition of VSV<sup>GFP</sup> replication was also observed after treatment of PK15 157 cells with a two-fold dilution of culture supernatants from cells transfected with Poly (I:C) (up to 158 dilution 1:8) and pCD2v-HA (up to dilution 1:4) but not with control plasmids (Fig. 4C). These 159 160 data indicate that expression of- or treatment with- CD2v in PK15 cells leads to induction of IFN- $\beta$ , ISGs and an antiviral state. 161

Induction of IFN-β by ASFV CD2v depends on NF-κB activation. NF-κB and IRF3 are two 162 important transcription factors involved in IFN- $\beta$  induction (28-32) that translocate to the nucleus 163 164 upon activation. To assess whether CD2v expression affects NF-kB-p65 and IRF3 nuclear 165 translocation, PK15 cells were transfected with pCD2v-HA or control plasmids pORFV120-Flag 166 and pORFV113-Flag, and examined by IFA at various times pt. Enhanced NF- $\kappa$ B-p65 nuclear translocation was observed in CD2v expressing cells at all times pt compared to controls (Fig. 5A 167 168 and B). In contrast, IRF3 nuclear translocation was not observed (data not shown). To confirm activation of the NF-kB pathway, PK15 cultures were transfected with pCD2v-HA or pORFV113-169 Flag (control), and the mean fluorescence intensity (MFI) of phosphorylated NF-KB (pNF-KB 170

171 S536) was examined at 3 h pt by flow cytometry. Consistent with the nuclear translocation results,

significantly increased MFI values (1.8-fold) were observed in CD2v-expressing cells comparedto the control (Fig. 5C).

To assess whether inhibition of NF-κB affects IFN-β induction by CD2v, PK15 cells were pretreated with the NF-κB inhibitor parthenolide (1 $\mu$ M) or vehicle control (DMSO) for one hour, transfected with pCD2v-HA in presence of the inhibitor (1 $\mu$ M) or vehicle, and assessed for NFκB-p65 nuclear translocation at 3 h pt by confocal microscopy. Percentage of NF-κB-p65 nuclear translocation in CD2v-expressing cells in presence of parthenolide was significantly reduced (49.3%) compared to DMSO (Fig. 6A and B).

To investigate the effect of NF-κB inhibition on CD2v-mediated IFN- $\beta$  induction, PK15 cells were pretreated for one hour with parthenolide (1µM) or DMSO (vehicle control), transfected with pCD2v-HA or pEmpty-HA (control) for 6 h in presence of parthenolide (1µM) or DMSO, and assessed for IFN- $\beta$  transcription by RT-PCR. Significant reduction in IFN- $\beta$  transcription was observed in cells transiently expressing CD2v in presence of parthenolide (1.5-fold) compared to cells expressing CD2v in presence of vehicle alone (2.6-fold) (Fig. 6C). Together, results above indicate that induction of IFN- $\beta$  by ASFV CD2v is mediated by NF- $\kappa$ B activation.

NF-κB activation and IFN-β induction by ASFV CD2v is mediated by interaction of CD2v
 with host CD58. CD2v contains all the domains present in cellular CD2 and some of the residues
 involved in binding to the CD58, the natural CD2 ligand (56, 57). To study the potential interaction
 between CD2v and CD58, PK15 cells were co-transfected with pEmpty-HA vector or pCD2v-HA
 and porcine pCD58-Flag, and cell lysates were prepared at 8 h pt for reciprocal co immunoprecipitation with anti-Flag or anti-HA antibodies as described in materials and methods.
 Figure 7A shows that CD2v and porcine CD58 reciprocally co-immunoprecipitate.

To confirm CD2v-CD58 interaction, PK15 cells were co-transfected with pCD2v-HA and 194 pCD58-Flag, and co-localization of proteins was examined using confocal microscopy. Strong 195 overlap of signals was observed (Fig. 7B). Similarly, strong reciprocal co-immunoprecipitation 196 197 and co-localization of CD2v with endogenous human CD58 was observed in 293T cells transfected with pCD2v-HA, using anti-HA or anti- hu-CD58 antibodies (Fig. 8A and B). Given that: 1) CD2v 198 interacts with CD58 (Fig. 7 and Fig. 8) and 2) host CD2-CD58 interaction activates downstream 199 200 cellular kinases (58-67), we examined the involvement of CD2v-CD58 interaction in CD2vmediated NF- $\kappa$ B activation and IFN- $\beta$  activation. 201

To evaluate the effect of CD58 downregulation on CD2v-mediated NF-kB activation, PK15 202 cells were transfected with siRNAs targeting porcine CD58 or control siRNA as described in 203 materials and methods. CD58 transcript knock-down of approximately 55% was routinely 204 205 obtained when PK15 cells were transfected with CD58 siRNA compared to the negative control 206 (Fig. 9A). Twenty-four hours after siRNA treatment, cells were transfected with pCD2v-HA for 3 207 h, and NF-kB-p65 nuclear translocation was assessed by confocal microscopy. Significant 208 reduction in NF-kB-p65 nuclear translocation (60%) in CD2v expressing cells was observed in 209 PK15 cells with reduced CD58 transcript compared to the control (Fig. 9B, C and D).

To investigate involvement of CD58-CD2v interaction in CD2v-mediated IFN- $\beta$  induction, siRNA knock down experiments and CD2v-HA/control transfections were performed as described above. Significant reduction of IFN- $\beta$  transcription was observed 6 h pt with CD2v-HA in PK15

#### 214 Purified CD2v induces NF-κB-p65 nuclear translocation and IFN-β transcription in swine

215 PBMC cultures. Uncharacterized soluble factors released by ASFV-infected macrophages have 216 been shown to inhibit proliferation of swine lymphocytes in response to lectins (54) and CD2v has 217 been shown to be involved in inhibition of mitogen-induced proliferation of bystander lymphocytes in virus-infected swine PBMC cultures (55). This and our observations that CD2v-218 219 expressing PK15 cells secrete CD2v into the culture supernatant, and that cells treated with- or expressing CD2v upregulate IFN-β and ISGs led us to hypothesize that 1- secreted CD2v induces 220 IFN-β and ISGs expression in lymphocytes and 2- induction involves NF-κB-p65 nuclear 221 translocation. 222

- To examine NF- $\kappa$ B-p65 nuclear translocation in swine lymphocytes, swine PBMC were treated with purified CD2v or purified control for 1.5 or 2 h, processed for IFA and deposited onto glass slides with a cytospin. Confocal microscopy analysis showed enhanced NF- $\kappa$ B-p65 nuclear translocation in swine PBMC treated with purified CD2v (2.8-fold at 1.5 h; 1.6-fold at 2 h) compared to control treatment (Fig. 10A and B).
- To investigate whether IFN- $\beta$  transcription in lymphocytes is affected by CD2v, swine PBMCs were treated as above and assessed for IFN- $\beta$  transcription by RT-PCR at various times posttreatment. We found that IFN- $\beta$  transcription was significantly induced in CD2v-treated PBMCs at 4 h (3.3-fold) and 6 h (3.2-fold) post treatment compared to purified control (Fig. 10C).
- Supernatants from CD2v- treated swine PBMCs exhibit antiviral activity. To examine the 232 233 functional significance of CD2v-induced IFN-β expression in swine lymphocytes, swine PBMCs were treated with purified CD2v or purified control for 24 h, and the supernatants collected, 234 diluted, and used to treat fresh PK15 cells. Twenty-four hours after treatment the cells were 235 infected with VSV<sup>GFP</sup> and examined for virus replication at 16 h post infection by IFA. Supernatant 236 collected from PK15 cell culture transfected with Poly I:C was used as positive control. Significant 237 inhibition of VSV<sup>GFP</sup> replication was observed in PK15 cells treated with supernatant from 238 PBMCs. We observed 32.1% inhibition with undiluted supernatant, 24.4% inhibition with 1:2 239 240 dilution and 28.1% inhibition with 1:4 dilution (Fig. 11A and B).
- CD2v induces apoptosis in swine PBMCs. ASF is characterized by severe destruction of lymphoid tissue and massive lymphocyte depletion due to apoptosis (3, 7, 8, 16, 19, 20). An explanation for this critical pathogenic event is lacking. ASFV replicates in cells of the monocyte lineage, most notably macrophages, but not in lymphocytes, thus apoptosis in bystander lymphocytes is most likely due to proteins or factors secreted by infected macrophages. Based on our data, we hypothesize that CD2v released by infected macrophages induces IFN expression in bystander lymphocytes leading to apoptosis.
- To investigate the effect of CD2v on PBMC apoptosis, swine PBMC cultures were treated with purified CD2v, purified control, or staurosporine (positive control) as described in the materials and methods, and caspase-3 activation and PARP1 cleavage assessed by Western blot at various times post treatment. As shown in Fig. 12 A-C, treatment of swine PBMCs with purified CD2v led to significant induction of caspase-3 activation at 18 h post treatment (1.9-fold) and PARP1 cleavage at 18 h (1.9-fold) and 24 h (1.6-fold) post treatment compared to purified control. This result indicates that CD2v treatment induces apoptosis in swine PBMC.
- Treatment with monoclonal antibodies against ASFV CD2v inhibits CD2v- induced NF κB activation and IFN-β transcription in swine PBMCs. Monoclonal antibodies against ASFV

CD2v were generated and screened as described in materials and methods. Four anti-CD2v 257 258 antibodies (A4, C4, C3 and F2) were used as a mixture to examine their reactivity against CD2v. 259 The antibodies mix detected the full length 100 kDa CD2v species in Western blot (Fig. 13A). To 260 confirm reactivity of antibodies, lysates from 293T cells transfected with pCD2v-HA were incubated overnight with the anti-CD2v monoclonal antibodies. Immunoprecipitation products 261 262 were assessed by Western blot using anti-HA antibodies. Strong pull down of CD2v by anti-CD2v 263 monoclonal antibodies was observed in CD2v- but not in control-transfected cells (Fig. 13B). The results show that the anti-CD2v monoclonal antibodies react with CD2v. 264

To investigate the effect of the anti-CD2v antibodies on CD2v-induced NF-kB activation in 265 swine PBMC, purified CD2v or purified control were incubated overnight with the monoclonal 266 antibody mix, anti-ORFV086 monoclonal antibody, or anti-IgG mouse isotype antibody control. 267 Swine PBMCs were then treated with purified CD2v or purified control pre-incubated with the 268 269 various antibodies, processed with NF- $\kappa$ B nuclear translocation assay as above, and analyzed by 270 confocal microscopy. As a control for the experiment, purified CD2v or purified control without pre-incubation with antibodies was used to treat swine PBMCs. Significant inhibition of NF-κB-271 p65 nuclear translocation (approximately 50% reduction) was observed in PBMCs treated with 272 CD2v previously incubated with anti-CD2v monoclonal antibodies compared to controls (Fig. 14A 273 274 and B). This result supports a role of secreted CD2v in induction of NF-kB-p65 nuclear translocation in swine PBMCs. 275

276 The effect of the anti-CD2v monoclonal antibodies on CD2v-induced IFN- $\beta$  expression in swine PBMC, was investigated by pre-incubating purified CD2v or purified control with the 277 monoclonal antibodies as above, followed by treatment of PBMC. As a control for the experiment, 278 purified CD2v or purified control without pre-incubation with antibodies was used to treat swine 279 PBMC. Total RNA was extracted 6 h post treatment and IFN-β transcription was assessed by RT-280 PCR. Significant inhibition in IFN-β transcription was observed when swine PBMCs were treated 281 with purified CD2v pre-incubated with anti-CD2v antibody mix (1-fold) as compared to purified 282 CD2v pre-incubated with anti-ORFV086 monoclonal antibody (1.5-fold) or without pre-283 incubation (1.8-fold). These results confirm a role of CD2v in the induction of IFN- $\beta$  transcription 284 in swine PBMCs (Fig. 15). 285

## 287 **DISCUSSION**

286

288 A landmark of acute ASF is the severe lymphoid tissue destruction and massive lymphocyte 289 depletion in infected pigs, which occurs as a result of bystander lymphocyte apoptosis (3, 8, 19). Since lymphocytes do not support ASFV replication, factors secreted by infected macrophages 290 291 have been implicated in triggering lymphocyte apoptosis (16, 19-22). ASFV CD2v is a glycoprotein with homology to host CD2, an adhesion molecule expressed by T and NK cells (56, 292 293 57, 72). CD2v has been shown to be involved in host immunomodulation, virulence and induction 294 of protective immune responses (55, 75, 88). Here, we show that expression of ASFV CD2v in swine cells induces NF-kB-mediated IFN expression through interaction with CD58 (Fig. 3, Fig. 295 5 and Fig. 9). We observed that treatment of swine PBMC with purified CD2v leads to significant 296 IFN-β induction, NF-κB-p65 nuclear translocation (Fig. 10A and B), and caspase-3 and PARP1 297

cleavage, thus providing a mechanism for CD2v-induced lymphocyte apoptosis (Fig. 12).

