1 Inflammasome-mediated antagonism of type I interferon enhances *Rickettsia* pathogenesis

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10 Summary

11	Inflammasomes and interferons constitute two critical arms of innate immunity. Most facultative bacterial
12	pathogens that inhabit the host cell cytosol avoid activating inflammasomes and are often resistant to
13	killing by type I interferon (IFN-I). We report that the human pathogen Rickettsia parkeri, an obligate
14	intracellular pathogen that resides in the cytosol, is sensitive to IFN-I. The mechanism of IFN-I-
15	dependent restriction requires the transcription factor IRF5, which upregulates anti-rickettsial factors
16	including guanylate-binding proteins and iNOS. However, R. parkeri curtails cGAS-dependent IFN-I
17	production by causing caspase-11-dependent pyroptosis. In vivo, inflammasome activation antagonizes
18	IFN-I production, enhancing <i>R. parkeri</i> abundance in the spleen. Mice lacking either IFN-I or IFN-γ
19	signaling are resistant to infection, but mice lacking both rapidly succumb, revealing that both interferons
20	are required to control R. parkeri. This study illuminates how an obligate cytosolic pathogen exploits the
21	intrinsic trade-off between cell death and cytokine production to escape killing by innate immunity.
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23	Highlights
24	Rickettsia killed by GBPs activates caspase-11 and GSDMD, promoting pyroptosis
25	Rickettsia exploits pyroptosis to avoid cGAS-dependent type I interferon
26	• IRF5, GBPs, and iNOS contribute to controlling <i>R. parkeri</i> infection
27	• Ifnar ^{-/-} Ifngr ^{-/-} mice succumb to infection, uncovering a mouse model to study <i>R. parkeri</i>
28	
29	Keywords
30	Inflammasome; type I interferon; IFN-γ; <i>Rickettsia parkeri</i> ; IRF5; guanylate-binding proteins; GBP2;
31	GBP5; caspase-11; cGAS
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37 Introduction

38 The innate immune response to microbial pathogens depends on upregulation of antimicrobial 39 factors, secretion of cytokines, and activation of host cell death pathways (Jorgensen and Miao, 2015; 40 Meunier and Broz, 2016; Mitchell and Isberg, 2017; Randow et al., 2013). Innate immune responses 41 have been characterized during infection with facultative intracellular bacterial pathogens such as 42 Listeria monocytogenes and virulent Francisella species, as well as viruses, which inhabit the host cell 43 cytosol (Brubaker et al., 2015; McNab et al., 2015; Wallet et al., 2016). However, less well understood 44 are the innate immune responses to obligate intracellular bacterial pathogens. Spotted fever group 45 (SFG) *Rickettsia* spp. are tick-borne pathogens that cause spotted fever diseases worldwide (Walker 46 and Ismail, 2008). As obligate intracellular bacteria that replicate exclusively in the host cell cytosol, 47 SFG *Rickettsia* spp. must continually interface with host innate immune sensors. Therefore, they 48 presumably have evolved sophisticated mechanisms to avoid or even exploit innate immune responses.

49 Following infection and subsequent detection of pathogen associated molecular patterns 50 (PAMPS) by host cell sensors (Takeuchi and Akira, 2010), cytokines including type I interferon (IFN-I) 51 are upregulated. For example, the detection of cytosolic DNA by cyclic GMP-AMP synthase (cGAS) 52 results in activation of the downstream adaptor stimulator of IFN genes (STING) and transcription 53 factors including IFN regulatory factor 3 (IRF3) to upregulate IFN-I expression and secretion (Sun et al., 54 2013). Binding of IFN-I to the interferon- α/β receptor (IFNAR) then activates a signaling cascade that 55 upregulates the expression of hundreds of interferon-stimulated genes (ISGs), mobilizing the cytosol to 56 an antimicrobial state (MacMicking, 2012; Meunier and Broz, 2016).

Previous studies revealed that *L. monocytogenes* is resistant to the killing effects of IFN-I in macrophages (Reutterer et al., 2008; Woodward et al., 2010), and in fact, this pathogen actively secretes STING agonists that stimulate IFN-I production (Burdette et al., 2011; Woodward et al., 2010). IFN-I enhances *L. monocytogenes* as well as *Francisella novicida* pathogenicity *in vivo* (Auerbuch et al., 2004; Carrero, 2013; Henry et al., 2010; O'Connell et al., 2004; Storek et al., 2015), demonstrating that these bacterial pathogens benefit from IFN-I signaling. In contrast, IFN-I has a critical role in restricting viral replication, and therefore the effects of IFN-I signaling on pathogens that inhabit the

cytosol are often regarded as anti-viral but not anti-bacterial (Boxx and Cheng, 2016; McNab et al.,
2015; Stetson and Medzhitov, 2006).

