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Analysis of Coronavirus Temperature-Sensitive Mutants Reveals an Interplay between the Macrodomain and Papain-Like Protease Impacting Replication and Pathogenesis

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1	Analysis of coronavirus temperature-sensitive mutants reveals an interplay
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5 6	Xufang Deng ¹ , Robert C. Mettelman ¹ , Amornrat O'Brien ¹ , John A. Thompson ¹ , Timothy E. O'Brien ² , and Susan C. Baker ^{1,a}
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17 Running title: CoV macrodomain and PLP2 interplay impacts replication

19 Analysis of temperature-sensitive (ts) mutant viruses is a classic method allowing researchers to identify genetic loci involved in viral replication and pathogenesis. Here, 20 we report genetic analysis of a ts strain of mouse hepatitis virus (MHV), tsNC11, 21 22 focusing on the role of mutations in the macrodomain and the papain-like protease 2 (PLP2) domain of nonstructural protein 3, a component of the viral replication complex. 23 Using MHV reverse genetics, we generated a series of mutant viruses to define the 24 25 contribution of macrodomain- and PLP2-specific mutations to the ts phenotype. Viral 26 replication kinetics and efficiency of plating analysis performed at permissive and nonpermissive temperatures revealed that changes in the macrodomain alone were both 27 necessary and sufficient for the ts phenotype. Interestingly, mutations in the PLP2 28 domain were not responsible for the temperature sensitivity but did reduce the 29 frequency of reversion of macrodomain mutants. Co-immunoprecipitation studies are 30 31 consistent with an interaction between the macrodomain and PLP2. Expression studies 32 of the macrodomain-PLP2 portion of nsp3 indicate that the ts mutations enhance the 33 proteasome-mediated degradation of the protein. Furthermore, we found that during virus infection, the replicase proteins containing the MAC and PLP2 mutations were 34 more rapidly degraded at the non-permissive temperature, as compared to the wild-type 35 proteins. Importantly, we show that the macrodomain- and PLP2-mutant viruses trigger 36 37 production of type I interferon in vitro and are attenuated in mice, further highlighting the 38 importance of the macrodomain-PLP2 interplay in viral pathogenesis.

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41 Importance

42 Coronaviruses are emerging human and veterinary pathogens with pandemic potential. Despite the established and predicted threat these viruses pose to human health, there 43 are currently no approved countermeasures to control these infections in humans. Viral 44 macrodomains, enzymes that remove post-translational ADP-ribosylation of proteins, 45 and viral multifunctional papain-like proteases, enzymes that cleave polyproteins and 46 remove polyubiquitin chains via deubiquitinating (DUB) activity, are two important 47 48 virulence factors. Here, we reveal an unanticipated interplay between the macrodomain and the PLP2 domain that is important for replication and antagonizing the host innate 49 immune response. Targeting the interaction of these enzymes may provide new 50 therapeutic opportunities to treat CoV disease. 51

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53 Introduction

Coronaviruses (CoVs) are enveloped, positive-sense, single-stranded RNA 54 viruses that primarily infect the respiratory or gastrointestinal tract. CoVs can emerge 55 from an animal reservoir, such as bats, to infect a new species and cause epidemic or 56 pandemic disease with high mortality. Recent emergence events exemplified by Severe 57 Acute Respiratory Syndrome coronavirus (SARS-CoV) and Middle East Respiratory 58 59 Syndrome coronavirus (MERS-CoV) in humans (1), and Swine Acute Diarrhea Syndrome coronavirus (SADS-CoV) in domestic pigs (2), have demonstrated how 60 devastating these viruses can be within naïve populations. To date, there are no 61 62 approved antivirals or effective vaccines that protect humans from coronavirus

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system to study coronavirus replication and pathogenesis in mice. The replication of the 67 virus initiates with the engagement of the spike glycoprotein with a host cell receptor 68 and the release of the positive-sense RNA into the cytoplasm of the cell. The large (~32 69 70 Kb) viral genomic RNA is translated to produce two long polyproteins, pp1a and pp1ab, 71 which are processed by viral proteases, including the papain-like proteases (PLP1 and/or PLP2), and the 3C-like protease (3CLpro or Mpro), into 16 nonstructural proteins 72 (nsp1-16, Figure 1A). To generate the viral replication complex, the coronaviral nsps 73 sequester host endoplasmic reticulum (ER) to generate convoluted membranes and 74 double-membrane vesicles (DMVs), which are the sites of viral RNA synthesis (3, 4). 75 76 The viral replication complex generates a nested-set of dsRNA intermediates to 77 produce copious amounts of mRNAs, which are then translated to produce the 78 structural (spike, envelope, membrane and nucleocapsid) and virus-specific accessory proteins. The genomic RNA and structural proteins assemble in the ER-Golgi 79 intermediate compartment to generate infectious virus particles that are released from 80

to facilitate the development of effect vaccines and antivirals.

diseases. Therefore, identifying viral factors that contribute to pathogenesis and

characterizing novel targets for therapeutic interventions are two important approaches

The murine coronavirus, mouse hepatitis virus (MHV), is widely used as a model

the cell (5, 6). 81

CoV replication induces profound rearrangement of the host ER, and generates 82 viral dsRNA intermediates, processes that can be sensed by the host to activate the 83 84 innate immune response. As a result, CoVs have evolved multiple strategies to counteract and delay activation of these host immune responses and establish an 85

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environment amenable to virus replication. These strategies include: expressing 86 species-specific accessory proteins as modulators of innate immune responses 87 [reviewed in (7)]; encoding highly-conserved nonstructural proteins that serve as 88 interferon antagonists (8-10) and sequestering viral RNA in DMVs (3, 4) to prevent 89 detection by host pattern recognition receptors. A key component in the assembly of 90 the DMVs is nsp3 (11, 12). To date, eleven distinct nsp3 domains have been identified 91 92 using either bioinformatic approaches or enzymatic studies (Figure 1A). Here, we focus on two multifunctional components encoded within nsp3, the macrodomain and the 93 papain-like protease 2. 94

