







# Structural mechanism of CRL4-instructed STAT2 degradation via a novel cytomegaloviral DCAF receptor

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## **Abstract**

Human cytomegalovirus (CMV) is a ubiquitously distributed pathogen whose rodent counterparts such as mouse and rat CMV serve as common infection models. Here, we conducted global proteome profiling of rat CMV-infected cells and uncovered a pronounced loss of the transcription factor STAT2, which is crucial for antiviral interferon signalling. Via deletion mutagenesis, we found that the viral protein E27 is required for CMV-induced STAT2 depletion. Cellular and in vitro analyses showed that E27 exploits host-cell Cullin4-RING ubiquitin ligase (CRL4) complexes to induce poly-ubiquitylation and proteasomal degradation of STAT2. Cryoelectron microscopy revealed how E27 mimics molecular surface properties of cellular CRL4 substrate receptors called DCAFs (DDB1- and Cullin4-associated factors), thereby displacing them from the catalytic core of CRL4. Moreover, structural analyses showed that E27 recruits STAT2 through a bipartite binding interface, which partially overlaps with the IRF9 binding site. Structurebased mutations in M27, the murine CMV homologue of E27, impair the interferon-suppressing capacity and virus replication in mouse models, supporting the conserved importance of DCAF mimicry for CMV immune evasion.

**Keywords** cullin-RING ubiquitin ligases; cytomegalovirus; interferon; ubiquitin–proteasome system; viral DCAF

Subject Categories Microbiology, Virology & Host Pathogen Interaction;
Post-translational Modifications & Proteolysis; Structural Biology
DOI 10.15252/embj.2022112351 | Received 11 August 2022 | Revised 15
December 2022 | Accepted 21 December 2022 | Published online 10 February

The EMBO Journal (2023) 42: e112351

# Introduction

Human cytomegalovirus (HCMV; Human herpesvirus 5) is the prototypic member of the *Betaherpesvirinae* and a ubiquitous human pathogen. More than three quarters of the global population are latently infected with HCMV (Virgin *et al*, 2009). HCMV infections usually progress subclinically in healthy adults, but cause morbidity and mortality in individuals with immature, impaired or senescent immunity, such as congenitally infected newborns, AIDS patients, transplant recipients and the elderly (Griffiths & Reeves, 2021). During host adaptation, cytomegaloviruses became strictly speciesspecific. Therefore, meaningful *in vivo* experiments cannot be performed with HCMV in unmodified small animal models. This roadblock for basic research and preclinical development is circumvented by studying closely related betaherpesviruses such as mouse cytomegalovirus (MCMV; Murid herpesvirus [MuHV] 1) and rat cytomegaloviruses (RCMVs; MuHV-2 and MuHV-8; Voigt *et al*, 2007;

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Geyer *et al*, 2015; Brizic *et al*, 2018) in their corresponding host species. Since MuHV infections represent some of the few genuine virus infection models in which the animal host is infected with its natural pathogen (Becker *et al*, 2007), they became well-established and broadly applied models in virology and immunology, enabling insights regarding general phenomena of virus pathogenesis and host immunity (Reddehase & Lemmermann, 2018).

Upon encounter of pathogens such as cytomegaloviruses, cytokines called interferons (IFNs) are expressed (Panne et al, 2007; Schneider et al, 2008; Ivashkiv & Donlin, 2014), which elicit potent antiviral activity (Isaacs & Lindenmann, 1957) by changing the cellular transcriptome and proteome (Trilling et al, 2013; Megger et al, 2017). In addition to the antiviral activity executed within infected host cells, IFNs also orchestrate cellular immune responses (Baranek et al, 2012). IFNs bind to cell surface receptors that are associated with janus kinases (JAK), which then phosphorylate intracellular receptor chains, generating docking sites for the signal transducer and activator of transcription (STAT; Darnell et al, 1994). After phosphorylation by JAKs, STATs homo- and heterodimerise and form transcription factor complexes that translocate into the nucleus where they induce the expression of IFN-stimulated genes (ISGs; Levy & Darnell, 2002). Type I IFNs (IFN-I) and type III IFNs (IFN-III) signal through STAT1/STAT2 heterodimers, which recruit IFN regulatory factor 9 (IRF9) to generate the active IFN-stimulated gene factor 3 (ISGF3) that binds IFN-stimulated response elements (ISRE) in the nucleus (Qureshi et al, 1995; Horvath et al, 1996; Kotenko et al, 2003). IFNγ, the only known type II IFN (IFN-II), mainly signals via phosphorylated STAT1 homodimers that recognise gammaactivated DNA sequences (GAS) and express a distinct set of ISGs. Accordingly, STAT2 is crucial for IFN-I and IFN-III signalling. Furthermore, STAT2 directly and indirectly affects IFN-II signalling, for example as part of IFNγ-induced ISGF3 complexes (Matsumoto et al, 1999; Zimmermann et al, 2005; Trilling et al, 2011, 2013; Le-Trilling et al, 2018), as constituent of unphosphorylated IRF9-STAT2 heterodimers (Blaszczyk et al, 2015; Platanitis et al, 2019), and by regulating STAT1 expression levels (Park et al, 2000; Hambleton et al, 2013; Ho et al, 2016). Accordingly, animals and humans lacking STAT2 are severely immunosuppressed and show an exaggerated susceptibility to infections (Park et al, 2000; Hambleton et al, 2013; Gowen et al, 2017; Le-Trilling et al, 2018).

The pivotal role of STAT2 for IFN signalling and innate immunity shaped virus evolution. Several relevant pathogens escape from an IFN-induced antiviral state by targeting STAT2. For example, Zika virus (ZIKV; Grant *et al*, 2016), Dengue virus (DENV; Jones *et al*, 2005; Ashour *et al*, 2009), paramyxoviruses (Parisien *et al*, 2001, 2002b), MCMV (Zimmermann *et al*, 2005; Trilling *et al*, 2011) and HCMV (Le-Trilling *et al*, 2020) induce STAT2 degradation along the ubiquitin–proteasome pathway (UPS). Intriguingly, the responsible virus proteins act as adapters that place STAT2 into the sphere of influence of cellular E3 ubiquitin ligases (E3s; Ulane & Horvath, 2002; Le-Trilling & Trilling, 2020; Barik, 2022).

The MCMV protein M27 was the first CMV STAT2 antagonist to be described (Zimmermann *et al*, 2005). MCMV mutants lacking *M27* are severely attenuated *in vivo* (Zimmermann *et al*, 2005; Le-Trilling *et al*, 2018) due to the inability to interfere with STAT2-dependent IFN signalling as evident from the restored replication in STAT2-deficient mice (Le-Trilling *et al*, 2018). Mechanistically, M27 binds DDB1, a component of host Cullin4-RING E3s (CRL4),

to induce poly-ubiquitination of STAT2, resulting in its rapid proteasomal degradation (Trilling *et al*, 2011; Landsberg *et al*, 2018). While the aforementioned complex formation has been confirmed in several models and by different techniques, direct binding of M27 to DDB1 and/or STAT2 in absence of other proteins has to our knowledge not been documented yet. This is relevant since paramyxoviral IFN antagonists bind one STAT and induce degradation of another, for example STAT1 is needed by human parainfluenza virus 2 to target STAT2, while STAT2 is required for both simian virus 5 (SV5) and mumps V proteins to target STAT1 (Ulane *et al*, 2005). Structural details regarding the ternary interactions between CRL4, STAT2 and CMV antagonists are currently not available.

CRL4, similar to other CRLs, are modular complexes, comprising a core formed by the cullin scaffold (CUL4A or CUL4B) and a catalytic RING domain subunit (RBX1). By the adapter protein DDB1, the core is connected to various exchangeable substrate receptors called DCAFs, forming various cellular CRL4 E3s with different specificities (Zimmerman et al, 2010). CRLs are regulated by cycles of activation through modification of the cullin C terminus with NEDD8 ("neddylation"), followed by de-neddylation of substratefree CRLs and substrate receptor exchange (Goldenberg et al, 2004; Cavadini et al, 2016; Baek et al, 2020; Reichermeier et al, 2020; Harper & Schulman, 2021). Among CRLs, CRL4 exhibits unique features like the presence of the co-adaptor DDA1, which stabilises at least certain substrate receptors on CRL4 by bridging them to DDB1, as in the case of DCAF15 (Olma et al, 2009; Shabek et al, 2018; Du et al, 2019; Bussiere et al, 2020; Faust et al, 2020), and the existence of a flexible hinge in the DDB1 adapter that allows for rotation of the catalytic subunit around the substrate receptor, thus creating a broad ubiquitylation zone to accommodate substrates of different size and shape (Fischer et al, 2011, 2014; Banchenko et al, 2021). Most likely, the latter property makes CRL4 a frequent target for viral exploitation to steer antiviral host factors efficiently into the ubiquitin-proteasome pathway. In addition to CMV M27, there are further examples of viral CRL4 binders that induce host protein degradation such as CMV RL1 (Nobre et al, 2019; Nightingale et al, 2022), UL35 (Salsman et al, 2012), UL145 (Le-Trilling et al, 2020), hepatitis B virus X (HBx; Decorsiere et al, 2016), paramyxovirus V—the first viral CRL4 hijacker to be structurally analysed (Precious et al, 2005; Li et al, 2006), and the accessory proteins Vpx and Vpr from immunodeficiency viruses (Schwefel et al, 2014, 2015; Greenwood et al, 2019; Banchenko et al, 2021). Mechanistic studies of these viral CRL4 subversion processes yield direct molecular insight into virus adaptation to host replication barriers, advancing the understanding of virus pathology and potentially paving the way for therapeutic opportunities (Le-Trilling et al, 2016; Becker et al, 2019).

Here, we pursued a proteomic approach to assess RCMV-dependent protein regulation and identified STAT2 as one of the most thoroughly downregulated factors. The RCMV protein E27 was necessary for the reduction of STAT2 protein levels by a process sensitive to the inhibition of the proteasome or CRLs. Moreover, biochemical reconstitution demonstrated that E27 was sufficient for the recruitment of STAT2 to the DDB1 adapter protein of host CRL4 complexes. Cryogenic electron microscopy (cryo-EM) structural analyses revealed that E27 replaces endogenous CRL4 substrate receptors, resulting in proper positioning of STAT2 for poly-ubiquitylation and subsequent proteasomal destruction to inhibit host IFN signalling.

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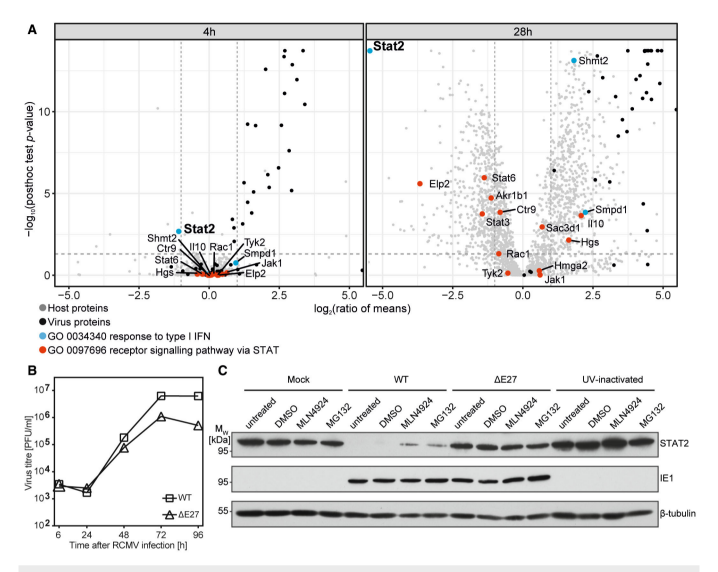


Figure 1. Host protein regulation during RCMV-E infection and the role of E27.

- A Mass spectrometric analyses of protein abundance changes upon RCMV-E infection, compared with mock-infected samples. Host and virus proteins are indicated as grey and black dots, respectively. Host proteins involved in the type I IFN response and the STAT signalling pathway are highlighted in blue and red.
- B Replication analysis of WT and  $\Delta$ E27-RCMV-E in REF. Symbols indicate mean values (pooled data of two biological replicates each titrated in triplicates).
- C Western blot analysis of STAT2 protein levels in mock-, WT-RCMV-E-,  $\Delta$ E27-RCMV-E- or UV-inactivated RCMV-E-infected REF. 24 h p.i., cells were either left untreated or treated for 5.5 h with DMSO, 2.5  $\mu$ M MLN4924 for blocking CRL activity or 10  $\mu$ M MG132 for proteasome inhibition. IE1 and  $\beta$ -tubulin served as infection and loading controls, respectively.