66 Cytosolic PAMPs can also be recognized by inflammasomes, resulting in host cell death. For 67 example, sensing of PAMPs such as microbial DNA by nucleotide-binding domain and leucine-rich 68 repeat containing gene family (NLR) proteins causes their oligomerization and subsequent activation of 69 the protease caspase-1 (Lamkanfi and Dixit, 2014; Strowig et al., 2012). Similarly, direct binding of 70 bacterial lipopolysaccharide (LPS) to the non-canonical inflammasome caspase-11 causes its 71 oligomerization and activation (Aachoui et al., 2013; Hagar et al., 2013; Kayagaki et al., 2013; Shi et 72 al., 2014). Active caspases-1 and -11 cleave the pore-forming protein gasdermin D (GSDMD), 73 promoting pyroptosis, a rapid lytic host cell death (Kayagaki et al., 2015; Shi et al., 2015). Pyroptosis 74 curtails microbial replication and exposes microbes to the extracellular environment where they can be 75 targeted by immune factors such as phagocytes and antibodies. To avoid inflammasome activation, F. 76 novicida modifies its LPS, enabling it to avoid caspase-11 (Hagar et al., 2013; Wallet et al., 2016). 77 Additionally, L. monocytogenes and F. novicida minimize bacteriolysis to prevent the release of DNA 78 (Peng et al., 2011; Sauer et al., 2010), and L. monocytogenes downregulates expression of flagellin (an 79 activator of the inflammasome) during infection (Shen and Higgins, 2006). These virulence strategies 80 reveal that pathogens can benefit from circumventing inflammasome activation. Interestingly, 81 inflammasome activation antagonizes IFN-I production (Banerjee et al., 2018; Corrales et al., 2016; 82 Jabir et al., 2014; Liu et al., 2018), and IFN-I can antagonize inflammasome activation (Guarda et al., 83 2011; Inoue et al., 2012). This suggests that the immune system may flip a switch between cell-intrinsic 84 anti-viral and anti-bacterial responses.

⁸⁵ Despite these advances, the role for IFN-I, inflammasomes, and other innate immune responses ⁸⁶ during infection by obligate intracellular bacteria is not fully understood. IFN-I has modest anti-rickettsial ⁸⁷ effects in immortalized endothelial cells *in vitro* (Colonne et al., 2011; 2013; Turco and Winkler, 1990). ⁸⁸ In regards to the inflammasome, caspase-1 is required for the production of IL-1 β during *R. australis* ⁸⁹ infection (Smalley et al., 2016). Many studies have also established a critical role for interferon- γ (IFN-⁹⁰ γ) and nitric oxide as anti-rickettsial factors (Feng et al., 1994; Li et al., 1987; Turco and Winkler, 1993;

91 Turco et al., 1998; Walker et al., 1997). However, the role for pyroptosis, IFN-I, and their relationships
92 to IFN-γ during *Rickettsia* infection remain unknown. Since *Rickettsia* obligately reside within the host
93 cytosol, we hypothesize that they have evolved sophisticated and distinctive measures to manipulate
94 the inflammasome and IFN-I response to their benefit.

95 Here, we investigated the roles for host cell death, IFN-I signaling, and antimicrobial factors in 96 controlling R. parkeri infection at the molecular, cellular, tissue, and organismal levels. We observed 97 that, unlike facultative cytosolic bacterial pathogens, R. parkeri intracellular growth is restricted by IFN-98 I in macrophages in vitro. If R. parkeri is killed in the host cytosol by guanylate-binding proteins (GBPs), 99 the inflammasome is activated, thus curtailing IFN-I production. Consistently, we discovered that R. 100 parkeri antagonized IFN-I production in vivo via inflammasome activation, which has tissue-specific 101 effects on controlling bacterial replication. We also found that both IFN-I and IFN- γ are critical for 102 controlling *R. parkeri* infection in mice, and propose that mice lacking interferon signaling can serve as 103 a robust animal model for investigating *Rickettsia* pathogenesis. Our data suggest an unprecedented 104 strategy whereby a bacterial pathogen exploits the inherent antagonism between the inflammasome 105 and IFN-I response to promote its pathogenesis.

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- 107 Results

Inflammasome activation promotes *R. parkeri* growth in macrophages by antagonizing the IFN I response

110 Although previous studies have investigated the roles of IFN-I and inflammasomes during 111 infection by facultative intracellular pathogens, little is known for obligate intracellular pathogens 112 including *R. parkeri*. Previous reports suggested that IFN-I causes a ~50% reduction in *Rickettsia* spp. 113 growth in immortalized endothelial cells (Colonne et al., 2011; 2013; Turco and Winkler, 1990). In vivo, 114 both endothelial cells (Feng et al., 1993; Sahni and Rydkina, 2009) and other cell types including 115 macrophages (Banajee et al., 2015; Feng et al., 1993; Osterloh et al., 2016; Papp et al., 2016; Walker 116 et al., 1999) are targeted by *Rickettsia* spp. during infection. However, the effect of IFN-I on *Rickettsia* 117 spp. replication in primary macrophages was unknown. We infected wild type (WT) primary bone

118 marrow-derived macrophages (BMDMs) with R. parkeri or L. monocytogenes, treated them with 119 recombinant IFN- β , and measured the impact on bacterial growth (by counting plaque-forming units 120 (PFU) for *R. parkeri* or colony-forming units (CFU) for *L. monocytogenes*). IFN-β caused a dose-121 dependent restriction of *R. parkeri* growth (Figure 1A), whereas it had no effect on *L. monocytogenes* 122 (Figure 1B). We further assessed whether each of these pathogens elicited an IFN-I response during 123 infection by testing supernatants from infected BMDMs for their ability to stimulate expression of an 124 IFN-responsive luciferase reporter. R. parkeri infection did not induce appreciable IFN-I expression, 125 whereas L. monocytogenes elicited a robust response (Figure 1C). These data suggest that, although 126 R. parkeri is sensitive to IFN-I, it elicits low amounts of IFN-I and therefore grows robustly in 127 macrophages.

