

1 Exploiting Bacterial Effector Proteins to Uncover Evolutionarily Conserved Antiviral Host 2 Machinery

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10 Abstract

11 Arboviruses are a diverse group of insect-transmitted pathogens that pose global public health
12 challenges. Identifying evolutionarily conserved host factors that combat arbovirus replication in
13 disparate eukaryotic hosts is important as they may tip the balance between productive and
14 abortive viral replication, and thus determine virus host range. Here, we exploit naturally abortive
15 arbovirus infections that we identified in lepidopteran cells and use bacterial effector proteins to
16 uncover host factors restricting arbovirus replication. Bacterial effectors are proteins secreted by
17 pathogenic bacteria into eukaryotic hosts cells that can inhibit antimicrobial defenses. Since
18 bacteria and viruses can encounter common host defenses, we hypothesized that some bacterial
19 effectors may inhibit host factors that restrict arbovirus replication in lepidopteran cells. Thus, we
20 used bacterial effectors as molecular tools to identify host factors that restrict four distinct
21 arboviruses in lepidopteran cells. By screening 210 effectors encoded by seven different bacterial
22 pathogens, we identify six effectors that individually rescue the replication of all four arboviruses.
23 We show that these effectors encode diverse enzymatic activities that are required to break
24 arbovirus restriction. We further characterize *Shigella flexneri*-encoded IpaH4 as an E3 ubiquitin
25 ligase that directly ubiquitinates two evolutionarily conserved proteins, SHOC2 and PSMC1,
26 promoting their degradation in insect and human cells. We show that depletion of either SHOC2
27 or PSMC1 in insect or human cells promotes arbovirus replication, indicating that these are
28 ancient virus restriction factors conserved across invertebrate and vertebrate hosts. Collectively,
29 our study reveals a novel pathogen-guided approach to identify conserved antimicrobial
30 machinery, new effector functions, and conserved roles for SHOC2 and PSMC1 in virus
31 restriction.

32 Author Summary

33 Microbial pathogens such as viruses and bacteria encounter diverse host cell responses during
 34 infection. While viruses possess antagonists to counter these responses in natural host species,
 35 their replication can be restricted in unnatural host cells where their antagonists are ineffective.
 36 Bacteria also employ a diverse repertoire of immune evasion proteins known as “effectors” that
 37 can inhibit antimicrobial responses found in invertebrate and vertebrate hosts. In this study, we
 38 hypothesized that some bacterial effectors may target host immunity proteins that restrict both
 39 bacteria and viruses. To test this hypothesis, we screened a bacterial effector library comprising
 40 210 effectors from seven distinct bacterial pathogens for their ability to rescue the replication of
 41 four viruses in insect cells that are normally non-permissive to these viruses. Though numerous
 42 effectors were identified to rescue the replication of each virus, the uncharacterized IpaH4 protein
 43 encoded by the human pathogen *Shigella flexneri* was able to rescue all four viruses screened.
 44 We discovered that IpaH4 enhances arbovirus replication in both restrictive insect and permissive
 45 human cells by directly targeting two novel, evolutionarily conserved antiviral host proteins,
 46 SHOC2 and PSMC1, for degradation. Our study establishes bacterial effectors as valuable tools
 47 for identifying critical antimicrobial machinery employed by eukaryotic hosts.

48 Introduction

49 Arboviruses comprise a diverse group of arthropod-borne viruses that are transmitted by
 50 dipteran (fly and mosquito) vectors to animal and human hosts. For example, vesicular stomatitis
 51 virus (VSV) is a negative-sense single-stranded (ss)RNA virus belonging to the *Rhabdoviridae*
 52 family that is the leading cause of vesicular disease in livestock in the United States, resulting in
 53 costly animal quarantines and trade embargoes [1]. Of the ~500 arboviruses that have been
 54 identified, ~150 are known to cause disease in humans [2]. Consequently, in 2022, the “Global
 55 Arbovirus Initiative” was launched by the World Health Organization to monitor and control
 56 arboviral disease [3]. Notable among arboviruses causing disease in humans are the positive-
 57 sense ssRNA viruses belonging to the *Togaviridae* family. This family includes chikungunya virus,
 58 the second-most prevalent arbovirus infecting humans worldwide [2]. However, the need for
 59 biosafety level-3 facilities to culture wild-type strains of chikungunya virus poses significant
 60 challenges to studying this togavirus. In contrast, other less pathogenic togaviruses [e.g. Ross
 61 River virus (RRV), O'nyong'nyong virus (ONNV), Sindbis virus (SINV)], can be cultured under
 62 BSL-2 conditions and thus have become important models for understanding togavirus-host
 63 interactions [4, 5]. However, we still lack vaccines and antiviral drugs to combat most human
 64 arbovirus infections, including those caused by togaviruses [5]. Thus, the identification of immune

65 mechanisms that restrict arbovirus replication may provide additional avenues for the
66 development of effective strategies to combat arboviral disease.

67 While genome-wide CRISPR-Cas9 and RNA interference (RNAi) screening platforms
68 have been used to identify host immunity factors affecting arbovirus replication [6-8], these assays
69 can be difficult, time-consuming, and cost prohibitive to set up, and are not easily applicable to
70 non-model host systems. Moreover, these assays cannot provide insight into the strategies used
71 by pathogens to combat host antiviral factors identified in these screens. Identification of
72 pathogen-encoded immune evasion proteins (IEPs) targeting host immunity factors is important
73 for several reasons. First, the existence of such IEPs is strong evidence for the physiologic
74 importance of these interactions during the “molecular arms race” between pathogen and host.
75 Second, while some IEPs simply bind/sequester host factors to inhibit their function, others can
76 alter post-translation modifications to modify stability or function [9]. Thus, IEPs can be used as
77 “tools” to both identify the host immunity factors they target and uncover molecular mechanisms
78 that regulate host factor function. Third, IEPs often drive virulence and thus their characterization
79 can reveal pathogenesis mechanisms [10, 11]. It is paramount to develop simplistic, functional
80 assays that can both identify key antiviral factors restricting viral replication and that provide
81 molecular tools to mechanistically dissect the function of such immunity factors.

82 Although arboviruses are well-adapted to replicate in dipteran and mammalian hosts, we
83 have previously shown that several arboviruses, such as VSV and SINV, undergo abortive
84 infections in cells derived from lepidopteran (moth and butterfly) hosts [12, 13]. For example, in
85 *Lymantria dispar* (spongy moth)-derived LD652 cells, VSV and SINV undergo abortive infections
86 post-entry after limited gene expression. However, their replication can be rescued by global
87 inhibition of host transcription or by expression of mammalian poxvirus-encoded IEPs termed
88 “A51R proteins”, suggesting that innate antiviral defenses block VSV and SINV replication in
89 LD652 cells [12]. However, the host immune responses that are at play during restricted arboviral
90 infections in LD652 cells remain poorly defined. More recently, we have reported the full genomic
91 sequence of *L. dispar* and the LD652 cell transcriptome [14], making virus-LD652 cell systems
92 more amenable to uncovering pathogen-host interactions at the molecular level. Our finding that
93 mammalian poxviral IEPs can retain immunosuppressive function in LD652 cells suggests that
94 some pathogen-encoded IEPs target host machinery conserved between insects and mammals.
95 Thus, we were interested in identifying IEPs from other mammalian pathogens that promote
96 arbovirus replication in LD652 cells. Such IEPs might be useful molecular tools in identifying the
97 conserved host immunity factors they target.

98 Bacterial pathogens encode a wide array of IEPs that can manipulate eukaryotic immune
99 responses. Many of these bacterial IEPs are “effector” proteins that are injected into eukaryotic
100 host cells through bacterial secretion systems [10, 15]. These effectors can manipulate, usurp,
101 and/or inhibit a variety of cellular processes once inside the host cell cytoplasm including
102 cytoskeletal dynamics, host signaling cascades, and innate immune responses [10, 15].
103 Interestingly, some bacterial effectors inhibit innate immune pathways that are also antagonized
104 by viruses, such as the Type I interferon (IFN) response [16], suggesting that bacterial and viral
105 pathogens may need to evade common eukaryotic defense mechanisms. Although significant
106 advances have been made towards understanding effector biology, the function of many effectors
107 remains unknown. Understanding bacterial effector function is important because these proteins
108 can be critical drivers of bacterial pathogenesis [10, 15]. However, defining the role of individual
109 effectors during infection can be challenging due to functional redundancy among independent
110 effectors encoded by a single bacterial pathogen [17]. Therefore, experimental strategies to study
111 effector functions outside of bacterial infections may be useful for determining their role during
112 natural infection.

113 Here, we further explore the restricted infections of arboviruses in lepidopterans by
114 infecting moth cells with the rhabdovirus, VSV, and the togaviruses: SINV, RRV, and ONNV. We
115 develop a simple, yet innovative approach to uncover evolutionarily conserved antiviral factors
116 through the identification of bacterial effectors that rescue arbovirus replication in LD652 cells. By
117 expressing a library of 210 effector proteins encoded by seven distinct bacterial pathogens, we
118 identify six effectors capable of rescuing all four restricted arboviruses in LD652 cells: SopB, lpgD,
119 HopT1-2, HopAM1, Ceg10, and lpaH4. Using mutagenesis, we demonstrate the importance of
120 diverse enzymatic functions for SopB, lpgD, HopAM1, and lpaH4 in breaking arbovirus restriction.
121 Moreover, crystallography and cell cultures studies reveal Ceg10 to encode a putative cysteine
122 protease function that is required for arbovirus rescue. By focusing on the *Shigella flexneri*-
123 encoded effector lpaH4, we reveal this novel bacterial E3 ubiquitin ligase to directly target two
124 conserved host proteins, SHOC2 and PSMC1, for degradation in moth and human cells. To our
125 knowledge, roles for these host factors in virus restriction had not been reported in any eukaryotic
126 system. However, we show that depletion of intracellular SHOC2 or PSMC1 levels in moth or
127 human cells promotes arbovirus replication, suggesting they have ancient roles in combating viral
128 infection across diverse eukaryotic host species. Together, our findings demonstrate the utility of
129 using naturally abortive arbovirus infections in lepidopteran cells for the interrogation of arbovirus-
130 host interactions and establish it as a model for identifying conserved host immunity proteins
131 targeted by pathogens.

Results

Inhibition of Host Transcription Rescues Restrictive Arbovirus Replication in LD652 Cells

Previously, we showed that the normally abortive infection of VSV and SINV in LD652 cells can be rescued by treatment of cultures with actinomycin D (ActD), an inhibitor of transcription [12]. ActD globally blocks transcription by host DNA-dependent RNA polymerases by intercalating into GC-rich regions of cellular DNA and thus does not impede viral RNA-dependent RNA polymerase-mediated transcription [18, 19]. The relief of arbovirus restriction by ActD treatment suggests that VSV and SINV undergo abortive infections in LD652 cells due to cellular antiviral responses that require active transcription [12]. To confirm these previous results and to determine if ActD treatment could relieve restriction of additional togaviruses related to SINV, such as RRV and ONNV, LD652 cells were infected with GFP reporter viruses (VSV-GFP [12], SINV-GFP [12], RRV-GFP [20], and ONNV-GFP [21]) in the absence or presence of ActD. Cells were then stained with CellTracker™ Orange Dye and imaged 72 h post-infection (hpi). Representative GFP fluorescence images and quantitative GFP signals (normalized to cell number with CellTracker™ signals) were used as a readout for viral replication and are shown in **Figure 1AB**. As expected, 0.05 µg/mL ActD treatment increased GFP signal in VSV-GFP and SINV-GFP infections by ~10,000- and 100-fold, respectively (**Figure 1B**). Additionally, ActD treatment during VSV-GFP and SINV-GFP infection increased their viral titer by ~1,000-fold for both viruses (**Figure 1C**). During ONNV-GFP and RRV-GFP infections, ActD increased GFP signal and viral titer by ~100-fold and ~10-fold, respectively (**Figure 1ABC**). Importantly, we have previously shown that LD652 cells treated with this dose of ActD retain ~90% viability [12], and thus enhanced virus replication is not due to a general decrease in cell viability. Together, these findings suggest that arbovirus infection of LD652 cells induces a restrictive immune response that requires active host transcription.

Specific Bacterial Effectors Relieve Arbovirus Restriction in LD652 Cells

We have previously shown that poxvirus-encoded A51R proteins are IEPs that rescue restricted arbovirus replication when expressed from plasmids transfected into LD652 cells [12, 22]. Therefore, we asked if bacterial effector proteins, which often function as IEPs, could also rescue arbovirus replication. To do this, we adapted and expanded a previously described effector library for expression in insect cells [23]. Briefly, 210 secreted bacterial effectors from seven pathogens (*Shigella flexneri*, *Salmonella enterica* serovar Typhimurium, *Pseudomonas syringae*, Enterohemorrhagic *E. coli* O157:H7, *Yersinia pseudotuberculosis*, *Legionella pneumophila*, and

164 *Bartonella henselae*) were cloned into the pIB/V5-His insect expression vector and screened for
 165 their ability to alleviate arbovirus restriction in LD562 cells. Cells were transiently transfected for
 166 48 h and then infected with either GFP reporter viruses (RRV-GFP and ONNV-GFP) or luciferase
 167 reporter strains (SINV-LUC and VSV-LUC [12, 22]) for 72 h (**Figure 2A**). Effector proteins that
 168 enhanced viral GFP signals by >2.5-fold or luciferase signals by >4-fold above empty vector-
 169 transfected cells, were considered “hits” in our screen (**Figure 2B-E**). These cutoffs were chosen
 170 to avoid false positives stemming from experimental noise within our screening system and
 171 allowed us to focus on effectors that robustly rescued arbovirus replication. Of the 210 effector
 172 proteins screened, 10 effectors rescued RRV-GFP, 11 rescued ONNV-GFP, 18 rescued SINV-
 173 LUC, and 10 rescued VSV-LUC (**Figure 2B-F** and **Table S1**). Interestingly, effectors generally
 174 rescued in a virus-specific manner. For instance, 21 effectors only rescued one of the four
 175 arboviruses screened (**Figure 2F**). This suggests that these effectors may relieve virus-specific
 176 restrictions to replication. In contrast, seven effectors rescued three or more arboviruses: IpaH4,
 177 SopB, HopT1-2, HopAM1, Ceg10, EspK, and SidM (**Figure 2F**), suggesting that these effectors
 178 may target host restriction mechanisms that are active against a broader range of viral pathogens.

179 *Diverse Effector Activities are Required for Viral Rescue*

180 Because our original effector library did not encode epitope-tagged effector genes, we
 181 wanted to confirm the expression and rescue functions of key hits from our screen. To do this, we
 182 chose five effectors: SopB, HopT1-2, HopAM1, Ceg10, and IpaH4 that rescued at least 3/4
 183 arboviruses screened. These five effectors are collectively encoded by four bacterial pathogens
 184 and have distinct known or putative structures and functions (**Figure 3A**). We cloned these five
 185 effectors into a novel expression vector, pDGOpIE2, along with Flag epitope tags (**Figure 3B**).
 186 The pDGOpIE2 vector uses the same baculovirus-derived OpIE2 promoter [24] to drive effector
 187 expression as in the pIB/V5-His vector used in our initial screen, but the former vector encodes
 188 unique restriction sites that facilitated restriction enzyme-based cloning (**Figure S1**). Using these
 189 constructs, we confirmed the expression of all five Flag-tagged effectors by immunoblot in LD652
 190 cells. We also confirmed expression of specific point mutants in some of these effectors
 191 (described below) predicted to inactivate effector enzymatic activities (**Figure 3B**). We next
 192 transfected either wild-type or mutant effector expression constructs into LD652 cells to
 193 determine: 1) if these Flag-tagged effectors could rescue arbovirus replication and 2) if known or
 194 putative enzymatic functions were required for viral rescue. Below we discuss each of these five
 195 effectors and their ability to rescue restricted arbovirus replication.