Source data are available online for this figure.

# Results

## RCMV-E E27 downregulates host STAT2

To uncover RCMV-induced proteome changes, rat embryo fibroblasts (REF) were infected with RCMV-England (RCMV-E). The protein content was analysed after 4 and 28 h by liquid chromatography coupled to mass spectrometry (LC–MS) and compared with mock-infected controls. In each condition, more than 2,300 proteins were quantified in six replicates (Appendix Fig S1A). To our knowledge, this represents the first proteome profiling of RCMV-infected cells. Gene ontology (GO) analysis of the data revealed that only very few ISGs

were expressed (Fig 1A). This under-representation of ISGs among RCMV-induced host proteins strongly suggested the expression of potent RCMV-encoded IFN antagonists. In order to obtain mechanistic insights, we further analysed the relative abundance of IFN signalling proteins. The expression levels of JAK1 and TYK2 kinases were similar to mock-infected cells at both time points (Fig 1A; Appendix Fig S1B). Conversely, STAT2 protein levels already started to diminish significantly 4-h postinfection and became virtually undetectable after 28 h of infection (Fig 1A; Appendix Fig S1B), indicating a rapid and sustained RCMV-induced loss of STAT2.

Next, we sought to identify the RCMV-E-encoded factor responsible for the reduction of STAT2 levels. Inspection of the RCMV-E

genome (Geyer et al, 2015) revealed 56% sequence identity of E27 to M27, the MCMV-encoded factor inducing STAT2 degradation in MCMV infection. To test whether E27 accounts for STAT2 downregulation, an RCMV-E mutant lacking E27 was constructed by CRISPR/ Cas9-mediated mutagenesis resulting in an E27-deficient RCMV-E ("ΔE27"). A virus replication analysis revealed a decreased titre of the  $\Delta$ E27 virus mutant in REF 48-h to 96-h postinfection (p.i.), compared with the wild type ("WT"), demonstrating reduced replication fitness upon E27 deletion (Fig 1B). Next, REF were infected with wild type and  $\Delta$ E27 virus and STAT2 expression was assessed by immunoblot analysis 24 h p.i. In accordance with the MS analysis, WT infection caused a complete loss of the STAT2 signal, whereas the  $\Delta E27$  virus failed to reduce STAT2 protein concentrations (Fig 1C). Moreover, the WT-mediated loss of STAT2 proved to be sensitive to treatment with MG132, a proteasome inhibitor, and MLN4924 (Fig 1C), an inhibitor of the NEDD8-activating enzyme, which shuts off cellular neddylation processes that are essential for CRL activity. These findings strongly suggested an involvement of the ubiquitin-proteasome system in E27-induced STAT2 downregulation and furthermore indicated the involvement of host CRLs.

#### E27 recruits STAT2 to CRL4 in vitro

These results prompted us to explore the physical association of E27 with STAT2 and DDB1. Most viral STAT2 antagonists are known to be species-specific (Parisien et al, 2002a; Ashour et al, 2010; Yoshikawa et al, 2019). Therefore, we cloned and purified Rattus norvegicus (rn) STAT2 for these studies. Conversely, hsDDB1 and rnDDB1 are extremely similar (1,128 of 1,140 amino acids are identical, and the few changes are located in the  $\beta$ -propeller domain B [BPB] of DDB1 or in disordered loops and regions not involved in intermolecular interactions). In analogy to various studies by us and others concerning DDB1 binders (Angers et al, 2006; Li et al, 2006; Fischer et al, 2011, 2014; Banchenko et al, 2021), we applied hsDDB1, and a variant that lacks the β-propeller (BP) domain B (DDB1 $_{\Delta BPB}$ ). The incubation of DDB1 $_{\Delta BPB}$  with equimolar amounts of STAT2, followed by analytical gel filtration (GF) chromatography, resulted in elution in separate peaks at the same elution volumes as the individual proteins in isolation (Fig 2A, blue, magenta and grey traces). This argues against an intrinsic association of DDB1 $_{\Delta BPB}$ and STAT2. By contrast, E27 formed stable binary complexes with STAT2 (Fig 2B, orange trace) and DDB1<sub>ABPB</sub> (brown trace). Incubation of E27 together with DDB1  $_{\Delta BPB}$  and STAT2 resulted in the elution of all three components in the same peak at an earlier elution volume than the isolated components (Fig 2A, purple trace). This showed the formation of a stable ternary DDB1<sub>ABPB</sub>/E27/ STAT2 protein complex, demonstrating that E27 is sufficient to bridge STAT2 to DDB1. Similar results were obtained upon incubation with full-length DDB1 (Appendix Fig S1C).

To obtain complementary information on the topology of the DDB1/E27/STAT2 complex, and to determine a possible involvement of the CRL4 co-adapter DDA1 in E27 and/or STAT2-binding, cross-linking mass spectrometry (XL-MS) was performed. DDB1\_ $\Delta$ BPB was assembled with hsDDA1 (98% identical to rnDDA1), E27, and STAT2 *in vitro* and purified by GF. Subsequently, the complex was cross-linked using the UV-activated cross-linker sulfo-SDA, followed by trypsin digestion and analysis of cross-linked peptides by XL-MS. Twenty-eight and 96 heteromeric cross links were identified

between DDB1 $_{\Delta BPB}$  and E27, and between E27 and STAT2, respectively, while only 1 cross link between DDB1 $_{\Delta BPB}$  and STAT2 was found (Fig 2C). These data are in line with a model where E27 serves as connector between DDB1 and STAT2, and further corroborate that the DDB1 BPB is dispensable for the DDB1/E27/STAT2 complex formation. Furthermore, 69 and 21 cross links were found between DDA1 and DDB1 $_{\Delta BPB}$  or E27, respectively, but no cross links extended from DDA1 to STAT2 (Fig 2C). These observations argue for a role of DDA1 in the stabilisation of the DDB1/E27 interaction, but against its involvement in the STAT2 recruitment.

#### Cryo-EM structure of the DDB1 ABPB/DDA1/E27/STAT2 complex

Cryo-EM imaging of a cross-linked DDB1<sub>ΔBPB</sub>/DDA1/E27/STAT2 complex (Appendix Fig S2) was pursued to gain structural insights. Single particle analyses yielded a density map at 3.8 Å resolution (map 1; Table EV1 and Appendix Fig S3). No experimental cryo-EM density corresponding to the expected position of DDA1 was observed (Appendix Fig S4A). Based upon distance restraints obtained from XL-MS (Fig 2C), the space accessible for interaction with the centre of mass of the DDA1 C-terminal helix was calculated using the DisVis algorithm (van Zundert & Bonvin, 2015). Visualisation of this space indicates that DDA1 preferably locates to a large, ill-defined DDB1 surface region around the edges of BPC and CTD. These data indicate that DDA1 is flexible and does not stably interact with E27, accounting for the lack of cryo-EM density (Appendix Fig S4B). Conversely, clear cryo-EM density corresponding to DDB1<sub>ABPB</sub> was present, allowing to unambiguously position both BP domains of the truncated DDB1 construct (Figs 3A and EV1A). Furthermore, assisted by XL-MS distance restraints and in silico structure prediction using AlphaFold2 (Jumper et al, 2021), a molecular model corresponding to E27 amino acid residues 105-570, folding into an elongated bundle containing 18  $\alpha$ -helices (the E27  $\alpha$ domain), could be fitted into a density segment on top of DDB1 $_{\Delta BPB}$ (Figs 3A and EV1B, see Materials and Methods for details). Prominent structural features were a protrusion from the edge of the  $\alpha$ domain, mainly formed by helix a8, inserting into a binding cleft formed by the DDB1 BPs, and a putative metal-binding motif composed of three cysteine residues and one histidine at the base of the protrusion. Taking into account the wealth of previous structural information (Laitaoja et al, 2013; Ireland & Martin, 2019), the tetragonal coordination geometry, the nature of the amino acid side chains coordinating the metal and the coordination bond lengths strongly argue in favour of a structural zinc-binding site. Accordingly, we modelled a zinc ion in the corresponding position (Fig EV1C). Moreover, cryo-EM map 1 showed extra density features consistent with the presence of a 4-helix bundle on top of E27, opposite of the DDB1-binding interface (Fig 3A). With the aid of an Alpha-Fold2 structure prediction of rnSTAT2, this density segment could be interpreted as a major portion of the STAT2 coiled-coil domain (CCD) encompassing residues 144–180 from helix αA, 196–231 (αB), 259–288 ( $\alpha$ C) and 293–317 ( $\alpha$ D; Fig EV1D). Twenty-one heteromeric cross links identified by XL-MS could be mapped on the DDB1<sub>ABPB</sub>/ E27/STAT2 CCD structure. In support of the cryo-EM model, 16 of these cross links are within the 25 Å distance threshold enforced by the cross-linker chemistry, and the remainder are only slight outliers (distances between 25 and 35 Å; Fig EV1E; Appendix Fig S5 and Table EV2).

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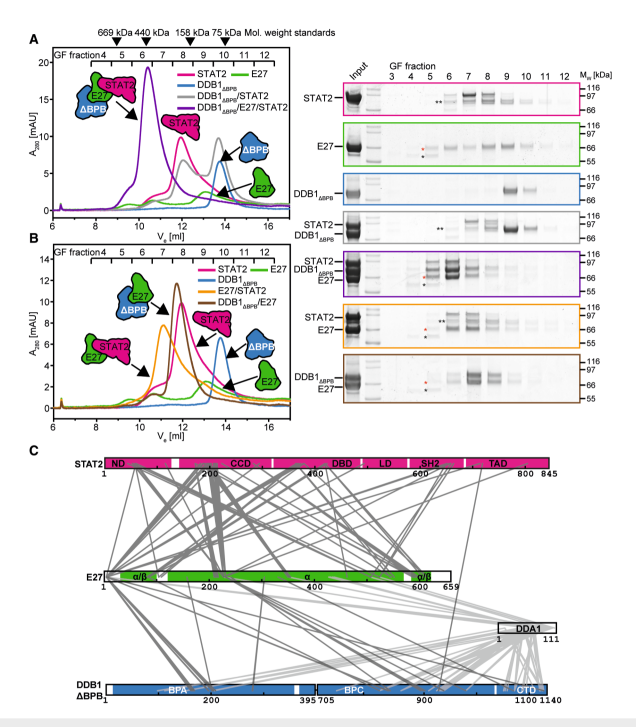


Figure 2. In vitro reconstitution and XL-MS analysis of DDB1<sub>ABPB</sub>(/DDA1)/E27/STAT2 protein complexes.

- A, B GF analysis of *in vitro* reconstitutions containing the indicated protein combinations. Coomassie blue-stained SDS–PAGE analyses of fractions collected during the GF runs are shown next to the chromatograms. The red \* indicates a contaminant, which we identified by cryo-EM analysis as *E. coli* ArnA, a notorious contaminant in *E. coli* Ni-NTA protein purifications (Andersen *et al*, 2013; Appendix Figs S2 and S3). Note that ArnA is present in GF fraction 6 (V<sub>e</sub> ~ 10.2 ml) of all runs containing E27 and migrates similarly to E27 on the SDS–PAGE. \*additional contaminant from E27 preparation; \*\*contaminant from STAT2 preparation.
- C Schematic of proteins used for XL-MS analysis with domains annotated. Heteromeric cross links obtained from XL-MS involving DDA1 are shown as light grey lines, otherwise as dark grey lines. ND, N-terminal domain; CCD, coiled-coil domain; DBD, DNA-binding domain; LD, linker domain; SH2, Src homology 2; TAD, transactivation domain; BP; β-propeller; CTD, C-terminal domain.

Source data are available online for this figure.