The relationship between ASFV infection and the IFN system is complex. ASFV has evolved 299 300 multiple strategies to counteract activation of the IFN and NF-kB signaling pathways (42-45). In 301 vitro infection of porcine macrophages with low virulence ASFV strains induced enhanced and 302 sustained IFN induction compared to virulent strains (46-51). However, acute ASFV infection of pigs is characterized by high level of systemic IFN production (52, 53). Viral factors involved in 303 304 IFN induction and sources of IFN during acute ASF infection remain to be identified. Here, we describe an immunomodulatory function of ASFV CD2v that, by targeting lymphocytes for 305 apoptosis, may contribute to the massive lymphocyte depletion observed in vivo. 306

CD58/LFA-3 is the natural ligand for host CD2 protein, and CD2-CD58 interaction initiates 307 cellular kinase signaling (58-67). We show that swine CD58 interacts with CD2v, and that the 308 interaction is involved in CD2v-mediated IFN-ß induction through NF-kB-p65 nuclear 309 translocation (Fig. 7 to 9). Since viral glycoproteins can also induce IFN-β through TLR-4/TLR-2 310 311 (76-79), involvement of these receptors cannot be ruled out. It will be interesting to determine whether the CD2v cytoplasmic domain, known to interact with the trans-Golgi network AP-1 312 factor and actin-binding adaptor protein SH3P7 (73, 80), plays a role in CD2v immunomodulation. 313 In ASFV-infected macrophages, CD2v is processed and thought to be secreted (72-74). Here, 314 we show for the first time that CD2v is released into the supernatant by CD2v-expressing cells 315 (Fig. 2B). Previous studies have described the involvement of soluble factor/s in the inhibition of 316 lymphocyte proliferation in PBMCs infected with ASFV or incubated with cell 317 318 extracts/supernatants free of virus (54, 55) and have defined an immunomodulatory role of CD2v in inhibition of mitogen-induced proliferation of bystander lymphocytes (55). Based on our 319 findings, secreted CD2v could be mimicking host CD2 by interacting with CD58, thus leading to 320 induction of IFN-β and inhibition of lymphocyte proliferation. Our observation that anti-CD2v 321 monoclonal antibodies inhibit CD2v-dependent NF-kB-p65 nuclear translocation and IFN-β 322

induction further strengthen the immunomodulatory role of secreted CD2v (Fig. 14). 323 The role of CD2v in ASFV virulence is not clearly understood. Infection of pigs with a CD2v 324 deletion mutant virus resulted in different infection phenotypes depending on the parental virus 325 strain. CD2v deletion in the Spanish strain BA71 resulted in virus attenuation (75), and the CD2v326 gene has been found interrupted in some attenuated ASFV strains (82) suggesting a role of CD2v 327 in virus virulence. In contrast, deletion of CD2v in virulent strains Malawi and Georgia 2007 did 328 not significantly affect virus virulence in pigs (55, 81). Although not essential for viral replication 329 in pigs, CD2v is critical for virus replication in the tick (83, 84). Results here support a role of 330 ASFV CD2v in viral pathogenesis and virulence by affecting lymphocyte survival. 331

CD2v protein has been implicated in protective immunity. CD2v expression is required for 332 333 partial protection conferred by specific vaccine constructs, and two predicted CD2v T-cell epitopes are speculated to affect protective immunity (85-87). Using ASFV inter-serotypic chimeric viruses 334 and vaccination/challenge experiments in pigs, serotype-specific CD2v and/or C-type lectin 335 336 proteins were shown to be important for protection against homologous ASFV infection (88). 337 Others, however, have found that inoculation of pigs with the BA71 virus lacking CD2v conferred protection against parental BA71 (75). This variability likely reflects the differences in the 338 339 virulence of the challenge strains used in the experiments. In conclusion, CD2v represents an 340 important factor contributing to lymphocyte depletion observed during ASFV infection in animals (Fig. 16). 341

342

## 343 MATERIALS AND METHODS

**Cells.** Porcine kidney cells (PK15) and monkey kidney cells (Vero) were obtained from American Type Culture Collection (ATCC) and were maintained at 37° C with 5% CO2 in minimal essential medium (MEM) supplemented with 10% fetal bovine serum (FBS) (Atlanta Biologicals, Flowery Branch, GA), 2mM L-glutamine, gentamicin (50  $\mu$ g/ml), penicillin (100 IU/ml), and streptomycin (100  $\mu$ g/ml). Human embryonic kidney (HEK 293T) were cultured in Dulbecco's modified essential medium (DMEM) supplemented as above.

Plasmids and transfection. ASFV Congo strain CD2v was synthesized, cloned into pUC57 350 NJ) and amplified with primers CD2v-Fw 351 (Genscript, Piscataway, (EcoRI) (5'-352 and CD2v-Rv (KpnI) (5'-CAGAATTCGCGGTACCAATAATTCTATCT ACATGAATAAGCG-3'). 353 354 The amplified full length CD2v gene was cloned into EcoRI and KpnI sites of pCMV-HA vector (Clontech, Mountain View, CA) to produce pCD2v-HA, which express C-terminally HA-tagged 355 CD2v protein (CD2v-HA). To enhance translation efficiency, a Kozak sequence (GCCACC) was 356 357 placed in front of the CD2v gene. The construct was sequenced to confirm integrity and fidelity. To construct expression plasmids pORFV120-Flag and pORFV113-Flag, ORFV120 and 358

ORFV113 coding sequences were PCR-amplified from orf virus strain OV-IA82 genome and cloned into p3xFlag-CMV-10 vector (pFlag) (Clontech).

For transfection of cells, Lipofectamine 2000 (Invitrogen) and plasmid DNA were separately diluted in Opti-MEM medium (Gibco) and incubated for 5 min. Diluted DNA was added to diluted lipofectamine 2000 (1:1ratio) and incubated for 20 min. Finally, DNA-lipid complex was added to cells and 5 h after incubation, Opti-MEM medium was replaced with 10% complete growth media.

Hemadsorption assay. PK15 cells grown in 6-well plates were mock transfected or transfected
with plasmid (p) CD2v-HA. At 24 h post-transfection (pt), culture media was removed and rinsed
two times with PBS. Transfected cultures were incubated with PBS-washed 1% swine RBC
overnight and observed with microscope (X100).

**Swine PBMCs isolation, freezing and culture.** Swine PBMCs were obtained from swine whole blood through density gradient centrifugation using Sepmate15 (Stem cell technologies) and lymphocyte separation media (Corning), and frozen in freezing media (50% RPMI-1640, 40% FBS, 10% DMSO) as described elsewhere (68, 69). Swine PBMC were maintained at 37° C with 5% CO2 in RPMI 1640 medium (Corning) supplemented with 10% fetal bovine serum (FBS) (Atlanta Biologicals, Flowery Branch, GA), 2mM L-glutamine, gentamicin (50 µg/ml), penicillin (100 IU/ml), streptomycin (100 µg/ml) and sodium pyruvate (1 mM).

CD2v purification. For CD2v protein purification, 293T cells were transfected with pCD2vHA or pEmpty-HA control vector for 30 h. Cell lysates were harvested using mammalian protein
extraction reagent (MPER) (Thermo Scientific) and incubated with anti-HA resin overnight using
spin columns (Thermo Scientific). Next day, CD2v was eluted using HA peptide (1 mg/ml)
(Thermo Scientific) in TBS buffer (Corning). Whole cell lysates obtained from two 6-well plates
were incubated with 100 µl of anti-HA resin, eluted in 200 µl buffer and used for downstream
process.

Monoclonal antibodies against CD2v. PK15 cells were transfected with pCD2v-HA, and 384 385 whole cell lysates obtained 30 h pt. Lysates were incubated overnight with anti-HA antibody (Cell 386 Signaling Technology) at 4° C, and immunoprecipitated using 50 µl of protein G agarose bead slurry (Millipore). Monoclonal antibodies recognizing CD2v-HA were generated by immunizing 387 BALB/c mice with immunoprecipitated CD2v-HA (100 µg/dose). To generate hybridomas, 388 389 splenocytes from immunized mice were harvested and processed according to the company's protocol (STEMCELL technologies/ClonalCell-HY Hybridoma Cloning Kit). Clones were 390 screened for reactivity against CD2v-HA by IFA using CD2v-HA-transfected PK15 cultures. 391 Monoclonal antibody titers were estimated by IFA using goat anti-mouse Alexa fluor 488 or 594 392 as secondary antibody. Mice were maintained at the University of Nebraska-Lincoln (UNL) in 393 accordance with the guidelines of UNL Institutional Animal Care and Use Committee (IACUC) 394 and used in accordance with the guidelines of the committees. 395

396 Western Blot. Fifty  $\mu g$  of whole protein cell extracts or 50  $\mu l$  of cleared culture supernatant were resolved by SDS-PAGE, blotted to nitrocellulose or PVDF membranes and probed with 397 primary antibody against HA (Cell Signaling Technology), caspase-3 (9662S; Cell Signaling 398 399 Technology), PARP1 (sc-53643; Santa Cruz) or glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (sc-25778; Santa Cruz). The blots were developed with appropriate HRP-conjugated 400 secondary antibodies and chemiluminescent reagents (Thermo Scientific), and imaged with 401 402 FluorChem R (Protein Simple). Densitometric analysis was performed using ImageJ software and 403 all readings were normalized to GAPDH values.

Immunoprecipitation. 293T cells were transfected with pCD2v-HA, and whole cell lysates in
 MPER lysis buffer (Thermo Scientific) were obtained at 30 h pt and incubated overnight with the
 anti-CD2v monoclonal antibody mix or anti-HA antibody (3724; Cell Signaling Technology). Pull
 down products were obtained by eluting in Laemmli buffer (Bio-Rad) and analyzed by Western
 blot using anti-HA antibody (2367; Cell Signaling Technology).

CD58 siRNA knock-down. To investigate involvement of CD58-CD2v interaction in CD2v-409 mediated IFN- $\beta$  induction, siRNA knock down experiments were performed using CD58 sense 410 (S) (CUUCCAGAGCCAGAACUAU) and anti-sense (AS) (AUAGUUCUGGCUCUG GAAG) 411 siRNA duplex (Sigma Aldrich). PK15 cultures were transfected with CD58 siRNA (15 nM) and 412 mission siRNA transfection reagent (Sigma) following the manufacturer's protocol, and 413 transfected 24 h later with pCD2v-HA or control pEmpty-HA for 6 h. CD58 knock-down was 414 assessed by comparing transcript levels between cultures transfected with one MISSION siRNA 415 Universal Negative control (Sigma aldrich) and CD58 siRNA-transfected cultures using RT-PCR. 416 SYBR primers for swine CD58 were sCD58 FW 5'-ACTTAAACACTGGGTCGGGC-3' and 417 418 sCD58 RV 5'-AAGCTGCAAGGATCAGGCAT-3'.