196 SopB is encoded by *Salmonella enterica* serovar Typhimurium and functions as a
197 phosphatase to generate phosphatidylinositol-5-phosphate (PI(5)P) from PI(4,5)P to activate
198 Phosphoinositide 3-kinase (PI3K) signaling [25]. This activity has been shown to promote
199 bacterial entry by inducing membrane ruffling [26] as well as altering endosomal maturation and
200 trafficking [27]. The phosphatase activity of SopB relies on a catalytic cysteine (C420) [25] and
201 mutants encoding a C420S substitution (SopB^{C420S}) have impaired enzymatic activity [28, 29].
202 When expressed in LD652 cells, we found wild-type SopB to significantly enhance the replication
203 of all four arboviruses, while SopB^{C420S} did not significantly affect arbovirus replication when
204 compared to empty vector control treatments (**Figure 3C-F**). Importantly, this lack of rescue by
205 the SopB^{C420S} mutant was not due to poor expression as it expressed to higher levels than wild-
206 type SopB (**Figure 3B**). Interestingly, the degree of arbovirus rescue was variable among the
207 viruses assayed. For instance, while SopB enhanced RRV-GFP by only ~three-fold over empty
208 vector controls, it increased the other three viruses by ~30-80-fold. Although the absolute fold
209 changes in arbovirus replication varied between our initial screen using pIB/V5-His and our
210 confirmatory screen with pDGOpIE2, the overall trends in SopB rescue were similar (**Figure 2B-**
211 **E** versus **Figure 3C-F**). Interestingly, the *Shigella flexneri*-encoded IpgD effector shares ~47%
212 amino acid identity to SopB and can also generate (PI(5)P) from PI(4,5)P reliant on a conserved
213 catalytic cysteine (C439) [30, 31]. Therefore, it was surprising that IpgD only rescued one
214 arbovirus in our initial screens. However, several immunoblot experiments failed to detect Flag-
215 tagged IpgD expression in LD652 cells when cloned into pDGOpIE2, suggesting that IpgD may
216 express more poorly in insect cells than SopB. Thus, we cloned a codon-optimized, Flag-tagged
217 form of IpgD into pDGOpIE2 vectors and re-tested its ability to rescue all four arboviruses.
218 Strikingly, we found the codon-optimized IpgD expressed in immunoblots (**Figure 3B**) and
219 enhanced the replication of all four arboviruses (**Figure 3C-F**). This illustrates that poor
220 expression of some effectors in our library may have produced false negatives. Importantly,
221 expression of a catalytically-inactive mutant IpgD^{C439S} [32] failed to rescue arbovirus replication,
222 despite robust expression (**Figure 3B-F**). These data suggest that a common phosphatase
223 activity encoded by effectors from *S. enterica* and *S. flexneri* can break arbovirus restriction in
224 LD652 cells.

225 The effector gene library used to screen for viral rescue included effectors encoded by the
226 plant pathogen *Pseudomonas syringae*. It was therefore notable that two effectors, HopAM1 and
227 HopT1-2, rescued arbovirus replication within the insect cells. These data suggest that host
228 substrates of effectors, highly conserved through to Plantae, can reveal novel points of viral
229 restriction. The HopAM1 effector possess a toll-like receptors and interleukin-1 receptors (TIR)

domain that suppresses pattern-triggered immunity in plant hosts [33]. HopAM1 was recently shown to catalyze the formation of a novel cyclic adenosine monophosphate (ADP)-ribose (cADPR) isomer, termed “v2-cADPR” [33], which is required for its immunosuppressive function and for *P. syringae* pathogenicity [34]. The hydrolase activity of HopAM1 requires a catalytic glutamate (E191) to hydrolyze nicotinamide adenine dinucleotide (NAD⁺) into v2-cADPR [33]. Expression of Flag-tagged HopAM1 constructs in LD652 cells significantly enhanced the replication of all four arboviruses, while a catalytically-inactive mutant (HopAM1^{E191A}) [35] did not significantly affect arbovirus replication (**Figure 3B-F**). HopT1-2 has also been implicated in suppressing plant immunity. Previous work has demonstrated that HopT1-2 antagonizes “nonhost resistance” defenses in plants [36] by suppressing expression of nonhost 1 proteins required for this immune response [37]. However, the molecular mechanism underlying this HopT1-2 function remains unknown. AlphaFold predicted structures (**Figure 3A**) and HHpred homology determination software [38] were unable to identify a putative catalytic activity for HopT1-2. Expression of Flag-tagged HopT1-2 was confirmed by immunoblot in LD652 cells and this construct rescued all four restricted arboviruses as we observed in our original screen (**Figure 3B-F**). Together, these results illustrate that two specific effectors encoded by a plant pathogen can antagonize antiviral responses in animal cells. Future studies will be needed to determine how HopAM1 generation of v2-cADPR breaks viral restriction, and the functional role of HopT1-2.

In addition to identifying effector proteins with known host substrates, our unbiased screen uncovered novel effector proteins involved in breaking host immunity. Ceg10 is an uncharacterized effector secreted by *Legionella pneumophila*, the causative agent of human Legionnaires’ disease [39]. BLAST homology searches identified proteins closely related to Ceg10 in different *Legionella* serovars, yet sequence comparisons did not reveal a putative function. We therefore took a structural biology approach to help assign a function to Ceg10. We were unable to obtain crystals suitable for X-ray crystallography with full-length Ceg10, possibly owing to flexible N- and C-terminal domains as determined by the AlphaFold prediction (**Figure 3A**). We then used limited proteolysis to produce a core Ceg10 central domain (T55-R287) that was amenable to structural determination by crystallography (**Figure 4A**). We determined the structure of the trypsin limited-proteolysis protected core (Ceg10^{TR}) from three different data sets to 1.7 Å, 1.4 Å and 1.5 Å resolution, respectively (**Table S2**).

Overall, Ceg10^{TR} is composed of 8 α -helices surrounding 7 β -sheets in a conformation resembling a cysteine proteinase with a putative catalytic triad consisting of Cys-159, His-196,

263 and Asp-204 (**Figure 4AB**). A structural search via the PDBeFold [40] reveals several homologs
 264 containing the Cys-His-Asp triad but with low sequence identity (**Figure 4A**). Interestingly, while
 265 the Cys-159 (C159) of Ceg10 is modeled in two roughly equal conformers in data set 1, in data
 266 sets 2 and 3, C159 is modeled in only one conformer as a nitrosylated cysteine, or S-nitrosothiol
 267 [41] (**Figure 4BC**). Despite this, both structural determinations overlap modestly (**Figure 4D**). All
 268 crystals used for data sets in this study were grown from the same batch of purified protein, but
 269 the crystallization condition for data set 1 was 2.0 M sodium, potassium phosphate whereas for
 270 data sets 2 and 3 the major precipitants were polyethylene glycols. The significant level of trace
 271 divalent metal ions present in commercial preparations of sodium or potassium phosphate, and
 272 the extended time between initiation of crystal growth and harvesting is likely the reason the
 273 protein crystallized in data set 1 has lost the S-nitrosothiol, as metal catalyzed decomposition has
 274 been well documented for this post-translational modification [42]. In the S-nitrosothiol structure,
 275 the carboxylate of Asp-110 is hydrogen bonded to both the nitrosyl oxygen of C159 and the
 276 sidechain of Trp-206 (**Figure 4B**).

277 The closest homolog of Ceg10, the *Legionella pneumophila* effector RavJ, is the only
 278 homolog that includes an analogous tryptophan (Trp-172) and aspartate (Asp-24), but the
 279 cysteine (Cys-101) is neither modified nor modeled in multiple conformations. Analysis of the
 280 electrostatic potential mapped to the surface of Ceg10 and RavJ reveals that the catalytic site of
 281 RavJ is buried while the S-nitrosothiol of Cys-159 in Ceg10 is solvent exposed (**Figure 4E**).
 282 These data together indicate that Ceg10 harbors a highly reactive catalytic cysteine that may be
 283 critical for regulating host responses to infection. Indeed, expression of Flag-tagged wild-type
 284 Ceg10 significantly enhanced the replication of all four arboviruses. Consistent with our structural
 285 analysis, conversion of the predicted catalytic cysteine to serine (C159S) generated a mutant
 286 (Ceg10^{C159S}) that was unable to rescue three of the four arboviruses screened, despite robust
 287 expression (**Figure 3B-F**). Although Ceg10^{C159S} significantly enhanced RRV-GFP replication
 288 compared to empty vector control, this rescue was significantly weaker than rescue observed with
 289 wild-type Ceg10 constructs (**Figure 3B-F**). Further studies will be needed to confirm Ceg10
 290 function and its regulation by S-nitrosylation, however, these data suggest that a putative cysteine
 291 protease encoded by *L. pneumophila* is capable of targeting antiviral immune pathways in insect
 292 cells.

293 Lastly, IpaH4 is secreted by *Shigella flexneri* which causes Shigellosis in humans [43].
 294 IpaH proteins are a family of bacterial E3 ubiquitin ligases that consist of two domains: a leucine
 295 rich repeat domain (LRR) used for substrate specificity and a novel E3 ligase domain (NEL)

296 containing the catalytic cysteine [15, 44, 45]. We confirmed expression of wild-type IpaH4 and
 297 demonstrated significant viral rescue in all cases, however, an IpaH4 mutant with a serine
 298 substitution at position 339 (IpaH4^{C339S}) was unable to rescue any arbovirus tested (**Figure 3B-**
 299 **F**). Our group [46-48] and others [49, 50] have shown that IpaH proteins antagonize various
 300 eukaryotic immune response pathways by targeting host defense proteins for degradation.
 301 However, IpaH4 has remained an uncharacterized member of the IpaH family. Thus, we were
 302 interested in both determining if IpaH4 was indeed an active E3 ubiquitin ligase and uncovering
 303 the putative host substrates that it targets.

304 *IpaH4 is an E3 ubiquitin Ligase that directly targets host PSMC1 and SHOC2 proteins*

305 To confirm that IpaH4 exhibits E3 ubiquitin ligase activity, we performed *in vitro*
 306 autoubiquitination assays. E3 ubiquitin ligase activity can be revealed by demonstrating: 1)
 307 autoubiquitination function; 2) polymerization of free ubiquitin chains; and/or 3) direct
 308 ubiquitination of a substrate [51, 52]. Given that the substrates of IpaH4 are unknown, we
 309 conducted *in vitro* autoubiquitination experiments with GST-tagged IpaH4 (GST-IpaH4) purified
 310 from *E. coli*. Ubiquitination is a post-translation modification that transfers ubiquitin molecules
 311 from E1 (ubiquitin-activating enzymes) to E2 (ubiquitin-conjugating enzymes) to E3 ubiquitin
 312 ligases, which ultimately covalently attach ubiquitin to target proteins [53]. When mixed with the
 313 necessary components of a ubiquitination reaction: E1, E2, ubiquitin, and ATP, GST-IpaH4
 314 displayed clear autoubiquitination as evidenced by the formation of higher molecular weight
 315 species that reacted with anti-GST and anti-ubiquitin antibodies (**Figure 5A**). However, when E1
 316 was removed from these *in vitro* reactions, these higher molecular weight species were not
 317 detected, as expected (**Figure 5A**). Interestingly, under these reaction conditions, we were unable
 318 to observe autoubiquitination activity for two additional IpaH family members, IpaH2.5 and
 319 IpaH9.8 (**Figure 5A**). This finding is consistent with prior observations that IpaH members typically
 320 display autoinhibition unless in the presence of substrates [54, 55]. Thus, the robust IpaH4
 321 autoubiquitination observed in the absence of substrates suggests an important distinction of
 322 IpaH4 autoregulation when compared with other IpaH members. GST-IpaH4 autoubiquitination
 323 activity could be detected at wide range of protein concentrations (0.2-1 μ M) and typically became
 324 more obvious at higher IpaH concentrations. However, the GST-IpaH4^{C339S} mutant predicted to
 325 inactivate the catalytic cysteine, was unable to display detectable autoubiquitination even at 1 μ M
 326 concentrations (**Figure 5B**). The robust autoubiquitination of wild-type IpaH4 may make it more
 327 prone to proteasomal degradation in host cells, and thus may contribute to the reduced

expression we typically observe when we transfect wild-type and C339S mutant IpaH4 vectors into LD652 cells (**Figure 3B**).

Next, we sought to identify the host substrates of IpaH4. As IpaH4 is encoded by a human pathogen and was able to rescue arbovirus replication in insect cells, we hypothesized that the natural substrates of IpaH4 may be highly conserved between mammals and invertebrates. Therefore, we employed two distinct approaches to identify putative targets of IpaH4 in both moth and human backgrounds. First, ubiquitin activated interaction trap (UBAIT) assays [53, 56] were conducted in LD652 whole cell extract (**Figure 5C**). We have used this approach previously to identify the host substrates of other IpaH members [46, 47]. Briefly, the human ubiquitin gene was cloned in frame with IpaH4 to generate a C-terminal fusion protein (IpaH4^{UBAIT}). This fusion allows IpaH proteins to bind their substrates through their N-terminal LRR domains and catalyze thiol-mediated ligation of the fused ubiquitin to a lysine on their substrates [56]. This results in the covalent linkage of IpaH proteins to their substrates, allowing their identification by mass spectrometry after affinity purification of IpaH-substrate complexes [56]. Using this approach, we identified 24 moth proteins that were enriched at least five-fold in IpaH4^{UBAIT} reactions compared to UBAIT reactions conducted with IpaH2.5^{UBAIT} constructs, which served as our control (**Figure 5D and Table S3**).

Because our goal was to identify evolutionarily conserved mechanisms of host immunity, we took advantage of an existing commercial human prey library and conducted two independent yeast two-hybrid (Y2H) screens using IpaH4 bait. An advantage of using a Y2H approach is that we can identify direct IpaH4-substrate interactions whereas UBAIT techniques tend to identify both direct and indirect IpaH-host interactions. Furthermore, by comparing the putative host substrates identified with UBAIT in the moth system with putative targets identified in human Y2H prey screens, we can detect substrate interactions common between two distinct approaches, and thus more likely to be bona fide interactors. Our Y2H screens with IpaH4 bait identified 12 [57] human proteins as putative IpaH4 interactors with >2 independent clones identified across two screens (**Figure 5D, Figure S2A and Table S3**). We then compared both UBAIT and Y2H screening results and found Leucine-rich repeat protein SHOC2 and proteasome 26S subunit, ATPase 1 (PSMC1; also known as Rpt2) to be the only two reproducibly identified putative substrates across moth and human backgrounds (**Figure S2A**).

To determine if IpaH4 directly ubiquitinates purified human SHOC2 or PSMC1 proteins, *in vitro* ubiquitination assays were conducted. Recombinant Flag-SHOC2 or PSMC1-His proteins were incubated with GST-IpaH4, E1, E2, ubiquitin, and ATP as described above for our

autoubiquitination assays. Wild-type GST-IpaH4, but not GST-IpaH4^{C339S}, was capable of ubiquitinating both host proteins in an E1-dependent manner, as evidenced by the formation of higher molecular weight Flag-SHOC2 and PSMC1-His species (**Figure 5E-F**). In contrast, incubation with GST-IpaH2.5 did not result in ubiquitination of either protein (**Figure 5E-F**). These results suggest that IpaH4 specifically ubiquitinates both SHOC2 and PSMC1 *in vitro*.

Lastly, given that other IpaH family members ubiquitinate substrates for proteasomal degradation [47, 49, 50], we hypothesized that intracellular SHOC2 and PSMC1 protein levels may be reduced in the presence of IpaH4. To assess this, degradation assays were conducted by co-expressing GFP-tagged IpaH4 with Flag-tagged human or moth SHOC2 and PSMC1 constructs in order to determine if IpaH4 activity altered intracellular levels of these putative substrates. Both human Flag-SHOC2 and Flag-PSMC1 levels were dramatically reduced when co-transfected with wild-type GFP-IpaH4 constructs, when compared to co-transfections with GFP- or GFP-IpaH4^{C339S}-expressing vectors in HEK293T cells (**Figure 5G**). Other putative targets of IpaH4 (**Figure S2A**) such as SLU7, PRPF6, CRTC1, and BRD7 that were identified in our Y2H screen with human prey but not in our moth UBAIT assays, mostly showed little-to-no reduction in their levels in HEK293T cells when in the presence of GFP-IpaH4 (**Figure S2B**). One exception to this trend was human RNF214, which was dramatically reduced in level when co-expressed with GFP-IpaH4 (**Figure S2B**). However, RNF214 lacks a discernable *L. dispar* ortholog [14] and therefore was unlikely to be involved in arbovirus restriction in LD652 cells.

We next sought to determine if SHOC1 and PSMC1 were targeted by IpaH4 in the insect cell background. We found that human Flag-SHOC2 and Flag-PSMC1 proteins were dramatically reduced in abundance when co-transfected with GFP-IpaH4 in LD652 cells, suggesting that IpaH4 can target these human proteins when expressed in either human or moth cells (**Figure 5H**). To determine if IpaH4 can also alter the levels of moth-encoded SHOC2 and PSMC1 proteins, we co-transfected Flag-tagged versions of these putative moth targets into LD652 cells in the absence or presence of GFP-IpaH4. These experiments demonstrated that GFP-IpaH4, but not GFP-IpaH4^{C339S} constructs, could dramatically reduce moth Flag-SHOC2 and Flag-PSMC1 levels (**Figure 5I**). Collectively, these results suggest that wild-type, but not catalytically-inactive IpaH4 proteins, can reduce the levels of SHOC2 and PSMC1 proteins encoded by both *L. dispar* and human hosts.