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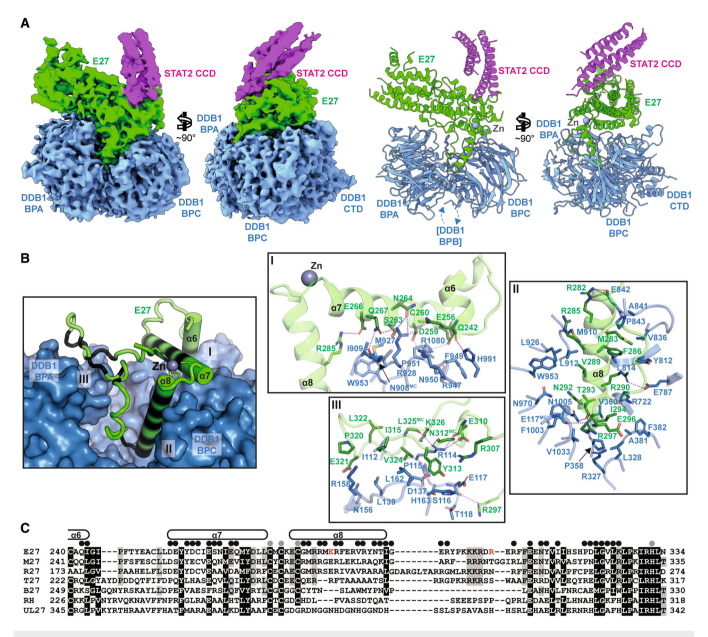


Figure 3. Cryo-EM structure of the DDB1  $_{\Lambda \rm BPB}$  /E27/STAT2 CCD protein complex.

- A (Left panel) Two views of cryo-EM map 1 contoured at threshold level 0.35 by the Chimerax software tool (Pettersen *et al*, 2021). Density segments corresponding to DDB1<sub>ABPB</sub> are coloured blue, to E27 green and to the STAT2 CCD magenta. (Right panel) Cartoon representation of the molecular model fitted in cryo-EM map 1, in the same orientation as in the left panel. DDB1<sub>ABPB</sub> is coloured blue and E27 green, and the zinc ion is shown as grey sphere.
- B (Left panel) Overview of the DDB1<sub>ABPB</sub>/E27 interaction interface. DDB1<sub>ABPB</sub> is shown in solvent-accessible surface representation and coloured blue, and E27 is shown as green cartoon with helices represented as cylinders. E27 residues in contact with DDB1<sub>ABPB</sub> are shaded black. The zinc ion is shown as grey sphere, and E27 side chains involved in zinc coordination are shown as sticks. (Right panels) Detailed views of interaction regions of interest I-III indicated in the left panel. Selected amino acid residues involved in intermolecular contacts are shown as sticks. Dashed lines represent hydrogen bonds or salt bridges.
- C Amino acid sequence alignment of E27 and homologues from other cytomegaloviruses. E27 residues contacting DDB1 are indicated by black dots and residues involved in zinc coordination as grey dots. E27 secondary structure elements are drawn above the alignment.

#### The DDB1<sub>ABPB</sub>/E27 protein complex

Initially, we focussed the structure analysis on the  $DDB1_{\Delta BPB}/E27$  interaction. E27 binds to a cleft lined by the inner surfaces of DDB1 BPA and BPC (Fig 3B). In this way, it is positioned on the opposite side of where DDB1 BPB would be located, which links to the CRL4

core. E27 helices  $\alpha 6$ ,  $\alpha 7$ ,  $\alpha 8$ , and the loop leading to  $\alpha 9$  are in direct contact with DDB1 and helix  $\alpha 8$  protrudes in the binding cleft (Fig 3B). The DDB1/E27 binding interface, covering an area of 2,131 Ų, is divided in three regions (Figs 3B and EV2A). The first involves E27 helices  $\alpha 6$  and  $\alpha 7$ , bordered by the zinc-coordinating cysteine triad, contacting residues located in DDB1 BPC WD40

repeats 6 and 7, and in the CTD. It is formed by a network of hydrogen bonds, additionally supported by salt bridges and hydrophobic interactions (Fig 3B, box I). A second, central interaction region includes the protrusion formed by helix  $\alpha 8$  and a subsequent polybasic loop, contacting residues lining the inner surface of DDB1 BPC and BPA WD40 repeats 3 and 7. This interface is mostly hydrophobic, with fewer polar and electrostatic interactions (Fig 3B, box II). A third region comprises the extended loop leading back to the E27 zinc-binding motif, adjacent to DDB1 BPA WD40 repeats 3 and 4. Again, mainly hydrophobic interactions constitute this interface, assisted by hydrogen bonding (Fig 3B, box III).

To evaluate the conservation of the DDB1/E27-binding surface, we mapped interface residues on a multiple sequence alignment of E27 homologues (Fig 3C). Twenty-eight out of 38 interfacing residues are type-conserved between RCMV-E E27 and MCMV M27, while 26 are type-conserved between E27 and R27, the homologue encoded by RCMV-Maastricht (RCMV-M). Furthermore, 16 and 12 interface residues are type-conserved in E27 homologues from bat and tupaiid CMV, respectively, and only 8 and 9 residues in rhesus macaque CMV and HCMV, respectively. Additionally, in the case of tupaiid, bat, rhesus macaque and human CMV, high sequence divergence of the central  $\alpha 8$  binding region is apparent. Together, this suggests that DDB1-binding functionality is restricted to rodent CMV E27, M27 and R27.

To validate the structure, an E27 construct was prepared, from which a significant part of the DDB1-binding region (residues E284-R305, marked red in Fig 3C) was deleted and replaced by a flexible linker (E27 $\Delta$ ). E27 $\Delta$  was purified and employed for *in vitro* reconstitution with DDB1 or STAT2. Subsequent gel filtration analysis showed that E27 $\Delta$ , in contrast to WT E27, was unable to bind DDB1 or DDB1<sub> $\Delta$ BPB</sub> (Fig EV2B and C). However, like WT E27, E27 $\Delta$  bound STAT2 and exhibited a similar elution profile as WT E27 in isolation, demonstrating that the overall integrity of E27 $\Delta$  was not compromised by mutagenesis (Fig EV2D and E). This suggests that the central binding region (Fig 3B, box II) is critical for E27/DDB1 interaction, while the peripheral regions (Fig 3B, boxes I and III) are not sufficient to sustain stable E27/DDB1 association on their own.

#### E27 mimics endogenous CRL4 substrate receptors

Next, the DDB1 $_{\Delta BPB}$ /E27 complex structure was compared with the DDB1-binding modes of DCAF1 and of the previously characterised viral DCAFs (vDCAFs) SV5 V and HBx. Overall, they are positioned similarly relative to DDB1 (Fig 4A). Analysis of their "molecular footprint" on DDB1 reveals that the E27 footprint is almost identical to the DCAF1-derived one, while SV5 V and HBx bind DDB1 in significantly smaller interaction interfaces, mostly restricted to DDB1 BPC (Fig 4B). To engage DDB1 surface regions, E27 utilises other structural elements than DCAF1, that is a short and two long helices ( $\alpha6$ –  $\alpha 8$ ) and an extended loop, in contrast to three short helices and the bottom side of four WD40 repeats in DCAF1 (Fig 4A, insets). Superposition of DDB1-bound E27 and cellular and viral DCAF structures showed that the only common secondary structure element is the Cterminal half of E27 a8, which aligns well with a short helical "Hbox" motif in DCAF-type receptors (Fig 4C). However, sequence conservation of this motif is low, with only four amino acid positions > 60% type-conserved (Fig 4D). Together, these data demonstrate how E27 emulates the DDB1-binding mode of endogenous DCAF substrate receptors using divergent structural principles. Furthermore, the analysis illustrates that all examined vDCAFs engage DDB1 via additional structural elements beyond the previously described "H-box" motif (Li *et al*, 2010). However, their structure and sequence are not conserved, and the only common characteristics are contact areas on the upper edge of DDB1 BPC. Accordingly, structural analyses of vDCAF/DDB1 interactions are necessary for a full mechanistic understanding of how they hijack CRL4.

Given the similar position and interface area of E27 and DCAF1 on DDB1, we hypothesised that E27 competes with endogenous DDB1/DCAF complexes for DDB1 binding. To gain biochemical insight, the preformed DDB1/DCAF1 complex was incubated with E27 and subjected to analytical GF. These analyses showed the formation of an elution peak containing DDB1 and E27, concomitant with the reduction of the DDB1/DCAF1 complex peak at later elution volume, in an E27 concentration-dependent manner, indicating that E27 is able to displace DCAF1 from DDB1 (Fig 4E).

#### Mechanism of STAT2 recruitment

Based on cryo-EM map 1, the binding between E27 and the STAT2 CCD was analysed in detail (Figs 5A and B, and EV3A). The interaction interface covers a surface area of 750  $\mathring{A}^2$ . It comprises hydrophobic residues from STAT2 CCD helices  $\alpha A, \, \alpha B$  and the C-terminal loop of  $\alpha C$ . These residues project into a hydrophobic pocket lined by E27 amino acids from the loop downstream of  $\alpha 4$  and side chains from a sequence stretch spanning  $\alpha 11$  to  $\alpha 12$ . In addition, several peripheral hydrogen bonds and a salt-bridge network strengthen the interface.

Conservation analysis of this binding interface revealed that out of 17 involved residues, 12 and 11 are type-conserved in rodent E27 homologues (M27 and R27, respectively), while only 8 are type-conserved in their tupaiid and bat counterparts and only 6 in macaque or human CMV (Fig 5B). Additionally, in the more distantly related, nonrodent CMV species, most of the VPVL loop downstream of  $\alpha 4$  is deleted, removing a significant portion of the STAT2 CCD-binding site. Thus, it is likely that STAT2 recruitment is restricted to rodent E27 homologues. Variation within this branch might be driven by species-specific adaptation to differences in the respective host STAT2 sequences, for example rnSTAT2 positions 169, 198 and 285 (Fig 5B).

To corroborate the structural data, an E27 R211E/R214E double mutant (E27 RE) was generated by site-directed mutagenesis to disrupt the intricate salt-bridge network between these residues and STAT2 D168 and D171 (Fig 5A). The E27 RE variant, alongside WT E27, was subjected to *in vitro* reconstitution with STAT2 or DDB1, followed by gel filtration analysis. The results demonstrate that E27 RE had lost the ability to form a stable complex with STAT2 (Fig EV3B), while it retains DDB1-binding activity (Fig EV3C and D) and exhibits a similar elution profile as WT E27 (Fig EV3E), thus validating the molecular model of STAT2 CCD recruitment.

Moreover, further 3D classification of the cryo-EM particle images yielded a second map at 5 Å resolution, exhibiting additional density features in the E27 and STAT2 regions (map 2, Figs 5C and EV4A; Appendix Fig S3 and Table EV1). This allowed the placement of a STAT2 AlphaFold2 model lacking the N-terminal and TAD domains and enabled fitting of the terminal E27  $\alpha/\beta$ -domain, revealing that at least a subset of the particles observed in the cryo-

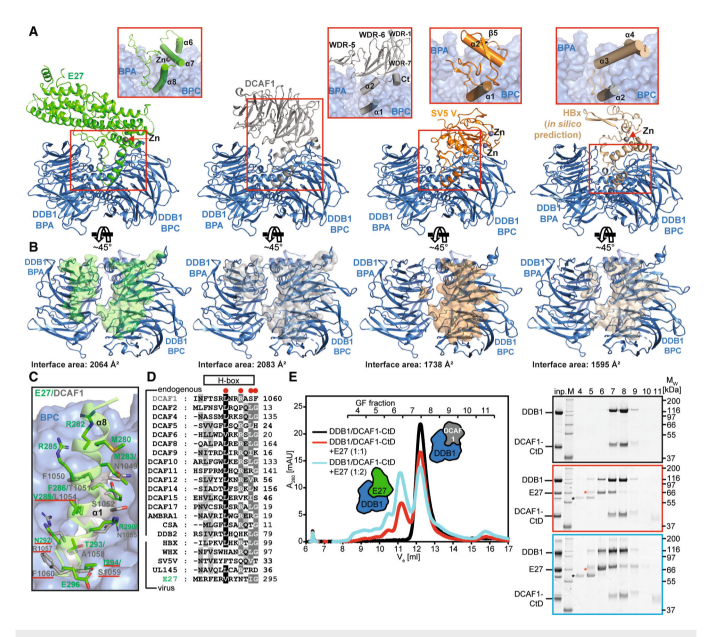


Figure 4. E27 replaces endogenous CRL4 substrate receptors.