128 We next investigated how R. parkeri escapes inducing IFN-I by examining the interaction 129 between R. parkeri and the inflammasome, leveraging the ability to generate BMDMs from mice 130 genetically deficient in inflammasome components. First, we focused on caspase-1, apoptosis-131 associated speck-like protein containing CARD (ASC), and the non-canonical inflammasome 132 component caspase-11. WT and mutant BMDMs were infected with *R. parkeri* and bacterial-induced 133 cell death, a readout of inflammasome-dependent pyroptosis, was monitored at 24 h post infection (hpi) using a lactate dehydrogenase (LDH) release assay. Infected Casp1^{-/-} and Asc^{-/-} mutant BMDMs 134 135 exhibited ~40% death, similar to WT cells (Figure 1D). In contrast, cell death was significantly reduced in Casp11^{-/-} mutant BMDMs, and little cell death was observed in Casp1^{-/-}Casp11^{-/-} double mutant 136 137 BMDMs (Figure 1D). Because caspases-1 and -11 activate pyroptosis by cleaving the pore-forming protein GSDMD, we next measured host cell death in Gsdmd^{-/-} BMDMs. Compared to Casp1^{-/-} or 138 Casp11^{-/-} single mutants, Gsdmd^{-/-} BMDM cell death was reduced, but it was not abolished as in the 139 Casp1^{-/-}Casp11^{-/-} double mutant cells (Figure 1D). These data reveal that the majority of *R. parkeri*-140 141 induced host cell death in macrophages depends on caspase-11 and GSDMD. To a lesser extent, R. 142 parkeri activates caspase-1-dependent and GSDMD-independent cell death.

As infections proceeded to 96 h, we observed that WT BMDMs became rounded and nonadherent, whereas many $Casp1^{-/-}Casp11^{-/-}$ double mutant cells remained adherent, similar to uninfected

145 cells (**Figure 1E**). When we measured the number of infectious bacteria in these cells by counting PFU, 146 we observed a reduction in the number of bacteria in $Casp1^{-/-}Casp11^{-/-}$ BMDMs, whereas the number 147 of bacteria increased over time in WT BMDMs (**Figure 1F**). There was also a reduction of infectious 148 bacteria in $Casp11^{-/-}$ and $Gsdmd^{-/-}$ single mutant BMDMs, albeit less pronounced than for the $Casp1^{-/-}$ 149 $Casp11^{-/-}$ double mutants (**Figure 1F**). Thus, a subpopulation of *R. parkeri* appears to exploit activation 150 of caspase-11 and GSDMD to promote replication of the remaining population.

151 Cytokines including IFN-I and TNF- α are produced by macrophages and can exert antimicrobial activity (Carrero. 2013: Weiss and Schaible, 2015). In addition, IFN-I is overproduced in cells lacking 152 153 the inflammasome (Corrales et al., 2016; Liu et al., 2018). Hence, we hypothesized that R. parkeriinfected Casp1^{-/-}Casp11^{-/-} BMDMs were secreting increased levels of IFN-I that restricted *R. parkeri* 154 growth. We therefore tested the bactericidal effects of adding supernatants from infected Casp1^{-/-} 155 Casp11^{-/-} cells to WT BMDMs that were infected with *R. parkeri*. We found that this supernatant 156 157 restricted bacterial growth in a dose-dependent manner (Figure 1G). Furthermore, IFN-I secretion was increased at least 15-fold in infected Casp1^{-/-}Casp11^{-/-} double mutant and Casp11^{-/-} and Gsdmd^{-/-} single 158 mutant cells (**Figure 1H**). To test whether IFN-I secreted from infected Casp1^{-/-}Casp11^{-/-} BMDMs was 159 necessary for bacterial killing, we transferred the supernatant from infected Casp1^{-/-}Casp11^{-/-} to infected 160 Ifnar^{-/-} BMDMs, which lack the IFN-I receptor and hence are not responsive to IFN-I. The supernatant 161 had no effect on bacteria growth in *Ifnar^{-/-}* cells (**Figure 1G**). Supernatants from infected Casp1^{-/-}Casp11^{-/-} 162 ^{-/-} cells were also treated with an anti-IFNAR antibody to prevent IFN-I signaling, and this similarly 163 164 restored bacterial growth (Figure 1I). To further test whether IFN-I was the inhibitory factor, we bred mice lacking *Ifnar*, *Casp1*, and *Casp11* (*Casp1^{-/-}Casp11^{-/-}Ifnar^{-/-}*), infected BMDMs from these mice, and 165 166 observed that growth of *R. parkeri* was restored (Figure 1I). A similar series of experiments failed to 167 show a role for TNF- α in killing *R. parkeri* (Figure S1). Together, these findings demonstrate that activation of the inflammasome allows R. parkeri to avoid killing by IFN-I. 168

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170 Pyroptosis masks cGAS-induced IFN-I