419 **Real-time PCR.** Interferon- $\beta$  (IFN- $\beta$ ) and interferon stimulated genes (ISGs) transcription 420 were assessed in PK15 cells transfected with pCD2v-HA or control plasmids, and in swine PBMCs 421 treated with purified CD2v or purified control. Total RNA was harvested at various times post 422 transfection/treatment with RNA-extraction kit (Zymo) and reverse transcribed with MLV-RT (Invitrogen) as previously described (70). IFN-β and ISGs mRNAs were quantified using ABI and 423 424 QuantStudio-3 Real time PCR system (Applied Biosystems, Foster city, CA), Power SYBR Green PCR Master Mix (Applied Bio) and primers sIFNBFw (5'-AGTGCATCCTCCAAATCGC T-3') 425 and sIFNBRv (5'-GCTCATGGAAAGAGCTGTGGT-3') for IFN-B mRNA, and sMX1Fw (5'-426

427 GGCGTGGGAATCAGTCATG-3'), sMX1Rv (5'-AGGAAGGTCTATGAGGGTCAGATCT-428 3'), sOAS1Fw (5'-GAGCTGCAGCGAGACTTCCT-3') and sOAS1Rv (5'-429 TGCTTGACAAGGCGGATGA-3') for ISGs *MX1* and *OAS*. Fold change was calculated by 430 comparison to Empty-HA for each time points. Experiments were conducted with biological 431 triplicates and at least three technical replicates.

432 NF-kB nuclear translocation assay. Involvement of NF-kB in CD2v-mediated upregulation of IFN-ß transcription was assessed by nuclear translocation assays in 1- PK15 cells transfected 433 with pCD2v-HA or control plasmids pORFV120-Flag and pORFV113-Flag, and 2- swine PBMCs 434 treated with purified CD2v or purified control. Cells were fixed at various times post-transfection 435 or-treatment, fixed with 2%-4% paraformaldehyde (PFA) (15 min), and permeabilized with 0.2% 436 triton X100 (10 min). PK15 cells were incubated overnight at 4° C with primary antibody against 437 HA (2367S; Cell Signaling Technology), control Flag (A00187; Genscript) and total NF-KB (8242; 438 439 Cell Signaling Technology), and PBMCs with antibody against total NF-KB. PK15 cells were washed with PBS and incubated with secondary antibodies, goat anti-mouse Alexa fluor 488 440 (A11029; Thermo Scientific) to detect CD2v, ORFV120 and ORFV113, and goat anti-rabbit Alexa 441 fluor 594 (A11012; Thermo Scientific) to detect total NF-κB; for PBMCs, goat anti-mouse Alexa 442 fluor 488. PBMCs were deposited on slides using Shandon cytospin 2 centrifuge (1500 rpm, 1 443 min). Nuclei were counterstained with DAPI. Images were obtained with a A1 Nikon confocal 444 microscope, and the number of cells exhibiting nuclear NF-kB staining was determined in 445 446 randomly selected fields and results were expressed as mean percentage of cells with nuclear NF- $\kappa B$  over three independent experiments. 447

To assess whether inhibition of NF-κB affects IFN-β induction by CD2v, PK15 cells were 448 pretreated with the NF-kB inhibitor parthenolide (InvivoGen) at 1µM final concentration or 449 vehicle control (DMSO) for one hour and transfected with pCD2v-HA in presence or absence of 450 parthenolide (1 µM). At 3 h pt, cultures were fixed, permeabilized, and incubated with primary 451 antibodies against HA and total NF-kB as above, washed, and incubated with secondary 452 antibodies, goat anti-mouse Alexa fluor 488 for CD2v and goat anti-rabbit Alexa fluor 594 for 453 454 total NF-κB. Cells were counterstained with DAPI and percentage NF-κB nuclear translocation in CD2v expressing cells in presence or absence of inhibitor were compared over three independent 455 experiments. 456

The role of CD2v-CD58 interaction in induction of NF- $\kappa$ B nuclear translocation was examined by transfecting PK15 cells with CD58 siRNA or siRNA universal negative control for 24 h, and with pCD2v-HA 24 h later for 3 h. Cultures were fixed, permeabilized and processed for HA and total NF- $\kappa$ B staining as described above. Percentage of NF- $\kappa$ B nuclear translocation in CD2vexpressing cells in which CD58 was knocked down were determined and compared in three independent experiments.

To investigate the effect of anti-CD2v monoclonal antibody mix in CD2v-induced NF- $\kappa$ B activation in swine PBMCs, purified CD2v or purified control were incubated overnight at 4° C with anti-CD2v monoclonal antibodies or anti-ORFV086 antibody or anti-IgG mouse isotype antibody control. PBMCs grown in 96-well plates were treated with pre-incubated CD2v or nonpreincubated CD2v control, processed as above for NF- $\kappa$ B nuclear translocation 1.5 h post treatment, cytospinned, and examined under the confocal microscope. Fields were randomly selected and scored for mean percentage of cells containing nuclear NF-κB over three independent
experiments.

471Interferon Bioassay. To investigate the functional significance of CD2v-induced IFN-β and472ISGs expression, the antiviral state of cells was examined using an IFN bioassay. PK15 cells were473transfected with pCD2v-HA or control plasmids pEmpty-HA or pORFV120-Flag, infected at 12474h, 24 h and 30 h pt with reporter vesicular stomatitis virus expressing GFP (VSVGFP; 50 PFU/well),475fixed with 4% PFA at 16 h post infection, and examined by IFA in a Leica DMI 4000B.

Supernatant obtained from PK15 cultures transfected with pCD2v-HA or control plasmids
pEmpty-HA or pORFV120-Flag or Poly I:C (+ control) were serially diluted and used to treat
PK15 cells. At 30 h post-treatment, cultures were infected with VSV<sup>GFP</sup> (50 PFU/well), and virus
replication was assessed 16 h post-infection.

The antiviral state was assessed in swine lymphocytes as follows. Supernatants from swine PBMCs treated with purified CD2v or purified control for 24 h or supernatant obtained PK15 culture transfected with Poly I:C (+ control) were serially diluted and used to treat fresh PK15 cells. PK15 cells were infected with VSV<sup>GFP</sup> (50 PFU/well) 24 h post treatment, and virus replication was examined 16 h post infection as described.

Flow cytometry. PK15 cells were transfected with pCD2v-HA or control plasmids, and then
infected with VSV<sup>GFP</sup> at 12 h, 24 h and 30 h pt. At 16 h post infection, cells were trypsinized, fixed
with 2% PFA, and washed. GFP mean fluorescence intensity (MFI) was measured using Cytek
Aurora flow cytometer (Cytek Biosciences, Fremont CA).

CD2v activation of the NF-κB pathway was further examined using flow cytometry assay. 489 PK15 cells were transfected with pCD2v-HA or pORFV113-Flag, fixed with 2% PFA at 3 h pt, 490 permeabilized with 0.2% Triton-X100 and incubated with anti-HA (2367S; Cell Signaling 491 Technology), anti-Flag (A00187; Genscript) or anti-phosphorylated NF-kB (S536) (3033S; Cell 492 Signaling Technology) for 45 min on ice. Secondary antibodies were goat anti-rabbit Alexa fluor 493 488 (A11008; Thermo Scientific) for pNF-kB (S536) and goat anti-mouse Alexa fluor 647 494 (A21236; Thermo Scientific) for CD2v and ORFV113. pNF-kB (S536) mean fluorescence 495 intensity (MFI) was examined in cells expressing CD2v or ORFV113. 496

**Co-immunoprecipitation.** To study the interaction between CD2v and CD58, PK15 cells were 497 co-transfected with pEmpty-HA and pCD58-Flag or pCD2v-HA and pCD58-Flag. Whole cell 498 extracts were prepared at 8 h pt using RIPA lysis buffer (Thermo Scientific). Reciprocal co-499 immunoprecipitations were performed using active motif Co-IP kit (Active Motif, Carlsbad, CA) 500 501 in moderate stringency buffer ( IP High buffer [Active Motif] and Protease inhibitor cocktail [Sigma], without detergent and salt) following the manufacturer's protocol. Whole cell extracts 502 503 were incubated with HA (3724S; Cell Signaling Technology) and Flag (A00187; Genscript) 504 antibodies overnight at 4° C, and then pre-washed with 50 µl of protein G agarose bead slurry 505 (Millipore) at 4° C for 2 h. Beads were washed four times with moderate stringency buffer and 506 bound proteins were eluted in Laemmli buffer. Whole cell protein extracts and immunoprecipitated 507 products were examined by SDS-PAGE-Western blot with the appropriate antibodies.

Interaction of CD2v with endogenous human CD58 was evaluated in 293T cells transfected with pCD2v-HA. Reciprocal co-immunoprecipitation was performed using anti-HA (3724S; Cell Signaling Technology) and hu-CD58 (sc20009; Santa Cruz) antibodies following the manufacturer's protocol (Active motif Co-IP kit). Immunoprecipitated products were examined by SDS-PAGE-Western blot with the appropriate antibodies (7074; Cell Signaling Technology,
7076; Cell Signaling Technology).

514 **Co-localization assays.** CD2v-CD58 interaction was examined by confocal microscopy. PK15 515 cells were co-transfected with pCD2v-HA and pCD58-Flag, fixed with 4% PFA 24 h pt, 516 permeabilized with 0.2% Tritron-X 100 and treated with anti-HA (3724S; Cell Signaling 517 Technology) or anti-Flag (A00187; Genscript) primary antibodies. Cells were then washed with PBS and incubated with goat anti rabbit Alexa fluor 594 (A11012; Cell signaling Technology) and 518 anti-mouse Alexa fluor 488 goat (A11029; Cell Signaling Technology) secondary antibodies for 519 1 h. Nuclei were stained with DAPI, and images obtained using A1 Nikon confocal microscope. 520 Similarly, co-localization of CD2v with endogenous human CD58 was investigated in 293T cells. 521 Cells were transfected with pCD2v-HA and IFA was performed using anti-HA (3724S; Cell 522 Signaling Technology) and anti-CD58 (sc20009; Santa Cruz) primary antibodies, and goat anti 523 524 rabbit Alexa fluor 594 (A11012; Cell Signaling Technology) and anti-mouse Alexa fluor 488 goat (A11029) secondary antibodies. 525

- 526 **Statistics.** All statistical analyses were performed using student's t test. Statistically significant 527 difference were indicated as \*, P < 0.05; \*\*, P < 0.01 and NS, not significant.
- 528

## 529 ACKNOWLEDGEMENTS

This work was funded by National Pork Board (grant 13-102) and by the USDA National
Institute of Food and Agriculture (grant 2013 – 67015-21335).

- We thank Hiep Vu (Nebraska Center for Virology, University of Nebraska-Lincoln) for his advice and assistance in generating monoclonal antibodies, Fernando A. Osorio (Nebraska Center for Virology, University of Nebraska-Lincoln) for laboratory support, James F. Lowe (College of Veterinary medicine at University of Illinois at Urbana-Champaign) and Anna Carol Dilger (Department of Animal Sciences at University of Illinois at Urbana-Champaign) for providing swine blood.
- 538
- 539
- 540