- A Comparison of the DDB1/E27-binding mode with endogenous and viral CRL4 substrate receptors. Structures of DDB1 (blue) in complex with E27 (green), DCAF1 (grey, PDB 6zue; Banchenko et al, 2021), SV5 V (orange, PDB 2hye; Angers et al, 2006) and HBx (wheat, AlphaFold2 prediction, Appendix Fig S6) are shown as cartoons. Regions of the AlphaFold2 HBx model with very low confidence have been removed. The insets display enlarged views of the boxed interaction areas, with DDB1 shown as semitransparent blue surface. Zinc ions are shown as grey spheres. The position of the HBx zinc ion was modelled based on information from (Ramakrishnan et al. 2019).
- B "Molecular footprint" of substrate receptors from A on the DDB1 surface. DDB1 is shown in the same representation as in A, but slightly rotated to allow an unobstructed view of the receptor-binding cleft. The receptors themselves have been removed for clarity. Parts of the DDB1 surface in contact with the respective substrate receptor are coloured according to the scheme in A.
- C Detailed comparison of a helical DDB1-binding motif in E27 (green) and the DCAF1 "H-box" (grey). DDB1 is shown as semitransparent blue surface, E27 and DCAF1 as semitransparent cartoon. Selected amino acid residues involved in intermolecular contacts are shown as sticks.
- D Amino acid sequence alignment of the "H-box" DDB1-binding motif in endogenous substrate receptors and viral factors hijacking DDB1. Black boxes: > 80% type-conserved; dark grey: > 60%; light grey: > 40%. > 60% type-conserved residues are marked by red dots and underlined in (C).
- E Preformed DDB1/DCAF1 protein complex was incubated with indicated amounts of E27 and analysed by GF. SDS—PAGE analyses of GF fractions are shown next to the chromatogram. The red \* indicates the ArnA contaminant (see Fig 2). \*additional contaminant from E27 preparation.

Source data are available online for this figure.

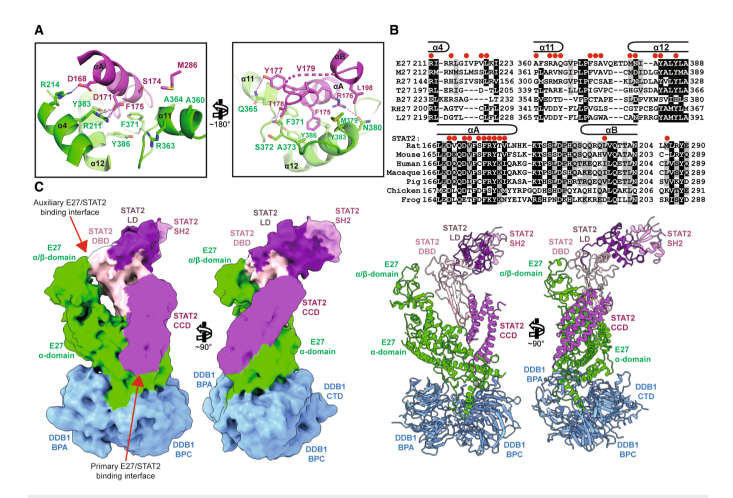


Figure 5. Mechanism of STAT2 recruitment.

- A Detailed view of the E27/STAT2 CCD interaction, derived from the cryo-EM map 1 molecular model. Proteins are represented as cartoons; E27 is coloured green and STAT2 magenta. Selected side chains involved in intermolecular interaction are shown as sticks. Dashed lines represent hydrogen bonds or salt bridges.
- B (Upper panel) Amino acid sequence alignment of E27 and homologues from other cytomegaloviruses. E27 residues contacting STAT2 are indicated by red dots. E27 secondary structure elements are drawn above the alignment. (Lower panel) Amino acid sequence alignment of STAT2 and homologues from other species. STAT2 residues contacting E27 are indicated by red dots. STAT2 secondary structure elements are drawn above the alignment.
- C (Left panel) Two views of cryo-EM map 2 contoured at threshold level 0.15 by the Chimerax software tool (Pettersen *et al*, 2021). Density segments corresponding to DDB1<sub>ΔBPB</sub> are coloured blue, to E27 green, to the STAT2 CCD magenta, to the STAT2 DBD light pink, to the STAT2 LD purple and to the STAT2 SH2 domain dark pink. (Right panel) Cartoon representation of the molecular model fitted in cryo-EM map 2, in the same orientation and colour code as in the left panel. The zinc ion is shown as grey sphere.

EM analysis exhibit a bipartite E27/STAT2-binding interface (Fig 5C). It comprises the primary interaction between the STAT2 coiled-coil domain (CCD) and the E27  $\alpha$ -domain, which has also been observed in cryo-EM map 1 (Figs 3 and 5A), as well as a second, auxiliary contact region involving the STAT2 DNA-binding domain (DBD) and the E27  $\alpha$ / $\beta$ -domain (Fig 5C; Appendix Fig S7). Thirty-nine out of 96 heteromeric cross links could be mapped on this E27/STAT2 complex, of which 22 lie within the 25 Å distance threshold and 9 are slight outliers (25–35 Å), supporting the structural data. Four severely overlength cross links are apparent (> 40 Å), all of which involve the E27  $\alpha$ / $\beta$ -domain or the STAT2 SH2 domain, indicating the flexibility of these distal components (Fig EV4B), rationalising the low local resolution of the corresponding cryo-EM map regions (Appendix Fig S3).

The auxiliary E27/STAT2 interaction area involves parts of the E27  $\alpha/\beta$ -domain, located between strand  $\beta A$  and helix  $\alpha A$ , between

 $\alpha B$  and  $\beta C,$  and a C-terminal stretch upstream of  $\beta D$  (Appendix Fig S7). These regions form a shallow surface, which contacts STAT2 DBD protrusions comprising amino acid stretches 377–383 and 406–416. However, due to the low local resolution of the cryo-EM map, we refrained from modelling individual side chain interactions.

# E27 positions STAT2 appropriately for CRL4-catalysed ubiquitin transfer

The structure of the DDB1 $_{\Delta BPB}/E27/STAT2$  complex enabled modelling of STAT2 in the context of a complete CRL4 assembly (Fig 6). The model shows that STAT2 is placed centrally in the ubiquitylation zone created by rotation of the CUL4/RBX1 scaffold arm. This rotation is enabled by a flexible hinge connecting DDB1 BPB, where CUL4/RBX1 is attached, to DDB1 BPC, which together with the BPA

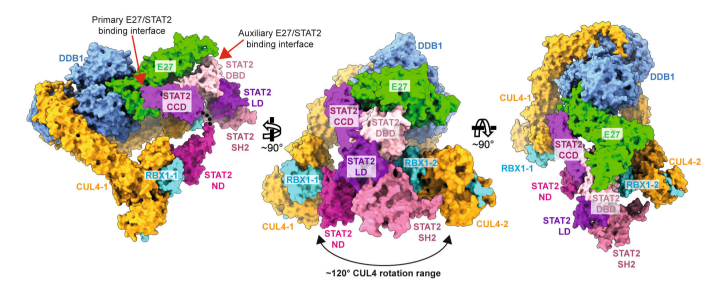


Figure 6. Model of the complete CRL4<sup>E27</sup>/STAT2 assembly.

A model of how E27/STAT2 is positioned relative to the remainder of CRL4 was assembled by superposition of DDB1<sub>ABPB</sub>/E27/STAT2 with DDB1 BPA/BPC from two CRL4<sup>DCAF1</sup> cryo-EM structures, which represent the two outermost positions of the CUL4 scaffold (CUL4-1, CUL4-2), thus visualising the rotation range of CUL4/RBX1 with respect to DDB1 (Banchenko *et al*, 2021). RBX1 (cyan), CUL4 (orange), DDB1 (blue), E27 (green) and STAT2 (coloured as in Fig 5) are shown in surface representation. The STAT2 N-terminal domain (magenta cartoon) was modelled in an arbitrary position, since it is connected to the STAT2 CCD via a flexible linker (indicated by the dotted line).

forms the binding platform for substrate receptors. Furthermore, the model also provides a rationale for the existence of the second, auxiliary E27/STAT2 DBD-binding interface, which nudges the STAT2 molecule into the ubiquitylation zone. In such a way, surface-exposed lysines on all STAT2 domains are within reach of ubiquitin-charged E2, which is positioned for ubiquitin transfer by a catalytic assembly comprising the neddylated distal end of CUL4 and RBX1 (Baek *et al*, 2020), guaranteeing efficient STAT2 ubiquitylation and subsequent degradation.

# The E27 structure reveals a conserved zinc-binding motif of functional importance

The cryo-EM density (map 1) revealed a cryptic zinc-binding motif in E27 involving three cysteine residues (C273, C275 and C278) and H332 from a loop upstream of helix  $\alpha 9$  (Figs 7A and EV1C), anchoring the protrusion, which inserts into the binding cleft on DDB1, to the E27  $\alpha$ -domain bundle. Site-directed mutagenesis of the E27 zinc-coordinating cysteines (AxAxxA variant), or the histidine (H332A), to alanine residues abrogated the ability to coimmunoprecipitate endogenous DDB1, demonstrating the importance of the zinc-binding motif for DDB1 binding (Fig 7B). The sequence conservation of this motif argues strongly in favour of its functional relevance (Fig 7C), which prompted us to investigate its function in closely related cytomegaloviruses. MCMV M27, the closest E27 homologue, exhibits 56 and 76% amino acid sequence identity and similarity, respectively (Appendix Fig S7). E27 and M27 AlphaFold2 structure predictions are almost indistinguishable (RMSD of 1.195 Å, 6,652 aligned atoms; Appendix Fig S8), and structural alignment of these predictions demonstrated conservation of side chain positions involved in zinc binding (Fig 7D). Thus, M27 represents a valid model to study functional consequences of

interference with the zinc coordination motif through site-directed mutagenesis and to test the relevance of our E27 structure model for the MCMV system.

First, in FLAG-tagged M27 constructs, individual residues and combinations in the zinc-binding site were mutated to alanine. The constructs were expressed in 293 T cells and their ability to bind endogenous DDB1 was assessed by co-immunoprecipitation. While WT-M27 efficiently co-precipitated DDB1, mutation of the three zinc-coordinating cysteine residues abrogated DDB1 binding (Fig 7E). Likewise, a mutation of the loop containing the zinc ligand H333 and of an adjacent LP motif interfered with DDB1 interaction. Serving as an additional control, the Y270F mutation upstream of the cysteine triad had no effect on DDB1 binding. Thus, the integrity of the zinc-binding motif is necessary for interaction with DDB1 in mouse and rat CMV-encoded STAT2 antagonists.

Next, MCMV mutants lacking M27 ("ΔM27") or carrying the AxAxxA triple cysteine mutation (M27<sub>AxAxxA</sub>) were generated. Immortalised mouse embryonic fibroblasts were infected with these viruses alongside WT-MCMV as a control. STAT2 protein levels of cell lysates were probed by western blot. As expected, the WT-MCMV infection depleted STAT2. By contrast, the M27<sub>AxAxxA</sub>-MCMV failed to reduce STAT2 protein levels, phenocopying the  $\Delta$ M27 virus (Fig 7F). In addition, to test the impact of these MCMV variants on ISG induction, NIH3T3 cells stably expressing an interferon-stimulated response element (ISRE)-dependent luciferase reporter were infected with WT,  $\Delta$ M27 and M27<sub>AxAxxA</sub> viruses. Cells were treated with IFN, and luciferase activity was quantified as reporter for IFN-driven gene expression. In accordance with the STAT2 degradation, IFN signalling was efficiently suppressed following WT-MCMV infection. Conversely, ΔM27-MCMV failed to counteract IFN-induced ISRE signalling. Intriguingly, the M27<sub>AxAxxA</sub>-MCMV infection exhibited strongly attenuated IFN inhibition

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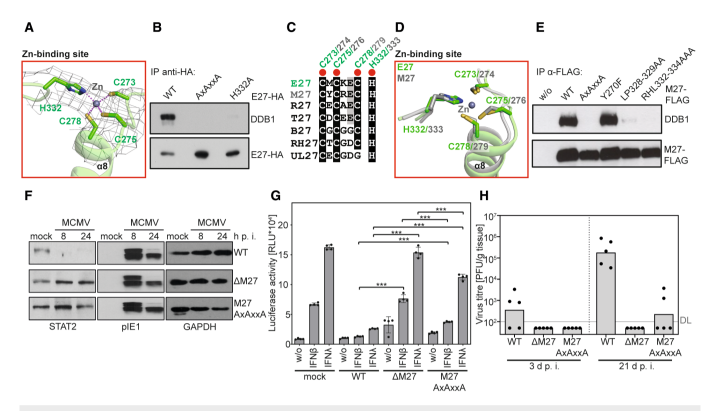


Figure 7. E27 structure reveals a conserved zinc-binding motif of functional importance in vivo.