Depletion of SHOC2 and PSMC1 Rescues Restrictive Arbovirus Replication in LD652 Cells

One prediction of our approach is that the host substrates of effector proteins secreted by mammalian and plant pathogens are key regulators of viral restriction in the moth. To then determine if endogenous SHOC2 proteins contributed to arbovirus restriction in moth cells as our data suggests, we adapted a previously described CRISPR-Cas9 system for disrupting gene expression in lepidopteran insects [58] to inhibit SHOC2 expression. We cloned two independent single-guide RNAs (sgRNAs) targeting *L. dispar* SHOC2 into pIE1-Cas9-SfU6-sgRNA-Puro [58], an “all-in-one” vector system that expresses Cas9 nuclease, sgRNA, and a puromycin resistance cassette for selection. As controls, cells were transfected with either empty vector or a sgRNA targeting the *L. dispar* *relish* gene, which encodes a Nuclear Factor- κ B-like transcription factor that we have shown to contribute to VSV and SINV restriction in LD652 cells [12, 22]. Transfected cells underwent three rounds of puromycin selection and were then challenged with reporter arboviruses. As expected, cells expressing sgRNA targeting *relish* were significantly more susceptible to VSV-GFP and SINV-GFP infection when compared to empty vector control treatments. RRV-GFP and ONNV-GFP replication was also elevated in cells expressing *relish*-targeted sgRNAs, indicating that these togaviruses are also restricted by Relish-dependent antiviral responses (**Figure 6A**). Interestingly, cells expressing either SHOC2 sgRNA-A or sgRNA-B were significantly more susceptible to infection with ONNV-GFP, SINV-GFP, and VSV-GFP infection (**Figure 6A**). While only cells expressing SHOC2 sgRNA-A displayed statistically-significant differences in RRV-GFP infection, SHOC2 sgRNA-B-expressing cells trended towards an increased susceptibility to this virus with an ~9-fold higher mean in GFP signal than empty vector controls (**Figure 6A**). These results indicate that SHOC2 is a broadly-acting restriction factor for multiple arboviruses in LD652 cells. However, the specific role of SHOC2 in arbovirus restriction requires further investigation.

PSMC1 has been reported to be an essential component of the 19S cap of the 26S proteasome [59]. Consistent with this, our attempts to knock out PSMC1 in LD652 cells with CRISPR-Cas9 techniques resulted in complete cell death after 1-2 rounds of puromycin selection. However, in mammalian systems, transient PSMC1 depletion has been achieved by siRNA knockdown [60]. Therefore, we sought to deplete PSMC1 in LD652 cells in an analogous manner. However, the application of siRNA-based RNAi in *L. dispar* and other lepidopteran cell types has not been well-established. Thus, we took advantage of a prior study that developed guidelines for designing siRNAs to achieve efficient knockdown in another moth species, *Bombyx mori*, as a basis for our siRNA design for use in LD652 cells [61]. To evaluate the efficiency of siRNA-mediated RNAi in LD652 cells, we designed siRNA targeting the coding sequences of *E. coli* *LacZ*

(negative control) and firefly luciferase. Cells were transfected with either an empty vector or a luciferase-encoding pDGOplE2 plasmid for 48 h and then subsequently transfected with siRNAs targeting transcripts encoding LacZ or luciferase. We then evaluated the relative expression of luciferase using luminescence assays 72 h later. Compared to luciferase signals observed in cells transfected with control LacZ siRNA, there was a significant ~75% reduction in luminescence signals in cells transfected with siRNA targeting transcripts encoding luciferase (**Figure 6B**), suggesting our siRNA design and transfection strategy was relatively efficient at reducing target gene expression.

We next designed three independent siRNAs targeting *L. dispar* PSMC1 sequence in LD652 cells [14] and assessed their relative impact on arboviral replication compared to treatments where control siRNAs targeting LacZ were transfected (**Figure 6C**). As a positive control for arbovirus rescue, siRNAs targeting transcripts encoding argonaute-2 (AGO2), which we have shown to restrict VSV and SINV replication in LD652 cells [62], were also transfected into cells. Compared to LacZ (negative control) siRNA treatments, at least 2/3 PSMC1-targeting siRNA transfections resulted in significant increases in viral replication for all four arboviruses (**Figure 6D**). These data indicate that, like SHOC2, PSMC1 may also play a role in restricting arbovirus replication in LD652 cells. Given that PSMC1 is a proteasome subunit, we asked if treatment of LD652 cells with the proteasome inhibitor bortezomib (Bort) would alter their susceptibility to arbovirus infection. Interestingly, addition of Bort to cell culture media 2 hpi resulted in significantly greater viral replication by 72 hpi (**Figure S3AB**). These data suggest that depletion of a proteasome subunit or inhibition of proteasome activity sensitizes LD652 cells to arbovirus infection. However, the mechanism(s) by which PSMC1 and proteasome activity restrict arbovirus replication will require additional studies in the future.

lpaH4 activity or SHOC2/PSMC1 depletion breaks the restriction of oncolytic virus replication in refractory human cancer cells

After using *lpaH4* as a molecular tool to uncover roles for SHOC2 and PSMC1 in viral restriction in insect hosts, we next wanted to examine if these host factors have conserved antiviral roles in mammalian hosts. However, the wild-type arboviruses used in our study are already well-adapted for robust replication in mammalian host cells. Therefore, we sought an alternative approach to examine whether SHOC2 and PSMC1 contribute to virus restriction in mammals.

Oncolytic virotherapy involves the use of viruses to replicate in, and destroy, cancerous cells. These viruses can also invoke anti-tumoral adaptive responses *in vivo*. For example, VSV strains encoding a deletion or arginine substitution of methionine 51 in the VSV matrix (M) protein (VSV Δ M51/M51R) are being intensively pursued as a potential oncolytic agent [63-68]. These strains display a relatively safe profile because their mutant M proteins are unable to block cellular gene expression and thus are highly susceptible to innate immune responses (e.g. IFN signaling) that are present in normal cells, but typically defective in transformed cells [63-68]. However, a wide array of human cancer cell lines and tumor types have been shown to be refractory to VSV Δ M51/M51R replication, presumably due restriction pathways that are still active in transformed cells [69-73]. This barrier to oncolytic VSV strain replication in refractory cancer types poses a significant challenge to the broad use of these strains for treating diverse malignancies [70, 74]. Given that IpaH4 expression could break wild-type VSV restriction in LD652 cells, we asked whether this bacterial effector could also break restriction of a VSV-M51R strain encoding GFP (VSV-M51R-GFP) in refractory 786-0 human adenocarcinoma cells [71, 75]. Interestingly, transfection of Flag-IpaH4 expression constructs into 786-0 cells led to a significant increase in VSV-M51R-GFP replication compared to empty vector (control) treatments. In contrast, Flag-IpaH4^{C339S} expression did not rescue VSV-M51R-GFP replication, indicating that IpaH4 E3 ubiquitin ligase activity was required to enhance the susceptibility of 786-0 cells to viral replication (**Figure 7AB**). These results also demonstrate that, as in moth cells, IpaH4 can sensitize human cells to VSV infection and demonstrate that bacterial effectors may be useful tools for breaking oncolytic virus restrictions in refractory human cancer cells.

Finally, we asked whether the targets of IpaH4 we identified (SHOC2 and PSMC1) may contribute to virus restriction in human 786-0 cells as observed in moth LD652 cells. To assess the impact of human SHOC2 and PSMC1 on VSV-M51R-GFP restriction, we used three independent siRNAs to deplete 786-0 cells of either SHOC2 or PSMC1 and then assessed VSV-M51R-GFP replication relative to non-targeting (scrambled) siRNA treatments. As a positive control for enhanced VSV-M51R-GFP replication, we also included siRNA treatments targeting *relA*, which encodes a human NF- κ B subunit, as NF- κ B signaling has been shown to contribute to oncolytic VSV strain restriction in these cells [71]. Compared to control RNAi treatments, knockdown of RelA significantly increased VSV-M51R infection as expected. Interestingly, at least 2/3 and 3/3 siRNAs targeting SHOC2 and PSMC1, respectively, sensitized 786-0 cells to VSV-M51R-GFP infection (**Figure 7CD**). Effective and specific knockdown was confirmed with immunoblotting 786-0 whole cell extracts. Notably, *SHOC2* siRNA-A was not as efficient at

knocking down human SHOC2 levels and may explain why we did not detect significant differences in VSV-M51R-GFP replication in those treatments (**Figure 7E**). Collectively, these data suggest that SHOC2 and PSMC1 contribute to virus restriction in human cells and suggest that they may be at least partly responsible for the refractory nature of 786-0 cells to VSV-M51R-GFP replication.

Discussion

Our study illustrates the utility of using abortive virus infections as a screening tool to identify IEPs and the host immunity factors they target. From this work, we identify a variety of bacterially-encoded proteins capable of rescuing individual arbovirus infections, as well as six that are capable of all four viruses screened. Additional analysis into the roles of these effectors showed that three of the effectors with known functions were dependent on their catalytic domains for their viral rescue phenotype. One effector of previously unknown function, Ceg10, was able to be briefly characterized through domain prediction, structure determination, and mutational analysis. Lastly, we established IpaH4 as a strong rescuer for all the arboviruses in our study, and uncovered two evolutionarily conserved targets for this E3 ubiquitin ligase: SHOC2 and PSMC1. These IpaH4 substrates were not only degraded in mammalian culture, but also in invertebrate cells upon expression of wild-type, but not catalytically-inactive, IpaH4. Both SHOC2 and PSMC1 proved important for restricting arbovirus replication in LD652 insect cells and in human 786-0 cells. We envision that our screening methodology can further identify unique IEPs targeting conserved pathways for a variety of pathogen-encoded proteins.

An exciting aspect of our work is that arbovirus replication in moth LD652 cells could be rescued by 30 different bacterial effector proteins, including six (SopB, IpgD, HopAM1, HopT1-2, Ceg10 and IpaH4) that could broadly rescue all four arboviruses examined. Initial characterization of two of these effectors revealed new facets of lipid signaling involved in viral restriction. For example, our studies suggest that acute changes in the phosphorylation status of phosphatidyl inositol (PI) lipids may play a key role in viral restriction in LD652 cells. Not only do both *S. enterica*-encoded SopB and the homologous *S. flexneri*-encoded IpgD rescue arbovirus infection, but inactivation of the phosphatidylinositol phosphatase activities of these bacterial effectors abolished the viral rescue phenotype (**Figure 3C-F**). There have been several reported consequences for the lipid phosphatase activity of SopB and IpgD during bacterial infection including inducing actin remodeling to promote membrane ruffling at the cell surface and bacterial entry [26, 76, 77] and activation of PI3K/AKT signaling to stunt programmed cell death pathways [78, 79]. Interestingly, actin remodeling pathways have been shown to be critical for late stages

of togavirus replication wherein these cytoskeletal arrangements are thought to mediate viral envelope protein transport to the cell surface [80]. Furthermore, many togaviruses activate PI3K/AKT signaling during infection and pharmacological inhibition of this signaling pathway has been shown to inhibit togavirus replication in vertebrate cells [81]. Although future mechanistic studies will be required to determine how exactly SopB and IpgD promote arbovirus replication in LD652 cells, it is possible that their phosphatase activity may complement a defect in the ability of these viruses to manipulate actin dynamics and/or regulate PI3K/AKT signaling pathways in these unnatural host cells.

Two unexpected hits from our screen, *P. syringae* effectors HopAM1 and HopT1-2, suggests the exciting possibility that innate immune responses may be conserved between plants and invertebrate animals. *P. syringae* is a plant pathogen that has been an important model for understanding plant immunity to bacterial infection. Recently, HopAM1 was shown to synthesize a novel cADPR variant, v2-cADPR, which has yet to be explored outside of prokaryotic and plant biology [33, 34]. Our data shows that the E191 residue responsible for the ability of HopAM1 to catalyze the v2-cADPR reaction was necessary for arbovirus rescue in LD652 cells (**Figure 3C-F**). This suggests that the v2-ADPR molecule produced by HopAM1 is countering an innate immunity pathway important for antimicrobial responses conserved across phyla. Interestingly, although HopT1-2 lacks clear homology to known enzymes, it has also been previously implicated in anti-bacterial immunity in plants. HopT1-2 can suppress plant nonhost resistance by inhibiting expression of nonhost 1 proteins required for activation of this immune response pathway, although how HopT1-2 achieves this function is unclear [36, 37]. Nonhost resistance responses are driven by nucleotide-binding leucine-rich repeat (NLR) proteins that recognize non-adapted pathogens [36, 37]. NLR proteins are conserved in animals and are also involved in innate immune responses to pathogens [82]. This raises the intriguing possibility that HopT1-2 may target a conserved component of NLR-mediated immunity pathways shared between plant and animal hosts.

By far the largest proportion of effector proteins used to screen arbovirus restriction are encoded by *Legionella pneumophila*, a bacteria species that utilizes a Type IV secretion system to deliver ~350 effector proteins into the host cells. While many of these effectors exhibit unique enzymatic activities and play different roles during infection, most remain uncharacterized [83]. We found the uncharacterized *L. pneumophila*-encoded Ceg10 protein to reproducibly break viral restriction in LD652 cells. Interestingly, the conserved catalytic cysteine, Cys-159 (C159), is nitrosylated in two of our three structural data sets, indicating that this cysteine is highly reactive

556 and particularly susceptible to oxidation by agents such as nitric oxide. Recent studies indicate
557 that S-nitrosylation of the *Vibrio cholera* virulence regulator AphB suppresses the enzymatic
558 activity and alters virulence gene expression [84]. To date, of the >215,000 structures available
559 on the Protein Data Bank, only 43 deposits display an S-nitrosothiol group, highlighting the unique
560 nature of this modification. Furthermore, to our knowledge, Ceg10 is the first structurally
561 characterized S-nitrosylated *Legionella* effector. We confirmed the importance of this residue by
562 mutating C159 to serine and demonstrated that mutant Ceg10 is not capable of breaking viral
563 restriction in L652 cells. These data suggest that Ceg10 targets a highly conserved immunity
564 evasion factor that may be important for controlling both viral and bacterial infections.
565 Furthermore, it reveals a potential mechanism for host defense against Ceg10 via S-nitrosylation
566 and subsequent inactivation.

567 While determining how SopB, IpgD, HopAM1, HopT1-2 and Ceg10 rescue virus
568 replication in moth cells will be an exciting avenue of future research, we sought to identify novel
569 points of viral restriction by exploring the activity of IpaH4 in greater detail given our prior interest
570 in IpaH family members [46-48]. Through independent UBAIT- and Y2H-based screening
571 methods, we identified host SHOC2 and PSMC1 proteins as conserved IpaH4 substrates in moth
572 and human cells. SHOC2 is a leucine-rich repeat scaffold protein that forms a ternary complex
573 with MRAS and PP1C that activates rapidly accelerated fibrosarcoma (RAF) kinases [85, 86].
574 Although the specific role of SHOC2 during virus infection is unclear, it has recently been shown
575 to activate ERK/STAT signaling in response to bacterial flagellin in shrimp hosts [87], implying
576 that SHOC2 may indeed have conserved roles in combating bacterial and viral pathogens.
577 Although it is unknown if SHOC2 restricts *S. flexneri* infection, this bacterium lacks flagella.
578 Moreover, recognition of flagella cannot explain how SHOC2 contributes to arbovirus restriction,
579 suggesting that different pathogens can activate SHOC2-mediated immunity through distinct
580 mechanisms. These findings highlight the strength of our system for identifying pathogen-
581 encoded modulators of key innate immunity pathways that are likely relevant to both bacterial and
582 viral pathogens.

583 PSMC1 is an essential component of the 19S cap of the proteasome functioning as one
584 of six ATPase subunits [88]. In yeast, PSMC1 (RPT2) controls the gate of the proteasome and
585 regulates the opening through an ATP-binding motif, allowing for substrates to enter [59].
586 Mutations within the ATP-binding motif in yeast PSMC1 was lethal and purified proteasomes
587 exhibited substantially less peptidase activity [89]. Data suggests a similar role of PSMC1 in
588 human proteasomes, although it remains unclear if other ATPase subunits are involved in gate-

589 regulating activity [88, 90]. We show that both knockdown of PSMC1 and suppression of
590 proteasome activity alleviate the restricted arbovirus replication seen in LD652 cells. It is attractive
591 to speculate that the proteasome degrades proteins required for the viral lifecycle in these cells,
592 and that IpaH4 suppresses this activity resulting in viral replication. However, it is entirely possible
593 that PSMC1-mediated virus restriction may be proteasome-dependent or -independent given the
594 growing evidence that proteasome subunits play roles outside the proteasome complex [57, 91,
595 92]. Future research will be needed to address the sophisticated interaction between viral
596 restriction and proteasome functions in the invertebrate host.