171 It remained unclear whether activation of host immune sensors was due to release of R. parkeri 172 ligands such as LPS or DNA. We therefore sought to identify the host sensor required for the increased IFN-I production observed in infected Casp1^{-/-}Casp11^{-/-} cells. We bred Cas^{-/-}, Sting^{gt/gt}, and Tlr4^{-/-} to 173 Casp1^{-/-}Casp11^{-/-} mice to generate triple mutants (Casp1^{-/-}Casp11^{-/-}Casp11^{-/-}Casp11^{-/-}Sting^{gt/gt}, 174 and Casp1^{-/-}Casp11^{-/-}Tlr4^{-/-}) and then infected BMDMs from these mice. As expected, due to their 175 176 inability to induce pyroptosis, all three triple mutants exhibited reduced host cell death, similar to levels seen for *Casp1^{-/-}Casp11^{-/-}* double mutants (**Figure 2A**). We next measured IFN-I secretion and found 177 that infected Casp1^{-/-}Casp11^{-/-}Tlr4^{-/-} cells maintained increased levels of IFN-I, similar to Casp1^{-/-} 178 Casp11^{-/-} cells (Figure 2B). In contrast, Casp1^{-/-}Casp11^{-/-}Casp1^{-/-}Casp11^{-/-}Casp11^{-/-}Sting^{Gt/Gt} 179 180 macrophages exhibited reduced IFN-I secretion, below levels of infected WT cells (Figure 2B). To 181 determine whether the amount of IFN-I production correlated with an effect on bacterial growth, we also measured PFU over time. Mutation of Cgas or Sting in the Casp1^{-/-}Casp11^{-/-} background rescued growth 182 183 of *R. parkeri* (Figure 2C) (for unknown reasons at 96 hpi, bacterial burdens were consistently lower in Casp1^{-/-}Casp11^{-/-}Cgas^{-/-} and Casp1^{-/-}Casp11^{-/-}Sting^{gt/gt} cells than in WT cells). In contrast, mutation of 184 185 TIr4 in the Casp1^{-/-}Casp11^{-/-} background did not rescue bacterial growth. Together, these data in 186 macrophages demonstrate that caspase-11 activation masks cGAS-dependent IFN-I production, and 187 are consistent with the notion that pyroptosis prevents IFN-I production (Corrales et al., 2016; Liu et al., 188 2018). This observation suggests that, in cells where bacteria release DNA to activate cGAS, they also 189 release LPS to activate caspase-11.

190

191 IRF5-regulates antimicrobial genes and is required for IFN-I-dependent killing of *R. parkeri*

The host factors that restrict *R. parkeri* cytosolic growth downstream of IFN-I are poorly understood, in part because IFN-I upregulates hundreds of ISGs, complicating the ability to identify specific antimicrobial factors. To narrow the list of potential ISGs that restrict *R. parkeri* growth, we first tested whether a specific IRF was required for killing *R. parkeri* upon IFN-I treatment. We analyzed bacterial growth over time following bacterial infection of WT, *Irf1^{-/-}*, *Irf3^{-/-}Irf7^{-/-}*, or *Irf5^{-/-}* mutant BMDMs and treatment with IFN-I. We observed that mutation of *Irf5* significantly rescued bacterial growth

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198 compared to WT cells, whereas mutation of *Irf1* or both *Irf3* and *Irf7* caused a modest increase in growth

199 (Figure 3A). This suggested that genes specifically regulated by IRF5 restrict *R. parkeri*.

200 To identify genes significantly and specifically regulated by IRF5, we performed high-throughput RNA-sequencing (RNAseq) on the different WT, *Irf1^{-/-}*, *Irf3^{-/-}Irf7^{-/-}*, or *Irf5^{-/-}* mutant BMDMs upon infection 201 202 and IFN-I treatment. To limit the identification of genes not directly responsible for bacterial killing, we 203 isolated RNA at 12 hpi, the earliest time when bacterial killing was observed (Figure 3B). Analysis of 204 the RNAseg data revealed that 136 genes were upregulated >4.0-fold in infected IFN-I-treated cells 205 when compared to infected untreated cells (**Table S1**). Among these, 82 were not as highly upregulated (>1.5-fold difference) in infected IFN-I-treated $lrf5^{-/-}$ cells when compared to infected IFN-I-treated WT 206 cells, suggesting that many were regulated by IRF5. We then compared the expression of these genes 207 between infected IFN-I-treated Irf5^{-/-} and Irf1^{-/-} and Irf3^{-/-}Irf7^{-/-} cells, and 36 had higher expression in Irf1^{-/-} 208 $^{\prime-}$ and Irf3 $^{\prime-}$ Irf7 $^{\prime-}$ cells than in Irf5 $^{\prime-}$ cells. From the analysis of the RNA-seq data, we conclude that 36 209 210 genes are significantly and specifically upregulated by IRF5 during R. parkeri infection and IFN-I 211 treatment (Table S1).

Many of the 36 genes encode known anti-viral and anti-bacterial proteins, including *Gbp2*, *Gbp5*, *Rsad2* (encoding Viperin), *Ifit1*, *Ifit2*, *Mx1*, and *Mx2* (**Figure 3C**). The RNA-seq also identified *Nos2*, encoding inducible nitric oxide synthase (iNOS), which is required for controlling *Rickettsia* spp. infection (Feng and Walker, 2000; Turco and Winkler, 1993; Turco et al., 1998; Walker et al., 1997), although its expression was also dependent on IRF1 (**Figure 3C**, right side). Together, these data demonstrate that IRF5 is critical for controlling *R. parkeri* upon IFN-I signaling, perhaps due to upregulating ISGs such as the GBPs and iNOS.