The region researchers now term the macrodomain was originally identified in 95 the 1990s as a highly-conserved domain of unknown function, termed the X domain, 96 contained within the replicase polyprotein of rubella virus, hepatitis E virus (HEV) and 97 coronaviruses (13–15). Structural and biochemical studies revealed that the X domain 98 99 exhibited structural similarity to the cellular histone MacroH2a and catalyzed measurable ADP-ribose-1"-phosphatase (ADRP) activity (16-18), although the 100 101 functional significance of this enzymatic activity was unclear. Using reverse genetics to 102 inactivate the catalytic site of the enzyme, researchers found that ADRP activity was not essential for CoV replication in cultured cells (19). However, further studies revealed 103 that an ADRP-catalytic mutant virus was attenuated in mice (20), and that ADRP activity 104 105 in SARS-CoV and human CoV-229E mediated resistance to antiviral interferon 106 responses (21). These findings were consistent with an essential role for enzymatic activity in vivo; although, the target for the ADRP activity was still unclear. A 107 breakthrough came in 2016 from a study revealing that the macrodomain of hepatitis E 108

virus acts as an ADP-ribose hydrolase (22). ADP-ribosylation is a known posttranslational modification that regulates cellular activities (23); therefore, viral enzymes
that reverse this process could interrupt host-cell signaling. For CoVs, nsp3
macrodomain activity was shown to promote MHV-induced encephalitis (24) and
increase virulence during SARS-CoV infection (25).

Another highly-conserved enzyme contained within nsp3 is the papain-like 114 protease 2 (PLP2). For MHV, PLP2 is responsible for processing the nsp3/4 junction 115 116 using a highly-conserved LXGG/X cleavage site (26). Studies using SARS-CoV revealed that the single papain-like protease encoded on nsp3 (termed PLpro) cleaves 117 all three sites at the amino-terminal end of the polyprotein (27). PLpro also functions as 118 a deubiquitinating enzyme (DUB), capable of removing polyubiquitin chains from 119 substrates (28, 29). Structural studies revealed that CoV PLpro/PLP2s are similar to 120 cellular DUBs (30). Enzymatic analysis revealed that CoV PLpro/PLP2s are 121 122 multifunctional with protease, deubiquitinating and deISGylating activity (30-33). The 123 viral DUB activity has been implicated as a modulator of the innate immune response to 124 viral infection (32, 34, 35), but the target(s) of the DUB activity have not yet been identified. Thus, both the PLP2 and macrodomains of nsp3 have been independently 125 identified as contributors to coronavirus virulence. 126

In this study, we characterized a temperature-sensitive MHV mutant virus containing mutations within both the macrodomain and PLP2 domain. We investigated the contribution of these mutations to the temperature-sensitive phenotype as well as the resulting effects on viral pathogenesis. The results presented here reveal a previously undescribed interplay between the macrodomain and PLP2 domain that

132 impacts replication, antagonizes the innate immune response, and contributes to viral pathogenesis. Modulating the macrodomain-PLP2 interaction may provide new 133 opportunities for therapeutic intervention. 134

Results 135

Identifying mutations associated with a temperature-sensitive phenotype. Murine 136 137 coronavirus strain tsNC11 was generated by chemical mutagenesis, plaque purified, and validated as a temperature-sensitive (ts) mutant defective in positive-sense RNA 138 synthesis at non-permissive temperatures (36). Complementation analysis indicated 139 that tsNC11 harbored mutation(s) in the ORF1a region of the replicase polyprotein, but 140 141 the specific mutations were unknown. To identify the nucleotide changes in tsNC11, we isolated the genomic RNA from viral supernatant, subjected it to deep sequencing then 142 aligned the reads to the genomic sequence of MHV-A59 (GenBank accession 143 #AY910861). In agreement with the complementation study by Schaad et al. (36), the 144 145 sequence analysis revealed 7 non-synonymous substitutions in the ORF1a of tsNC11. These substitutions resulted in 7 amino acid changes: two in nsp2 (I4V and T543I), four 146 147 in nsp3, and one in nsp10 (P23S). The four mutations within nsp3 are distributed between the macrodomain (K532E and G554D) and the PLP2 domain (D1026N and 148 D1071N) (Figure 1A). As noted above, previous studies documented the importance of 149 the macrodomain and PLP2 domain in virus replication and disease; therefore, we 150 focused our efforts on evaluating how these substitutions contributed to the ts 151 phenotype, the stability of the phenotype, and the pathogenesis of the virus in mice. 152

153 To evaluate the contributions of the macrodomain and PLP2 domain mutations to 154 the ts phenotype, three mutant viruses were generated using the MHV-A59 reverse

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155 genetics system (37). The first mutant virus, designated MACmut, contains the macrodomain mutations K532E and G554D. The second virus was engineered with the 156 D1026N and D1071N mutations within the PLP2 domain and is designated PLP2mut. 157 The third virus, MAC/PLP2mut, combines the mutations in the macro- and PLP2 158 domains into one virus. In addition, an isogenic wild-type MHV (icWT) was used as a 159 160 control. These viruses were recovered, plaque purified, and propagated in DBT cells at 161 a permissive temperature of 32°C. Deep-sequencing results confirmed the incorporation of the desired nucleotide changes in nsp3 and revealed no additional amino acid 162 changes within the ORF1 region. 163

First, we evaluated the one-step growth curves of all 5 viruses (tsNC11, icWT 164 and the 3 engineered mutants) at the permissive (32°C) and non-permissive (37°C and 165 40°C) temperatures. As expected, icWT replicates to high titer at all three temperatures, 166 whereas tsNC11 is impaired at both 37°C and 40°C, as reported by Schaad et al. (36) 167 168 (Fig. 1B). Analysis of the three engineered mutants revealed that the two substitutions 169 in the PLP2 domain were not sufficient to confer a temperature-sensitive phenotype, as the kinetics of replication mirrored those of the wild-type virus. In contrast, the MACmut 170 virus exhibited reduced virus replication at 40°C, but was only slightly impaired at 37°C. 171 The MAC/PLP2 mutant virus mirrored the kinetics of tsNC11 with impaired replication at 172 173 both 37°C and 40°C, implicating the mutations in both the macro- and PLP2 domains as contributors to the temperature-sensitive phenotype of tsNC11. 174

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175 We also evaluated the plaque size and efficiency of plating (EOP) of the viruses at permissive and non-permissive temperatures. As expected, all viruses replicated to 176 high titer and formed similarly-sized plaques at 32°C (Figure 2, upper panel). tsNC11 is 177