597 Given that the arboviruses in our study robustly replicate in human cells, we suspected
598 that it may be more difficult to identify roles for human PSMC1 and SHOC2 in the restriction of
599 these viruses. Thus, we sought to recapitulate a similar restricted arbovirus replication system as
600 found in LD652 cells but in a human cell background. To do this, we took advantage of the fact
601 that the oncolytic virotherapy strain, VSV-M51R, poorly replicates in human renal
602 adenocarcinoma 786-0 cells due to an inability of this mutant virus to inhibit innate immune
603 responses [63-68]. We showed that 786-0 cells became sensitize to VSV-M51R infection after
604 either overexpression of IpaH4 or knockdown of IpaH4 substrates, PSMC1 and SHOC2 (**Figure**
605 **6**). This raises the interesting possibility that either bacterial effector function or inactivation of the
606 host targets of effectors might enhance the replication of oncolytic viruses in cancer cell types
607 normally refractory to these viruses. Importantly, both SHOC2 and PSMC1 are often up-regulated
608 in human malignancies have been linked to cellular transformation and tumor metastasis [93-95].
609 Indeed, gain-of-function SHOC2 mutations have been linked to RASopathies, a group of clinical
610 pathologies associated with dysregulated RAS/MAPK signaling such as Noonan syndrome and
611 patients with these syndromes are predisposed to certain cancers [96-98]. Moreover, elevated
612 SHOC2 and PSMC1 expression has been associated with poorer clinical outcomes among
613 cancer patients [95, 98]. Given the common upregulation of SHOC2 and PSMC1 in transformed
614 cells, they may be particularly relevant to the reported refractory nature of some tumors to
615 oncolytic VSV strains replication [70, 74]. Our work opens up the exciting possibility that
616 expression of IpaH4 (or inhibition of the host factors IpaH4 targets) may provide a novel strategy
617 for increasing oncolytic VSV strain replication in refractory tumors that, in turn, may improve
618 virotherapy efficacy.

619 **Limitations of the study**

620 Our study utilizes a less developed lepidopteran cell system where reagents (e.g. antibodies) that
621 could be used to validate SHOC2 or PSMC1 knockouts/knockdowns in *L. dispar* cells are not

currently available. Furthermore, attempts to use antibodies raised against mammalian SHOC2 or PSMC1 failed to show cross reactivity in LD652 whole cell extract. Thus, we either used alternative methods for validation of knockdown procedures in LD652 cells (ex. Figure 5C) and/or multiple, independent sgRNAs or siRNAs to confirm SHOC2- and PSMC1-related phenotypes in insect cells. Furthermore, because our focus here was on identifying host factors that contribute to virus restriction, we have not yet determined if SHOC2 or PSMC1 are targets of IpaH4 during intracellular *S. flexneri* infection. These future studies will be important for understanding the impact of IpaH4-eukaryotic host interactions to bacterial replication and pathogenesis.

Materials and Methods

Cell Lines and Cell Culture

Mammalian cell lines were maintained at 37°C in 5% CO₂ atmosphere. HEK293T and BHK-21 cells were cultured in DMEM supplemented with 10% FBS. 786-0 cells were cultured in RPMI supplemented with 10% FBS. BSC-40 cells were cultured in MEM supplemented with 5% FBS. All media additionally contained 1% non-essential amino acids, 1% L-glutamine, and 1% antibiotic/antimycotic (Gibco). LD652 cells were cultured as described previously in a 1:1 mixture of Grace's Insect Media (Sigma) and Ex-Cell 420 (Sigma) at 27°C under normal atmospheric conditions [12].

Viruses

Stock preparation and culture of recombinant VSV and SINV was performed as previously described [12]. VSV-M51R-eGFP was obtained from Dr. Doug Lyles (Wake Forest University). RRV-GFP (strain T48), ONNV-GFP (strain SG650) constructs were obtained from Dr. John Schoggins (UTSW Medical Center). VSV, SINV, RRV, and ONNV stocks were amplified using low MOI conditions in BHK-21 cells. Viruses were collected from supernatants using ultracentrifugation (22,000 rpm, 2 h, 4°C) and titrated on BSC-40 using fluorescent plaque/foci assays as described [12].

Viral infections were incubated for 2 h in serum free media (DMEM for mammalian cells or Sf-900 II for invertebrate cells; **Table S4**) before the inoculum was replaced with complete media for the remainder of the infection. Where indicated, complete media containing 0.05 µg/mL ActD or 50 nM Bort was added for the remainder of the infection.

Plasmid Constructs for Mammalian and Insect Cell Expression

652 The bacterial gene library was assembled from a previously generated pENTR library of genes
653 [99]. Additional bacterial effector genes were added to the pENTR library by Gateway® Cloning
654 using BP Clonase II (Invitrogen; **Table S4**), following the manufacturer's instructions. All pENTR
655 effector gene were verified by Sanger sequencing. The entire pENTR effector library was then
656 transferred into the pIB/V5-His vector (Invitrogen; **Table S4**) using LR Clonase II (Invitrogen;
657 **Table S4**), following manufacturer's instructions.

658 N-terminal Flag-tagged versions of lpgD codon-optimized for expression in insect cells
659 [lpgD(CO)], SopB, Ceg10, HopT1-2, and HopAM1 and C-terminal Flag-tagged lpaH4 were
660 generated by PCR amplification using primers containing Flag sequences and flanking SacII /
661 PacI off pIB/V5-His templates using iProof DNA polymerase (Bio-Rad; **Table S4**) and cloned into
662 the pDGOpIE2 vector. The pDGOpIE2 vector was synthesized by Gene Universal and contains
663 an OpIE2 promoter, multiple cloning site, polyA sequence and puromycin resistance cassette. A
664 plasmid map of the pDGOpIE2 vector can be found in **Figure S1**. Catalytically-inactive mutants
665 of lpaH4, SopB, and lpgD were constructed using site-directed mutagenesis by PCR amplification
666 using Q5 DNA polymerase (NEB; **Table S4**).

667 N-Terminal GFP-lpaH4 was constructed through gateway cloning into pEGFP-C2 (for mammalian
668 expression), then PCR amplified using primers introducing SacII / PacI sites and cloned into
669 pDGOpIE2 (for insect cell expression).

670 N-terminal Flag human PSMC1 and SHOC2 were generated via Gateway cloning into pCDNA3,
671 (for mammalian cell expression), then PCR amplified using primers introducing SacII / PacI sites
672 and cloned into pDGOpIE2 (insect cell expression).

673 N-terminal Flag *L. dispar* PSMC1 and SHOC2 constructs were generated by RT-PCR
674 amplification from total RNA isolated from LD652 cells using Superscript III reverse transcriptase
675 (Thermo Fisher), iProof DNA polymerase, and primers containing SacII/PacI sites. SacII/PacI
676 constructs were cloned into a modified pCDNA3 vector containing SacII/PacI sites [12] or
677 pDGOpIE2.

678 *Immunoblotting*

679 Protein extracts were diluted in 5X SDS-PAGE loading buffer then boiled at 95°C for 10 min.
680 Samples were subjected to SDS-PAGE electrophoresis at 125 V for approximately 1.5 h.
681 Separated proteins were transferred to nitrocellulose membranes in either 1X transfer buffer
682 (BioRad; **Table S4**) at 1300 mA at 25°C, or in Towbin Buffer (BioRad; **Table S4**) at 100 V at

683 4°C for 100 min. Membranes were blocked in Odyssey Blocking Buffer (LI-COR; **Table S4**) for
 684 1 h at 25°C. Membranes were blotted with primary antibody overnight at 4°C, with actin serving
 685 as a loading control unless otherwise stated. After three, 5 min washes with PBS-T (PBS, 0.1%
 686 Tween; **Table S4**), membranes were incubated in secondary antibody conjugated to an IRDye
 687 (LI-COR; **Table S4**) for 1 h followed by two, 5 min washes in PBS-T and one 5 min PBS wash.
 688 Membranes were then imaged with an Odyssey Fc Imager (LI-COR; **Table S4**).

689 *Fluorescence Microscopy*

690 Cells were stained with 200 µL of serum free media containing CellTracker™ Orange (Invitrogen;
 691 **Table S4**) dye at specified concentrations for 30 min followed by replacement with 1 mL PBS.
 692 Cells were imaged using a 2X objective on an EVOS-FL fluorescence microscope (Thermo
 693 Fisher; **Table S4**) using RFP and GFP cubes. Each condition had 3 replicate wells, and two
 694 images/well were collected for analysis. Image analysis was conducted using Fiji (NIH) to quantify
 695 the percent area of each field of view containing GFP and RFP signal. Positive signal was
 696 determined by using uninfected wells lacking CellTracker™ stain to set a minimum threshold in
 697 both GFP and RFP channels that was applied to the remaining dataset. Only signal above the set
 698 threshold was highlighted and percent area of each field of view with fluorescent signal was
 699 measured using the Fiji analyze particles function. GFP signals were then divided by the percent
 700 area of RFP to give normalized GFP signal. Fold change in GFP signals were calculated by
 701 dividing normalized GFP values of experimental treatments with normalized GFP values in control
 702 treatments (indicated in each figure).

703 *Structural determination of Ceg10*

704 To determine the structured core of Ceg10 (NCBI accession number YP_094338.1), primers
 705 were designed to clone Ceg10 into pET28b using restriction enzymes NdeI and NotI by HiFi
 706 DNA Assembly kit (NEB; **Table S4**), producing the His_{6x}-TEV-Ceg10 vector. *E. coli* BL21 cells
 707 transformed with His_{6x}-TEV-Ceg10 were grown in LB medium supplemented with kanamycin
 708 (50 mg/ml). After cultures reached an OD₆₀₀ of 0.4-0.5, protein expression was induced with 0.2
 709 mM IPTG overnight at 18°C. Cells were lysed using an Emulsiflex C5 (Avestin) in buffer
 710 containing 50 mM Tris-base, 500 mM sodium chloride (NaCl), 10 mM imidazole, 1 mM
 711 phenylmethylsulfonyl fluoride (PMSF), 0.5 mM tris(2-carboxyethyl)phosphine (TCEP), and
 712 SIGMAFAST™ Protease Inhibitor Cocktail Tablets at pH 7.0. The fusion protein was affinity
 713 purified using Ni-NTA agarose (Qiagen; **Table S4**), eluted with 50 mM Tris-base, 150 mM NaCl,

250 mM imidazole, and 0.1 mM PMSF, and dialyzed overnight into 50 mM Tris-base, 150 mM NaCl, and 1 mM magnesium chloride (MgCl_2) at 4°C. Limited proteolysis of Ceg10 was carried by digesting 6.4 μM His_{6x}-TEV-Ceg10 with 0.89 μM Trypsin (Sigma) in PBS. After 30 min at 25°C, the reaction was quenched by the addition of 1 mM PMSF and the molecular weight of the core was determined by intact mass spectroscopic analysis (UTSW Proteomic Core), revealing a highly abundant peptide (T55-R287) corresponding to residues Thr-55-Arg287 (Ceg10^{TR}) in Ceg10. Primers were designed to clone Ceg10^{TR} pET28b as described above to produce the His_{6x}-TEV-Ceg10^{TR} vector. *E. coli* BL21 cells transformed with His_{6x}-TEV-Ceg10 were grown in LB medium supplemented with kanamycin (50 $\mu\text{g/ml}$). After cultures reached an OD₆₀₀ of 0.4-0.5, protein expression was induced with 0.2 mM IPTG overnight at 18°C. Cells were lysed using an Emulsiflex C5 (Avestin) in buffer containing 50 mM Tris-base, 500 mM NaCl, 10 mM imidazole, 1 mM PMSF, 0.5 mM TCEP, and SIGMAFAST™ Protease Inhibitor Cocktail Tablets at pH 7.4. The fusion protein was affinity purified using Ni-NTA agarose (Qiagen), eluted with 50 mM Tris-base, 150 mM NaCl, 250 mM imidazole, 0.5 mM TCEP and 0.1 mM PMSF, and dialyzed overnight into 50 mM Tris-base, 150 mM NaCl, and 0.5 mM TCEP at 4°C. The 6XHis-tag was removed in the presences of 1 mM EDTA by digestion with His-tagged Tobacco Etch Virus (TEV) protease (1:50 TEV:protein ratio). After 16 h at 25°C, MgCl_2 (5 mM final) was added and TEV protease was removed by passing the cleave reaction over 3 ml of Ni-NTA agarose, followed by washing with 20 ml of 10 mM Tris-base, 50 mM NaCl, and 1 mM TCEP at pH 8.0. Ceg10^{TR} was concentrated to 2.5 ml using a 10,000 MWCO Amicon 50 spin concentrator, subjected to centrifugation at 40,000 rpm (TLA55 rotor), and further purified using a Superdex highload 200 16/600 gel filtration chromatography column (Cytiva Life Sciences). Fractions containing Ceg10^{TR} were concentrated of 20 mg/ml and filtered with a 0.22 μm Durapore® membrane filter.

Native crystals were grown by the sitting drop vapor diffusion method at 4°C in 96-well Intelliplate trays using a 1:1 ratio of protein/reservoir solution containing 1.5 M sodium phosphate monobasic, 0.5 M potassium phosphate dibasic, 10 mM sodium phosphate dibasic/citrate buffer at pH 4.2 and were cryoprotected with 25% ethylene glycol. Native crystals diffracted to a minimum Bragg spacing (d_{min}) of 1.70 Å and exhibited the symmetry of space group C222₁ with cell dimensions of $a = 86.0$ Å, $b = 112.3$ Å, $c = 55.2$ Å, and contained one Ceg10⁵⁵⁻²⁸⁷ per asymmetric unit. Crystals of S-nitrosylated Ceg10⁵⁵⁻²⁸⁷ were grown by the sitting drop vapor diffusion method at 4°C in 96-well Intelliplate trays using a 1:1 ratio of protein/reservoir solution containing 20% PEG 1,000, 0.1 M Tris pH 7.0 and were cryoprotected

747 with 25% ethylene glycol. S-nitrosylated Ceg10⁵⁵⁻²⁸⁷ crystals diffracted to a minimum Bragg
748 spacing (d_{\min}) of 1.40 Å and exhibited the symmetry of space group P2₁2₁2 with cell dimensions
749 of $a = 103.4$ Å, $b = 115.8$ Å, $c = 40.9$ Å, and contained two Ceg10⁵⁵⁻²⁸⁷ per asymmetric unit.
750 Crystals of Ta₆Br₁₂²⁻ derivatized S-nitrosylated Ceg10⁵⁵⁻²⁸⁷ were grown by the sitting drop vapor
751 diffusion method at 4°C in 96-well Intelliplate trays using a 1:1 ratio of protein/reservoir solution
752 containing 10% PEG 8,000, 10% PEG 1,000, 0.2 M MgCl₂, 0.1 M sodium acetate pH 5.0 and
753 were cryoprotected with 25% ethylene glycol. Derivatized S-nitrosylated Ceg10⁵⁵⁻²⁸⁷ crystals
754 diffracted to a minimum Bragg spacing (d_{\min}) of 1.52 Å and exhibited the symmetry of space
755 group P2₁2₁2 with cell dimensions of $a = 103.6$ Å, $b = 116.4$ Å, $c = 40.8$ Å, and contained two
756 Ceg10⁵⁵⁻²⁸⁷ as well as two Ta₆Br₁₂²⁻ clusters per asymmetric unit. All diffraction data were
757 collected at beamline 19-ID (SBC-CAT) at the Advanced Photon Source (Argonne National
758 Laboratory, Argonne, Illinois, USA) and processed in the program HKL-3000 [100] with applied
759 corrections for effects resulting from absorption in a crystal and for radiation damage [101, 102],
760 the calculation of an optimal error model, and corrections to compensate the phasing signal for
761 a radiation-induced increase of non-isomorphism within the crystal [103, 104].

762 Phases were obtained from a single wavelength anomalous dispersion (SAD) experiment using
763 the Ta₆Br₁₂²⁻ derivatized S-nitrosylated Ceg10⁵⁵⁻²⁸⁷ with data to 1.52 Å collected at the LIII-edge
764 of tantalum. Twelve tantalum sites were located by the program AutoSol, part of the Phenix
765 package [105], and an initial model containing 78.5% of all Ceg10⁵⁵⁻²⁸⁷ residues was
766 automatically generated. This model was used for molecular replacement phasing of the native
767 Ceg10⁵⁵⁻²⁸⁷ and the S-nitrosylated Ceg10⁵⁵⁻²⁸⁷. Completion of these models was performed by
768 multiple cycles of manual rebuilding in the program Coot [106]. Positional and isotropic atomic
769 displacement parameter (ADP) as well as TLS ADP refinement was performed using the
770 program Phenix with a random 5.0% of all data set aside for an R_{free} calculation. Data collection
771 and structure refinement statistics are summarized in Table S2.

772 For determining the electrostatic potential of Ceg10 and Ceg10^{TR} the APBS plugin for PyMOL
773 (PyMOL Molecular Graphics System, Version 2.0 Schrödinger, LLC) was used.

774 *Luciferase Assay*

775 Luciferase assays were performed as described with minor modifications [12, 22]. Briefly, at
776 indicated times post infection/transfection, cells were washed in PBS, pelleted by brief
777 centrifugation, and lysed in reporter lysis buffer (Promega; **Table S4**). WCE was spotted in a 96-

well dish and luciferase activity was measured in arbitrary light units (LU) using a FLUOstar Omega plate reader (BMG Labtech).