## 541 **REFERENCES**

- Montgomery RE. 1921. On A Form of Swine Fever Occurring in British East Africa (Kenya Colony). J Comp Pathol Ther 34:159–191.
- Detray D. 1957. Persistence of viremia and immunity in African swine fever. Am J Vet Res 18:811–816.
- Survey State
  Survey State<
- Anderson EC, Hutchings GH, Mukarati N, Wilkinson PJ. 1998. African swine fever virus infection of the bushpig (Potamochoerus porcus) and its significance in the epidemiology of the disease. Vet Microbiol 62:1–15.
- 5515.Gómez-Villamandos JC, Bautista MJ, Sánchez-Cordón PJ, Carrasco L. 2013. Pathology552of African swine fever: The role of monocyte-macrophage. Virus Res 173:140–149.
- Moulton J, Coggins L. 1968. Comparison of lesions in acute and chronic African swine
  fever. Cornell Vet 58:364–388.
- 555 7. Blome S, Gabriel C, Beer M. 2013. Pathogenesis of African swine fever in domestic pigs

and European wild boar. Virus Res 173:122–130. 556 8. Ramiro-Ibáñez F, Ortega A, Brun A, Escribano JM, Alonso C. 1996. Apoptosis: A 557 mechanism of cell killing and lymphoid organ impairment during acute African swine 558 559 fever virus infection. J Gen Virol 77:2209-2219. Carrasco L, Bautista MJ, Gómez-Villamandos JC, De Las Mulas JM, De Lara FCM, 560 9. Wilkinson PJ, Sierra MA. 1997. Development of microscopic lesions in splenic cords of 561 pigs infected with African swine fever virus. Vet Res 28:93–99. 562 10. Wilkinson PJ, Wardley RC, Williams SM. 1981. African swine fever virus (Malta/78) in 563 564 pigs. J Comp Pathol 91:277–284. 11. Wardley RC, Hamilton F, Wilkinson PJ. 1979. The replication of virulent and attenuated 565 566 strains of African swine fever virus in porcine macrophages. Arch Virol 61:217–225. Mebus C. 1988. African swine fever. Adv Virus Res 35:251-269. 567 12. Carrillo C, Borca M V, Afonso CL, Onisk D V, Rock DL. 1994. Long-term persistent 13. 568 infection of swine monocytes/macrophages with African swine fever virus. J Virol 569 68:580-3. 570 14. Muñoz-Moreno R, Galindo I, Cuesta-Geijo MÁ, Barrado-Gil L, Alonso C. 2015. Host cell 571 572 targets for African swine fever virus. Virus Res 209:118–127. Carrasco L, Chacón-M De Lara F, Martín De Las Mulas J, Gómez-Villamandos JC, 573 15. Hervás J, Wilkinson PJ, Sierra MA. 1996. Virus association with lymphocytes in acute 574 575 African swine fever. Vet Res 27:305–312. 16. Carrasco L, Chacon-M. de Lara F, Martin de las Mulas J, Gomez-Villamandos JC, Perez 576 577 J, Wilkinson PJ, Sierra MA. 1996. Apoptosis in lymph nodes in acute African swine fever. J Comp Pathol 115:415–428. 578 17. Gómez-Villamandos JC, Hervás J, Mendéz A, Carrasco L, Villeda CJ, Wilkinson PJ, 579 Sierra MA. 1995. A pathological study of the perisinusoidal unit of the liver in acute 580 African swine fever. Res Vet Sci 59:146–151. 581 Gómez-Villamandos JC, Hervás J, Moreno C, Carrasco L, Bautista MJ, Caballero JM, 582 18. Wilkinson PJ, Sierra MA. 1997. Subcellular changes in the tonsils of pigs infected with 583 acute African swine fever virus. Vet Res 28:179–189. 584 19. Salguero FJ, Sánchez-Cordón PJ, Núñez A, Fernández de Marco M, Gómez-Villamandos 585 JC. 2005. Proinflammatory cytokines induce lymphocyte apoptosis in acute African swine 586 fever infection. J Comp Pathol 132:289–302. 587 588 20. Gomez-Villamandos JC, Hervas J, Mendez A, Carrasco L, De las Mulas JM, Villeda CJ, Wilkinson PJ, Sierra MA. 1995. Experimental African swine fever: Apoptosis of 589 lymphocytes and virus replication in other cells. J Gen Virol 76:2399–2405. 590 Gómez del Moral M, Ortuño E, Fernández-Zapatero P, Alonso F, Alonso C, Ezquerra A, 591 21. 592 Domínguez J. 1999. African swine fever virus infection induces tumor necrosis factor alpha production: implications in pathogenesis. J Virol 73:2173-2180. 593 594 22. Salguero FJ, Ruiz-Villamor E, Bautista MJ, Sánchez-Cordón PJ, Carrasco L, Gómez-Villamandos JC. 2002. Changes in macrophages in spleen and lymph nodes during acute 595 African swine fever: Expression of cytokines. Vet Immunol Immunopathol 90:11–22. 596 597 23. Fensterl V, Sen GC. 2009. Interferons and viral infections. BioFactors 35:14–20. 598 24. Fensterl V, Chattopadhyay S, Sen GC. 2015. No Love Lost Between Viruses and 599 Interferons. Annu Rev Virol 2:549–572. 600 25. McNab F, Mayer-Barber K, Sher A, Wack A, O'Garra A. 2015. Type I interferons in infectious disease. Nat Rev Immunol 15:87-103. 601

- Katze MG, He Y, Gale M. 2002. Viruses and interferon: A fight for supremacy. Nat Rev
  Immunol 2:675–687.
- Nan Y, Nan G, Zhang YJ. 2014. Interferon induction by RNA viruses and antagonism by
  viral pathogens. Viruses 6:4999–5027.
- 60628.Kim TK, Maniatis T. 1997. The mechanism of transcriptional synergy of an in vitro607assembled interferon- $\beta$  enhanceosome. Mol Cell 1:119–129.
- Lenardo MJ, Fan CM, Maniatis T, Baltimore D. 1989. The involvement of NF-κB in β interferon gene regulation reveals its role as widely inducible mediator of signal
   transduction. Cell 57:287–294.
- 30. Thanos D, Maniatis T. 1995. Identification of the rel family members required for virus induction of the human beta interferon gene. Mol Cell Biol 15:152–164.
- 61331.Sato M, Suemori H, Hata N, Asagiri M, Ogasawara K, Nakao K, Nakaya T, Katsuki M,614Noguchi S, Tanaka N, Taniguchi T. 2000. Distinct and essential roles of transcription615factors IRF-3 and IRF-7 in response to viruses for IFN- $\alpha/\beta$  gene induction. Immunity61613:539–548.
- 617 32. Honda K, Takaoka A, Taniguchi T. 2006. Type I Inteferon Gene Induction by the
  618 Interferon Regulatory Factor Family of Transcription Factors. Immunity 25:349–360.
- 619 33. Chen L, Greene WC. 2004. Shaping the nuclear action of NF-kappaB. Nat Rev Mol Cell
  620 Biol 5:392–401.
- 34. Perkins ND. 2007. Integrating cell-signalling pathways with NF- κ B and IKK function.
  Nat Rev Mol Cell Biol 8:49-62.
- Scheidereit C. 2006. IkappaB kinase complexes: gateways to NF-kappaB activation and transcription. Oncogene 25:6685–6705.
- Baeuerle PA, Baltimore D. 1988. IκB: A specific inhibitor of the NF-κB transcription
  factor. Science 242:540–546.
- 527 37. Li C-CH, Dai R-M, Longo DL. 1995. Inactivation of NF-kB Inhibitor IkB: Ubiquitin528 Dependent Proteolysis and its Degredation Product. Biochem Biophys Res Commun
  529 215:292-2301.
- Moschonas A, Ioannou M, Eliopoulos AG. 2012. CD40 Stimulates a "Feed-Forward" NF κB–Driven Molecular Pathway That Regulates IFN-β Expression in Carcinoma Cells. J
   Immunol 188:5521–5527.
- Balachandran S, Beg AA. 2011. Defining emerging roles for NF-κB in antivirus
  responses: Revisiting the interferon-β Enhanceosome paradigm. PLoS Pathog 7:1–4.
- 40. Bertolusso R, Tian B, Zhao Y, Vergara L, Sabree A, Iwanaszko M, Lipniacki T, Brasier
  AR, Kimmel M. 2014. Dynamic cross talk model of the epithelial innate immune response
  to double-stranded RNA stimulation: Coordinated dynamics emerging from cell-level
  noise. PLoS One 9.
- 41. Zhao J, He S, Minassian A, Li J, Feng P. 2015. Recent advances on viral manipulation of
  NF-κB signaling pathway. Curr Opin Virol 15:103-111.
- de Oliveira VL, Almeida SCP, Soares HR, Crespo A, Marshall-Clarke S, Parkhouse RM.
  2011. A novel TLR3 inhibitor encoded by African swine fever virus (ASFV). Arch Virol
  156:597–609.
- 644 43. Correia S, Ventura S, Parkhouse RM. 2013. Identification and utility of innate immune
  645 system evasion mechanisms of ASFV. Virus Res 173:87–100.
- 44. Wang X, Wu J, Wu Y, Chen H, Zhang S, Li J, Xin T, Jia H, Hou S, Jiang Y, Zhu H, Guo
  K. 2018. Inhibition of cGAS-STING-TBK1 signaling pathway by DP96R of ASFV China

648 2018/1. Biochem Biophys Res Commun 506:437–443.

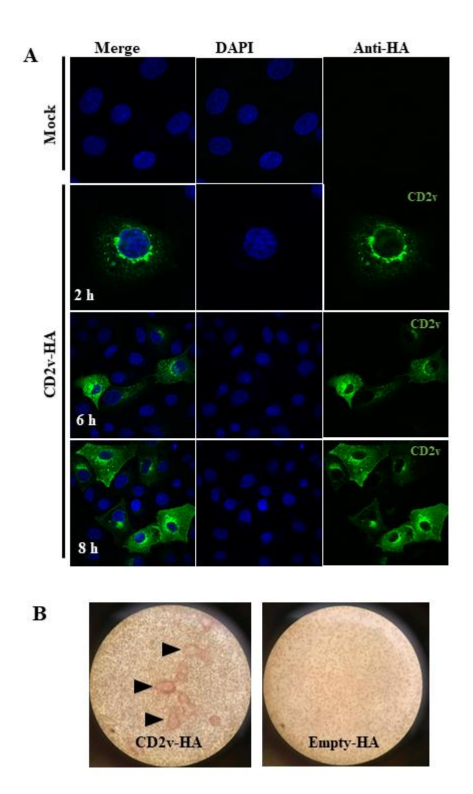
- 45. Revilla Y, Callejo M, Rodri JM, Culebras E, Nogal L, Salas L, Vin E, Fresno M. 1998.
  Inhibition of nuclear factor kappaB activation by a virus-encoded IkappaB-like protein. J
  Biol Chem 273:5405–5411.
- 46. Afonso CL, Piccone ME, Zaffuto KM, Neilan J, Kutish GF, Lu Z, Balinsky CA, Gibb TR,
  Bean TJ, Zsak L, Rock DL. 2004. African swine fever virus multigene family 360 and
  530 genes affect host interferon response. J Virol 78:1858-1864.
- Gil S, Sepúlveda N, Albina E, Leitão A, Martins C. 2008. The low-virulent African swine
  fever virus (ASFV/NH/P68) induces enhanced expression and production of relevant
  regulatory cytokines (IFNα, TNFα and IL12p40) on porcine macrophages in comparison
  to the highly virulent ASFV/L60. Arch Virol 153:1845–1854.
- 48. Zhang F, Hopwood P, Abrams CC, Downing A, Murray F, Talbot R, Archibald A,
  Lowden S, Dixon LK. 2006. Macrophage transcriptional responses following in vitro
  infection with a highly virulent African swine fever virus isolate. J Virol 80:10514-10521.
- García-Belmonte R, Pérez-Núñez D, Pittau M, Richt JA, Revilla Y. 2019. African Swine
  Fever Virus Armenia/07 Virulent Strain Controls Interferon Beta Production through the
  cGAS-STING Pathway. J Virol 93:1–17.
- 50. Portugal R, Leitão A, Martins C. 2018. Modulation of type I interferon signaling by
  African swine fever virus (ASFV) of different virulence L60 and NHV in macrophage
  host cells. Vet Microbiol 216:132–141.
- 668 51. Reis AL, Abrams CC, Goatley LC, Netherton C, Chapman DG, Sanchez-Cordon P, Dixon
  669 LK. 2016. Deletion of African swine fever virus interferon inhibitors from the genome of
  670 a virulent isolate reduces virulence in domestic pigs and induces a protective response.
  671 Vaccine 34:4698–4705.
- 52. Karalyan Z, Zakaryan H, Sargsyan K, Voskanyan H, Arzumanyan H, Avagyan H,
  Karalova E. 2012. Interferon status and white blood cells during infection with African
  swine fever virus in vivo. Vet Immunol Immunopathol 145:551–555.
- 675 53. Golding JP, Goatley L, Goodbourn S, Dixon LK, Taylor G, Netherton CL. 2016.
  676 Sensitivity of African swine fever virus to type I interferon is linked to genes within 677 multigene families 360 and 505. Virology 493:154–161.
- 678 54. González S, Mendoza C, Sánchez-Vizcaino JM, Alonso F. 1990. Inhibitory effect of
  679 African swine fever virus on lectin-dependent swine lymphocyte proliferation. Vet
  680 Immunol Immunopathol 26:71–80.
- 55. Borca M V, Carrillo C, Zsak L, Laegreid WW, Kutish GF, Neilan JG, Burrage TG, Rock
  DL. 1998. Deletion of a CD2-like gene, 8-DR, from African swine fever virus affects viral
  infection in domestic swine. J Virol 72:2881–2889.
- 684 56. Rodríguez JM, Yáñez RJ, Almazán F, Viñuela E, Rodriguez JF. 1993. African swine fever
  685 virus encodes a CD2 homolog responsible for the adhesion of erythrocytes to infected
  686 cells. J Virol 67:5312–5320.
- 57. Borca M V., Kutish GF, Afonso CL, Irusta P, Carrillo C, Brun A, Sussman M, Rock DL.
  1994. An African Swine Fever Virus Gene with Similarity to the T-Lymphocyte Surface
  Antigen CD2 Mediates Hemadsorption. Virology 199:463-468.
- 58. Bockenstedt LK, Goldsmith MA, Dustin M, Olive D, Springer TA, Weiss A. 1988. The
- 691 CD2 ligand LFA-3 activates T cells but depends on the expression and function of the 692 antigen receptor. J Immunol 141:1904–1911.
- 59. Satiimul T. 1988. Accessory Cell-Dependent T-Cell Activation via Involvement of CD2-