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220 GBPs and nitric oxide contribute to restricting *R. parkeri* growth in macrophages

We next assessed whether the IRF5-regulated genes that have previously known antimicrobial activity are important for inhibiting *R. parkeri* growth in the presence or absence of IFN-I. We derived BMDMs from the femurs of mice lacking *Ifit1, Ifit2, Rsad2, LipG* (the gene most highly upregulated by IFN-I), and the chromosome 3 cluster of *Gbps* (*Gbp^{chr3}*) (Yamamoto et al., 2012), which includes *Gbp2*,

225 Gbp5 and three other Gbp genes. Bacterial growth was then measured upon IFN-I treatment. In IFN-Itreated *Gbp^{chr3-/-}* macrophages, bacterial growth was increased 3-to-5-fold compared with that in WT 226 227 IFN-I-treated macrophages (Figure 4A). No change in bacterial growth was observed in the other 228 mutant macrophages upon IFN-I treatment (Figure S2B). When pyroptosis was assessed by measuring LDH-release, *Gbp^{chr3}*-deficient macrophages showed dramatically reduced cell death upon infection, 229 whereas the other mutant cells did not (Figure 4B). Gbp^{chr3-/-} macrophages did not exhibit increased 230 231 IFN-I production (Figure 4C), consistent with the notion that the GBPs are required for release of 232 bacterial LPS as well as DNA. These data suggested that GBPs contribute to both IFN-I-dependent and 233 IFN-I-independent killing of *R. parkeri*. Next, to assess the role for nitric oxide during *R. parkeri* infection. 234 we treated WT- and *Gbp^{chr3}*-infected cells with IFN-I and the specific iNOS inhibitor L-NIL and monitored bacterial growth. Upon iNOS inhibition, bacterial growth was increased in both WT and *Gbp^{chr3-/-}* cells 235 236 (Figure 4D, 4E). We also measured bacterial growth in IFN-I-treated Nos2^{-/-} BMDMs, but did not 237 observe any rescue in mutant versus WT cells (Figure S4A), but this result is complicated by the fact 238 that iNOS is a negative regulator of the NLRP3 inflammasome (Hernandez-Cuellar et al., 2012; Mishra et al., 2013) and we observed more host cell death in Nos2^{-/-} BMDMs compared to WT cells (Figure 239 **S4B**). All together, these experiments reveal that the GBPs and perhaps nitric oxide are IRF5-regulated 240 241 genes that non-redundantly restrict R. parkeri growth in macrophages upon IFN-I treatment.

242 GBP-mediated killing of other pathogens has also been examined downstream of IFN-y 243 treatment (Pilla-Moffett et al., 2016; Yamamoto et al., 2012). To assess the role of GBPs in IFN-ydependent restriction of *R. parkeri*, bacterial growth was measured in WT and *Gbp*^{chr3-/-} cells in the 244 245 presence of different concentrations of IFN- γ . At lower concentrations of IFN- γ (0.2 ng/ml), we observed a small but significant rescue in bacterial growth in *Gbp^{chr3-/-}* versus WT cells (Figure 4F). However, 246 higher concentrations of IFN- γ (4 ng/ml) ablated *R. parkeri* growth in both WT and *Gbp*^{chr3-/-} BMDMs 247 248 (**Figure 4F**). This revealed that IFN- γ potently stimulates macrophages to kill *R. parkeri* by a mechanism 249 that partially depends on GBPs.

250 The GBPs have been observed to localize to the surface of intracellular pathogens at steady-251 state conditions and upon the addition of interferons (Liu et al., 2018; Mitchell and Isberg, 2017; 252 Yamamoto et al., 2012). To determine if the GBPs localized to the surface of *R. parkeri*, we performed 253 immunofluorescence microscopy using a GBP2-specific antibody. GBP2 localized to the surface of ~1% 254 of bacteria in untreated WT cells at 3 hpi, and ~5% of total bacteria in IFN-I treated cells (Figure 4G, **H**). No colocalization was observed in infected *Gbp^{chr3-/-}* macrophages, demonstrating that the antibody 255 256 is specific for the GBPs (Figure S3). Together, these data demonstrate that the GBPs localize to R. 257 parkeri and restrict bacterial growth at steady-state conditions, which is enhanced by IFN-I.

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259 In spleens, the inflammasome antagonizes the anti-rickettsial effects of IFN-I

260 Our data from macrophages suggested a model for *R. parkeri* intracellular survival whereby 261 inflammasome activation antagonizes the anti-rickettsial activity of IFN-I. To determine whether this 262 pathway is important *in vivo*, we analyzed the role for the inflammasome, IFN-I, and IFN-y in spleens of 263 infected mice, where Rickettsia spp. proliferate upon systemic infection (Feng et al., 1994). In 264 agreement with the data from macrophages *in vitro*, IFN-β mRNA was significantly upregulated in spleens of infected Casp1^{-/-}Casp11^{-/-} mice when compared to WT mice, as measured by gPCR (Figure 265 266 **5A**). To determine if the increased IFN- β transcript abundance was dependent on cGAS, we also measured IFN-β mRNA in infected Casp1^{-/-}Casp11^{-/-}Cgas^{-/-} mice, and observed a reduction in IFN-β 267 268 transcript abundance (Figure 5A). This demonstrated that the inflammasome antagonizes cGAS-269 induced IFN-I in vivo, as it does in vitro.