RNAi and Viral Infection in Cell Culture

Each 21-nucleotide siRNA sequence was designed based on gene prediction using LD652 genome [14] following rules established by others RNAi experiments in *Bombyx mori* [61]. siRNAs were designed against target sites that were at least 75 nts downstream of the initiation codon of each target gene, were 19 nts in length, and had a GC content of 35-55% [61]. Targets were chosen such that the sense siRNA start nucleotide was preceded by “AA” sequence and the antisense siRNA ended on an A or U [61]. Transient siRNA knockdown was achieved by forward transfection of LD652 with Cellfectin II (Gibco; **Table S4**) according to the manufacturer’s protocol for 48 h prior to infection. Cells were then infected with virus for 72 h prior to imaging-based quantification of infection.

Y2H Screens

Y2H screening was conducted by Hybrigenics Services (Boston, MA). The coding sequence for IpaH4 was PCR amplified from Flag-IpaH4 pCDNA3 vectors and cloned into pB66 as a C-terminal fusion to the Gal4 DNA-binding domain (but without the Flag tag) creating Gal4-IpaH4 pB66. The construct was used as a bait for two independent screens with a random-primed human lung cancer cDNA library constructed into pP6. Clones were screened at 4-fold complexity of the library using mating approach with YHGX13 (Y187 ade2-101::loxP-kanMX-loxP, mata) and CG1945 (mata) yeast strains. His⁺ colonies were selected on a medium lacking tryptophan, leucine and histidine. Prey fragments of the positive clones were amplified by PCR and sequenced at their 5’ and 3’ junctions. The resulting sequences were used to identify the corresponding interacting proteins in the GenBank database (NCBI).

UBAIT

For UBAIT assay to capture unknown substrates from cell lysates, LD652 cells were grown to confluency in three 15 cm dishes. Cells were washed with cold PBS and lysed in 300 µL 1x Ubiquitination buffer (Boston Biochem; **Table S4**) containing 1% Triton X-100, 1x Protease inhibitor (Sigma), and 1mM DTT. The lysate was centrifuged for 10 min at 13,000 g and 4°C. The supernatant was transferred to a new tube before adding 25 µg of the GST-IpaH4-3xFlag-ub construct and incubating for 1 h at 4°C with rotation. His-UbE1 (100 nM), His-Ubch5b (2000 nM), ATP (1 mM), MgCl (1 mM) and 1x Ubiquitination buffer (Boston Biochem; **Table S4**) were added

809 and the reaction was incubated at 30°C for 10 min. Following incubation, 300 µL TBS buffer (25
810 mM Tris-HCl pH 7.4, 150 mM NaCl, 1 mM DTT) and 30 µL of washed GST beads were added.
811 The reaction was incubated at 4°C for 2 h with rotation. The reaction was transferred to a column
812 and the beads were washed four times in TBS containing 0.5% Triton-X 100 and two times in
813 TBS before eluting in 150 µL reduced glutathione pH 8.0. Beads were centrifuged and the
814 supernatant was transferred to a new tube followed by addition of 0.25% SDS (final concentration)
815 and 5 mM DTT (final concentration). The solution was boiled for 5 min at 95 °C before adding 1.2
816 mL TBS and 20 µL M2-Flag beads (Sigma; **Table S4**) and incubated at 4°C for 2 h with rotation.
817 Flag beads were then washed four times in TBS containing 0.5% Triton X-100 and twice with
818 TBS. Finally, 35 µL of 95°C SDS-PAGE loading buffer was added, and the beads were boiled an
819 additional 5 min at 95°C. Samples were subjected to SDS-PAGE and Coomassie stained. To
820 identify captured proteins, a lane of the gel above the unmodified IpaH4-UBAIT band was excised
821 and proteins were digested in-gel with trypsin and run on a Q Exactive MS platform at the
822 University of Texas Southwestern Medical Center Proteomics Core.

823 *Degradation Assay*

824 100,000 HEK-293T cells were co-transfected with 150 ng GFP/GFP-IpaH4/GFP-IpaH4^{C339S} and
825 350 ng target vectors for 48 h using FuGENE (Promega; **Table S4**). Cells were harvested in RIPA
826 buffer containing 100 µM PMSF and protease inhibitor cocktail (Abcam; **Table S4**). Protein
827 extracts were then subjected to SDS-PAGE and subsequent immunoblotting as indicated in each
828 figure.

829 *Protein Purification*

830 To obtain C-terminal 5XHis-tagged PSMC1, human *PSMC1* was cloned into pET21a by restriction
831 cloning with NdeI / XhoI and expressed in *E. coli* BL21 in the presence of ampicillin. After the
832 cultures reached OD₆₀₀ 0.7, protein expression was induced with 0.5 mM IPTG at 18 °C overnight.
833 Cells were lysed using an Emulsiflex C5 (Avestin) in buffer containing 25 mM HEPES, 100 mM
834 NaCl, 100 mM KCl, 10% glycerol, 10 mM MgCl₂, 0.5 mM EDTA, 1 mM DTT, and 10 mM Imidazole
835 (HBSi; **Table S4**) with the final pH at 7.5. The fusion protein was affinity-purified using TALON
836 Metal Affinity Resin (Takara Biosciences; **Table S4**) and eluted in buffer containing 25 mM
837 HEPES, 100 mM NaCl, 100 mM KCl, 10% glycerol, 10 mM MgCl₂, 0.5 mM EDTA, 1 mM DTT,
838 and 500 mM Imidazole. The protein was dialyzed using a 10,000 MWCO Slide-A-Lyzer™ G3
839 dialysis cassette (Thermo; **Table S4**) in 1 L of buffer containing 25 mM HEPES, 100 mM NaCl,
840 100 mM KCl, 10% glycerol, 10 mM MgCl₂, 0.5 mM EDTA, 1 mM DTT.

841 To obtain N-terminal 6XHis-MBP-TEV-Flag-tagged SHOC2, *E. coli* carrying pET28a-SHOC2
842 were grown in LB media supplemented with ampicillin. After the cultures reached OD₆₀₀ 0.7,
843 protein expression was induced with 0.5 mM IPTG at 17°C overnight. Cells were lysed by
844 sonication with 5 second pulses and 20 second intervals for 50 min at 4°C in buffer containing 50
845 mM Tris-HCl, 150 mM NaCl, 1 mM TCEP, 1 mM PMSF, and 20 mM Imidazole (HBSi) with the
846 final pH at 8. The lysate was then subjected to centrifugation at 10,000 x g for 30 min at 4°C. The
847 fusion protein was affinity-purified using TALON metal affinity resin (Takara Biosciences) and
848 eluted in buffer containing 50 mM Tris-HCl, 150 mM NaCl, 1 mM TCEP, 0.1 mM PMSF, and 250
849 mM Imidazole. The protein was concentrated using a 30,000 MWCO Amicon 50 spin concentrator
850 then dialyzed using a 10,000 MWCO Slide-A-Lyzer™ G3 dialysis cassette (Thermo) in buffer
851 containing 25 mM Tris-HCl and 75 mM NaCl. Flag-SHOC2 was cleaved from 6X-His-MBP by
852 incubating 37 mg purified protein with 0.65 mg TEV protease at 4°C overnight followed by affinity
853 purification using Nickel affinity resin (Qiagen) where cleaved Flag-SHOC2 was collected in the
854 flow-through.

855 Recombinant GST-IpaH4-3XFlag for UBAIT experiments was obtained as previously described
856 for other IpaH proteins [107, 108]. Briefly, GST-IpaH4-3XFlag was cloned into pGEX6P-1 with a
857 3XFlag peptide followed by the coding sequence for ubiquitin using Gibson Cloning (NEB; **Table**
858 **S4**). Sequence-verified constructs were transformed into *E. coli* BL21 cells, induced, and
859 recombinant protein was purified using glutathione sepharose beads (GE Healthcare Life
860 Sciences).

861 Recombinant GST-IpaH4 for *in-vitro* experiments was expressed and purified as previously
862 described for other IpaH proteins [107, 108]. Briefly, GST-IpaH4 was cloned into pGEX6P-1,
863 transformed into *E. coli* BL21 cells, induced, and recombinant protein was purified using
864 glutathione sepharose beads (GE Healthcare Life Sciences; **Table S4**).

865 *In-Vitro Ubiquitination Assay*

866 *In vitro* ubiquitination reactions were performed in 50 mM HEPES pH 7.5, 150 mM NaCl, 20 mM
867 MgCl₂, and 10 mM ATP (47). Components were mixed as indicated at the following concentrations
868 1 μM UbE1 (E1), 5 μM UbcH5b (E2), 500 nM-10 μM GST-IpaH or GST-IpaH4^{C339S}, 50 μM
869 ubiquitin, and 5 μM His-tagged PSMC1, or 5 μM Flag-tagged SHOC2. Reactions were incubated
870 for 18 h at 30°C before the addition of 2X loading buffer containing β-mercaptoethanol (BME).

871 Samples were then boiled for 10 min at 95°C, subjected to SDS-PAGE, and immunoblotted with
872 anti-GST, anti-Ubiquitin, anti-His or anti-Flag antibodies.

873 *Statistical Analysis*

874 Graphs and charts were presented as mean values \pm standard error of mean (SEM) with individual
875 data points for each independent experiment shown. At least three independent experiments were
876 conducted for all quantitative experiments shown. All statistical analyses were performed with
877 Prism software v10.0.2 (GraphPad) and statistical tests used are indicated in respective figure
878 legends. Statistical significance ($P < 0.05$) between compared groups is indicated in figures as
879 either: ns (not significant), $* = P < 0.05$, $** = P < 0.01$, $*** = P < 0.001$, $**** = P < 0.0001$.

880 **Acknowledgements**

881 We would like to thank Drs. John Schoggins and Andrew Sandstrom (UT Southwestern Medical
882 Center) and members of the Gammon and Alto laboratories for helpful discussions. We also thank
883 Dr. Don Jarvis (University of Wyoming) for providing the pIE1-Cas9-SfU6-sgRNA-Puro vector.
884 We thank the UT Southwestern Medical Center Proteomics Core for assistance with aspects of
885 experimental execution. This work was supported by grants to DBG from the NIH
886 (1R35GM137978-01 and 1R21AI169558-01A1) and by funding to DBG from the UTSW Endowed
887 Scholars Program. NMA was supported by NIH NIAID R01AI083359, The Welch Foundation (I-
888 1704) and The Burroughs Wellcome Fund (1011019). This research was also supported with
889 training grant funding from the NIH to AE, NSB, DBH (T32 AI007520).

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904 Editing.

905 **Competing Interests**

906 The authors declare that there are no competing interests.

907 **Data Availability**

908 All relevant data are within the manuscript or supporting data. Ceg10 unmodified and S-nitrolylated
909 structures (.pdb files) are included with this manuscript.

910 **Figure Legends**

911 **Figure 1. Abortive arbovirus replication in LD652 cells can be relieved with ActD treatment.**

912 **A.** Representative fluorescence microscopy images (GFP channel) of LD652 cells treated with
913 DMSO (vehicle) or 0.05 µg/mL ActD and infected with the indicated GFP reporter strains for 72
914 h. **B.** Fold-change in normalized GFP signals in ActD-treated cultures relative to DMSO
915 treatments. Cells were stained 72 hpi with CellTracker™ Orange dye (not shown) and imaged in
916 GFP and RFP channels to calculate fold-change in GFP signal after normalization of cell number
917 using CellTracker™ (RFP) channel signals. **C.** Fold-change in titer of supernatants from LD652
918 cell cultures treated as in **A-B** 72 hpi relative to input inoculum (dotted line). Data in B-C are
919 means ± SD; n=3. Statistical significance was determined with unpaired student's t-test;
920 ns=P>0.1234, *=P<0.0332, **=P<0.0021, ***=P<0.0002, ****=P<0.0001.

921 **Figure 2. Specific Bacterial Effectors Relieve Arbovirus Restriction in LD652 Cells. A.**

922 Schematic outlining screen for bacterial effectors that rescue arbovirus restriction in LD652 cells.
923 Cells were transfected with expression plasmids from a library consisting of 210 different effector
924 proteins. After 48 h, cells were infected with either GFP or luciferase reporter strains. At 72 hpi,
925 viral replication was quantified using fluorescence microscopy (RRV-GFP and ONNV-GFP) or
926 luciferase assays (VSV-LUC and SINV-LUC). Image was created with Biorender.com. **B-E.** Fold-
927 change in reporter readout, normalized to empty vector controls for all four screens. The cutoff
928 for fold-change in GFP-based assays was set to >2.5, while the cutoff for luciferase reporters was
929 set to >4-fold (represented by dotted horizontal lines). Data points are means. RLU = relative light
930 units. **F.** Summary of bacterial effector proteins that rescued at least one virus. Green blocks
931 indicate the effector rescued the virus indicated in the column header. The bacterium encoding
932 each effector is noted to the right: *Shigella flexneri* (*S. flexneri*), *Pseudomonas syringae* (*P.*
933 *syringae*), *Salmonella enterica* (*S. enterica*), *Legionella pneumophila* (*L. pneumo.*)

934 *Enterohemorrhagic Escherichia coli* 0157:H7 (EHEC). Additional effector proteins from *Yersinia*
935 *pseudotuberculosis* and *Bartonella henselae* were also screened but did not rescue arbovirus
936 replication.

937 **Figure 3. Validation and characterization of top hits from bacterial effector screens. A.**
938 AlphaFold predicted structures, and known or predicted enzymatic functions for indicated effector
939 proteins identified as hits in arbovirus screens. Effector catalytic residues where substitution
940 mutations were made are highlighted in red in AlphaFold structures. **B.** Representative
941 immunoblots of Flag-tagged bacterial effector expression in LD652 cells 48 h post transfection.
942 **C-F.** Fold-change in normalized viral GFP signal relative to empty vector (EV) controls 72 hpi with
943 after transfection with indicated Flag-tagged effector constructs. Wild-type (WT) effectors are
944 compared to their mutants (E/A or C/S). Cells were stained 72 hpi with CellTracker™ dye and
945 imaged to calculate fold-change in normalized GFP signal over signals in EV treatments. Data in
946 C-F are means \pm SD; n=3. Statistical significance was determined with unpaired student's t-test;
947 ns=P>0.1234, *=P<0.0332, **=P<0.0021, ***=P<0.0002, ****=P<0.0001.

948 **Figure 4. Structural analysis of *Legionella pneumophila* effector Ceg10. A.** Six structural
949 homologs of Ceg10, shown in the same orientation with the putative active site near the top of
950 the figure. Top row, from left: Ceg10 (this study), *L. pneumophila* RavJ, *L. pneumophila* LapG.
951 Bottom row, from left: *S. enterica* Ssel, *S. flexneri* Ospl, *P. savastanoi* AvrPphB. Table shows
952 these structural homologs as determined via the Dali Lite server and their PDB ID. **B.** The putative
953 active site of Ceg10 with residues shown in stick representation and hydrogen bonds are shown
954 as dotted yellow lines. The catalytic Cys (C159), Asp (D204) and His (H192) are labeled as well
955 as residues Asp110 and Trp 206 which are hydrogen bonded to each other in both structures. In
956 the S-nitrosylated structure, Asp110 also hydrogen bonds to the nitrosylated-C159 and van der
957 Waals interactions occur between the aromatic ring of Trp206 and the nitrosylation moiety. **C.**
958 Electron density for C159 in the native (left) and S-nitrosylated (right) Ceg10 structures. The final
959 refined 2Fo-mDFc electron density map, contoured at the 1 σ level, is shown superimposed on
960 each residue, as well as a 180° rotation of this region. **D.** Superposition of native (blue) and S-
961 nitrosylated (brown) Ceg10 structures. **E.** Electrostatic surface potential of both Ceg10 structures
962 and RavJ. All structures are orientated with the putative catalytic cysteine residue in
963 approximately the center of the surface, and the orientations between Ceg10 and RavJ
964 correspond to protein alignments. The displayed surface is colored by electrostatic potential from
965 -10 kT (red) to + 10 kT (blue), as calculated by the APBS plugin in PyMOL.