178 profoundly temperature-sensitive, with a low number of plaques detected at the 10⁻¹ 179 dilution plate incubated at the non-permissive temperature. The tsNC11 plaques that were detected at the non-permissive temperature exhibited a large-plaque phenotype 180 suggesting that these viruses may be revertants. We found that the PLP2mut virus 181 formed large plagues at 40°C, which is consistent with the results of the kinetic analysis 182 183 and indicates that the mutations in the PLP2 domain are not sufficient to cause the ts 184 phenotype. Analysis of the MACmut virus revealed a mixed population of small and large plaques at the non-permissive temperature, the majority of which displayed the 185 small-plaque phenotype. The MAC/PLP2 mutant virus mirrored the plaque size and 186 plating efficiency of tsNC11. We calculated the EOP values, which represent the ratio of 187 viral titers obtained at 40°C and 32°C (Figure 2B). Again, both the PLP2mut and icWT 188 viruses had similar titers at both temperatures, resulting in an EOP of ~1. In contrast, 189 the MACmut virus exhibited titers that were significantly lower at 40°C compared to 190 titers obtained at 32°C (EOP=10⁻²). These results indicate that the MACmut virus, but 191 not the PLP2mut or icWT viruses, has a defect in plaque formation at the non-192 193 permissive temperature. Taken together, these data demonstrate that the mutations in the macrodomain, but not those in the PLP2 domain, are the major determinants of the 194 195 ts phenotype of tsNC11. Additionally, these results are consistent with a critical role of the macrodomain in viral replication. Interestingly, we found that the MAC/PLP2 mutant 196 virus mirrored the plaque size and low reversion frequency of tsNC11 (EOP = 10^{-5}), 197 198 supporting a role for the PLP2 domain as a genetic enhancer of the ts phenotype. A 199 genetic enhancer, as defined by genetic studies of eukaryotic organisms, is a mutation 200 in one gene that intensifies the phenotype caused by a mutation in another gene (38).

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202 noticed that, in addition to the majority population having the small-plaque phenotype, a 203 subpopulation of large plaques were also present at 40°C. The large plaques consistently appeared even after several rounds of plague purification of the small-204 205 plaque isolates. Therefore, we asked if the small plaques were formed by temperature-206 sensitive viruses, while the large plagues were due to revertant viruses. To address this 207 question, we selectively isolated plaques with different sizes and propagated them at 32°C to obtain viral stocks for subsequent analysis (Figure 3). We found that the small-208 209 plaque isolates recapitulated the phenotype of the parental MACmut virus: small plaques and similar EOP values (Figure 3A). In contrast, the large-plaque isolates 210 exhibited a phenotype similar to icWT. Sequencing results of PCR amplicons, 211 representing the region spanning the macro- and PLP2 domains, revealed that small-212 213 plague isolates had no additional mutations in either the macrodomain or PLP2. In 214 contrast, the large-plague isolates had either a true reversion (D554-to-G), or harbored 215 putative suppressive mutations located within the macrodomain, or the adjacent, downstream sequence (Figure 3B). Among seven large-plaque revertants, all 216 maintained the K532E mutation, indicating it was not associated with the ts phenotype. 217 218 Three isolates had the D554-to-G reversion, suggesting that it may be sufficient for the 219 ts phenotype of the MACmut virus. We found that isolates 4-7 maintained the engineered mutations, but had also acquired additional, potentially suppressive, 220 221 mutations in the downstream region. Together, these results indicate that altering the coding sequence of either the macrodomain or the downstream region is likely sufficient 222 to revert or suppress the ts phenotype caused by the G554D mutation. We also 223

Evaluating revertants of the MACmut virus. While generating the MACmut virus, we

evaluated the MAC-PLP2 region of tsNC11 large-plaque revertant viruses and found that all three isolates had the D554-to-G reversion (Figure 3C), consistent with our findings with the MACmut revertants.

Mutations in PLP2 enhance the ts phenotype by reducing reversion frequency. 227 228 We determined that the macrodomain mutations are the major contributors to the ts phenotype; however, we noticed that the MACmut virus did not completely phenocopy 229 tsNC11. We found that the replication of the MACmut virus was defective at 40°C, but 230 not at 37°C. In addition, the MACmut virus exhibited a higher EOP value (10⁻²) 231 compared to that of tsNC11 (10⁻⁴) (Fig 2B), indicating a relatively high reversion 232 frequency. These data imply that mutations outside the macrodomain may enhance the 233 ts phenotype by stabilizing the replication defect, thereby preventing reversion to the 234 235 wild-type phenotype (38). Therefore, we asked if the addition of the PLP2 mutations observed in tsNC11 could enhance the ts phenotype of the MACmut virus and reduce 236 237 reversion. We found that the MAC/PLP2mut virus exhibits a severe replication defect at 238 both 37°C and 40°C (Figure 1B), and only replicated under permissive conditions, 239 similar to tsNC11. The MAC/PLP2mut and tsNC11 viruses exhibited similar EOP values (~10⁻⁴) (Figure 2B). Of note, the low titer of the MAC/PLP2mut virus at 40°C indicates a 240 low level of reversion to the wild-type phenotype, suggesting that the PLP2 mutations 241 stabilize the MACmut virus. Taken together, these data demonstrate that while the 242 243 PLP2 mutations are not sufficient to cause the ts phenotype, they act to enhance the ts 244 phenotype caused by the mutation in the macrodomain. Enhancing phenotypes have been described for other coronavirus interacting proteins (39, 40), which motivated us to 245

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246 determine if the enhancement phenotype we detected here is due to an interaction between the macro and PLP2 domains. 247

Evaluating macrodomain interaction with the PLP2 domain. The structures of 248 several domains of nsp3 have been solved individually [reviewed in (41)] or in 249 combination (42). However, owing to the size and complexity this protein, the complete 250 structure of nsp3 remains unsolved. The capacity of the PLP2 mutations to enhance the 251 ts phenotype in the presence of the macrodomain mutations raises the possibility of 252 253 domain-domain interaction between the macrodomain and PLP2. To test this 254 hypothesis, we generated plasmids that express either an epitope-tagged macrodomain (HA-MAC) or PLP2 domain (PLP2-V5) (depicted in Figure 4A). When these plasmids 255 were co-transfected into HEK-293T cells, the expression of both the macrodomain and 256 257 PLP2 were detectable by the cognate epitope antibodies (Figure 4B). We detected HA-MAC in lysates immunoprecipitated with anti-V5, and inversely, PLP2-V5 was detected 258 259 when HA-MAC was immunoprecipitated from the lysates. These results indicate that the 260 ectopically expressed macrodomain associates with PLP2 in cell lysates, consistent with either a direct or indirect interaction. 261