To determine if the increased IFN-I observed in $Casp1^{-/-}Casp11^{-/-}$ mice restricted *R. parkeri*, we compared bacterial burdens between $Casp1^{-/-}Casp11^{-/-}$ and $Casp1^{-/-}Casp11^{-/-}Ifnar^{-/-}$ mice. The triple $Casp1^{-/-}Casp11^{-/-}Ifnar^{-/-}$ mutant showed a significantly increased bacterial burden in comparison to the double $Casp1^{-/-}Casp11^{-/-}$ mutant (**Figure 5B**). To ascertain if the increased bacterial growth was due to cGAS-dependent IFN-I, we also measured bacterial burdens in $Casp1^{-/-}Casp11^{-/-}Cgas^{-/-}$ mice, and similarly observed increased bacterial burdens compared with the $Casp1^{-/-}Casp11^{-/-}$ mutant (**Figure 5B**). It remains unclear why spleens of $Casp1^{-/-}Casp11^{-/-}$ mice, which produce more IFN- β , do not have

reduced bacterial burdens when compared to WT (as in macrophages); nevertheless, these data reveal
that the mechanisms of *R. parkeri* growth in spleens is similar to macrophages *in vitro*, where bacterial
activation of the inflammasome limits the antimicrobial effects of cGAS-induced IFN-I.

280 In macrophages, we also observed a role for IFN- γ in restricting *R. parkeri* growth; however, the 281 role for IFN- γ and its relationship to the inflammasome during *Rickettsia* infection *in vivo* is unknown. 282 We therefore next assessed if IFN- γ restricted *R. parkeri* growth in the spleens of infected mice. Neutralization of IFN- γ using an anti-IFN- γ antibody increased bacterial burdens in Casp1^{-/-}Casp11^{-/-} 283 mice 2.5-fold when compared to untreated $Casp1^{-/-}Casp11^{-/-}$ mice, suggesting that IFN- γ restricts 284 285 bacterial growth (**Figure 5B**, right side of dotted line). To reveal if IFN-γ and IFN-I both control infection in the spleen, we neutralized IFN- γ in infected Casp1^{-/-}Casp11^{-/-}Ifnar^{-/-} mice. This increased bacterial 286 burdens 51-fold over untreated Casp1^{-/-}Casp11^{-/-} mice (**Figure 5B**, right side of dotted line). These data 287 288 demonstrate that in spleens of WT mice, inflammasome activation limits protective IFN-I production, 289 and that IFN-I and IFN- γ both contribute to protection.

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In livers, the inflammasome antagonizes IFN-I, which in turn antagonizes the anti-bacterial effects of IFN-γ

293 We also measured bacterial burdens in the livers, where Rickettsia and other bacterial 294 pathogens also reside during infection (Feng et al., 1994; Rayamajhi et al., 2010). In contrast to what was observed in spleens, infected Casp1^{-/-}Casp11^{-/-} mice had 15-fold higher bacterial burdens than WT 295 296 mice in the liver (Figure 5C). Also in contrast to what was observed in spleens, the bacterial burden in infected Casp1^{-/-}Casp11^{-/-}Ifnar^{-/-} mice, as well as Casp1^{-/-}Casp11^{-/-}Cgas^{-/-}mice, was indistinguishable 297 298 from WT (Figure 5C), suggesting that the 15-fold increase observed in Casp1^{-/-}Casp11^{-/-} mice was 299 dependent on IFN-I. We hypothesized that this effect might be due to IFN-I antagonizing the anti-300 bacterial effects of IFN- γ , as it was previously demonstrated that IFN-I antagonizes the effects of IFN- γ 301 in the liver during L. monocytogenes infection (Rayamajhi et al., 2010). To test whether IFN-I antagonized the anti-rickettsial effects of IFN- γ , infected Casp1^{-/-}Casp11^{-/-} and Casp1^{-/-}Casp11^{-/-}Ifnar^{-/-} 302

mice were treated with the IFN- γ neutralizing antibody and bacterial burdens were measured. The difference in bacterial burdens between these mice was erased upon IFN- γ neutralization (**Figure 5C**, right), demonstrating that IFN-I was responsible for antagonizing IFN- γ . Together, these results suggest that the inflammasome protects against *R. parkeri* infection in the liver, in part by limiting IFN-I antagonism of IFN- γ (**Figure 5D**).

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Both IFN-I and IFN-γ are required to control *R. parkeri* infection in mice

310 Because we observed that bacterial burdens in organs increased when both IFN-I and IFN- γ 311 were removed, we next tested whether these cytokines were protective at the whole animal level. We 312 infected C57BL/6 mice deficient for both IFN-I and IFN- γ receptors (*Ifnar*^{-/-}*Ifngr*^{-/-}) via the intravenous route. Strikingly, whereas WT, *Ifnar^{-/-}*, and *Ifngr^{-/-}* mice survived an infectious dose of 10⁷ bacteria with 313 no severe symptoms, the majority of *Ifngr^{-/-}Ifnar^{-/-}* mice succumbed within 6 d (**Figure 6A**), and lost body 314 315 weight and had reduced body temperature over time (Figure 6B,C). Symptoms of infection were dosedependent. At a dose of 10⁶ bacteria, mice lost weight, their temperature decreased, and many 316 317 succumbed to infection, although some eventually recovered to their original body weight and temperature. At dose of 10⁵ bacteria, none of the mice succumbed to infection, but the animals lost 318 319 weight and their body temperature decreased, which then recovered to pre-infection levels. We also 320 performed these experiments using AG129 mice, which are similarly mutated for Ifngr and Ifnar, but are 321 in a different genetic background. These mice were also highly susceptible to infection with 10^7 bacteria, 322 with 100% fatality by 6 dpi (Figure 6D). Together, these data demonstrate that both IFN-I and IFN- γ 323 potently control R. parkeri growth in animals. The discovery of the sensitivity of this mouse genotype 324 reveals it as robust animal model for further investigations into pathogenesis of the SFG Rickettsia.