Figure 5. Identification of host SHOC2 and PSMC1 as conserved targets of IpaH4. **A.** Representative immunoblot of *in vitro* ubiquitination assay performed with indicated GST-IpaH proteins in the absence of substrates. **B.** Representative immunoblot of *in vitro* ubiquitination assay performed with indicated concentrations of wild-type GST-IpaH4 or GST-IpaH4^{C339S} mutant proteins. **C.** Schematic outlining UBAIT protocol [53, 56]. **D.** Venn diagram showing conserved putative substrates (overlapping region) of IpaH4 across UBAIT experiments (n=3) in LD652 cell lysates and Y2H screens (n=2) against a human prey library. **E.** Representative immunoblot of *in vitro* ubiquitination assay showing IpaH4-mediated ubiquitination of human Flag-SHOC2 proteins. **F.** Representative immunoblot of *in vitro* ubiquitination assay showing IpaH4-mediated ubiquitination of human PSMC1-His proteins. **G.** Representative immunoblot of degradation assays using indicated Flag-tagged human proteins in transfected HEK293T cells co-expressing GFP, IpaH4 (WT) or catalytic mutant GFP-IpaH4^{C339S} (C339S). **H.** Representative immunoblot of degradation assays using indicated Flag-tagged human proteins in transfected LD652 cells co-expressing GFP, IpaH4 (WT) or catalytic mutant GFP-IpaH4^{C339S} (C339S). **I.** Representative immunoblot of degradation assays of Flag-tagged moth (*L. dispar*) protein in LD652 cells expressing GFP, IpaH4 (WT) or catalytic mutant GFP-IpaH4^{C339S} (C339S). Image was created with Biorender.com.

Figure 6. Depletion of IpaH4 substrates SHOC2 and PSMC1 enhances arbovirus replication in LD652 cells. **A.** Fold-change in normalized viral GFP signals in cells expressing gRNA targeting Relish or SHOC2 relative to empty vector controls 72 hpi. Cells were stained with CellTracker™ dye 72 hpi and imaged to calculate fold-change in normalized GFP signal over empty vector (control) treatments. Data are means ± SD; n=3. Statistical significance was determined with unpaired student's t-test. **B.** Relative Light Units (RLU) of LD652 lysates from cells transfected with empty vector (EV) or luciferase (Luc)-expressing vectors for 48 and then transfected with siRNA targeting LacZ (negative control) or Luc sequences for 48 h. Data are means ± SD; n=3. Statistical significance was determined with unpaired student's t-test. **C.** Schematic detailing siRNA knockdown protocol to assess impact of PSMC1 on restricted arbovirus replication in LD652 cells. Image was created with Biorender.com. **D.** Fold-change in normalized viral GFP signals relative to LacZ siRNA (control) treatments. Cells were stained with CellTracker™ dye 72 hpi and imaged to calculate fold-change in normalized GFP signal over LacZ (control) siRNA treatments. Data are means ± SD; n=3. Statistical significance was determined with unpaired student's t-test; ns=P>0.1234, *=P<0.0332, **=P<0.0021, ***=P<0.0002, ****=P<0.0001.

Figure 7. Bacterial effector expression or depletion of effector targets enhances oncolytic

virus replication in human cancer cells. A. Representative fluorescence microscopy images (GFP channel) of human 786-0 renal adenocarcinoma cell line infected with VSV-M51R-GFP at 16 hpi. **B.** Fold-change in normalized viral GFP signal relative to empty vector (EV) control from experiments as in **A**. Cells were stained with CellTracker™ Orange dye 16 hpi and imaged to calculate fold-change in normalized GFP signal over EV treatments. Data are means \pm SD; n=3. Statistical significance was determined with unpaired student's t-test. **C.** Representative fluorescence microscopy images (GFP channel) of 786-0 cells infected with VSV-M51R-GFP at 16 hpi after transfection with indicated siRNAs. **D.** Fold-change in normalized viral GFP signal relative to scrambled (control) treatments as in **C**. Cells were stained with CellTracker™ dye 16 hpi and imaged to calculate fold-change in normalized GFP signal over control treatments. **E.** Representative immunoblot of 786-0 whole cell lysate 72 h post-transfection with indicated siRNAs. For **B** and **D**: ns=P>0.1234, *=P<0.0332, **=P<0.0021, ***=P<0.0002, ****=P<0.0001.

Supplementary Tables

Supplemental Table 1. Bacterial effector library screening results. 1A. Fold-change in GFP or luciferase reporter signals for each reporter arbovirus after transfection of indicated expression construct.

Supplemental Table 2. Data collection and refinement statistics, Ceg10 structures.

Supplemental Table 3. Identification of IpaH4 substrates. Screening putative substrates of IpaH4.

Supplemental Table 4. Key resources and reagents.

Supplementary Figure Legends

Figure S1. Generation of insect expression vector pDGOpIE2. A. Snapgene vector map of pDGOpIE2 vector and features of interest. **B.** Complete sequence of pDGOpIE2 vector.

Figure S2. Identification of putative targets of bacterial E3 ubiquitin ligase IpaH4. A. Table summarizing results of two independent Y2H screens using a human prey library and three independent ubiquitin-activated interaction trap (UBAIT) assays using LD652 cell lysate. Hits were then analyzed via Blastp to determine percent identity to their *Homo sapiens* ortholog. N.D. = Not determined; either no ortholog found or no significant homology (as determined by BLAST). (B) Representative immunoblot of degradation assays for Flag-tagged human proteins following 48

h co-expression in HEK293T cells with GFP, lpaH4 (WT) or catalytic mutant GST-lpaH4C339S (C339S).

Figure S3. Proteasomal activity plays a role in restricting arbovirus replication in LD652 cells. **A.** Representative fluorescence microscopy images (GFP channel) of LD652 cells 72 hpi with the indicated GFP reporter strains that were treated with DMSO (vehicle) or 50 nM Bortezomib (Bort). DMSO or Bort was added 2 hpi. **B.** Fold-change in normalized GFP signals in Bort-treated cultures relative to DMSO treatments. Cells were stained 72 hpi with CellTracker™ Orange dye (not shown) and imaged in GFP and RFP channels to calculate fold-change in GFP signal after normalization of cell number using CellTracker™ (RFP) channel signals. Data in B are means ± SD; n=3. Statistical significance was determined with unpaired student's t-test; ns=P>0.1234, *=P<0.0332, **=P<0.0021, ***=P<0.0002, ****=P<0.0001.

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1302 PubMed Central PMCID: PMC4816540 and YC and JHK are employed by Tunnell Government
1303 Services, Inc. There are no patents, products in development or marketed products to declare. These
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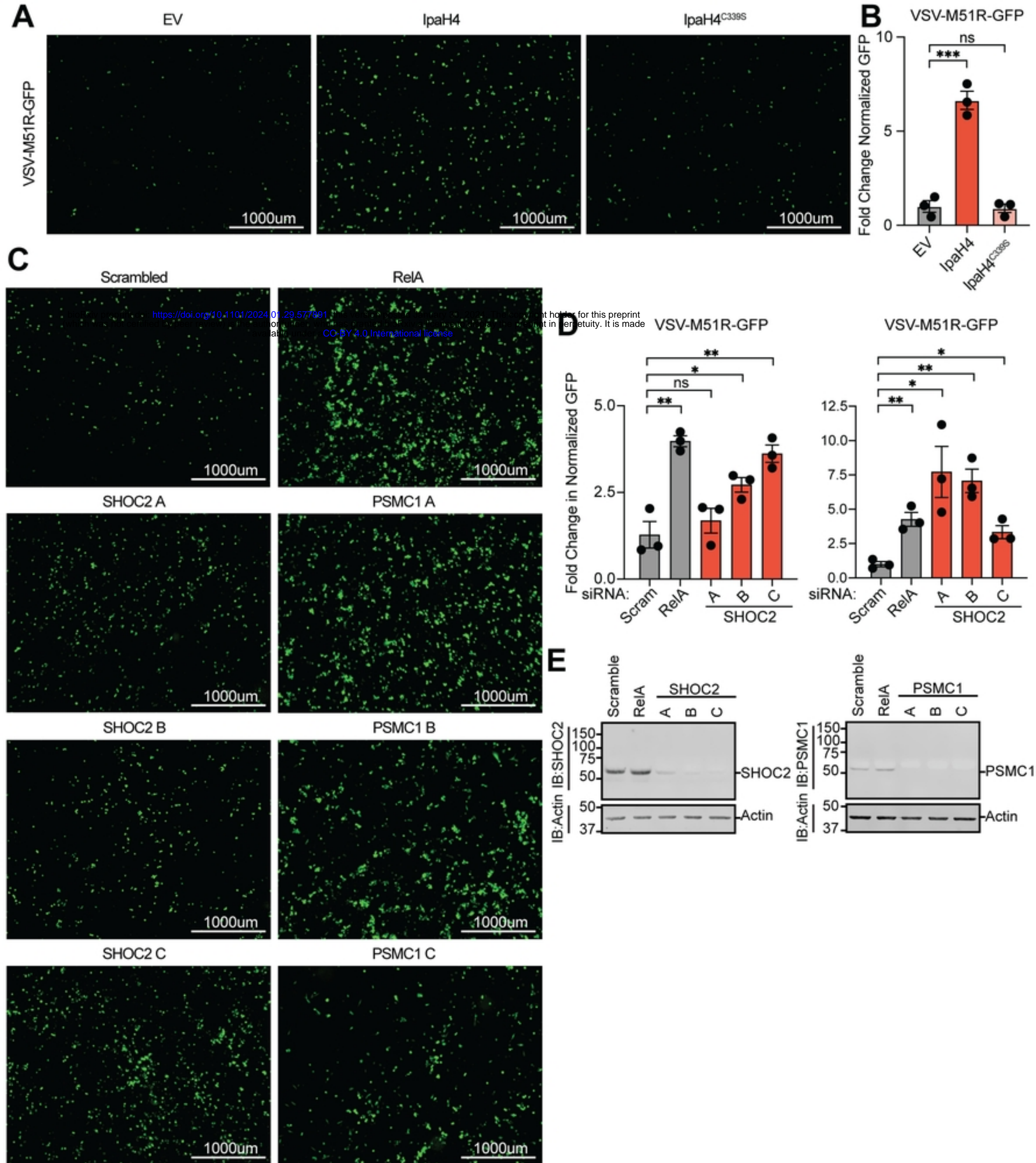