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Mutations in the macrodomain and PLP2 domain affect protein stability. Because 262 we found that mutation in the macrodomain (G554D) is the major ts determinant and the 263 264 PLP2 mutations enhance the ts phenotype, we reasoned that these mutations might alter protein folding, thereby rendering the protein unstable and susceptible to 265 proteasome-mediated degradation. To determine if the mutations in the macrodomain 266 267 and/or the PLP2 domain alter protein stability, plasmid DNA expressing wild-type or 268 mutant forms of MAC/PLP2 polypeptide (Figure 5A) were transfected into HEK-293T

269 cells. The cells were maintained at 37°C throughout the experiment. We added cycloheximide (CHX) at 16 hours post-transfection to block translation, and harvested 270 cell lysates at the indicated times. The level of expressed proteins was determined by 271 immunoblotting (Figure 5B and C). The MAC/PLP2 (WT) protein was maintained at 272 levels comparable to those prior to treatment, up to 5 hours post-treatment with CHX. In 273 274 contrast, we detected rapid reductions in the levels of all of the mutant forms of the 275 protein. Addition of the proteasome inhibitor MG132 blocked the degradation of the proteins (Figures 5B and 5C). These results indicate that mutations in both the MAC 276 and PLP2 domains affect the protein folding and stability, rendering the proteins more 277 susceptible to proteasome-mediated degradation. 278

279 To determine if these MAC/PLP2 mutations affect the stability of the replicase proteins during virus replication at the non-permissive temperature, we performed 280 temperature shift experiments as outlined in Figure 6. We infected cells with either WT 281 282 or MAC/PLP2mut virus and incubated at the permissive temperature for 9.5 h. At this 283 point, we added CHX to block translation and shifted the infected-cells to the non-284 permissive temperature. Cell lysates were collected every 30 minutes and evaluated using immunoblotting for the level of nonstructural intermediate nsp2-3 and product 285 nsp3. We found that WT nsp2-3 and nsp3 were relatively stable, with loss of detection 286 occurring at 3 hours after the temperature shift and addition of CHX (Fig 6C, lanes 2-8). 287 288 In contrast, the levels of nsp2-3 and nsp3 in the MAC/PLP2mut-infected cells 289 diminished more rapidly, with reduced levels at 1.5 hours after the temperature shift and addition of CHX (Fig 6C, lanes 9-15). These results support the finding that the MAC 290

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291 and PLP2 mutations destabilize the replicase protein at the non-permissive temperature. 292

ts mutant viruses induce interferon in macrophages and are attenuated in mice. 293 Previous studies have shown that the papain-like protease domains of MHV, SARS-294 295 CoV, and MERS-CoV antagonize the IFN response, likely through the deubiguitinating activity of these enzymes (32, 34, 35, 43-46). In addition, coronaviral macrodomains 296 have been shown to suppress IFN production both in vitro and in vivo (20, 21, 24, 25). 297 298 We asked if the mutations in the macrodomain and PLP2 modulate the type I IFN 299 response during infection of macrophages. As shown in Figure 7A, infection of mouse 300 bone marrow-derived macrophages (BMDMs) with mutant viruses at permissive temperature produced significantly more IFN- α during infection compared to the icWT 301 virus infection. At 12 hours post-infection, the MACmut virus induced 2-fold more IFN- α 302 than icWT virus. Furthermore, the level of N gene transcript, which reveals the 303 304 abundances of all viral mRNAs, was reduced in the MACmut-infected cells, compared 305 to the wild type virus. We found that the PLP2mut virus elicited dramatically more IFN- α 306 than WT virus, while the level of N gene expression was similar. The MAC/PLP2mut 307 virus exhibited the most robust IFN- α and the lowest level of N gene expression. These results show that mutations in the macrodomain and the PLP2 domain result in elevated 308 309 levels of type I IFN mRNA expression during infection of macrophages, further supporting the role of macrodomain and PLP2 in modulating host innate immunity. 310

Because the ts mutant viruses had reduced replication efficiency and elicited 311 type I IFN production during infection of macrophages, we were interested in evaluating 312 313 the pathogenicity of these viruses. To this end, C57BL/6 mice were intracranially 14

inoculated with 600 plaque-forming units (PFUs) of virus and monitored for weight loss and mortality. As shown in Figure 7B, all WT virus-infected mice lost weight rapidly and succumbed to infection by day 11 post-infection. In contrast, the mutant virus-infected mice exhibited transient or no weight loss during the infection period and all mice survived. These results demonstrate that the ts mutant viruses are attenuated *in vivo*, and those mutations adjacent to the catalytic sites of the macro- and PLP2 domains can modulate viral pathogenesis.

321

322 Discussion

Identifying viral factors that modulate the immune response to viral infection 323 provides new opportunities for developing novel antiviral interventions. Here, we 324 described an unanticipated interplay between two previously characterized virulence 325 factors, the macrodomain and the papain-like protease, of coronaviruses. The 326 enzymatic activities of these domains have been implicated in removing post-327 translational modifications: macrodomains remove mono- or poly-ADP-ribose from 328 proteins (18, 22, 23); deubiquitinating activity of viral papain-like proteases removes 329 330 mono- or poly-ubiquitin chains from signaling proteins (30, 32, 47). Our study stems 331 from characterizing a temperature-sensitive mutant virus that harbored mutations within both the macrodomain and the PLP2 domain of nsp3. We found that the mutation within 332 333 the macrodomain (G554D) was associated with the most significant temperaturesensitive phenotype, but that this alteration of the macrodomain reverted to the wild-334 type phenotype at high frequency. However, viruses containing mutations in both the 335 macrodomain and PLP2 domain reverted less frequently, consistent with the PLP2 336 15

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domain having an enhancing effect on the ts phenotype. Although these two enzymes reside within the same nsp3 polypeptide (Figure 1A), to our knowledge, this is the first suggestion of an interplay between these domains. By expressing the macrodomain and papain-like protease 2 domain on independent expression plasmids, we were able to evaluate and detect co-immunoprecipitation of the proteins, consistent with either a direct or indirect interaction. Furthermore, we report that the mutations identified in the macrodomain and PLP2 domain destabilize the proteins, as revealed by proteasomedependent degradation. Lastly, we demonstrate that these mutant viruses promote type I IFN production from macrophages in tissue culture and are attenuated in mice. This 345 work confirms and extends previous studies that independently identified the 346 macrodomain and the papain-like protease 2 domain as modulators of the innate 347 immune response and virulence factors [reviewed in (31, 48, 49), (35)]. 348