LFA-3 Interactions. Scand J Immunol 28:277-284. 694 60. Karmann K, Hughes CCW, Fanslow WC, Pober JS. 1996. Endothelial cells augment the 695 expression of CD40 ligand on newly activated human CD4+T cells through a CD2/LFA-3 696 697 signaling pathway. Eur J Immunol 26:610–617. Diaz-Sanchez D, Chegini S, Zhang K, Saxon A. 2018. CD58 (LFA-3) stimulation 698 61. provides a signal for human isotype switching and IgE production distinct from CD40. J 699 Immunol 153:10-20. 700 62. King PD, Sadra A, Han A, Liu XR, Sunder-Plassmann R, Reinherz EL, Dupont B. 1996. 701 702 CD2 signaling in T cells involves tyrosine phosphorylation and activation of the Tec family kinase, EMT/ITK/TSK. Int Immunol 8:1707-1714. 703 704 63. Lin H, Hutchcroft JE, Andoniou CE, Kamoun M, Band H, Bierer BE. 1998. Association 705 of p59 fyn with the T Lymphocyte Costimulatory Receptor CD2. Binding of the Fyn Src homology (SH)3 domain is regulated by the Fyn SH2 domain. J Biol Chem 273:19914-706 19921. 707 708 64. Le Guiner S, Le Dréan E, Labarrière N, Fonteneau JF, Viret C, Diez E, Jotereau F. 1998. LFA-3 co-stimulates cytokine secretion by cytotoxic T lymphocytes by providing a TCR-709 710 independent activation signal. Eur J Immunol 28:1322-1331. 65. Meinl E, Lengenfelder D, Blank N, Pirzer R, Barata L, Hivroz C. 2000. Differential 711 Requirement of ZAP-70 for CD2-Mediated Activation Pathways of Mature Human T 712 713 Cells. J Immunol 165:3578-3583. Ariel O, Levi Y, Hollander N. 2009. Signal transduction by CD58: the transmembrane 714 66. 715 isoform transmits signals outside lipid rafts independently of the GPI-anchored isoform. Cell Signal 21:1100-1108. 716 Skånland SS, Moltu K, Berge T, Aandahl EM, Taskén K. 2014. T-cell co-stimulation 717 67. through the CD2 and CD28 co-receptors induces distinct signalling responses. Biochem J 718 719 460:399-410. Meier WA, Galeota J, Osorio FA, Husmann RJ, Schnitzlein WM, Zuckermann FA. 2003. 720 68. Gradual development of the interferon-y response of swine to porcine reproductive and 721 respiratory syndrome virus infection or vaccination. Virology 309:18–31. 722 723 69. Parida R, Choi IS, Peterson DA, Pattnaik AK, Laegreid W, Zuckermann FA, Osorio FA. 2012. Location of T-cell epitopes in nonstructural proteins 9 and 10 of type-II porcine 724 reproductive and respiratory syndrome virus. Virus Res 169:13–21. 725 726 70. Diel DG, Delhon G, Luo S, Flores EF, Rock DL. 2010. A Novel Inhibitor of the NF-κB 727 Signaling Pathway Encoded by the Parapoxvirus Orf Virus. J Virol 84:3962–3973. Malogolovkin A, Burmakina G, Tulman ER, Delhon G, Diel DG, Salnikov N, Kutish GF, 728 71. Kolbasov D, Rock DL. 2015. African swine fever virus CD2v and C-type lectin gene loci 729 730 mediate serological specificity. J Gen Virol 96:866-873. Goatley LC, Dixon LK. 2011. Processing and Localization of the African Swine Fever 731 72. 732 Virus CD2v Transmembrane Protein. J Virol 85:3294–3305. 73. Kay-Jackson PC, Goatley LC, Cox L, Miskin JE, Parkhouse RM, Wienands J, Dixon LK. 733 2004. The CD2v protein of African swine fever virus interacts with the actin-binding 734 735 adaptor protein SH3P7. J Gen Virol 85:119-130. 736 74. Ruíz-Gonzalvo F, Coll JM. 1993. Characterization of a soluble hemagglutinin induced in 737 African swine fever virus-infected cells. Virology 196:769-777. 738 75. Monteagudo PL, Lacasta A, López E, Bosch L, Collado J, Pina-pedrero S, Correa-fiz F, 739 Accensi F. 2017. A71ACD2: a New Recombinant Live Attenuated African Swine Fever

740 Virus with Cross-Protective Capabilities. J Virol 91:1–17.

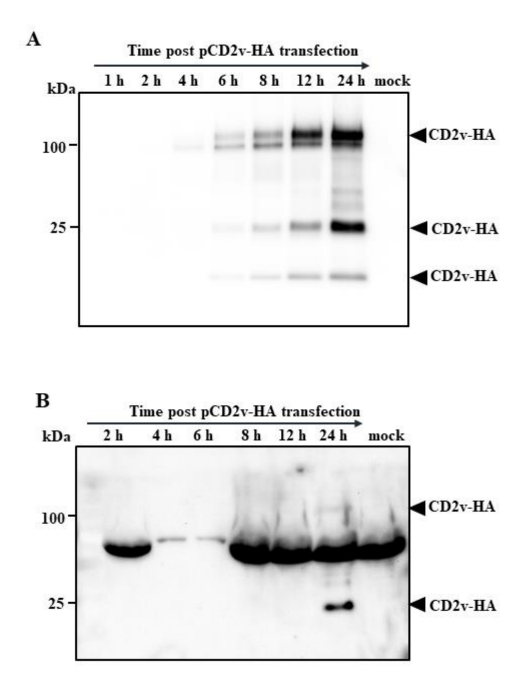
- 741 76. Barbalat R, Lau L, Locksley RM, Barton GM. 2009. Toll-like receptor 2 on inflammatory monocytes induces type i interferon in response to viral but not bacterial ligands. Nat
  743 Immunol 10:1200–1209.
- 744 77. Bauernfeind F, Hornung V. 2009. Tlr2 joins the interferon gang. Nat Immunol 10:1139–
  1141.
- 746 78. Kurt-Jones EA, Popova L, Kwinn L, Haynes LM, Jones LP, Tripp RA, Walsh EE,
  747 Freeman MW, Golenbock DT, Anderson LJ, Finberg RW. 2000. Pattern recognition
  748 receptors TLR4 and CD14 mediate response to respiratory syncytial virus. Nat Immunol
  749 1:398–401.
- 750 79. Georgel P, Jiang Z, Kunz S, Janssen E, Mols J, Hoebe K, Bahram S, Oldstone MBA,
  751 Beutler B. 2007. Vesicular stomatitis virus glycoprotein G activates a specific antiviral
  752 Toll-like receptor 4-dependent pathway. Virology 362:304–313.
- Pérez-Núñez D, García-Urdiales E, Martínez-Bonet M, Nogal ML, Barroso S, Revilla Y,
  Madrid R. 2015. CD2v interacts with Adaptor Protein AP-1 during African swine fever
  infection. PLoS One 10:1–19.
- 81. Borca MV, O'Donnell V, Holinka LG, Risatti GR, Ramirez-Medina E, Vuono EA, Shi J,
  Pruitt S, Rai A, Silva E, Velazquez-Salinas L, Gladue DP. 2020. Deletion of CD2-like
  gene from the genome of African swine fever virus strain Georgia does not attenuate
  virulence in swine. Sci Rep 10:1-8.
- 760 82. Chapman DAG, Tcherepanov V, Upton C, Dixon LK. 2008. Comparison of the genome sequences of non-pathogenic and pathogenic African swine fever virus isolates. J Gen Virol 89:397–408.
- Rowlands RJ, Duarte MM, Boinas F, Hutchings G, Dixon LK. 2009. The CD2v protein
  enhances African swine fever virus replication in the tick vector, Ornithodoros erraticus.
  Virology 393:319–328.
- Kleiboeker SB, Burrage TG, Scoles GA, Fish D, Rock DL. 1998. African swine fever
  virus infection in the argasid host, Ornithodoros porcinus porcinus. J Virol 72:1711–1724.
- Argilaguet JM, Pérez-Martín E, Nofrarías M, Gallardo C, Accensi F, Lacasta A, Mora M,
  Ballester M, Galindo-Cardiel I, López-Soria S, Escribano JM, Reche PA, Rodríguez F.
  2012. DNA Vaccination Partially Protects against African Swine Fever Virus Lethal
  Challenge in the Absence of Antibodies. PLoS One 7:1–11.
- Takamatsu HH, Denyer MS, Lacasta A, Stirling CMA, Argilaguet JM, Netherton CL,
  Oura CA, Martins C, Rodríguez F. 2013. Cellular immunity in ASFV responses. Virus
  Res 173:110–121.
- 87. Burmakina G, Malogolovkin A, Tulman ER, Xu W, Delhon G, Kolbasov D, Rock DL.
  2019. Identification of T-cell epitopes in African swine fever virus CD2v and C-type
  lectin proteins. J Gen Virol 100:259–265.
- 88. Burmakina G, Malogolovkin A, Tulman ER, Zsak L, Delhon G, Diel DG, Shobogorov
  NM, Morgunov YP, Morgunov SY, Kutish GF, Kolbasov D, Rock DL. 2016. African
  swine fever virus serotype-specific proteins are significant protective antigens for African
  swine fever. J Gen Virol 97:1670–1675.
- 782
- 783
- 784
- 785

bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



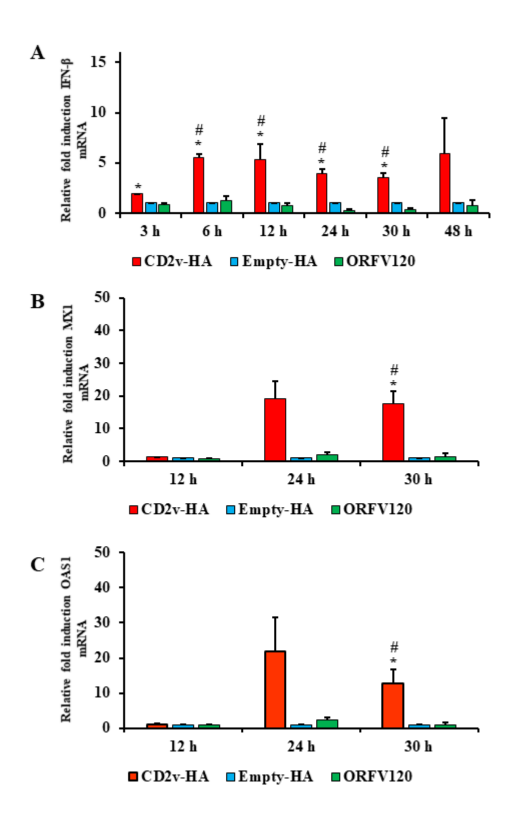
#### **FIG 1. Subcellular localization and characterization of ASFV CD2v protein.** (A) PK15 cells