325

326 **Discussion**

327 Avoiding innate immunity is critical for microbial pathogens to survive and cause disease. 328 Whereas some facultative pathogens that inhabit the cytosol resist killing by IFN-I and avoid the

inflammasome, the innate immune response to obligate intracellular bacteria has remained largely unexplored. We report the unexpected discovery that the innate immune response to the obligate intracellular human pathogen *R. parkeri* is distinctive among other cytosolic pathogens, as the bacteria are sensitive to IFN-I-mediated killing, but avoid stimulating a robust IFN-I response by exploiting the inherent trade-off between inflammasome activation and IFN-I production.

We observed that *R. parkeri* were restricted by IFN-I in macrophages and in mice. Furthermore, IFN-I much more potently restricted *R. parkeri* in macrophages than was previously seen for other *Rickettsia* spp. in immortalized endothelial cells and fibroblasts (Colonne et al., 2011; Turco and Winkler, 1990). In contrast, facultative cytosolic pathogens such as *L. monocytogenes* are resistant to the killing effects of IFN-I in primary myeloid cells (Bauler et al., 2011; Woodward et al., 2010) and in mice (Auerbuch et al., 2004; Rayamajhi et al., 2010). Given their sensitivity to IFN-I, we propose that *Rickettsia* must have evolved mechanisms to avoid stimulating IFN-I production.

341 Our results suggest that one critical mechanism for avoiding IFN-I production is that R. parkeri-342 induces pyroptosis, which dampens IFN-I production, protecting the remaining bacterial population that 343 has successfully infected other cells (Figure 7, left). In support of this conclusion, we observed that a 344 subpopulation of *R. parkeri* activated caspase-11 and GSDMD to cause pyroptosis. Because pyroptosis 345 is a rapid post-translational signaling event (Kayagaki et al., 2015), activation of this cell death pathway 346 prohibits IFN-I production, which is slower as it requires transcription, translation, and secretion. We 347 found that macrophages that are deficient for inflammasome signaling restricted R. parkeri because 348 bacterial ligands instead activated robust IFN-I production via cGAS, leading to bacterial killing (Figure 349 7, right). We also observed that the GBPs are required for pyroptosis and for IFN-I production in the 350 absence of host cell death. From these observations, we conclude that bacteriolysis is likely responsible 351 for the release of LPS and DNA that stimulate caspase-11 and cGAS. In contrast with what we observed 352 for *R. parkeri*, several facultative bacterial pathogens have evolved different mechanisms to avoid the 353 inflammasome (Jorgensen and Miao, 2015), including modification LPS by F. novicida to avoid 354 caspase-11 (Hagar et al., 2013; Wallet et al., 2016), downregulation of flagellin by L. monocytogenes 355 to avoid NAIP5 (Miao et al., 2010; Shen and Higgins, 2006), and infrequent lysis by F. tularensis and L.

356 *monocytogenes* to avoid AIM2 (Sauer et al., 2010; Ulland et al., 2010). We propose that inflammasome 357 activation is a trade-off that allows for growth of a pathogen that is sensitive to IFN-I.

358 Downstream of IFN-I production, we observed that the transcription factor IRF5 is critical for 359 restriction of *R. parkeri* in macrophages in vitro. IRF5 in turn upregulates the expression of anti-microbial 360 factors including the GBPs and iNOS, which protect against R. parkeri. These data agree with a 361 previously established role for nitric oxide in protecting against infection by other *Rickettsia* spp. in 362 mouse and human cells (Feng and Walker, 1993; 2000; Feng et al., 1994; Turco and Winkler, 1993; 363 Woods et al., 2005), although it remains unclear if nitric oxide directly or indirectly kills these pathogens. 364 These data also reveal an unappreciated role for the GPBs in rickettsial killing. We observed that iNOS 365 and GBPs acted additively and non-redundantly, and only partially account for the killing effects of IFN-366 I. We suspect that additional IRF5-regulated genes contribute to controlling *R. parkeri* infection, and 367 identifying these factors will be critical for understanding how interferons control infection by obligate 368 pathogens.

369 Consistent with our observations in macrophages in vitro, we found that the inflammasome 370 antagonizes IFN-I production in vivo. However, we observed tissue-specific differences in the spleen 371 and liver with regard to the relative protective role for IFN-I. In the spleen, IFN-I protects against 372 infection. These data are in agreement with previously known roles for the inflammasome in the spleen. 373 where the major protective pathway elicited by the inflammasome is pyroptosis (Maltez et al., 2015). 374 Our observations expand on these previous findings, by revealing that an inherent trade-off of 375 pyroptosis is that it limits protective IFN-I in the spleen. However, in the liver, IFN-I enhances infection 376 due to its effects of antagonizing IFN- γ . These observations are also in agreement with a previous study 377 demonstrating that cytokine production is the more important facet of inflammasome activation in the 378 liver (Maltez et al., 2015), and expand on these studies by demonstrating that the inflammasome also 379 acts by antagonizing IFN-I. Our observations are also consistent with a study demonstrating that IFN-I 380 antagonizes IFN- γ in the liver, as observed during *L. monocytogenes* infection (Rayamajhi et al., 2010). 381 Moreover, our observations in both organs are consistent with earlier studies establishing a protective 382 role for IFN-γ during *Rickettsia* infection *in vivo* (Feng et al., 1994; Li et al., 1987). Based on our findings

383 that IFN-I and IFN- γ act differently in the spleen and liver, we speculate that interferons may pressure 384 pathogens to evolve organ-specific tropisms to accommodate their intracellular lifestyle.