Macrodomains have been shown to play a role in the virulence of positive-sense 349 350 RNA viruses including hepatitis E virus (HEV), alphaviruses, and coronaviruses 351 [reviewed in (48, 49)]. Studies of the alphavirus Chikungunya virus (CHIKV) revealed 352 that the macrodomain at the N terminus of nsP3 hydrolyzes ADP-ribose groups from mono-ribosylated proteins and that this de-ribosylating activity is critical for CHIKV 353 replication in vertebrate and insect cells, and for virulence in mice (50). Interestingly, 354 viruses engineered to encode a mutation of the CHIKV macrodomain catalytic site 355 356 rapidly reverted to the wild-type sequence (51), similar to the high frequency reversion 357 we reported for the MHV MACmut virus (Figure 3). Studies of the role of the macrodomain during coronavirus replication indicate that catalytic activity is not required 358 for virus replication in interferon non-responsive cell lines (19, 20, 24). However, 359

catalytic activity is important for replication in primary cells and in mice, implicating the macrodomain in evading the innate immune response and promoting viral pathogenesis (20, 24, 25). Identifying the ribosylated substrates that are targeted by the viral enzymatic activity is an important future direction for this work.

364 Our study implicated an adjacent viral domain, the papain-like protein 2 domain, as an interacting partner with the macrodomain. Interestingly, the helicase domain 365 adjacent to the macrodomain of hepatitis E virus (HEV) was found to modulate 366 367 macrodomain activity. Biochemical assays revealed that the presence of the HEV 368 helicase domain in cis enhanced the binding of the macrodomain to ADP-ribose and stimulated the hydrolase activity (22). Furthermore, we previously found that the 369 370 mutations in the Ubl-2 domain could cause a ts phenotype and destabilize the PLP2 371 domain (52). Here, we found that the mutations in the macro- and PLP2 domains destabilized the replicase proteins, as shown by the more rapid degradation of the 372 373 proteins after temperature shift. We speculate that there may be a dynamic interaction 374 between adjacent domains within the nsp3 polyproteins.

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375 As a multidomain protein, nsp3 must hold a sophisticated architecture to function properly and precisely. To date, four essential functions have been documented for this 376 multidomain protein: 1) interaction of the Ubl-1 domain with the nucleocapsid (N) protein 377 is important for genomic RNA synthesis and encapsidation (39, 40); 2) proteolytic 378 processing of the N-terminal region of pp1a and pp1ab to release nsp1, nsp2, and nsp3 379 (26, 27); 3) hijacking the cellular reticular network in concert with other membrane-380 381 associated proteins (nsp4 and nsp6) to form virus-specific membrane structures for 382 RNA synthesis (11, 12); and 4) antagonizing the innate immune response through the 383 actions of the de-ADP-ribosylating activity of the macrodomain and the deubiguitinating activity of the PLP2 domain [reviewed in (31, 48)]. The removal of post-translational 384 modifications such as ADP-ribosylation and poly-ubiquitination could be directed either 385 at cellular proteins to redirect them for use during viral replication, or to subvert 386 signaling of innate immune responses. Ultimately, structural and biochemical studies 387 will be needed to fully investigate the multiple *cis* and *trans* interactions of nsp3 and to 388 389 determine if there is a dynamic interplay that modulates the stability, substrate specificity and/or affinity of the enzymes and substrates. 390

391 We found that the MAC/PLP2mut virus recapitulated the ts phenotype of tsNC11 (Figure 1B). However, it is possible that some or all of the other mutations we identified 392 by deep sequencing (I4V and T543I in nsp2 and P23S in nsp10) may contribute in a 393 subtle way to the phenotype of tsNC11. Nsp2 was shown to be dispensable for MHV 394 and SARS-CoV replication, but the deletion of the nsp2 coding sequence resulted in 395 396 decreased viral replication and RNA synthesis (53). For nsp10, previous studies 397 revealed that this protein plays critical roles in the 3C-like protease-mediated 398 polyprotein processing and viral RNA synthesis (54, 55). The results from these studies indicate that the mutations in nsp2 and nsp10 may also contribute to a ts phenotype. 399 While our study focuses on the contribution of the macrodomain and PLP2, further 400 studies are needed to fully evaluate the impact of other ORF1a mutations on the 401 402 replication and pathogenesis of coronaviruses.

In summary, we report what is, to our knowledge, the first indication of an interplay between the macrodomain and papain-like protease 2 domain of CoV nsp3. We found that this interplay impacts virus replication efficiency, innate immune

406 antagonism and virulence in mice. A detailed understanding of the relationship between 407 the macro- and PLP2 domains will require further structural and enzymatic studies. We anticipate that the genetic analysis, co-immunoprecipitation and in vivo pathogenesis 408 outcomes reported here will facilitate these future studies. 409 410

Materials and Methods 411

Virus and cells. Human embryonic kidney (HEK) 293T cells (CRL-11268, ATCC) were 412 cultured in Dulbecco's modified Eagle medium (DMEM) with 10% fetal bovine serum 413 (FBS) and 2% L-glutamine. Delayed brain tumor (DBT) cells were grown in minimal 414 essential media (MEM) (catalog no. 21800-0400; Gibco) supplemented with 10% 415 tryptose phosphate broth (TPB) media, 5% heat-inactivated FBS (Atlanta Biological), 416 2% penicillin/streptomycin (pen/strep; HyClone), and 2% L-glutamine. The BHK-MHVR 417 cell line was kindly provided by Mark Denison at Vanderbilt University Medical Center 418 and cultured in Dulbecco's modified Eagle medium (DMEM) (catalog no. 12100-046, 419 Gibco) supplemented with 10% heat-inactivated FBS and G418 (0.8 mg/mL; HyClone). 420 Differentiated BMDMs were maintained in bone marrow macrophage media containing 421 422 DMEM (catalog no. 10-017-CV, Corning) supplemented with 30% L929 cell 423 supernatant, 20% FBS, 1% L-glutamine, 1% sodium pyruvate, and 1% pen/strep. HeLa-MHVR cells (56) were grown in DMEM (catalog no. 12100-046, Gibco) supplemented 424 with 10% FBS, 1% L-glutamine, 0.5% HEPES, and 1% pen/strep. Temperature-425 sensitive MHV strain tsNC11 was propagated in DBT cells at 32°C. The infectious clone 426 MHV-A59 strain (GenBank accession no. AY910861) serves as wild-type (icWT) virus 427 for this study. 428