- A Detailed view of the zinc-binding motif. E27 is shown as green cartoon, residues involved in zinc binding as sticks. Zinc coordination bonds are indicated as dashed lines. Cryo-EM density, contoured at 6σ, is shown as grey mesh.
- B 293 T cells were transfected with expression plasmids encoding E27-HA or indicated variants. 24 h after transfection, cells were lysed and E27-HA was precipitated using an HA-specific antibody. Precipitates were separated by SDS—PAGE, blotted and probed using the indicated antibodies.
- C Amino acid sequence alignment of E27 and homologues from other CMV species. Residues involved in zinc coordination are marked by red dots.
- D Superposition of the E27 zinc-binding site (green, structure from this work) with the one from M27 (grey, AlphaFold2 prediction, see Appendix Fig S8).
- E 293 T cells were transfected with expression plasmids encoding M27-FLAG or the indicated M27-FLAG variants. 24 h after transfection, cells were lysed and M27-FLAG was precipitated using a FLAG-specific antibody. Precipitates were separated by SDS-PAGE, blotted and probed using the indicated antibodies.
- F Mouse fibroblasts were infected with MCMV or the indicated MCMV mutants. At the indicated time points, cells were lysed and proteins were separated by SDS–PAGE and blotted. Blots were probed using the indicated antibodies. pIE1 and GAPDH were used as infection and loading controls.
- G NIH3T3 reporter cells stably expressing ISRELuc were infected with the indicated MCMV strains, treated with IFN $\beta$  or IFN $\lambda$ , and luciferase activity was measured. The error bars represent the standard deviation, N=4 biological replicates, \*\*\* $P \leq 0.001$  (ANOVA).
- H BALB/c mice were infected i.p. with WT-MCMV and indicated MCMV mutants. At 3 and 21 days p.i., salivary glands were collected and frozen. Virus titres were determined from organ homogenates by plaque titration. Titrations were performed in quadruplicates. Bars depict the geometric mean, and dots show titres of individual mice (N = 5 biological replicates). DL, detection limit.

Source data are available online for this figure.

(Fig 7G). These data demonstrate that the disruption of the M27 zinc-binding site interferes with M27-dependent STAT2 degradation and alleviates suppression of IFN responses.

To verify the application of the MCMV model for challenging and extending our structural E27 data, we engineered a  $\Delta M27\text{-MCMV}$  variant expressing HA-tagged E27 (" $\Delta M27\text{-E27HA}$ "). Infection of primary rat fibroblasts with  $\Delta M27\text{-E27HA}$ , followed by immunoblotting of the cell lysates, revealed complete loss of STAT2 signal, similar to WT-MCMV infection, demonstrating functional interchangeability of M27 and E27 under these experimental conditions (Fig EV5).

Lastly, to assess the relevance of these observations *in vivo*, mice were infected with WT-,  $\Delta$ M27-, or M27<sub>AxAxxA</sub>-MCMVs, and virus titres in salivary glands were determined. Virus titre in  $\Delta$ M27- and M27<sub>AxAxxA</sub>-MCMV infection were reduced strongly in comparison

with WT-MCMV (Fig 7H). This highlights the importance of M27 for MCMV replication and shows that the M27  $_{\rm AXAXXA}$  mutation severely impairs the IFN inhibitory capacity in the mouse model. In conclusion, the disruption of the M27 zinc-binding motif causes a loss of DDB1 binding and STAT2 degradation, an impairment of IFN antagonism, and ultimately a virus attenuation  $in\ vivo$ . This establishes the predictive power of our E27 structure also for the MCMV infection model by connecting it with the MCMV  $in\ vivo$  replication capacity.

# Discussion

We identified STAT2, the central hub driving the antiviral IFN response, as one of the most drastically downregulated host factors during RCMV-E infection and elucidated E27 as causative factor.

The structure determination of E27 uncovered a zinc-binding motif close to the DDB1-binding site, highly conserved in homologous proteins from related cytomegaloviruses. Disruption of this motif in the closely related E27 homologue M27 resulted in impairment of DDB1 binding, STAT2 degradation, and ISG suppression, and finally to inhibition of virus replication in vivo. Similarly, the integrity of conserved zinc-coordinating motifs has been shown to be crucial for CRL4 association and function of SV5 V (Lin et al, 1998) and the retroviral accessory proteins Vpx and Vpr (Wang et al, 2017; Yamamoto et al, 2017). However, HBx side chains involved in zinc coordination seem to be dispensable for CRL4 interaction, but critical for substrate recruitment (Ramakrishnan et al, 2019). These data, together with structure-based sequence alignments (Fig 3C; Appendix Fig S7), strongly suggest that RCMV-E E27 and MCMV M27 employ a conserved mechanism to integrate into CRL4 via DDB1 binding and that this mechanism depends on the integrity of a conserved zinc-binding motif.

With UL27, HCMV encodes a positional and sequence homologue of M27 and E27. HCMV reduces STAT2 levels (Le et al, 2008; Nightingale et al, 2018) and-like MCMV-exploits the UPS and CRLs to this end (Le et al, 2008; Le-Trilling et al, 2016). However, UL27 is neither essential nor sufficient for the HCMV-induced STAT2 degradation (Le et al, 2008), and at best interacts very weakly with DDB1 (Reitsma et al, 2011; Trilling et al, 2011; Landsberg et al, 2018). Our structure-based sequence alignment shows exchange of the third zinc-coordinating cysteine to a glycine in UL27, alongside considerable sequence divergence in the critical DDB1-binding helix  $\alpha 8$ , thereby rationalising these observations. In addition, only few of the E27 residues mediating the critical interaction with the STAT2 CCD are conserved in UL27. In accordance with these observations, HCMV UL27 induces the degradation of the unrelated host Tip60 acetyltransferase, eventually via direct interaction with proteasome components (Reitsma et al, 2011).

The HCMV-encoded STAT2 antagonist was recently identified as UL145 (Le-Trilling et al, 2020). Like M27, UL145 interacts with DDB1 and CRL4 (Nightingale et al, 2018) and induces proteasomal degradation of STAT2 (Le-Trilling et al, 2020). Accordingly, an unbiased single-cell sequencing approach combined with a genomewide CRISPR screen confirmed the central importance of the IFNAR2/STAT2/IRF9 signalling axis as well as neddylation and Cullin-RING E3s in defining HCMV infection propensity (Hein & Weissman, 2022). Sequence alignments from previous and from our present study indicate the presence of an H-box motif in the N terminus of UL145 (residues 25-36, Fig 4D), suggesting that this UL145 region is positioned similarly in the DDB1-binding cleft as endogenous DCAF substrate receptors and the C-terminal half of E27 helix α8. Indeed, site-directed mutagenesis of several putative H-box residues in UL145 abrogated DDB1 interaction in coimmunoprecipitation experiments (Le-Trilling et al, 2020), and a crystal structure of the UL145 H-box peptide in complex with DDB1 very recently supplied the ultimate proof of this hypothesis (Wick et al, 2022). However, future structural studies are necessary to clarify how the whole of UL145 integrates into CRL4, in order to pinpoint additional DDB1-interacting motifs and to decipher the mechanisms of STAT2 recruitment. These will be highly informative to fully understand the different mechanistic principles pursued by HCMV and rodent CMV for STAT2 antagonism, both converging on CRL4.

We observed some residual IFN inhibitory capacity of the DDB1binding-deficient M27<sub>AxAxxA</sub>-MCMV variant (Fig 7G). A comparison of the E27/STAT2 CCD-binding mode discovered here to the IRF9/ STAT2 CCD complex (Rengachari et al, 2018) demonstrates an overlap of the binding interfaces (Appendix Fig S9A). Previous structural studies of ZIKV STAT2 antagonism established that ZIKV NS5 competes against IRF9 for STAT2 binding by occupying the IRF9 binding site on the STAT2 CCD (Wang et al, 2020). Therefore, we wondered whether E27 might inhibit IRF9 recruitment to STAT2 by a similar steric exclusion mechanism. To test this, we performed in vitro competition assays by analytical GF. However, addition of E27 in equimolar amounts did not to disassemble preformed rnIRF9/STAT2 protein complexes, and rnIRF9 readily disassembled preformed E27/STAT2 complexes (Appendix Fig S9B). These data suggest that, at least under the experimental in vitro conditions, IRF9 possesses a higher binding affinity for STAT2 than E27. Additional factors, not reflected in our experimental in vitro set-up, that explain the residual ISG suppression upon M27AXAXA-MCMV infection remain to be discovered.

The comparison of the DDB1-binding modes of E27 and other CRL4 substrate receptors revealed that E27 engages similar surface regions in the DDB1 binding cleft as DCAF-like receptors. However, structure and sequence determinants in E27 and other viral and cellular DCAFs mediating DDB1 binding are highly divergent. In fact, the only common element in these DDB1 binders with structural homology, and at least a certain degree of sequence conservation is the short H-box motif (Fig 4C and D; Li et al, 2010), interacting with the bottom of the binding cleft formed by DDB1 BPA and BPC. The physico-chemical properties of this DDB1 cleft seem to select for helical secondary structure, for an L or V side chain at position 7 of the sequence alignment in Fig 4D (albeit with exception of the viral DCAFs HBx and SV5V), and with less stringency for R at position 10, L/I/V at position 12 and G at position 13. The latter G residue, having a very low propensity to be located in  $\alpha$ -helices (Pace & Scholtz, 1998), might be necessary as helix-breaking residue to lead over into the loop segment exiting the binding cleft.

Our biochemical analysis indicated that E27 replaces the endogenous DCAF1 substrate receptor on DDB1. This is in agreement with recent results demonstrating that H-box peptides of the vDCAFs UL145 and HBx exhibit significantly higher DDB1-binding affinities than the corresponding H-box peptides derived from endogenous DCAF1 and DDB2 receptors (Wick *et al*, 2022). Accordingly, it is tempting to speculate that vDCAFs, in addition to marking antiviral substrates for degradation, might block a plethora of cellular DCAF-dependent ubiquitylation processes. Further research is needed to delineate if and how virus replication would benefit from these putative ubiquitylation inhibition events.

Interestingly, another CRL4 substrate receptor, CRBN, possesses a unique DDB1-binding mechanism involving three  $\alpha$ -helices, which do not structurally align to the H-box motif at all (Chamberlain et~al,~2014; Fischer et~al,~2014). This pronounced plasticity of the DDB1 substrate receptor-binding cleft impedes sequence-based prediction and suggests the existence of further, yet unidentified viral and endogenous DDB1 binders. In line with these observations, recent HCMV interactome studies discovered several more DDB1- and CRL4-associated HCMV factors (Nobre et~al,~2019; Nightingale et~al,~2022). Accordingly, future studies of DDB1-binding mechanisms, together with proteomics and novel in

*silico* structure prediction methods (Humphreys *et al*, 2021; Jumper *et al*, 2021), might greatly expand the spectrum of known DDB1 binders and extend our understanding of CRL4 biology and its viral exploitation.