- 791 were mock transfected or transfected with a plasmid expressing HA-tagged ASFV CD2v, fixed at
- various times post transfection (pt), incubated with anti-HA primary antibody, sequentially
- stained with Alexa fluor 488-labeled secondary antibody and DAPI, and examined by confocal
- 794 microscopy. Results are representative of two independent experiments. (B) CD2v-HA-
- expressing PK15 cells hemadsorbed swine red blood cells (RBCs). PK15 cells were transfected
- with plasmid pCD2v-HA for 24 h, incubated with swine RBCs overnight, and observed with the
- microscope (X100). CD2v-dependent rosetting is indicated by the arrowheads. Results are
- representative of two independent experiments.
- 799



801 802

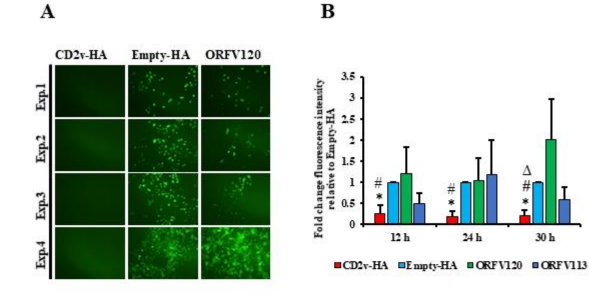
FIG 2. Expression kinetics and secretion of CD2v. (A) PK15 cells mock transfected or 803 804 transfected with plasmid pCD2v-HA were harvested at the indicated times post transfection. Total cell protein extracts were resolved by SDS-PAGE, blotted and incubated with antibodies 805 against HA. Results are representative of two independent experiments. (B) Detection of CD2v 806 in culture supernatant of PK15 cells transiently expressing CD2v. PK15 cells were transfected as 807 above, and supernatants harvested at various times pt. Cleared supernatants (50 µl) were resolved 808 by SDS-PAGE and analyzed by Western blot using antibodies against HA. Results are 809 representative of two independent experiments. 810



816 FIG 3. Induction of IFN-β and ISGs in CD2v-expressing cells. PK15 cells were transfected

- 817 with plasmids pCD2v-HA, pEmpty-HA or pORFV120-Flag, and transcription of IFN-  $\beta$  and
- selected ISGs were assessed by real-time PCR. (A) Fold changes of IFN- $\beta$  levels in cells
- transfected with pCD2v-HA relative to pEmpty-HA and pORFV120-Flag at various times post
- 820 transfection (pt). Results are average mRNA levels from three independent experiments. P-
- values relative to Empty-HA and ORFV120 were 0.007 and 0.1 (3 h); 0.04 and 0.01 (6 h); 0.016
- and 0.017 (12 h); 0.015 and 0.006 (24 h); and 0.019 and 0.006 (30 h), respectively. (B and C)
- Fold changes of ISG mRNA levels at 12 h, 24 h and 30 h pt. Results are average mRNA levels
- from three independent experiments. *P*-values relative to Empty-HA and ORFV120 at 30 h pt
- were 0.02 and 0.018 for MXI (B) and 0.04 and 0.037 for OAS1 (C). \* and # denote statistical
- significance compared to Empty-HA and ORFV120, respectively.
- 827

bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



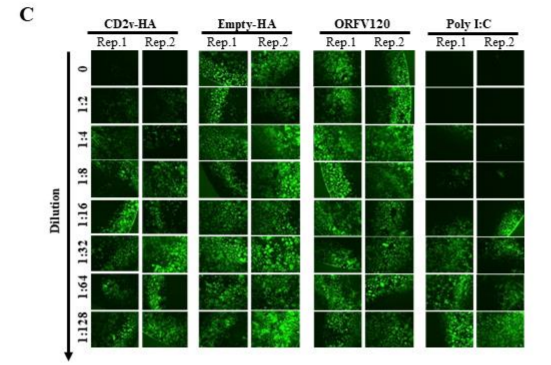


FIG 4. Induction of antiviral state in CD2v-expressing cells. PK15 cells were transfected with

- pCD2v-HA, pEmpty-HA, pORFV120-Flag or pORFV113-Flag. At 12 h, 24 h or 30 h post
- transfection, cultures were infected with vesicular stomatitis virus expressing GFP (VSV<sup>GFP</sup>, 50
- 835 PFU/well). (A) Fluorescence microscopy images taken at 16 h post infection. Note decreased
- 836 VSV replication in cells transfected with pCD2v-HA relative to controls. Results representative
- of four independent experiments. Exp., denotes experimental replicates. (B) Mean GFP
- 838 fluorescence measured by flow cytometry at 16 h post infection. Results are mean values of four
- 839 independent experiments. *P*-values relative to transfection with plasmids pEmpty-HA,
- pORFV120-Flag and pORFV113-Flag were 0.0064, 0.042 and 0.096 (12 h); 0.0002, 0.041 and
- 841 0.07 (24 h); and 0.0002, 0.02 and 0.02 (30 h), respectively. \*, # and  $\Delta$  denote statistical
- significance compared to Empty-HA, ORFV120 and ORFV113, respectively. (C) PK15 cells
- 843 were treated with supernatants obtained from cultures transfected with pCD2v-HA, pEmpty-HA
- or pORFV120-Flag, or with poly I: C, and infected with VSV<sup>GFP</sup> (50 PFU/well) 30 h post
- treatment. Fluorescence microscopy images taken at 16 h post infection are shown. Results
- representative of two independent experiments. Rep., denotes technical replicates.

bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

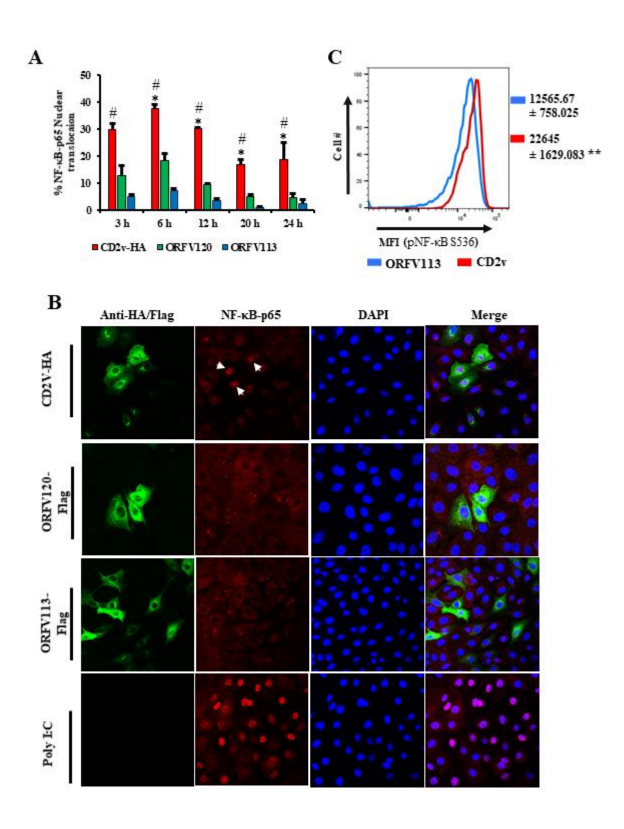
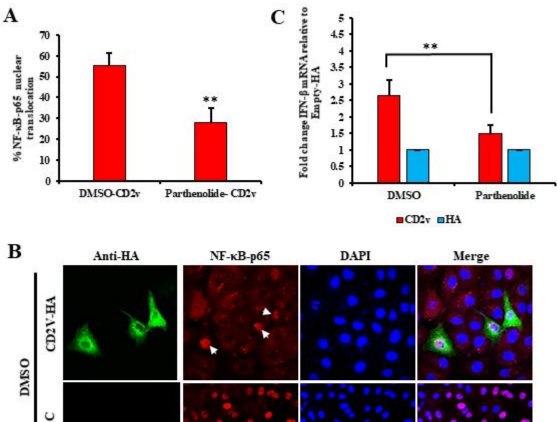


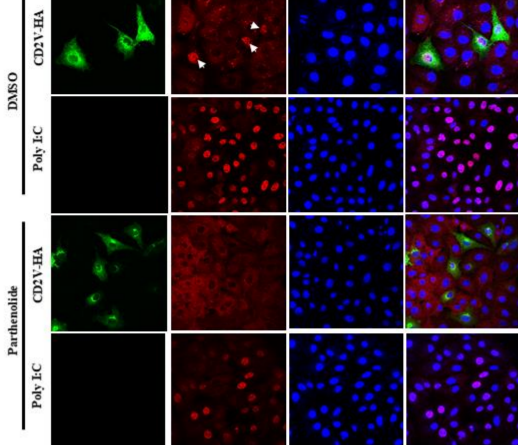


FIG 5. Induction of NF-κB-p65 nuclear translocation in CD2v-expressing cells. PK15 cells
 were transfected with pCD2v-HA, pORFV120-Flag or pORFV113-Flag, processed for

- immunofluorescence using primary antibodies against HA or Flag and NF- $\kappa$ B-p65, and
- secondary antibodies Alexa fluor 488 to detect CD2v, ORFV120 or ORFV113 and Alexa fluor
- 594 to detect NF-κB-p65, and counterstained with DAPI. Cells were counted from 15 random
- 858 fields/slide (approximately 100 cells/slide) and results, shown as percentage of cells with nuclear
- NF- $\kappa$ B-p65, are mean values from three independent experiments. (A) Percentage of NF- $\kappa$ B-p65
- 860 nuclear translocation following the different treatments. *P*-values relative to ORFV120 and
- 861 ORFV113 were 0.056 and 0.009 (3 h); 0.01 and 0.0006 (6 h); 0.019 and 0.02 (12 h); 0.005 and
- 862 0.004 (20 h); and 0.016 and 0.009 (24 h), respectively. \* and # denote statistical significance
- compared to ORFV120 and ORFV113, respectively. (B) Confocal microscopy images showing
   NF-κB-p65 nuclear translocation at 3 h post transfection (pt; arrows). Green, CD2v or ORFV120
- or ORFV113; Red, NF- $\kappa$ B-p65; Blue, DAPI. (C) Mean fluorescence intensity (MFI) of
- phosphorylated NF- $\kappa$ B (S536) fluorescence measured by flow cytometry in CD2v and ORFV113
- expressing cells at 3 h pt. Results representative of three independent experiments. *P*-value
- 868 relative to ORFV113 is 0.0082.
- 869

bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.





#### FIG 6. CD2v expression induces IFN-β transcription in NF-κB dependent manner in PK15

**cells.** PK15 cells pretreated with the NF- $\kappa$ B inhibitor parthenolide (1  $\mu$ M) or DMSO (vehicle

control) for one hour, were transfected with pCD2v-HA, fixed at 3 h post transfection and

processed for immunofluorescence with antibodies against HA and NF- $\kappa$ B-p65. Cells were counted from 15 random fields/slide (approximately 100 cells/slide) (A) Percentage of CD2v-

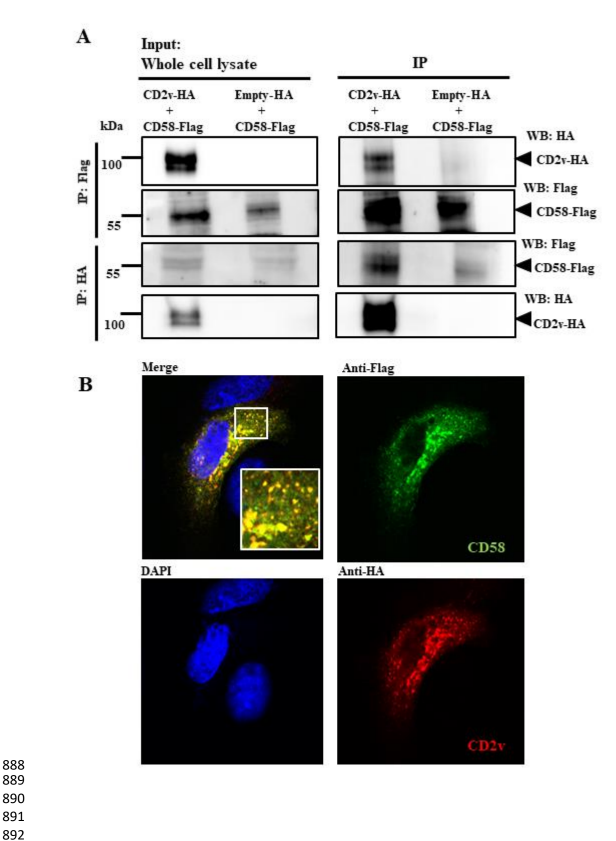
expressing cells with nuclear NF- $\kappa$ B-p65. Results are expressed as mean values from three

independent experiments (P = 0.001). (B) Representative confocal images of cells treated as

above. Green, CD2v; Red, NF- $\kappa$ B-p65; Blue, DAPI. Arrows indicate nuclear NF- $\kappa$ B-p65. (C)