429 Deep sequencing and bioinformatic analysis. Viral RNA was extracted from the supernatant of tsNC11-infected DBT cells incubated at 32°C. Isolated RNA was sent to 430 GENEWIZ, Inc. for cDNA library preparation and Illumina Miseg high-throughput 431 sequencing. Raw reads were subject to pairing and trimming and aligned to the genome 432 sequence of the synthetic construct of MHV A59 strain (GenBank accession no. 433 434 AY910861) using Geneious software (Geneious R7, https://www.geneious.com). A 435 medium-low sensitivity and an iteration of up to 5 times were chosen. A total of 195,824 sequences with a mean coverage of 898.8 were aligned to the MHV Synthetic Construct 436 template. Polymorphisms were detected using the "find variations/SNPs" tool. 437 Parameters included a minimum coverage of 5 with a minimum variant frequency of 438 25% in order for a variation to be called. The maximum variant p-value was set at 10^{-6} 439

and the minimum strand bias p-value was set at 10⁻⁵ when exceeding 65% bias. We
focused our analysis on the first 12 kb of the replicase gene, since previous studies
reported the ts phenotype was associated with changes in this region (36, 57).

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443 Generation of mutant viruses. All infectious clones were generated using the reverse 444 genetics system previously established for MHV-A59 (37). Mutations identified by deep sequencing within the macrodomain and PLP2 domain were introduced into plasmids A 445 and B, respectively, then verified by sequencing of the plasmid DNA. DNA fragments 446 were ligated together and used for in vitro transcription of viral RNA. In vitro transcribed 447 genomic RNA and N gene RNA was electroporated into BHK-MHVR cells, which were 448 449 overlaid onto DBT cells in a T75 flask. These cells were incubated at the permissive temperature of 32°C to facilitate the replication of ts mutant viruses. Supernatants were 450 collected at the time when cytopathic effect was evident, usually between 36 and 48 h 451

452 post-electroporation. All infectious clone mutant viruses were plaque purified, propagated on DBT cells, and subjected to full genome sequencing to validate the 453 genotype. These infectious clones were designated MACmut, PLP2mut, and 454 MAC/PLP2, according the locations of introduced mutations as shown in Figure 1. 455

456 Temperature-sensitive assay and one-step growth kinetics. To determine the temperature sensitivity of mutant viruses, the efficiency of plating (EOP = titer 40°C/ titer 457 32° C) of virus was measured. DBT cells were seeded into two 6-well plates at 5.0 x 10^{5} 458 459 cells/well a day prior to infection. Each viral stock supernatant was serially diluted and 460 inoculated onto the DBT cells. After 1 h incubation at 37°C, inoculum was removed, and cells were subsequently overlaid with 0.8% 2x MEM/agar mixture. One plate was 461 incubated at 32°C for 60 h, and the second plate was incubated at 40°C for 48 h. 462 Agarose-covered cells were fixed using 4% formaldehyde for 1 h and stained using 463 0.1% crystal violet solution after removal of agarose. Plaques were counted and titers 464 465 were calculated.

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To evaluate the kinetics of virus replication, a one-step growth curve was generated at 466 467 each temperature. Briefly, DBT cells were infected with the designated virus at multiplicity of infection of 5 for 1 h at 37°C, then plates were incubated at the specified 468 temperatures. The supernatants were collected at indicated time points and titrated on 469 470 DBT cells incubated at 32°C for 60 h.

Isolation and characterizations of ts revertants. To isolate ts revertants, plaque 471 assays were performed at 40°C. Viruses from single plaques were isolated and 472 propagated in DBT cells at 32°C to obtain viral stocks. To determine the ts phenotype of 473 the isolates, the isolates were titrated at both 32°C and 40°C and the EOP values were 474 21

475 calculated as described above. To identify any mutations within the region of macro-476 and PLP2 domains of the revertants, viral genomic RNA was extracted using TriReagent (MRC, Inc.) according to the manufacturer's instruction and subsequently 477 subject to cDNA synthesis. A genomic region (3976-6101 nt) containing the macro- and 478 PLP2 domain was amplified by PCR using specific primers (Sense: 5'- CAA GAA AGG 479 TCT TTA GGG CTG CTT -3'; anti-sense: 5'- GAC ACC ATC AAC CTT CTC AAA TG -480 481 3'). The PCR products were sequenced and the sequencing results were compared to the tsNC11 sequence. 482

483 MAC and PLP2 expression plasmids. Nucleotide sequences encoding the macrodomain [467-622 amino acids (aa) of nsp3] were amplified from a codon-484 optimized MHV nsp3 gene (sequence available upon request) and cloned into pCAGGS 485 vector with an HA epitope tag, designated as HA-MAC. The pCAGGS-PLP2 plasmid 486 (PLP2-V5) was generated in a previous study (52). The coding sequence of the 487 488 macrodomain through PLP2 domain (467-1085 aa) was inserted into pcDNA3.1 and fused with a c-terminal V5 epitope tag (pMP-WT). Mutations were introduced into these 489 490 constructs using site-directed mutagenesis PCR or Gibson Assembly technique to pMP-GD (G554D), pMP-2DN (D1026N/D1071N) and pMP-GD/2DN 491 generate (G554D/D1026N/D1071N), which all contain a c-terminal V5 tag. 492

493 **Co-immunoprecipitation.** HA-MAC and PLP2-V5 plasmids were co-transfected into 494 HEK-293T cells in 35 mm dishes. Cells were harvested using 500 μ L lysis buffer (20 495 mM Tris pH 7.5, 150 mM NaCl, 1 mM EGTA, 1 mM EDTA, 1% Triton X-100, 2.5 mM 496 sodium pyrophosphate, 1 mM β -glycerophosphate, 1 mM sodium ortho-vanadate, 1 497 μ g/mL leupeptin, 1 mM PSMF) and 200 μ g of whole cell lysates were used for immunoprecipitation with 1 μ g primary mouse anti-V5 (R96025, Invitrogen) or -HA (MMS-101R-200, Biolegend) monoclonal antibody (Ab). Protein-Ab mixtures were rotated at 4°C overnight and then added 15 μ L magnetic protein G beads (LSKMAGA02, Millipore) for 1 h incubation. Beads were washed three times with washing buffer (lysis buffer composition except 450 mM NaCl) and eluted with 40 μ L 2x sample buffer (10% glycerol, 5% β -ME, 3% SDS, 7.5 mg/mL Trizma-base, bromophenol blue). Eluted products and 5% of cell lysates as input were subject to SDS-PAGE gel electrophoresis and immune-blotting with anti-V5 or anti-HA antibodies.