Figure 7

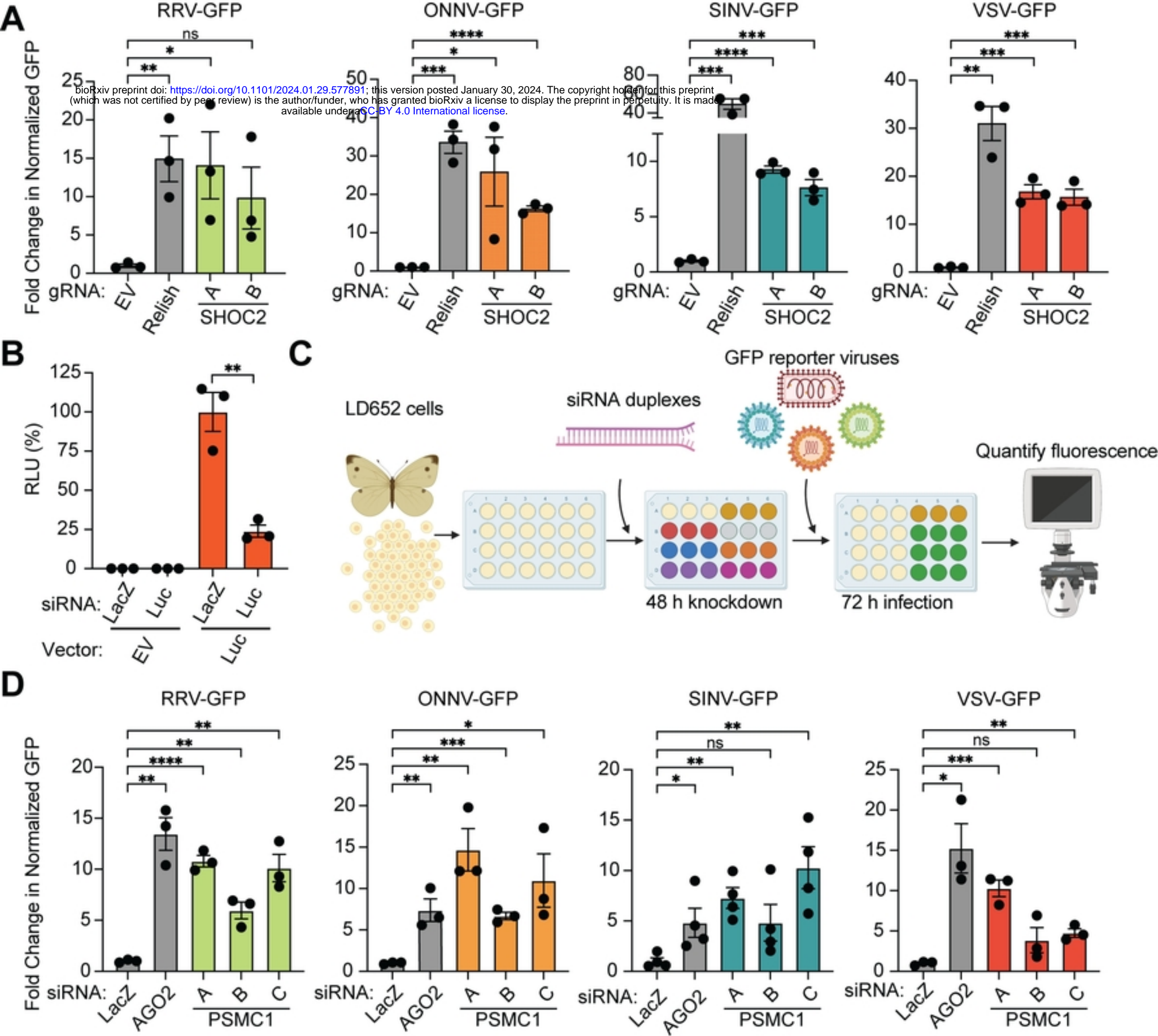


Figure 6

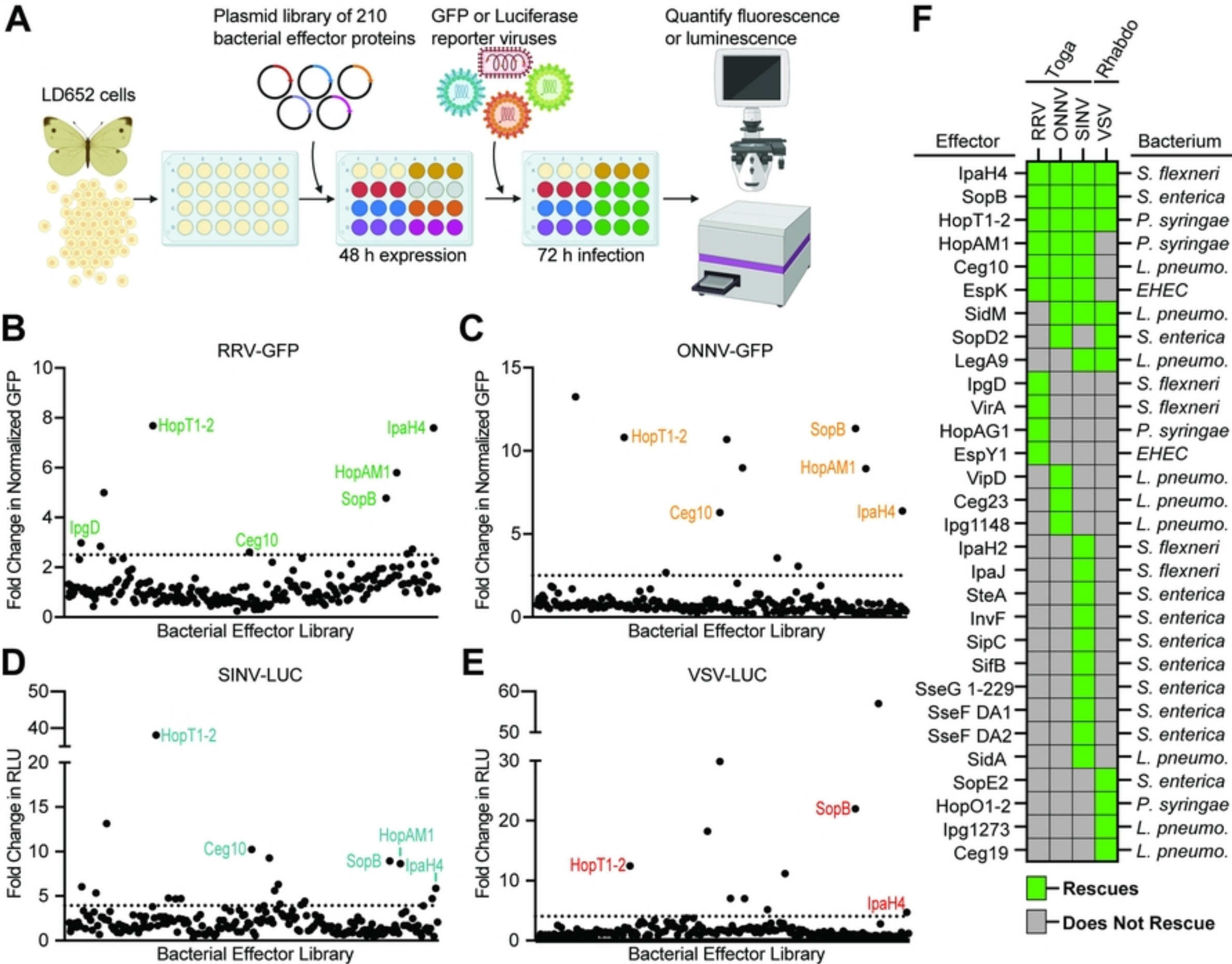


Figure 2

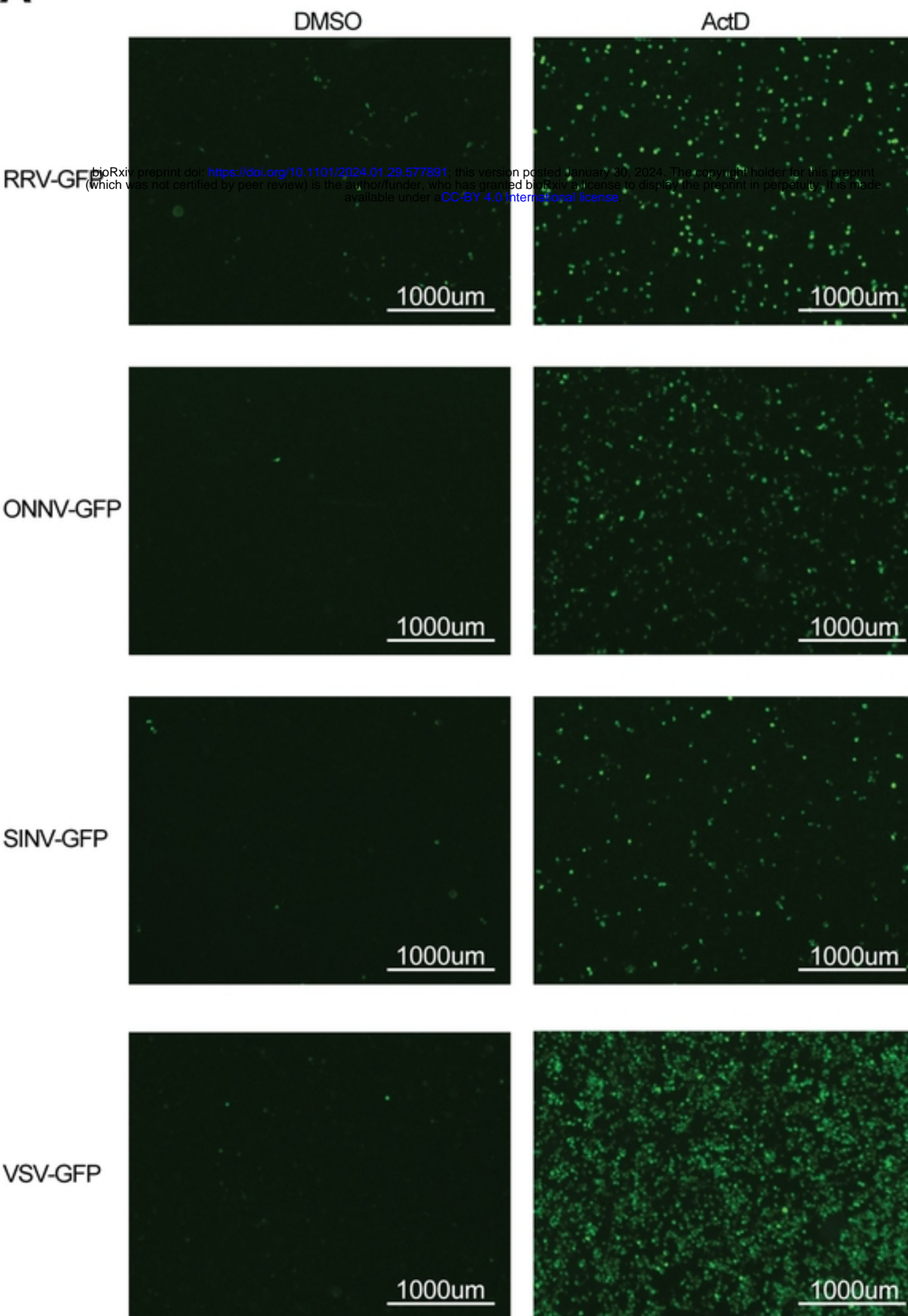
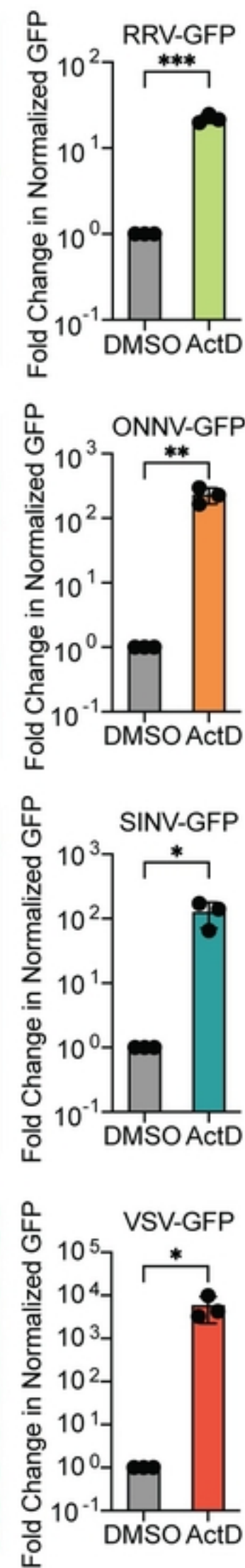
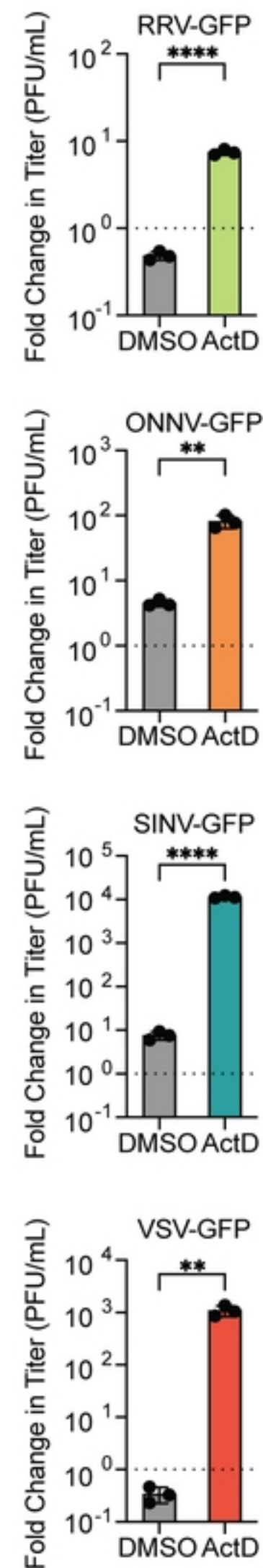
A**B****C**

Figure 1

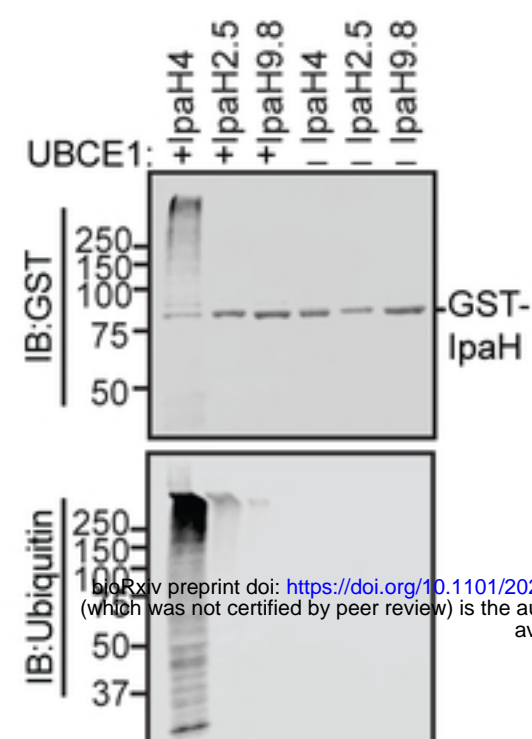
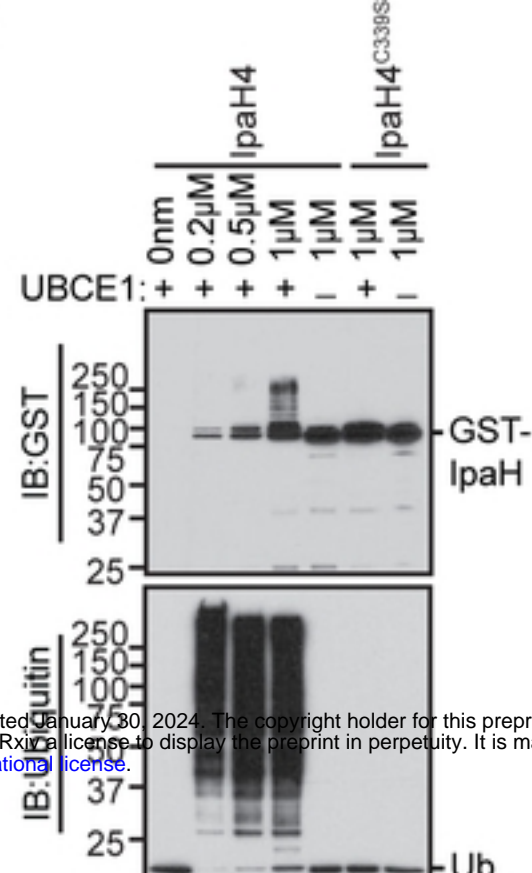
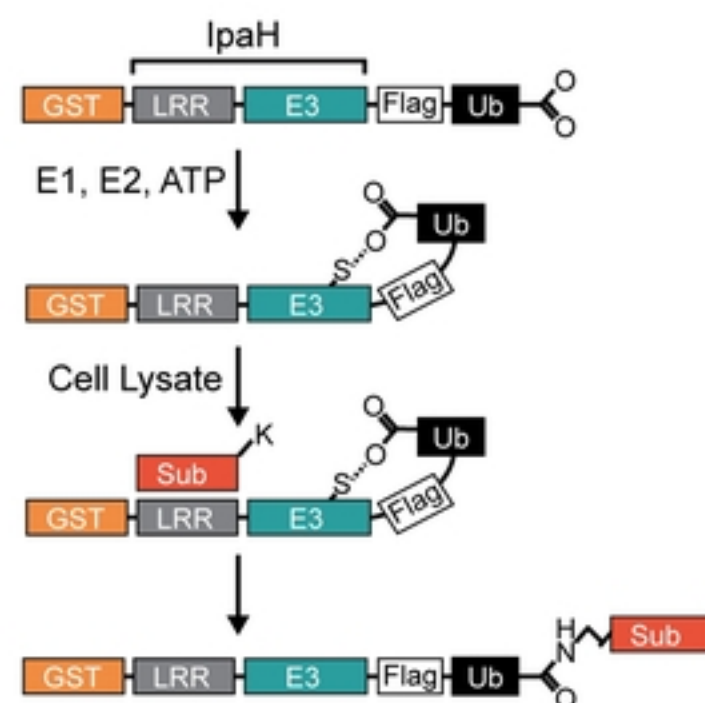
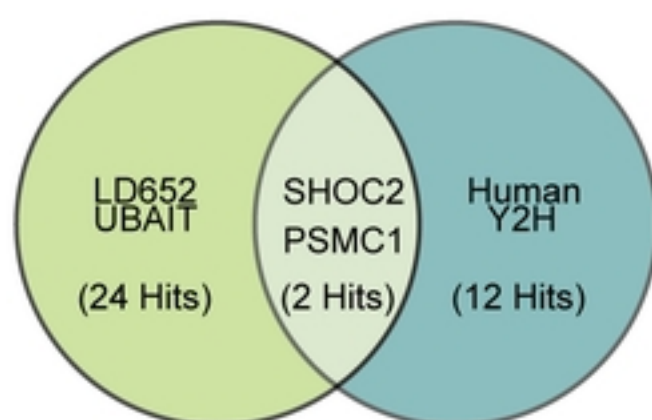
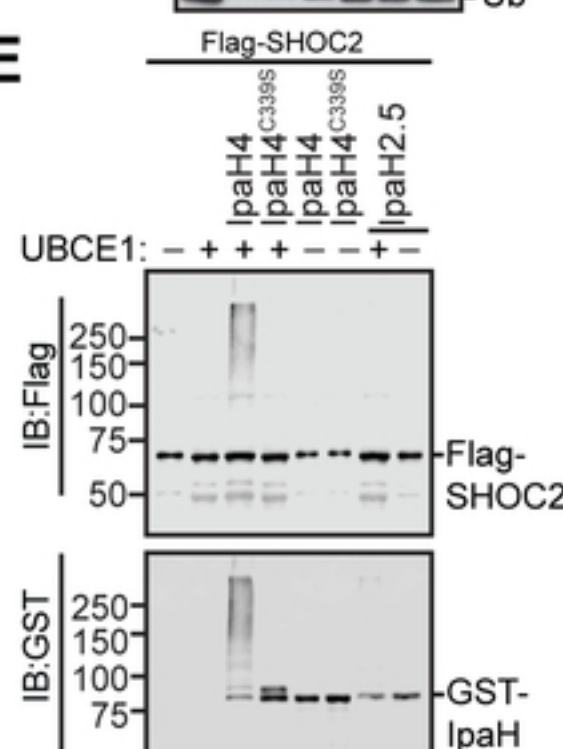
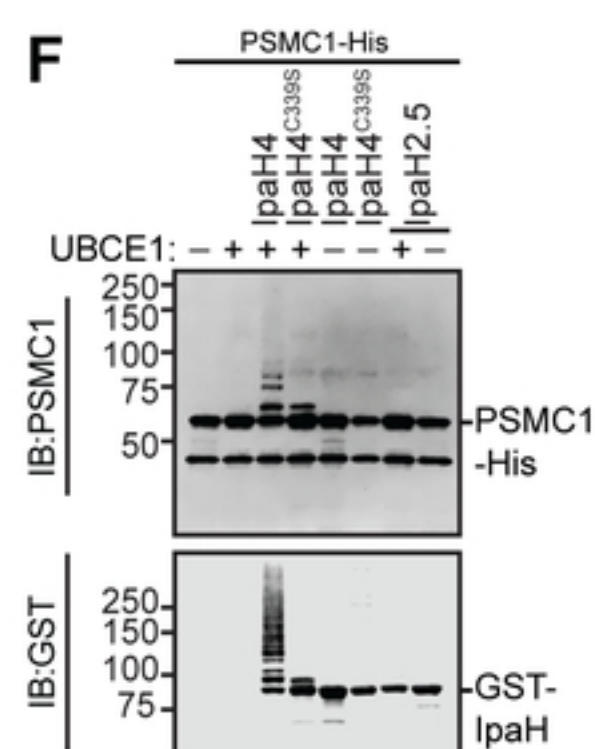
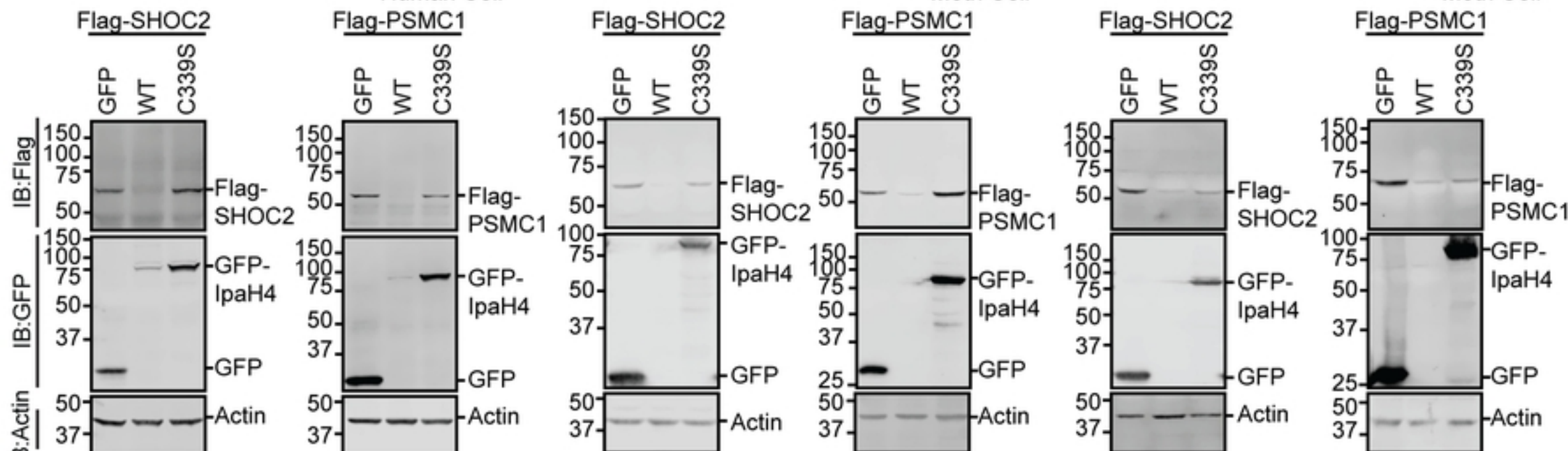
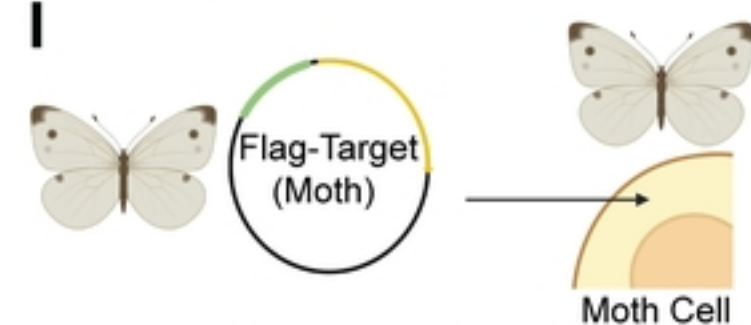
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Figure 5