Our study has limitations. We employed human DDB1 for in vitro reconstitution and structure determination of the complex with the RCMV factor E27 and rat STAT2. It is important to note that DDB1 is highly conserved across eukaryotes (Tang & Chu, 2002), because it is critical for development and cell viability as an essential component of the nucleotide excision DNA repair machinery (Cang et al, 2006; Wakasugi et al, 2007). As a consequence, out of 1,140 DDB1 amino acids, there are only 12 positions exchanged between human and rat DDB1, of which four are typeconserved (Appendix Fig S10A). We have mapped these exchanges on a model of full-length human DDB1 bound to E27 and the rat STAT2 CCD, to show that none of these are involved in intermolecular interaction, demonstrating the validity of our approach (Appendix Fig S10B and C). In support of these observations, there is precedence in studying interspecies DDB1 protein complexes, for example the SV5V, WHX, SIV Vpx and Vpr, zebrafish DDB2 and chicken CRBN proteins have been in vitro reconstituted with human DDB1 for structural characterisation, and valid biological conclusions have been drawn from these studies (Angers et al, 2006; Li et al, 2006, 2010; Fischer et al, 2011, 2014; Schwefel et al, 2014, 2015; Wu et al, 2015; Banchenko et al, 2021).

We used a DDB1 mutant lacking the entire BPB for XL-MS and structure determination. This had technical reasons, since protein complex preparations including full-length DDB1, E27 and STAT2 did not yield cryo-EM reconstructions at appropriate resolution for model building, most likely due to the flexibility of the DDB1 BPB, impeding proper alignment of the cryo-EM particle images. Importantly, the structure model including full-length DDB1 (Appendix Fig S10B) clearly demonstrates that the BPB is located on the opposite side of the E27 binding interface on DDB1, where it is situated to flexibly connect to the CUL4 scaffold to create the ubiquitylation zone around immobilised substrates (Fig 6). This rules out any involvement of DDB1 BPB in E27- and/or STAT2-binding processes, thus validating the use of DDB1 $_{\Delta BPB}$  in experiments aimed at studying E27- and/or STAT2-binding mechanisms. Along these lines, several previous studies have employed DDB1ABPB to facilitate structure determination and have successfully applied the resulting structural models for functional follow-up experiments in the context of complete CRL4 assemblies (Petzold et al, 2016; Bussiere et al, 2020; Slabicki et al, 2020).

Lastly, it should be noted that we used a cross-linked protein complex sample for cryo-EM structure determination. Again, this had technical causes, since the equivalent non-cross-linked sample failed to produce cryo-EM specimens suitable for single particle analysis, likely because of aggregation and/or denaturation of the particles in the plunge-freezing process. Even though cross linking is a common procedure to optimise cryo-EM samples (Stark, 2010; Drulyte *et al*, 2018) and XL-MS has been proven to be a suitable tool to inform modelling of native structures (O'Reilly & Rappsilber, 2018; Schneider *et al*, 2018; Piersimoni *et al*, 2022), we cannot exclude that the cross-linking procedure stabilised a rare or non-native conformation of the protein sample.

Our data, to our knowledge for the first time, reveal that RCMV (like MCMV) exploits host CRL4 complexes to induce

proteasomal degradation of STAT2. With UL145, HCMV relies on an analogous, but evolutionary unrelated factor, also exploiting DDB1 and CRL4 for STAT2 degradation and evasion from the antiviral activity of IFNs. In clear contrast to HCMV, RCMV and MCMV are both amenable to in vivo experiments in small animal models. Over 40 years ago, HCMV has been called the troll of transplantation (Tx) due to its grim impact on immunosuppressed individuals after Tx (Balfour, 1979). Despite tremendous efforts, HCMV still harms and kills graft recipients. While MCMV allows various experiments regarding general principles of cytomegaloviral pathogenesis and antiviral immunity, one shortcoming is that mice are rather small animals in which chirurgical and solid organ transplantation (SOT) procedures are difficult. In clear contrast, various SOT procedures that are indispensable for human medicine can be recapitulated in bigger rat models. We hope that our work provides the molecular foundation for future studies addressing the relevance of STAT2-dependent IFN signalling, CMV-induced STAT2 degradation and viral DDB1-CRL4 exploitation in rat SOT models.

# Materials and Methods

#### Animals

BALB/c mice were obtained from Harlan and bred and housed in the animal facility of the Institute for Virology of the University Hospital Essen. Mice were maintained under selective pathogen-free conditions. Age- (8–16 weeks) and sex-matched (male and female) groups of BALB/c mice were infected intraperitoneally with  $2\times 10^5$  PFU of WT-MCMV or MCMV mutants per mouse. All procedures were performed in accordance with the guidelines of the University Hospital Essen, Germany, the national animal protection law (Tierschutzgesetz [TierSchG]) and the recommendations of the Federation of European Laboratory Animal Science Association (FELASA). The study was approved by the North Rhine-Westphalia State Office for Nature, Environment and Consumer Protection (LANUV NRW, Düsseldorf, North Rhine-Westphalia, Germany; permit #84–02.04.2016.A441). No blinding was done. No animals were excluded.

## Microbe strains

Escherichia coli NovaBlue and Rosetta™ 2(DE3) were cultivated in LB medium at 37°C in a shaker incubator (150 rpm; Table EV3).

#### **Cell lines**

Immortalised mouse fibroblasts were generated from primary C57BL/6 and BALB/c MEF by crisis immortalisation (Rattay et al, 2015). NIH3T3-ISREluc:IFNLR cells were described in (Le-Trilling et al, 2018). Mouse fibroblasts, REF and HEK293T (ATCC CRL-11268, female) cells were grown in Dulbecco's minimal essential medium (DMEM) supplemented with 10% (v/v) foetal calf serum (FCS), penicillin, streptomycin and 2 mM glutamine at 37°C in 5% CO<sub>2</sub>. Sf9 cells (originally derived from the ovaries of *Spodoptera frugiperda*) were cultivated in Insect-XPRESS<sup>TM</sup> medium at 28°C in a shaker incubator (180 rpm; Table EV3).

#### **Primary cultures**

Organs of infected mice were harvested, snap-frozen in liquid nitrogen and stored at  $-80^{\circ}\text{C}$  until titrations were performed. Plaque titrations were conducted by use of primary mouse fibroblasts (mouse embryonic fibroblasts [MEF] and mouse newborn cells [MNC]), which were isolated from mouse embryos and newborns, respectively, according to described protocols (Le-Trilling & Trilling, 2017). Fibroblasts were grown in DMEM supplemented with 10% (v/v) FCS,  $100~\mu\text{g/ml}$  streptomycin, 100~U/ml penicillin and 2 mM glutamine. All cell culture media and supplements were obtained from Gibco/Life technologies.

#### Viruses

WT-RCMV-E has previously been described (Ettinger et al, 2012). ΔE27-RCMV-E was generated by CRISPR/Cas9-mediated genome editing. REF were transduced with a mixture of four CRISPR/Cas9 lentiviral vectors carrying sgRNAs that target viral genome up and downstream of the E27 gene (see Table EV3). Successfully transduced cells were selected with 2.5 µg/ml puromycin prior to infection with WT-RCMV-E. This resulted in an E27-knockout virus ( $\Delta$ E27) with a complete deletion of the 1970-bp E27 gene. E27 deletion was verified by PCR and sequencing. Mutant virus was screened by limiting dilution and was double plaque-purified. WT-MCMV was reconstituted from the BAC described in (Jordan et al, 2011). ΔM27-MCMV has been described previously (Le-Trilling et al, 2018). The M27<sub>AxAxxA</sub> mutant was generated according to previously published procedures (Wagner et al, 2002; Tischer et al, 2006) using the primers MCMV-M27-QC1-3\_1 and MCMV-M27-QC1-3\_2 (Table EV4). For the construction of the  $\Delta$ M27-E27HA virus, a DNA fragment harbouring one frt site, the ZeoR gene, the EF1 promoter and E27HA was introduced into the ΔM27-MCMV-BAC (harbouring an frt site instead of the M27) by FLP-mediated homologous recombination in E. coli DH10B using the FLPexpressing plasmid pCP20. Correct mutagenesis was confirmed by restriction digest analysis, Southern blot and PCR analysis. MCMV mutants were reconstituted by transfection of BAC DNA into permissive cells. Viral titres were determined by standard plaque titration on primary MEF, REF or MNC.

#### Plasmids

For CRISPR/Cas9 viral genome editing, crRNA sequences targeting the viral genome up and downstream of the E27 gene were designed manually or using the CRISPOR website (http://crispor.tefor.net); crRNA sequences are presented in Table EV4 and were cloned as sgRNA downstream a human U6 promoter in the pSicoR-CRISPR-PuroR vector (van Diemen et al, 2016). The pHisSUMO-E27 and pHisSUMO-Rattus norvegicus (rn)STAT2 expression plasmids were prepared using standard restriction enzyme cloning. Full-length E27 and rnSTAT2 open reading frames, and the sequence stretch corresponding to amino acid residues 180–387 from rnIRF9, were PCR-amplified from cDNA templates using oligonucleotide primers XmaI-E27-fw and SacI-E27-rev, XmaI-STAT2-fw and NotI-STAT2-rev, or XmaI-IRF9-fw and SacI-IRF9-rev, and ligated in the pHisSUMO plasmid, to generate N-terminal 6xHis-SUMO-fusion proteins. The rnSTAT2 and rnIRF9 cDNA templates were ordered as *E. coli* codon-

optimised Strings DNA fragments from ThermoFisher. To generate pAcGHLT-B-3C-Homo sapiens (hs)DDB1, the thrombin protease site between the GST purification tag and DDB1 in pAcGHLT-B-hsDDB1 was changed to a 3C protease site using insertion/deletion PCR according to (Hansson et al, 2008), using the oligonucleotide primers 3C-DDB1-A and 3C-DDB1-B. To generate pAcGHLT-B-3C-hsDDB1 ΔBPB (amino acid residues 396-705 replaced with a GNGNSG sequence), and HisSUMO-E27 $\Delta$  (residues 284–305 replaced with a GNGNSG linker), the same method was used, with the oligonucleotide primers  $\triangle BPB-A$  and  $\triangle BPB-B$  or  $E27\Delta-A$  and E27  $E27\Delta-B$ , and pAcGHLT-B-3C-hsDDB1 or pHisSUMO-E27 as template. Point mutations were introduced by site-directed mutagenesis using the primers E27-R211E/R214E-fw and E27-R211E/R214E-rev, and pHisSUMO-E27 as template. The M27-FLAG expression plasmid was described previously (Trilling et al, 2011). Site-directed mutations were introduced using the E27-QC and M27-QC primers and the QuikChangeII kit (Agilent) according to the manufacturer's instructions, using pIRES2-EGFP-E27HA and pIRES2-EGFP-M27FLAG as template. For primer sequences, see Table EV4. All constructs were verified by Sanger sequencing.

#### LC-MS/MS analyses

Liquid chromatography coupled to mass spectrometry analyses were performed as described previously (Megger et al, 2017). Briefly, cells were harvested and dissolved in 50 mM ammoniumbicarbonat containing 0.1% (w/v) Rapigest SF Surfactant (Waters, Boston, MA, USA). Four µg proteins per sample were reduced with dithiothreitol for 30 min at 60°C and thereafter alkylated with 15 mM iodoacetamide for 30 min at room temperature. Subsequently, the samples were digested using trypsin (SERVA Electrophoresis, Heidelberg, Germany) in a ratio of 1:20 (w/w) for 16 h at 37°C. The reaction was stopped by adding trifluoroacetic acid (TFA) to final 1% (v/v), and the precipitated surfactant was removed by centrifugation at 16,000 g for 10 min. The supernatants were dried by vacuum centrifugation and dissolved in 0.1% TFA. An amount of 300 ng tryptic peptides per sample was used for LC-MS/MS analysis and injected into an Ultimate 3000 RSLC nanoLC online coupled to an Orbitrap Elite mass spectrometer (both Thermo Scientific). The peptides were concentrated for 7 min on the trap column (Acclaim PepMap 100, 75  $\mu$ m  $\times$  2 cm, C18, 5  $\mu$ m particle size, 100 Å pore size) using a flow rate of 30 µl/min. Subsequently, the samples were separated on the analytical column (Acclaim® PepMap RSLC, 75  $\mu m \times 50$  cm, nano Viper, C18, 5 µm particle size, 100 Å pore size) by applying a gradient from 5 to 40% solvent B over 98 min (solvent A: 0.1% (v/ v) formic acid; solvent B: 0.1% (v/v) formic acid, 84% acetonitrile) at a flow rate of 400 nl/min. The column oven temperature was 60°C. Full-scan MS spectra were acquired in the Orbitrap analyser at a resolution of 60,000 (400 m/z) with a mass range from 350 to 2,000 m/z. Tandem mass spectra of the 20 highest abundant precursors were acquired in the linear ion trap after fragmentation by collision-induced dissociation.