Evaluating protein stability after addition of cycloheximide. To determine the 506 507 steady-state level of protein, 0.5 µg of the specified plasmid DNA was transfected into 508 HEK-293T cells with transfection reagent TransIT-LT1 (MIR2300, Mirus) according to 509 the manufacture's recommendation. At 16 h post-transfection, cells were treated with 20 µg/mL of cycloheximide (CHX) (5087390001, Sigma Aldrich) or a combination of 20 510 511 µg/mL CHX and 10 µM MG132 (474790, Calbiochem), a proteasome inhibitor, and harvested at the indicated time points. Equal amounts of cell lysate were subjected to 512 immunoblotting with anti-V5 or anti- β -actin (A00702, Genscript) antibodies. The relative 513 514 intensity of MAC/PLP2 bands (relative to β -actin) were measured and calculated with AlphaView software (Protein Simple). To assess the rate of decay of the protein amount 515 516 over time for the four viral protein types, we fit the two-parameter simple exponential nonlinear regression function, $y = \theta_1 e^{-\theta_2 x}$, using the NLIN procedure in SAS 9.4 517 software package and verified using Minitab software version 18. In this regression 518 equation, θ_1 is the initial viral amount parameter at time zero and θ_2 is the slope or rate 519

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520 of decay parameter. The slope parameters (θ_2) were each compared with the WT slope using NLIN's provided two-sided t-tests and p-values < 0.05 were deemed significant. 521

To evaluate the steady level of replicase proteins nsp2-3 and nsp3, we 522 performed a temperature shift experiment. Briefly, HeLa-MHVR cells were infected with 523 either WT or MAC/PLP2mut virus (moi = 5) and incubated at the permissive 524 temperature for 9.5 h, when the cells were shifted to 40°C and treated with 20 ug/mL of 525 CHX. Whole cell lysates were prepared at 30 min intervals by the addition of lysis 526 527 buffer A (4% SDS, 3% DTT, 40 % glycerol and 0.065 M Tris, pH 6.8). The lysates were passed through a 25-gage needle to break up aggregates, incubated at 37°C for 30 min 528 529 and loaded onto a 6% SDS-PAGE, followed by transfer to a nylon membrane. The membrane was incubated with a 1:2,000 dilution of rabbit polyclonal anti-nsp2-3 530 antibody (anti-D3) (58), followed by horseradish peroxidase (HRP)-conjugated donkey 531 anti-rabbit IgG (H+L) (Southernbiotech), and developed with Western Lightening Plus-532 533 ECL reagents (PerkinElmer). The membrane was stripped and re-probed using a 1:2,000 dilution of mouse anti-calnexin antibody, followed by HRP-conjugated goat anti-534 mouse IgG (H+L), and developed as above. 535

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Reverse transcription quantitative PCR (RT-qPCR). The protocol of RT-qPCR was 536 described previously (9) with slight modification. Briefly, BMDMs were mock-infected or 537 infected with wild-type or mutant MHVs at a multiplicity of infection (MOI) of 1 and 538 incubated at a permissive temperature of 32 °C. At indicated time points, cells were 539 harvested for RNA extraction using an RNeasy Mini Kit (74104, Qiagen). An equal 540 amount of RNA was used for cDNA synthesis using Rt2 HT First Strand Kit (330401, 541 Qiagen). To determine IFN- α 11, β -actin, or MHV-A59 N gene mRNA production, qPCR 542

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was performed with specific primers for mouse IFN- α 11 (PPM03050B-200, Qiagen), mouse β-actin (PPM02945B-200, Qiagen) or MHV-A59 N gene (Sense; 5'- AGC AGA CTG CAA CTA CTC AAC CCA ACT C -3'; anti-sense: 5'- GCA ATA GGC ACT CCT TGT CCT TCT GCA -3') using RT2 SYBR Green qPCR Mastermix (330502, Qiagen) in the Bio-Rad CFX96 system. The thermocycler was set as follows: one step at 95 °C (10 min), 40 cycles of 95 °C (15 s), 60 °C (1 min) and plate read, one step at 95 °C (10 s), and a melt curve from 65 °C to 95 °C at increments of 0.5 °C/0.05 s. Samples were evaluated in triplicate and data are representative of three independent experiments. 550 The levels of mRNA were relative to β -actin mRNA and expressed as $2^{-\Delta CT}$ [ΔCT = 551 $C_{T(gene of interest)} - C_{T(\beta-actin)}].$ 552

Evaluating viral pathogenesis. The protocol for evaluating pathogenesis of MHV was 553 554 approved by the Lovola University Chicago IACUC and previously described (59). Briefly, six-week-old C57BL/6 female mice were purchased from the Jackson 555 556 Laboratory. Mice were intracranially inoculated with 600 PFU in 20 μL PBS and monitored daily for changes in body weight. Infected mice were euthanized when weight 557 loss was over 25% according to the protocol. Statistical analysis of survival rate was 558 559 evaluated using the log-rank test.

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Author contributions 567

568 X.D. and S.C.B. conceived the concept, planned the experiments, and wrote the manuscript with contributions from all authors. X.D., R.C.M., and A.O. performed 569 570 specific experiments and analyzed the data. J.A.T performed the bioinformatic analysis. T.E.O. conducted the statistical analysis. Current contact information for J.A.T. is 571 jtho46@lsuhsc.edu 572

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774 Figure legends

Figure 1. Evaluating the replication kinetics of coronavirus temperature-sensitive mutants at permissive and non-permissive temperatures. (A) Schematic diagram of the MHV genome and the domains of nsp3. Abbreviations: Ubl1, ubiquitin-like domain 1; Ac, acidic region; PLP1, papain-like protease 1; MAC, Macrodomain; DPUP, domain

779 proceeding Ubl2 and PLP2; Ubl2, ubiquitin-like domain 2; PLP2, papain-like protease 2; 780 NAB, nucleic acid-binding domain; G2M, coronavirus group 2 marker domain; TMDs, 781 transmembrane domains; Y, coronavirus highly-conserved domain. Representative structures of the macrodomain with ribose (229E: PDB: 3EWR) and PLP2 (MHV: PDB: 782 4YPT) are shown in cyan and green with catalytic pockets circled and the residues 783 784 involved in catalysis shown in magenta. The mutations described in this study are 785 shown in red. (B) Growth kinetics of MHV and mutants at three temperatures. DBT cells were inoculated with the indicated virus (MOI of 5) for 1 h at 37°C and then shifted to 786 787 the indicated temperatures. Culture supernatant was collected at the indicated hours post-infection and titrated in DBT cells at 32°C. The data are representative of two 788 independent experiments. Error bars indicate ±SD. 789

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Figure 2. Analysis of plaque size and efficiency of plating at the permissive and non-permissive temperatures. (A) Representative plaque assays at 32° C and 40° C for icWT, tsNC11 and engineered mutant viruses. The dilution of the viral stock is indicated and selected to visualize ~20-50 plaques per plate. (B) Efficiency of plating (EOP) = average titer at 40° C/ average titer at 32° C.