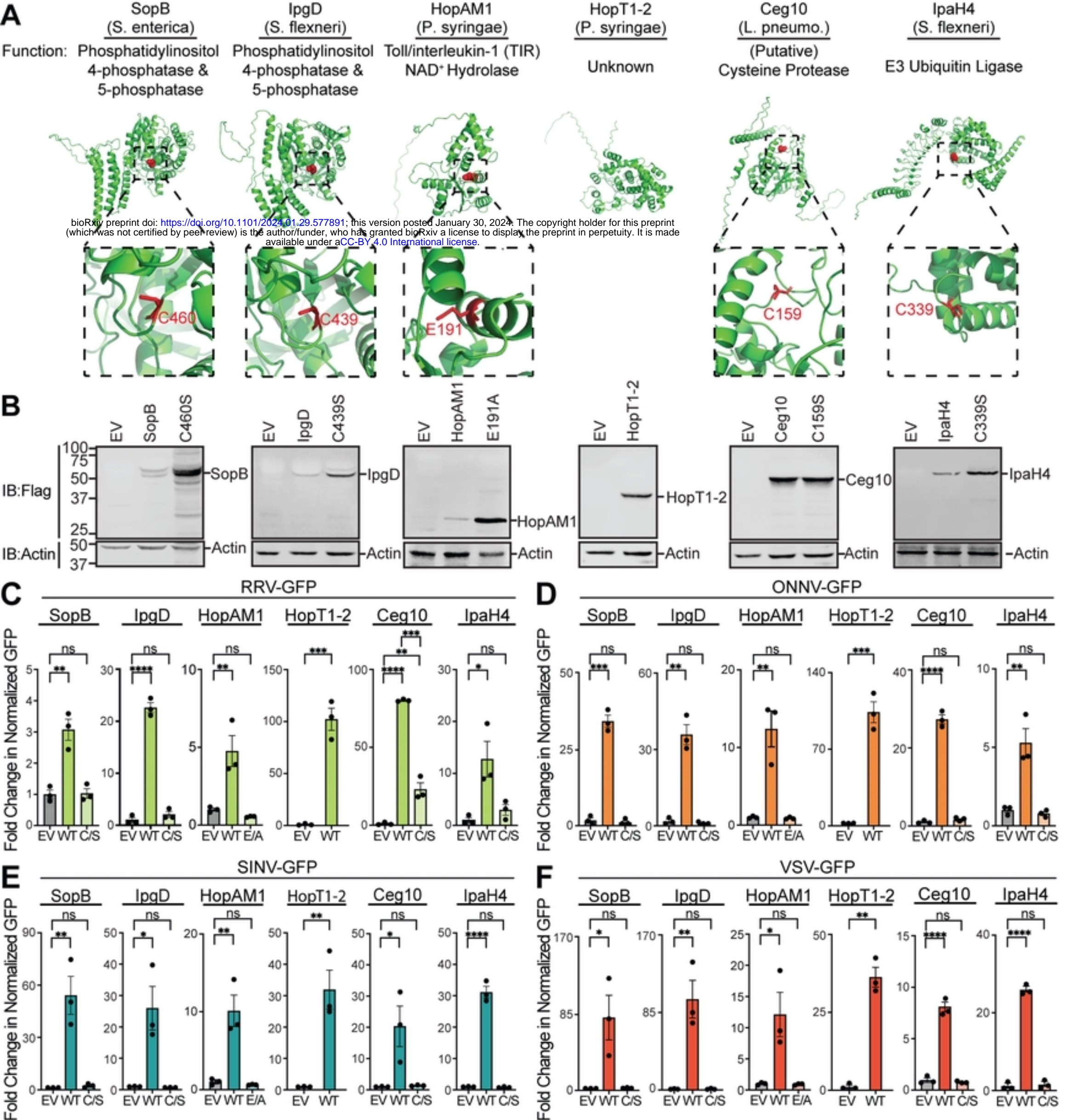


Figure 3

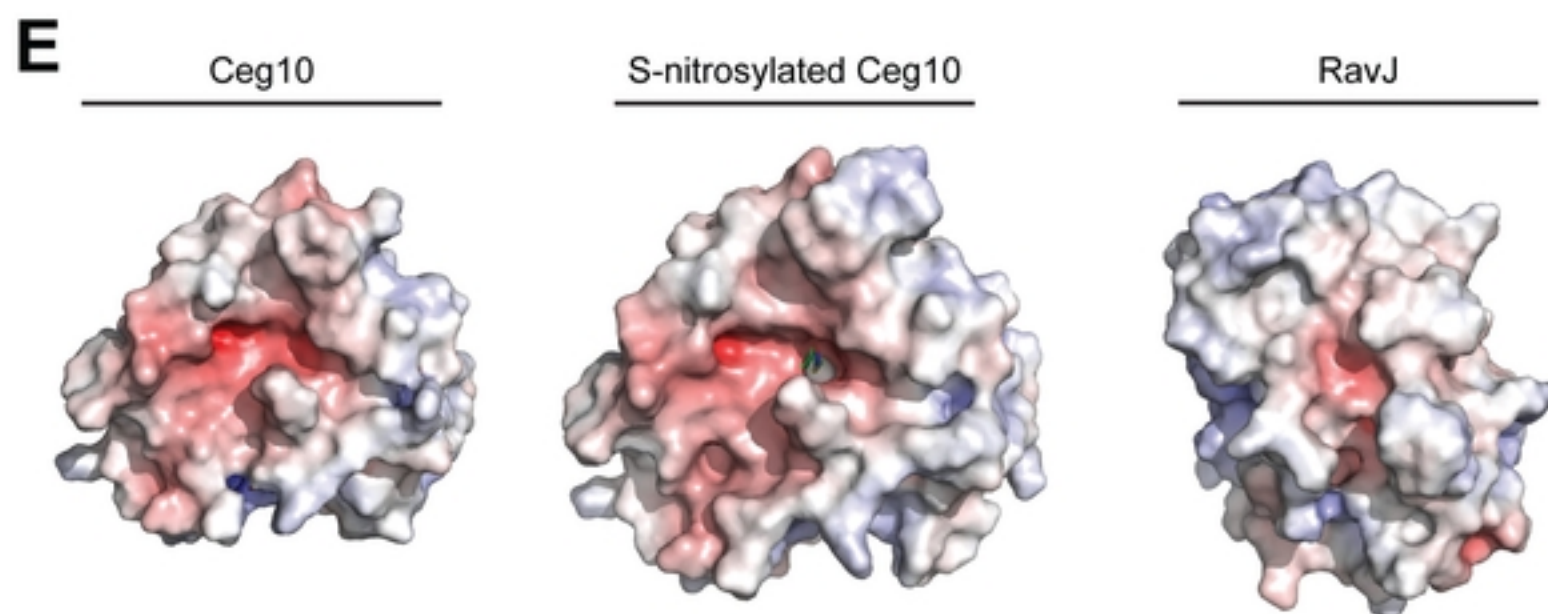
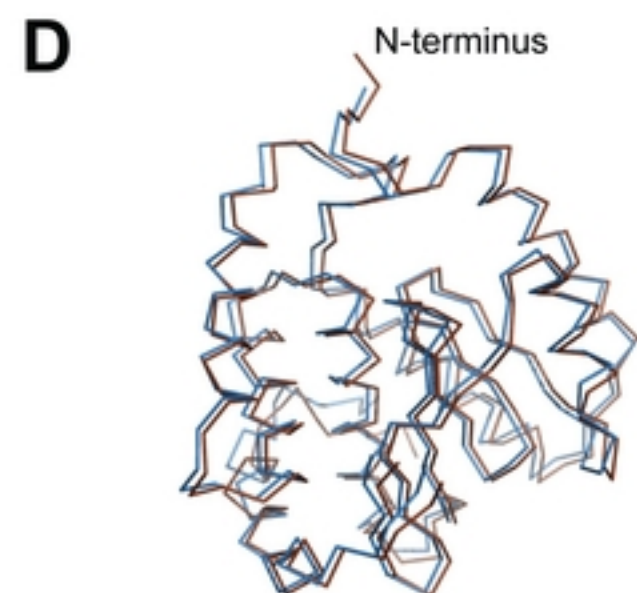
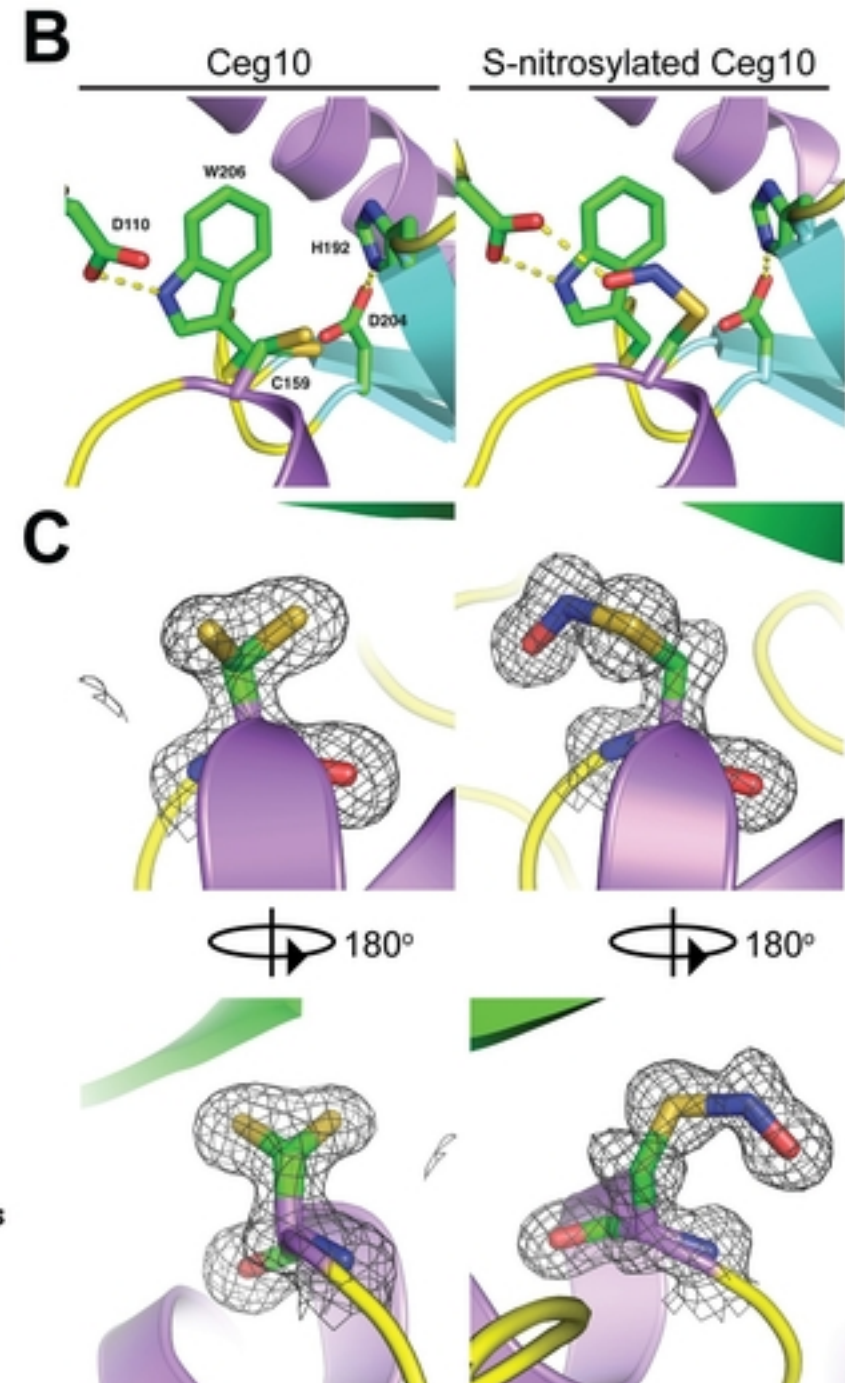
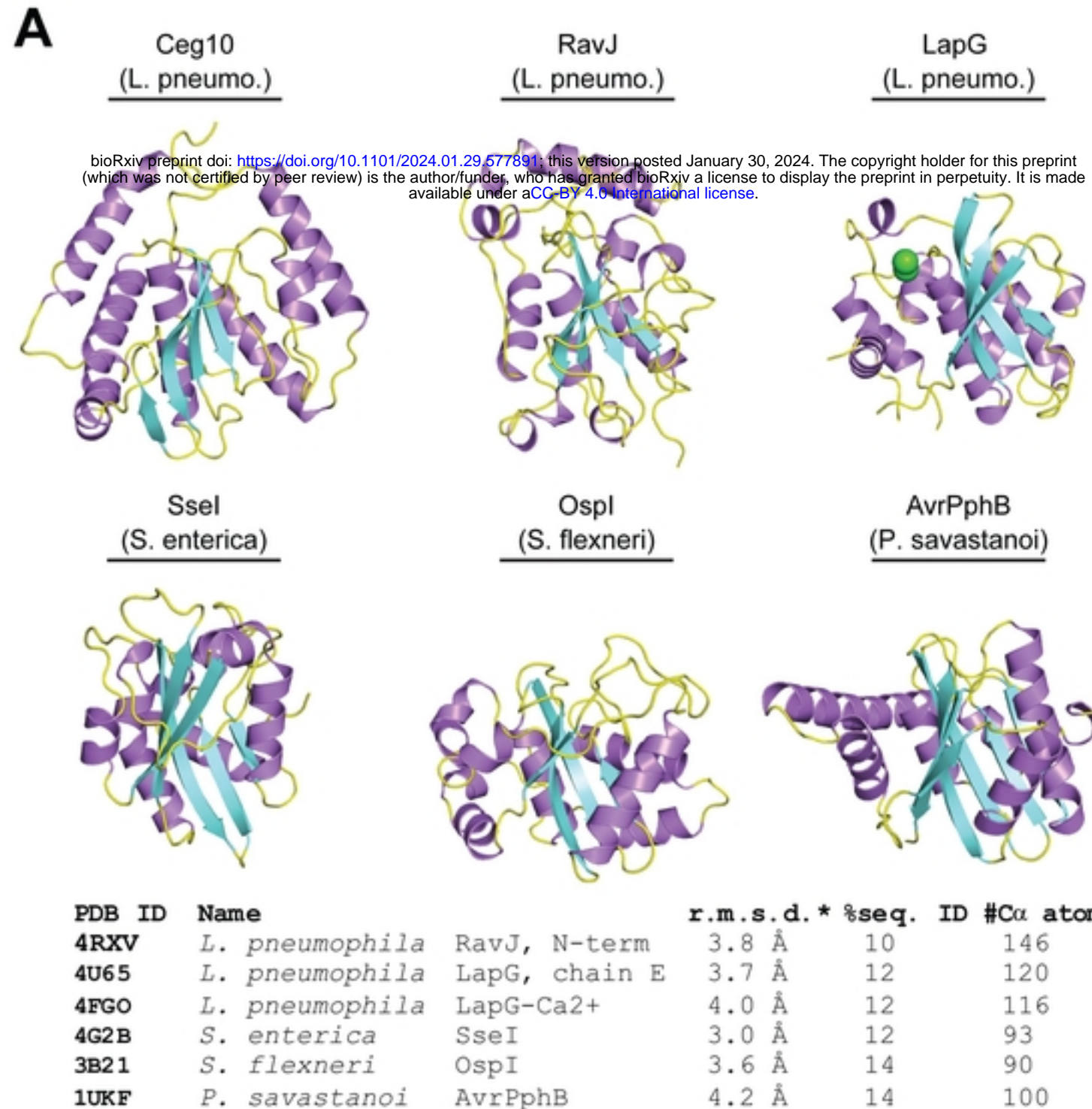


Figure 4