Peptide identification was carried out using Proteome Discoverer 1.4 (Thermo Scientific, Bremen, Germany). The acquired spectra were searched against the UniProt reference proteome for *Rattus norvegicus* (UP000002494, release 2015\_11, 29,880 entries) and UniProt/TrEMBL sequences for *Rat cytomegalovirus* (RCMV-E, isolate England; release 2015\_11, 208 entries) using Mascot (v.2.5;

Matrix Science, London, UK). Trypsin was set as digesting enzyme and one missed cleavage site was considered as well as dynamic oxidation of methionine and static carbamidomethylation of cysteine residues. The mass tolerance was set to 5 ppm for precursor ions and 0.4 Da for fragment ions. The false discovery rate (FDR) was calculated using the percolator function, and only identifications passing an FDR threshold of 1% were considered. Labelfree quantification was carried out using Progenesis QI for Proteomics (ver.2.0.5, Nonlinear Dynamics, Newcastle upon Tyne, UK), and the peptide spectrum matches derived from Proteome Discoverer were used for protein identification. Normalised protein abundancies were exported and further processed using R (R Foundation for Statistical Computing, Vienna, Austria). Statistical significance was calculated using acrsinh-transformed data by ANOVA followed by Tukey's honest significant difference method. Ratios of means were calculated using nontransformed data. Proteins passing an ANOVA p<sub>FDR</sub> value ≤0.05 (adjusted according to Benjamini-Hochberg), a post hoc test P-value ≤ 0.05 and an absolute ratio of means  $\geq 2$  were considered as significantly differentially abundant.

#### **Immunoblotting**

Immunoblotting was performed according to standard procedures. Briefly, cells were lysed as described previously (Trilling *et al*, 2009) and equal amounts of protein were subjected to SDS–PAGE and transferred to nitrocellulose membranes. Immunoblot analysis was performed using the specific antibodies recognising the indicated proteins (Table EV3). Proteins were visualised using peroxidase-coupled secondary antibodies (Dianova) and the ECL chemiluminescence system (Cell Signaling).

#### Co-immunoprecipitation

Transfected HEK293T cells were washed with ice-cold PBS and subsequently lysed as described previously (Le et~al,~2011; Trilling et~al,~2011) by incubation for 1 h on ice in the following buffer: 0.1 mM EDTA, 150 mM NaCl, 10 mM KCl, 10 mM MgCl<sub>2</sub>, 10% (v/v) glycerol, 20 mM HEPES (pH 7.4), 0.5% (v/v) NP-40, 0.1 mM PMSF, 1 mM DTT, 10  $\mu$ M pepstatin A, 5  $\mu$ M leupeptin, 0.1 mM Navanadate and complete protease inhibitor (Roche). Lysates were cleared by centrifugation for 30 min at 4°C and 16,000 g. The IP antibody (Table EV3) was added to the supernatant and incubated overnight at 4°C in a rotating device. Immune complexes were precipitated with Protein-G-Sepharose (GE Healthcare) for 1 h at 4°C in a rotating device and washed with 150, 250 and 400 mM NaCl-containing lysis buffer. Samples were separated by SDS-PAGE for immunoblot analysis of indicated proteins.

#### Luciferase assays

NIH3T3 reporter cells were infected with MCMV, treated with IFN $\beta$  or IFN $\lambda$  for 6 h and lysed for quantification of reporter gene expression (Le-Trilling *et al*, 2018). Luciferase activity was measured according to the manufacturer's instructions (pjk) using a microplate multireader (Mithras2 LB 943; Berthold Technologies). Quantification of luminescence was performed by use of the microplate multireader Mithras2 LB 943 and MicroWin software (Berthold Technologies). The resulting data were analysed using GraphPad

Prism software. The values are reported as mean  $\pm$  standard deviation (SD) as described in the figure legends.

#### Protein production and purification

Constructs in the pHisSUMO vector, containing an N-terminal HisSUMO-tag, followed by a 3C protease cleavage site, were expressed in Escherichia coli Rosetta™ 2(DE3). In a typical expression, 2 L LB medium containing 100 μg/ml Ampicillin and 34 μg/ml Chloramphenicol were inoculated with 20 ml of an overnight culture and grown under shaking in a Multitron HT incubator (Infors) at 37°C, 150 rpm, until  $OD_{600} = 0.7$ . Then, temperature was lowered to 18°C and protein expression was induced by addition of 0.2 mM Isopropyl β-d-1-thiogalactopyranoside (IPTG). Cultures were grown for further 20 h. Bacteria were pelleted by centrifugation at 4,000 rpm, 4°C for 15 min using a JLA-9.1000 centrifuge rotor (Beckman). The pellet was resuspended in 70 ml buffer containing 50 mM Tris-HCl, pH 7.8, 500 mM NaCl, 4 mM MgCl<sub>2</sub>, 0.5 mM tris-(2-carboxyethyl)-phosphine (TCEP), mini-complete protease inhibitors (1 tablet per 50 ml; Merck) and 30 mM imidazole. 5 µl Benzonase (Merck) was added and the suspension was stirred thoroughly. Cells were lysed by passing the suspension at least twice through a Microfluidiser (Microfluidics). The lysate was clarified by centrifugation at 48,000 g for 45 min at 4°C using a JA-25.50 rotor (Beckman) and applied on an XK 16/20 chromatography column (Cytiva) containing 10 ml Ni Sepharose® High Performance beads (Cytiva), using an Äkta pure FPLC (Cytiva) at 4°C. The column was washed with 250 ml buffer containing 50 mM Tris-HCl, pH 7.8, 500 mM NaCl, 4 mM MgCl<sub>2</sub>, 0.5 mM TCEP and 30 mM imidazole. Protein was eluted with the same buffer supplemented with 0.3 M imidazole. Eluent fractions were analysed using SDS-PAGE, and appropriate fractions were pooled and concentrated to 5 ml with centrifugal filter devices (Vivaspin 20, Sartorius). In pHisSUMO-rnSTAT2 preparations, 30 μg 3C protease per mg total protein was added and the sample was incubated for 12 h on ice to cleave off the HisSUMO-tag. In pHisSUMO-E27 purifications, the tag was not removed, because tag-free E27 precipitated out of solution at higher concentration in the absence of additional binding partners. Subsequently, gel filtration chromatography (GF) was performed on an Äkta prime plus FPLC (Cytiva), using a Superdex 200 16/600 column (Cytiva), equilibrated in 10 mM Tris-HCl pH 7.8, 150 mM NaCl, 4 mM MgCl<sub>2</sub>, 0.5 mM TCEP buffer, at a flow rate of 1 ml/min. GF fractions were analysed by SDS-PAGE, appropriate fractions were pooled and concentrated to approx. 20 mg/ml, flash-frozen in liquid nitrogen in small aliquots and stored at -80°C. Protein concentration was determined with a NanoDrop spectrophotometer (ND 1000, Peqlab), using theoretical absorption coefficients calculated based upon the amino acid sequence by Prot-Param on the ExPASy webserver (Wilkins et al, 1999).

Constructs in the pAcGHLT-B vector, encoding an N-terminal GST-His-tag, followed by a 3C protease cleavage site, were expressed by infecting Sf9 cells with recombinant baculoviruses (Autographa californica nucleopolyhedrovirus clone C6), which were generated as described previously (Zhao et al, 2003). Sf9 cells were cultivated in Insect-XPRESS medium (Lonza) at 28°C, 180 rpm in an Innova 42R incubator shaker (New Brunswick). 1 L Sf9 cells at a cell density of  $3\times10^6$  cells/ml were infected with 0.5 ml of high titre recombinant baculovirus for 72 h. Cells were pelleted by

centrifugation at 1,000 rpm, 4°C for 30 min using a JLA-9.1000 centrifuge rotor (Beckman). The pellet was resuspended in 500 ml hypotonic lysis buffer containing 10 mM Tris—HCl pH 7.8, 4 mM MgCl<sub>2</sub>, 30 mM imidazole, 0.5 mM TCEP, mini-complete protease inhibitors (10 tablets; Merck) and stirred at 4°C for 30 min. The lysate was cleared by centrifugation at 4,000 g, 4°C for 30 min. Subsequent purification steps were performed as described previously for proteins expressed in E. coli.

#### Analytical gel filtration (GF)

A sample containing 10  $\mu$ M of each protein component was incubated with 30  $\mu$ g 3C protease in a volume of 150  $\mu$ l buffer containing 50 mM Tris–HCl, pH 7.8, 500 mM NaCl, 4 mM MgCl<sub>2</sub> and 0.5 mM TCEP, for 12 h on ice. To remove the cleaved purification tags and GST-tagged 3C protease, 20  $\mu$ l GSH-Sepharose FF beads (Cytiva) were added and the sample was rotated at 4°C for 1 h. GSH-Sepharose beads were removed by centrifugation at 4°C, 4,000 rpm for 5 min, and 120  $\mu$ l of the supernatant was loaded on an analytical GF column (Superdex 200 Increase 10/300 GL, Cytiva), equilibrated in 10 mM Tris–HCl pH 7.8, 150 mM NaCl, 4 mM MgCl<sub>2</sub> and 0.5 mM TCEP, at a flow rate of 0.5 ml/min, using an Äkta pure FPLC (Cytiva). 1 ml fractions were collected and analysed by SDS–PAGE.

#### Cryo-EM sample preparation

A sample containing 7  $\mu$ M HisSUMO-E27, 7  $\mu$ M GST-DDB1<sub>ABPB</sub> and 7  $\mu$ M rnSTAT2 was incubated with 0.5 mg 3C protease in a volume of 1 ml buffer containing 10 mM HEPES-NaOH pH 7.8, 150 mM NaCl, 4 mM MgCl<sub>2</sub>, 0.5 mM TCEP, for 12 h on ice. Subsequently, the sample was subjected to GF chromatography on an Äkta prime plus FPLC (Cytiva), using a Superdex 200 16/600 column (Cytiva), connected in line to a 1 ml Glutathione Sepharose<sup>®</sup> 4 Fast Flow column (Cytiva), at a flow rate of 1 ml/min, using the same buffer as eluent. GF fractions were analysed by SDS–PAGE. Fractions corresponding to the elution peak containing E27, DDB1<sub>ABPB</sub> and rnSTAT2 were pooled and concentrated to 70  $\mu$ l (protein concentration 4.1 mg/ml).

Suberic acid bis(N-hydroxysuccinimide ester; DSS) cross-linker (Merck) was added 1:1 (w/w) from a 100 µg/ml stock solution in DMSO, and the sample volume was adjusted to 400 µl with buffer containing 10 mM HEPES-NaOH pH 7.8, 150 mM NaCl, 4 mM MgCl<sub>2</sub> and 0.5 mM TCEP. The sample was incubated for 1 h at 22°C, and the cross-linking reaction was quenched by the addition of 20 µl 1 M Tris–HCl pH 7.8 for 10 min at 22°C. Precipitate was removed by centrifugation at 16,000 g, 22°C for 5 min. The supernatant was loaded on a Superdex 200 Increase 10/300 GL GF column, equilibrated in 10 mM Tris–HCl pH 7.8, 150 mM NaCl, 4 mM MgCl<sub>2</sub> and 0.5 mM TCEP, at a flow rate of 0.5 ml/min, using an Äkta pure FPLC (Cytiva). 0.5 ml fractions were collected and analysed by SDS–PAGE. Fractions containing the cross-linked DDB1<sub>ABPB</sub>/E27/STAT2 complex were pooled, concentrated to 1.3 mg/ml, flash-frozen in 5 µl aliquots in liquid nitrogen and stored at  $-80^{\circ}$ C.

R1.2/1.3400 mesh Cu holey carbon grids (Quantifoil) were glow-discharged for 30 s using a Harrick plasma cleaner with technical air at 0.3 mbar and 7 W. 3.5  $\mu$ l solution containing 0.5  $\mu$ M (0.13 mg/ml) cross-linked DDB1\_ $\Delta$ BPB/E27/STAT2 complex were applied to the grid, incubated for 45 s, blotted with a Vitrobot Mark IV device

(Thermo Fisher) for 1–2 s at 4°C and 99% humidity and plunged in liquid ethane. Grids were stored in liquid nitrogen until imaging.