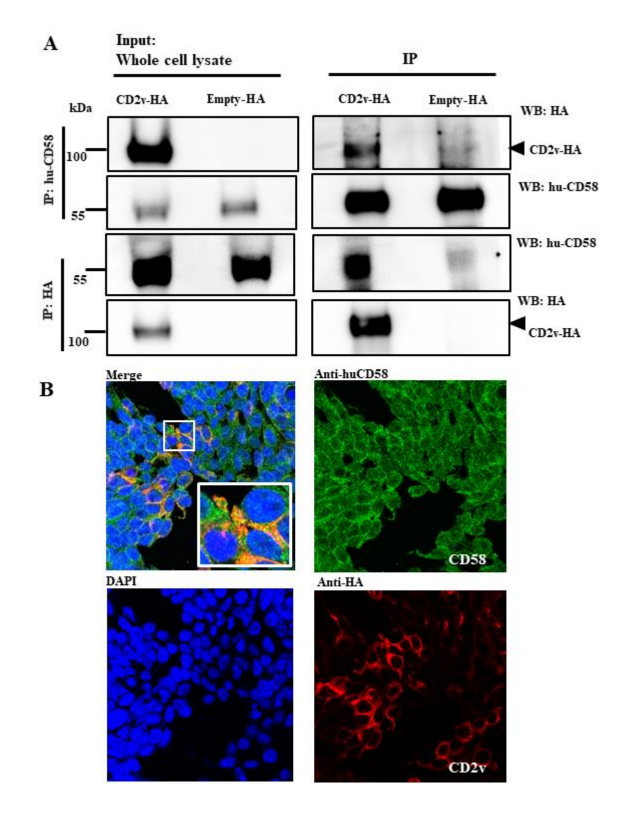
- PK15 cells treated with parthenolide  $(1 \mu M)$  or DMSO for one hour, were transfected with
- pCD2v-HA in presence or absence of parthenolide. Total RNA was extracted at 6 h pt, cDNA
- prepared and IFN-  $\beta$  transcription assessed by RT-PCR. Fold changes relative to Empty-HA and
- data are means from four independent experiments (P = 0.008). (\*, P < 0.05; \*\*, P < 0.01).

bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



#### **FIG 7. Interaction between CD2v and porcine CD58.** (A) For co-immunoprecipitation

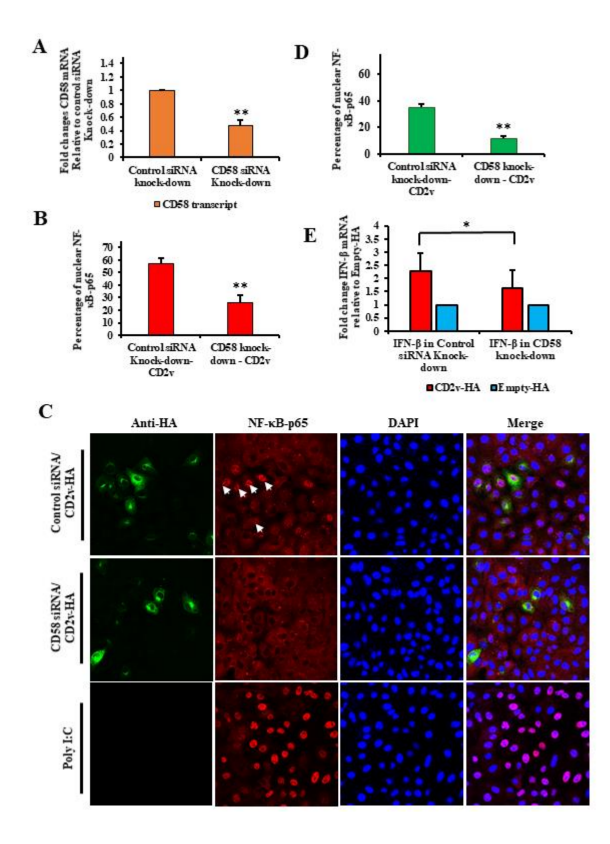
- experiments, PK15 cells were co-transfected with plasmids pCD58-Flag and pCD2v-HA, or
- pCD58-Flag and pEmpty-HA (control) and harvested at 8 h post transfection (pt). Whole cell
- lysates (left) and extracts immunoprecipitated with anti-Flag antibodies (IP, two top panels) or
- anti-HA (IP, two bottom panels) were examined by Western blotting with antibodies directed
- against proteins indicated on the right. Results are representative of three independent
- 899 experiments. (B) For co-localization studies, PK15 cells were co-transfected with pCD58-Flag
- and pCD2v-HA, fixed at 24 h pt, incubated with mouse anti-Flag and rabbit anti-HA primary
- antibodies, washed, and incubated with secondary antibodies (Alexa-fluor 488-labeled anti-
- mouse and Alexa-fluor 594-labeled anti-rabbit). Cells were counterstained with DAPI, and
- examined with the confocal microscope. Results are representative of three independent
- 904 experiments. Insets show magnified areas of the field.



**FIG 8. Interaction between CD2v and human CD58.** (A) For co-immunoprecipitation, 293T

- cells were transfected with pCD2v-HA or pEmpty-HA (control) and harvested at 8 h pt. Whole
- cell lysates (left) and extracts immunoprecipitated with mouse anti-huCD58 antibodies (IP, two
- top panels) or anti-HA (IP, two bottom panels) were examined by Western blotting with
- antibodies directed against proteins indicated on the right. Results are representative of three
- 916 independent experiments. (B) For co-localization, 293T cells were transfected with pCD2v-HA,
- 917 fixed at 24 h pt, incubated with mouse anti-huCD58 and rabbit anti-HA primary antibodies,
- 918 washed, and incubated with Alexa-fluor 488-labeled anti-mouse and Alexa-fluor 594-labeled
- anti-rabbit secondary antibodies. Cells were counter stained with DAPI, and examined with the
- 920 confocal microscope. Results are representative of three independent experiments. Insets show
- 921 magnified areas of the field.
- 922

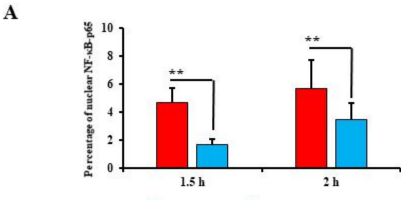
bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



#### 928 FIG 9. CD2v-CD58 interaction affects CD2v-mediated NF-kB-p65 nuclear translocation

- **and IFN-\beta induction.** (A) siRNA knock down of CD58. PK15 cells were transfected with CD58
- siRNA or siRNA universal negative control, total RNA was extracted at 24 h pt, cDNA prepared
- and CD58 transcription assessed by RT-PCR. Results are mean of five independent experiments
- 932 (P = 0.0002). (B) NF- $\kappa$ B-p65 nuclear translocation. PK15 cells were sequentially transfected
- 933 with CD58 siRNA or siRNA universal negative control and pCD2v-HA or pEmpty-HA, fixed at 3 h pt, and processed for detection of NF- $\kappa$ B-p65 by immunofluorescence. Cells were counted
- 3 h pt, and processed for detection of NF- $\kappa$ B-p65 by immunofluorescence. Cells were counted from 15 random fields/slide (approximately 100 cells/slide) and results are shown as percentage
- 936 of CD2v-expressing cells with nuclear NF- $\kappa$ B-p65. Results are mean of three independent
- experiments (P = 0.0035). (C) Percentage of CD2v-expressing cells with strong nuclear NF- $\kappa$ B-
- p65 translocation (P = 0.0015). (D) Representative images of NF- $\kappa$ B-p65 nuclear translocation
- under conditions outlined in A. Green, CD2v; Red, NF-κB-p65; Blue, DAPI. Arrows indicate
- 940 nuclear NF- $\kappa$ B-p65. (E) IFN- $\beta$  induction. PK15 cells were sequentially transfected with CD58
- siRNA or siRNA universal negative control, and pCD2v-HA or pEmpty-HA. Total RNA was
- extracted at 6 h pt, cDNA prepared and transcription of IFN-β assessed by RT-PCR. Fold
- 943 changes are relative to Empty-HA and data are mean mRNA levels from eight independent
- 944 experiments (P = 0.025). (\*, P < 0.05 and \*\*, P < 0.01).

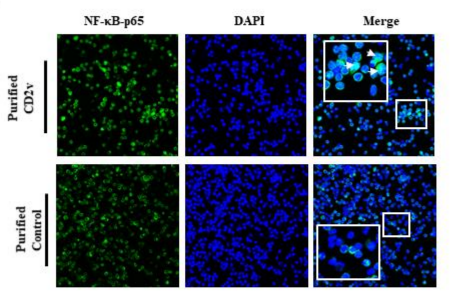
bioRxiv preprint doi: https://doi.org/10.1101/2020.11.06.372417; this version posted November 8, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

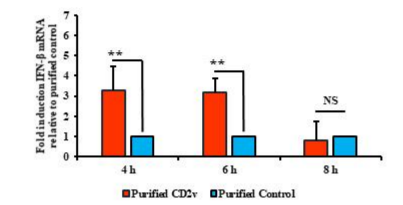






С



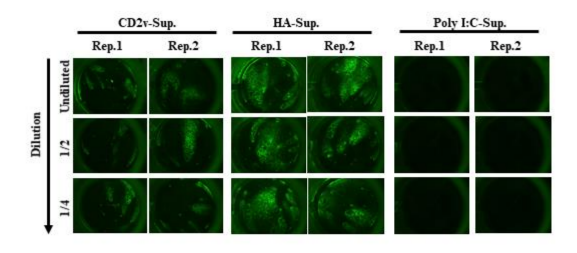




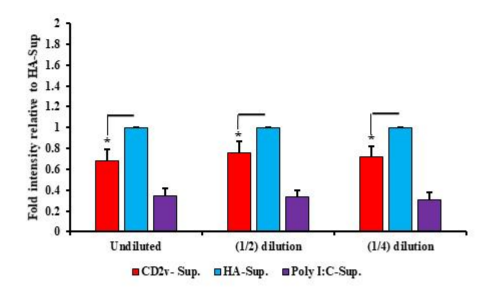
#### 951 FIG 10. Purified CD2v induces NF-kB-p65 nuclear translocation and IFN-β transcription

- 952 in swine PBMCs. Purified CD2v, purified control, and swine PBMCs were obtained as indicated
- in materials and methods. (A) Swine PBMCs were treated with purified CD2v or purified
- control, fixed at 1.5 h and 2 h post treatment, sequentially incubated with anti-NF-κB-p65
- primary antibody and Alexa fluor 488-labeled secondary antibody, stained with DAPI,
- 956 cytospined, and examined by confocal microscopy. Cells were counted from 15 random
- 957 fields/slide (approximately 2000 cells/slide) and results are shown as percentage of cells with
- 958 nuclear NF- $\kappa$ B-p65. Results are mean of four independent experiments (1.5 h, P = 0.01; 2 h, P =
- 959 0.012). (B) Representative confocal images of NF-κB-p65 nuclear translocation under conditions
- outlined in A. Green, NF-κB-p65; Blue, DAPI. Insets show magnified areas of the field. Arrows
- 961 indicate nuclear NF- $\kappa$ B-p65. (C) IFN- $\beta$  induction. Total RNA was harvested at 4 h, 6 h and 8 h
- post treatment and IFN- $\beta$  transcription was assessed by RT-PCR. Fold changes are relative to
- purified control and data are mean mRNA levels of seven independent experiments (4 h, P =
- 964 0.016; 6 h, *P* =0.002). (\*, *P*<0.05 and \*\*, *P*<0.01).