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Figure 3. Analysis of small- and large-plaque variants in the MACmut virus
 population. (A) MACmut isolates with distinct plaque sizes were evaluated for a ts
 phenotype. Sequence analysis of individual plaque-purified revertant isolates identified

800 mutations in the macrodomain and the adjacent downstream region in the large-plaque 801 variants of the MACmut (B) and tsNC11(C) viruses.

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803 Figure 4. Evaluating co-immunoprecipitation of the macrodomain and the PLP2 domain. (A) Schematic diagram of the individual constructs used to evaluate potential 804 805 interactions between the macrodomain and PLP2. (B) Western blotting to identify 806 expression and co-immunoprecipitation of HA-MAC and PLP2-V5. HEK-293T cells were transfected with the indicated plasmid DNAs, lysates were prepared at 18 hours post-807 transfection, subjected to immunoprecipitation with the indicated antibody and the 808 809 products analyzed by SDS-PAGE and immunoblotting. The data represent the results of 810 three independent experiments. Astersks indicate the cross detection of IgG chains by 811 secondary antibody.

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813 Figure 5. Mutations in macrodomain and PLP2 enhance degradation of the polypeptide. (A) Schematic diagram of constructs used to evaluate protein stability. (B 814 and C) Western blotting detecting wild-type or mutant forms of MAC-PLP2 polypeptide 815 in the presence of cycloheximide (CHX) or a combination of CHX and a proteasome 816 inhibitor MG132. HEK-293T cells were transfected with the indicated expression 817 818 plasmid of wild-type (WT) or mutant forms of MAC-PLP2. At 16 h post-transfection, cells were treated with 20 µg/mL of CHX or a combination of 20 µg/mL CHX and 10 µM 819 MG132 and harvested at the indicated time points. Equal amount of cell lysate were 820 subjected to immunoblotting with anti-V5 or anti-β-actin antibodies. The relative intensity 821

of MAC/PLP2 bands (relative to β-actin) were measured and calculated with AlphaView
software. The experiment was repeated two times and the representative immunoblots
(B) and the curves of relative intensity (C) are shown. The slope parameters of the
decay curves were evaluated using non-linear regression and two-sided t-tests
compared to WT. **, P<0.005; ****, P<0.0001.

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Figure 6. Mutations in the macrodomain and PLP2 alter the stability of replicase 828 protein nsp3. HeLa-MHVR cells were infected with either icMHV-WT or 829 830 MAC/PLP2mut virus (MOI of 5) and incubated at 32°C for 9.5 h, then 20 µg/mL of cvcloheximide (CHX) was added and cells were shifted to the non-permissive 831 temperature. Lysates were prepared every 30 min, and the proteins separated by SDS-832 PAGE, and nonstructural proteins nsp2-3 and nsp3 were visualized by immunoblotting. 833 A) Schematic diagram of MHV replicase polyprotein indicating the processing pathway 834 835 and the region identified by the anti-nsp2-3 antibody. B) Outline of the experiment. C) Western blot evaluating the level of nsp2-3 and nsp3 proteins detected after shift to the 836 non-permissive temperature. This is representative data of two independent 837 experiments. Arrowhead indicates detections of cellular protein in all lysates. Asterisk 838 indicates degradation products detected by anti-nsp2-3 antibody in the MAC/PLP2mut 839 virus-infected cells. 840

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Figure 7. Macrodomain mutant viruses induce type I interferon in primary macrophages and are attenuated in mice. (A) Mouse bone marrow-derived

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844 macrophages were infected with the indicated virus (MOI of 1) at 32°C. Total RNA was 845 extracted at the indicated time points and subjected to RT-qPCR. The mRNA levels of IFN- α (left) and N gene (right) are presented relative to β -actin. The results are 846 representative of three independent experiments and subjected to a two-tailed, unpaired 847 t-test. Error bars indicate ±SD. ***, P<0.001; ****, P<0.0001. n.s.: not significant. N.D.: 848 not detected. (B) Six-week-old mice were injected intracranially with either icWT or the 849 indicated ts mutant virus (600 PFU per mouse) and monitored for weight loss. Viral 850 pathogenicity was evaluated by body weight loss (left) and percent survival (right). The 851 number (n) of infected mice is indicated in parentheses. Error bars indicate ± SEM. 852 Differences in survival rates were calculated using a log-rank test. 853

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	Titer (P		
Virus Strain	32°C	40°C	EOP
icWT	2.0 x 10 ⁷	1.3 x 10 ⁷	0.7 x 10 ⁰
tsNC11	4.0 x 10 ⁶	7.0 x 10 ²	1.8 x 10 ⁻⁴
PLP2mut	5.0 x 10 ⁷	6.0 x 10 ⁷	1.2 x 10 ⁰
MACmut	1.3 x 10 ⁷	2.3 x 10 ⁵	1.8 x 10 ⁻²
MAC/PLP2mut	4.0 x 10 ⁶	2.6 x 10 ²	0.7 x 10 ⁻⁴

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Α		32°C	40°C				
MAC 40°C	Small						
	20			Representative	e Titer (F	PFU/mL)	
E.		Barris Contraction	e	isolate	32°C	40°C	EO
				Small plaque	1.3 x 10 ⁷	2.3 x 10 ⁵	1.8 x 1
Ne.		a hanne a	R	Large plaque	5.0 x 10 ⁷	2.0 x 10 ⁷	0.4 x 1
В	Large plaque	C-PLP2)	487 622 69 (MAC)	93 755 827 DPUP(Ubl2) PL	1085 P2		
	WT MACmut #2 #3 Rev #4 #5 #6 #7	⁵³⁰ AEK AG Q . E . E	VGEC	615 Macro DNDVSLTY	GVVTKN	IVILVSNNQ	DDFD
С	WT	530 AEKAG	VGEC ⁵⁹⁵ NKCI	₆₁₅ Macro DNDVSLTY		VILVSNNO	DDFD
_	tsNC11	E	.D		• • • • • • • •		••••
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