#### Cryo-EM data collection and processing

Data were collected on a 300 kV Tecnai Polara cryo-EM (Thermo Fisher) equipped with a K2 Summit direct electron detector (Gatan) in super-resolution mode, at a nominal magnification of 31,000×, with a pixel size of 0.63 Å/px and the following parameters: defocus range of -1 to  $-2.5 \mu m$ , 50 frames per movie, 200 ms exposure per frame and electron dose of 6.2 e/ $Å^2/s$ , leading to a total dose of 62 e/Å<sup>2</sup> per movie. In four independent sessions, data from four grids were collected using Leginon (Suloway et al, 2005) yielding 8.069 movie stacks. Movies were aligned and dose-weighted using MotionCor2 (Zheng et al, 2017). Initial estimation of the contrast transfer function (CTF) was performed with the CTFFind4 package via the Relion 3.0.7 GUI (Mindell & Grigorieff, 2003; Scheres, 2012; Zivanov et al, 2018). Power spectra were manually inspected to remove ice contamination and astigmatic, weak or poorly defined spectra. Particles were picked using Gautomatch 0.53 (https:// www2.mrc-lmb.cam.ac.uk/download/gautomatch-056/) or the Relion 3.0.7 autopicker as indicated in Appendix Fig S3. All subsequent image processing procedures were performed using Relion 3.0.7 or cryoSPARC 3.1 (Punjani et al, 2017, 2020; Punjani & Fleet, 2021) and are shown in Appendix Fig S3.

#### Model building and refinement

Guided by the cryo-EM density map 1 (Appendix Fig S3) and XL-MS distance restraints (Fig 2C), a partial molecular model of E27 was generated de novo using the program Coot 0.9.5 (Emsley & Cowtan, 2004; Emsley et al, 2010). In addition, in silico AlphaFold2 structure prediction of E27 was performed (Jumper et al, 2021) using the ColabFold web server (https://colab.research.google.com/ github/sokrypton/ColabFold/blob/main/AlphaFold2.ipynb; Mirdita et al, 2022). The AlphaFold2 output coordinates were in very good agreement with the experimentally determined E27 structure (RMSD of 1.489 Å, 2764 aligned atoms). Accordingly, the model was completed by merging the initial model with the AlphaFold2 template and adjusting backbone and side chain positions where necessary using Coot 0.9.5. DDB1 coordinates were taken from a previously solved DDB1/DCAF1-CtD complex structure (Banchenko et al, 2021), rigid body-fitted and adjusted manually using Coot 0.9.5. The coordinates of the rnSTAT2 CCD were extracted from the AlphaFold protein structure database (https://alphafold.ebi.ac.uk/ entry/Q5XI26; Varadi et al, 2022), rigid body-fitted and adjusted manually using Coot 0.9.5. As final step, automated real space refinement was performed using the program Phenix 1.19.2-4158 (Liebschner et al, 2019). The refined DDB1/E27/STAT2 CCD structure was then fitted as rigid body in cryo-EM density map 2 (Appendix Fig S3). Full-length rnSTAT2 coordinates (https:// alphafold.ebi.ac.uk/entry/Q5XI26) were placed by structural superposition of the CCDs. The STAT2 NTD and TAD were removed, and flexible loops in the SH2 domain were trimmed, because they could not be assigned to the experimental density, indicating that they are flexibly attached and mobile relative to the other STAT2 domains. Lastly, a chain break was introduced between STAT2 residues 315 and 317, and the STAT2 portion encompassing residues 317-701

(DBD, LD and SH2) was once more separately fitted as rigid body, to account for a small movement relative to the CCD. Subsequently, the model was subjected to rigid body and grouped b-factor real space refinement using Phenix 1.19.2–4158, followed by molecular dynamics flexible fitting using the Namdinator server (Kidmose *et al*, 2019), applying default parameters.

#### **Cross-linking mass spectrometry**

#### Photo-cross-linking

The cross-linker sulfo-SDA (sulfosuccinimidyl 4,4'-azipentanoate; Thermo Scientific) was dissolved in cross-linking buffer (10 mM HEPES-NaOH, pH 7.8, 150 mM NaCl, 4 mM MgCl, 0.5 mM TCEP) to 100 mM before use. The labelling step was performed by incubating 23 μg aliquots of the DDB1<sub>ΔBPB</sub>/E27/DDA1/STAT2 complex at 1 mg/ml with 0.2, 0.3, 0.6 and 0.8 mM sulfo-SDA, added, respectively, for an hour. The samples were then irradiated with UV light at 365 nm using a Luxigen LZ1 LED emitter (Osram Sylvania Inc.), to form cross links, for 10 s and quenched with 50 mM Tris-HCl pH 7.5 for 20 min. All steps were performed on ice. Reaction products were separated on a Novex Bis-Tris 4-12% SDS-PAGE gel (Life Technologies) in order to identify suitable cross-linker concentrations. Samples corresponding to 0.2 and 0.3 mM SDA concentrations were precipitated in 80% acetone at -20°C. All subsequent steps were carried out at room temperature. Protein pellets were resuspended in denaturing buffer (50 mM NH<sub>4</sub>HCO<sub>3</sub>, 8 M urea), reduced with 2 mM DTT for 30 min and alkylated with 5 mM iodoacetamide for 30 min. Urea concentration was then reduced to 1.5 M by dilution with 50 mM NH<sub>4</sub>HCO<sub>3</sub> and samples were digested with trypsin (Thermo Scientific Pierce; Shevchenko et al, 2006) overnight. The resulting tryptic peptides were extracted, desalted using C18 StageTips (Rappsilber et al, 2003) and pooled. Eluted peptides were fractionated on a Superdex Peptide 3.2/300 increase column (GE Healthcare) at a flow rate of 10 µl/min using 30% (v/ v) acetonitrile and 0.1% (v/v) trifluoroacetic acid as mobile phase. 50 µl fractions were collected and vacuum-dried.

#### XL-MS acquisition

Samples for analysis were resuspended in 0.1% (v/v) formic acid, 3.2% (v/v) acetonitrile. LC-MS/MS analysis was performed on an Orbitrap Fusion Lumos Tribrid mass spectrometer (Thermo Fisher) coupled online with an Ultimate 3000 RSLCnano HPLC system (Dionex, Thermo Fisher). Samples were separated on a 50 cm EASY-Spray column (Thermo Fisher). Mobile phase A consisted of 0.1% (v/v) formic acid and mobile phase B of 80% (v/v) acetonitrile with 0.1% (v/v) formic acid. Flow rates were 0.3 μl/min using gradients optimised for each chromatographic fraction from offline fractionation, ranging from 2% mobile phase B to 55% mobile phase B over 90 min. MS data were acquired in data-dependent mode using the top-speed setting with a 2.5 s cycle time. For every cycle, the full-scan mass spectrum was recorded using the Orbitrap at a resolution of 120,000 in the range of 400 to 1,500 m/z. Ions with a precursor charge state between 3+ and 7+ were isolated and fragmented. Analyte fragmentation was achieved by stepped higherenergy collisional dissociation (HCD) and a decision tree strategy (Kolbowski et al, 2017). Fragmentation spectra were then recorded in the Orbitrap with a resolution of 60,000. Dynamic exclusion was enabled with single repeat count and 60-s exclusion duration.

#### XL-MS processing

A recalibration of the precursor m/z was conducted based on high-confidence (< 1% false discovery rate [FDR]) linear peptide identifications. The recalibrated peak lists were searched against the sequences and the reversed sequences (as decoys) of cross-linked peptides using the Xi software suite (v.1.7.6.3) for identification (Mendes *et al*, 2019). Final cross-link lists were compiled using the identified candidates filtered to < 2% FDR on link level with xiFDR v.2.1.5.5 (Fischer & Rappsilber, 2017) imposing a minimum of 5 observed fragments per peptide and at least one cross-linker-containing fragment. FDR settings were optimised by boosting heteromeric cross links based on delta score, minimum fragments and minimum sequence coverage, and lower result levels.

#### DisVis

Centre of mass position of the DDA1 C-terminal region relative to the complex was determined using DisVis (van Zundert & Bonvin, 2015; van Zundert *et al*, 2017). The DDB1/STAT2/E27 complex structure from this study was selected as the fixed chain and residues 49–76 of DDA1 from PDB 6ue5 (Bussiere *et al*, 2020) were selected as the scanning chain. Forty six SDA cross-linking restraints were entered from  $C\alpha$  to  $C\alpha$  with allowed distance between 2.5 and 22 Å. The rotational sampling was set to 15°.

#### Sequence alignments

Sequence alignments were calculated using the EBI ClustalOmega server (Madeira *et al*, 2019; https://www.ebi.ac.uk/Tools/msa/clustalo/) and adjusted manually using the program GeneDoc (Nicholas *et al*, 1997).

#### Virus taxonomy IDs (TaxID)

HCMV, 10359; MuHV-1, 10366; MuHV-2, 79700; MuHV-8, 1261657.

#### Amino acid sequences used for multiple sequence alignments

E27, NCBI YP\_007016432; M27, NCBI AWV68118; R27, NCBI NP\_064132; T27, NCBI NP\_116370; B27, NCBI AFK83839; rhesus macaque UL27 (RH27), NCBI AAZ80544; UL27, NCBI ACN63126; DCAF1, UniProt Q9Y4B6; DCAF2, UniProt Q9NZJ0; DCAF4, UniProt Q8WV16; DCAF5, UniProt Q96JK2; DCAF6, UniProt Q58WW2; DCAF8, UniProt Q5TAQ9; DCAF9, UniProt Q8N5D0; DCAF10, UniProt Q5QP82; DCAF11, UniProt Q8TEB1; DCAF12, UniProt Q5T6F0; DCAF14, UniProt Q8WWQ0; DCAF15, UniProt Q66K64; DCAF17, UniProt Q5H987; AMBRA1, UniProt Q9C0C7; CSA, UniProt Q13216; DDB2, UniProt Q92466; HBX, NCBI AMH41022; WHX, NCBI AAA46776; SV5V, UniProt P11207; UL145, UniProt F5HF44; rat STAT2, UniProt Q5X126; mouse STAT2, UniProt Q9WVL2; human STAT2, UniProt P52630; rhesus macaque STAT2, UniProt F6R1P6; pig STAT2, UniProt O02799; chicken STAT2, UniProt A0A1D5PIV4; frog STAT2, UniProt A0A1L8HHX5.

# Data availability

LC-MS data have been deposited at ProteomeXchange via the PRIDE partner repository (Perez-Riverol *et al*, 2019) with the dataset identifier PXD033639 (https://www.ebi.ac.uk/pride/archive/projects/PXD033639). XL-MS data have been deposited at the JPOST ProteomeXChange (Deutsch *et al*, 2020) with accession number

JPST001566 (PXD033366; http://proteomecentral.proteomexchange. org/cgi/GetDataset?ID = PXD033366). The EM maps have been deposited at the Electron Microscopy Data Bank (EMDB) with the accession numbers EMD-14802 (map 1, https://www.ebi.ac.uk/emdb/EMD-14802) and EMD-14812 (map 2, https://www.ebi.ac.uk/emdb/EMD-14812). Atom coordinates have been deposited at the Protein Data Bank (PDB) with the accession numbers 7zn7 (DDB1 $_{\Delta BPB}$ /E27/STAT2 CCD, https://www.rcsb.org/structure/7ZN7) and 7znn (DDB1 $_{\Delta BPB}$ /E27/STAT2 FL, https://www.rcsb.org/structure/7ZNN).

**Expanded View** for this article is available online.

#### **Acknowledgements**

This research was supported by the German Research Foundation (DFG) Emmy Noether Programme SCHW1851/1-1 (D.S.), by an EMBO Advanced grant aALTF-1650 (D.S.) and by an iNEXT grant 5419 (D.S.). M.T. receives funding from the DFG through RTG1949, TR1208/1-1 and TR1208/2-1. We thank the MPI-MG for granting access to the TEM instruments of the microscopy and cryo-EM service group; Dominik A. Megger for initial mass spectrometry analyses; Birgit Zülch and Kristin Fuchs for technical support. Open Access funding enabled and organized by Projekt DEAL.

#### **Author contributions**

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#### Disclosure and competing interests statement

The authors declare that they have no conflict of interest.

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