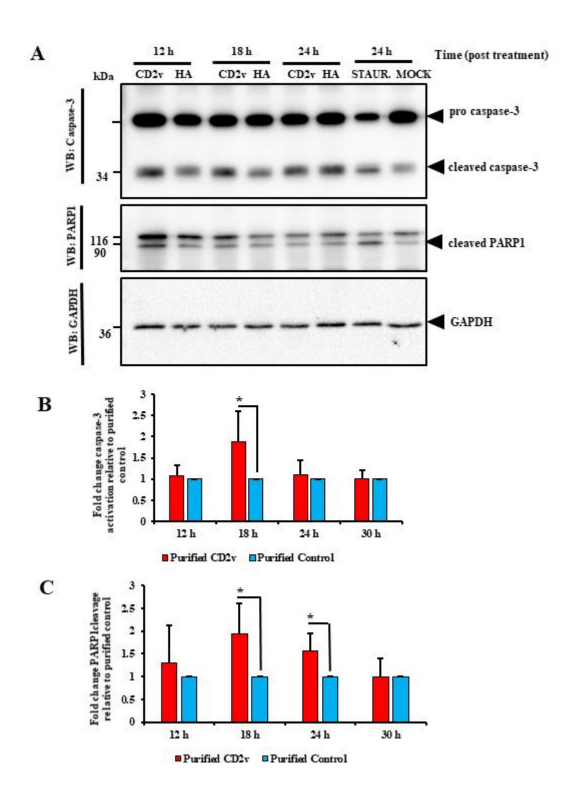


B



#### 971 FIG 11. Antiviral activity of supernatants from CD2v -treated swine PBMCs. Swine PBMCs

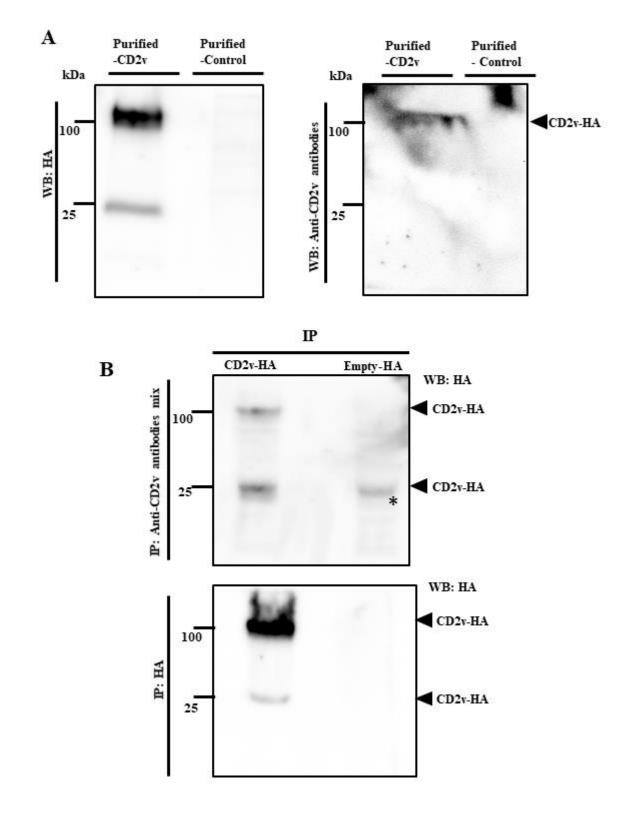
- were treated with purified CD2v protein or purified control, and supernatants collected 24 h post
- treatment. Supernatant collected from PK15 transfected with Poly I:C (positive control). PK15
- grown in 96 well plates were treated with different dilutions of PBMC supernatants for 24 h and
- 975 subsequently infected with VSV<sup>GFP</sup> (50 PFU/well) for 16 h. (A) Representative fluorescence
- 976 images of PK15 fixed at 16 h post VSV<sup>GFP</sup> infection. Green, VSV<sup>GFP</sup>. Rep., denotes technical
- 977 replicates. (B) Intensity (mean gray value) measured by ImageJ. Fold changes are relative to
- 978 purified control and data are mean of four independent experiments. *P*-values relative to purified
- control for undiluted, 1:2 diluted and 1:4 diluted supernatants were 0.014, 0.033 and 0.017,
- 980 respectively (\*, *P*<0.05).
- 981





**FIG 12. Purified CD2v induces apoptosis in swine PBMC** (A) Swine PBMC were treated with purified CD2v, purified control, or staurosporine (positive control) and whole cell lysates were obtained at 12 h, 18 h, 24 h and 30 h post treatment, resolved by SDS-PAGE, blotted, and probed with antibodies against Caspase-3, PARP1 and GAPDH. (B) Densitometric analysis showing the fold change in caspase-3 activation relative to purified control treatment, the caspase-3 results are mean values of six independent experiments (18 h, P = 0.042). (C) Densitometric analysis

- showing fold changes in PARP1 cleavage relative to purified control treatment. Results are mean
- values of six independent experiments (18 h, P = 0.026; 24 h, P = 0.018). (\*, P < 0.05).





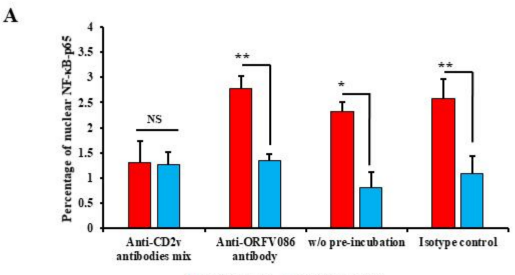
# 1001 FIG 13. Monoclonal antibodies generated against ASFV CD2v identifies CD2v in Western

**blots.** (A) Purified CD2v or purified control, were resolved by SDS-PAGE, blotted, and probed

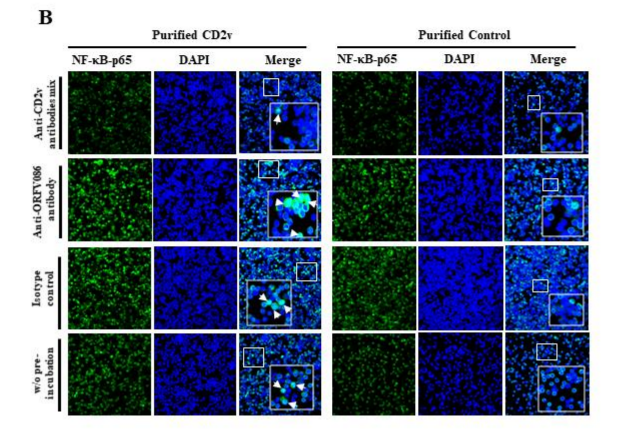
1003 with anti-CD2v monoclonal antibodies (A4, C4, C3 and F2; top right) or probed with anti-HA

antibody (control; top left). Results are representative of three independent experiments. (B)
 Whole cell extracts were immunoprecipitated with anti-CD2v monoclonal antibodies (upper

- 1006 blot) or anti-HA antibody (control; lower blot), resolved with SDS-PAGE, and probed with anti-
- 1007 HA antibody. Results are representative of three independent experiments. \* denotes light chain
- 1008 band.
- 1009

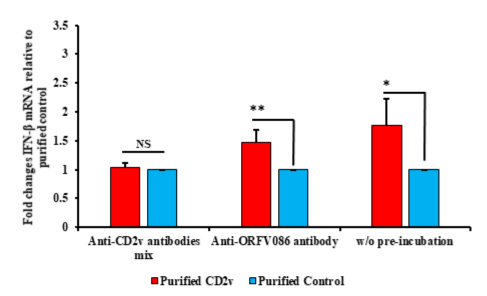






## 1015 FIG 14. Monoclonal antibodies against ASFV CD2v inhibit CD2v-induced NF-кВ

- 1016 activation in swine PBMCs. (A) Purified CD2v or purified control pre-incubated with anti-
- 1017 CD2v monoclonal antibody mix, anti-ORFV086 antibody or isotype control antibody were used
- to treat swine PBMCs for 1.5 h, and cells were processed for NF-κB-p65 staining by indirect
- 1019 immunofluorescence as explained in materials and methods. Control cells were treated with
- 1020 CD2v or purified control that have not been pre-incubated with antibodies. Cells were counted
- 1021 from 15 random fields/slide (approximately 2500 cells/slide). Results, shown as percentage of
- 1022 cells with nuclear NF- $\kappa$ B-p65, are mean values from three independent experiments (for %
- 1023 nuclear NF- $\kappa$ B-p65 for anti-CD2v antibody mix relative to anti-ORFV086, P = 0.012; without
- 1024 pre-incubation, P = 0.031 and for isotype control, P = 0.03. (\*, P < 0.05; \*\*, P < 0.01) (B)
- 1025 Representative confocal images of NF-κB-p65 nuclear translocation. Green, NF-κB-p65; Blue,
- 1026 DAPI. Arrows indicate nuclear NF- $\kappa$ B-p65. Insets show magnified areas of the field.
- 1027
- 1028
- 1029
- 1030
- 1031
- 1032
- 1032
- 1033



# 1035 Fig 15. Monoclonal antibodies against ASFV CD2v inhibits CD2v-induced IFN-β

**transcription in swine PBMC cultures**. Swine PBMCs were treated with purified CD2v or

1037 purified control pre-incubated with anti-CD2v monoclonal antibody mix or anti-ORFV086

antibody, or with the proteins without previous incubation with antibodies. Total RNA was
 harvested at 6 h post-treatment, and IFN-β transcription was assessed by RT-PCR. Fold changes

are relative to purified control and data are mean mRNA levels from five independent

1041 experiments. *P*-value for IFN- $\beta$  fold induction for anti-CD2v antibody mix compared to anti-

1042 ORFV086 is 0.014 and without pre-incubation is 0.028 (\*, *P*<0.05 and \*\*, *P*<0.01).

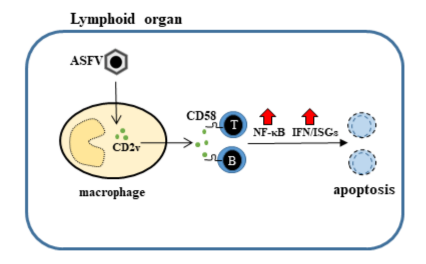
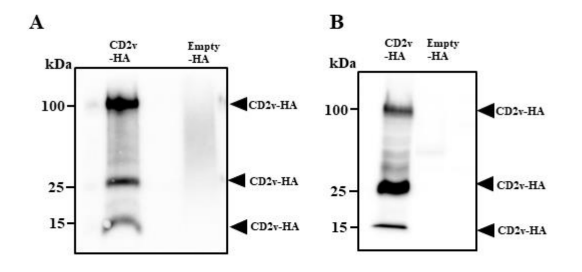


FIG 16. Proposed mechanism of action of ASFV CD2v. CD2v secreted from ASFV-infected
 macrophages interacts with by-stander lymphocytes via CD58. This interaction promotes NF-κB

1061 activation and induction of IFN- $\beta$  and ISGs, which lead to lymphocyte apoptosis.

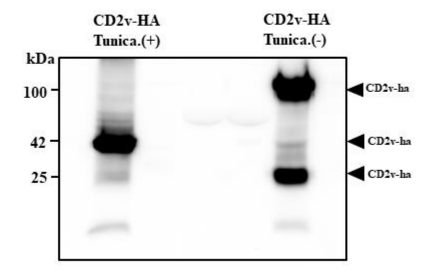


**FIG S1. Expression of CD2v in 293T and Vero cells.** 293T (A) and Vero cells (B) mock

1065 transfected or transfected with plasmid pCD2v-HA were harvested at 24 h post transfection.

1066 Total cell protein extracts were resolved by SDS-PAGE, blotted and incubated with antibodies

against HA. Results are representative of two independent experiments.



1069

1070 **FIG S2. CD2v expression in presence of tunicamycin**. Pk15 cells transfected with pCD2v-HA. 1071 Five hours post-transfection, media was replaced with complete growth media containing  $1\mu$ g/ml 1072 tunicamycin and incubated for 24 hr. Total cell extracts were resolved by SDS-PAGE, blotted 1073 and incubated with antibodies against HA. Results are representative of two independent 1074 experiments.