385 In keeping with our observations that IFN-I and IFN- γ restrict R. parkeri growth in macrophages 386 *in vitro*, we observed that *Ifnar^{-/-} Ifngr^{-/-}* mice rapidly succumbed to infection. These findings demonstrate 387 that both types of interferon play a critical role in protecting against R. parkeri infection. Our observations 388 are also of strong practical importance, as this mouse exhibits a pathology that makes it a useful animal 389 model for future studies examining the bacterial determinants that contribute to SFG Rickettsia 390 pathogenesis and for evaluating the impacts of infection at the organ, tissue and cellular levels in vivo. 391 Future studies will aim to characterize facets of infection including dissemination, vascular damage, as 392 well as the innate immune response.

393 Our data suggest that obligate cytosolic pathogens such as R. parkeri may occupy an 394 intermediate position in the spectrum between bacteria and viruses, both with regard to their 395 dependency on the host and their interplay with the innate immune system (Figure 7B). For example, 396 during infection with R. parkeri as well as with herpes simplex virus 1 or Vaccinia virus, activation of the 397 inflammasome antagonizes the protective IFN-I response (Wang et al., 2017). Additionally, IRF5 is 398 required for controlling *R. parkeri* and viral infection (Carlin et al., 2017; Proenca-Modena et al., 2016; 399 Thackray et al., 2014). Lastly, as with *R. parkeri*, many viruses have increased lethality in *Ifnar^{-/-}Ifngr^{-/-}* 400 mice (Milligan et al., 2017; Rossi et al., 2016), suggesting a non-redundant protective role for IFN-I and 401 IFN-y in both types of infection. These similarities are not true for facultative cytosolic bacterial 402 pathogens such as L. monocytogenes, for which IFN-γ is significantly more protective than IFN-I 403 (Rayamajhi et al., 2010). Our observations regarding the interaction between R. parkeri and innate 404 immune pathways underscores the notion that obligate pathogens have evolved different strategies 405 than facultative pathogens to fit their ecological niche. Their ability to tolerate inflammasome activation 406 to avoid IFN-I is an evolutionary strategy that may be shared among obligate microbes, whose complete 407 reliance on host cell metabolic and cellular pathways for survival may also dispose them to IFN-I 408 sensitivity. Continued investigation of the interaction between obligate intracellular bacteria and innate

immunity will enhance our understanding of how pathogens exploit host cell immunity to survive and tocause disease.

411

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424

425 **Author contributions**

Conceptualization: T.P.B; Methodology: T.P.B., P.E.; Investigation: T.P.B., P.E., and R.C.;
Visualization: T.P.B., P.E., R.E.V., and M.D.W.; Writing: Original Draft, T.P.B.; Writing – Review &
Editing, T.P.B., P.E., R.E.V., and M.D.W.; Funding Acquisition, R.E.V. and M.D.W.; Resources, R.E.V.
and M.W.D.; Supervision, R.E.V. and M.D.W.

430

431 **Declaration of interests**

432 The authors declare no conflicts of interest.

- 433
- 434 Figure Legends

18

Figure 1: Inflammasome activation promotes *R. parkeri* pathogenesis by antagonizing the IFN-I

436 response in macrophages

437 A) Measurement of Rickettsia parkeri (Rp) PFU in BMDMs over time. WT BMDMs were infected in a 438 24-well plate at an MOI of 0.2. MOI was calculated based on the ratio of the number of infectious 439 bacteria, as determined by plaque assay, to the number of BMDMs. At each time point, BMDMs were 440 lysed with water and lysates were plated over a confluent monolayer of Vero cells. Recombinant IFN-B 441 (PBL, 12405-1) was added at 0 hpi. U signifies units of recombinant IFN-β. Statistical comparisons were 442 made between each sample and untreated cells at each time point. Data are a compilation of at least 443 two separate experiments and are expressed as means + SEM. B) Measurement of Listeria 444 monocytogenes (Lm) CFU in BMDMs. WT BMDMs were infected in 24-well plate at an MOI of 1. MOI 445 was calculated based on the ratio of bacteria in liquid culture to BMDMs. 3,000 U of recombinant IFN-446 β (PBL, 12405-1) was added at 0 hpi. Data are the combination of at least two separate experiments 447 and are expressed as means + SEM. C) Measurement of IFN-I in supernatants of cells infected with R. 448 parkeri or L. monocytogenes. WT BMDMs were infected with R. parkeri or L. monocytogenes at an MOI 449 of 1 and supernatants were harvested at 8 hpi for L. monocytogenes and 24 hpi for R. parkeri. 450 Supernatants were used to stimulate a luciferase-expressing cell line and relative light units (RLU) were 451 measured and compared between each sample and uninfected cells. Statistical comparisons were 452 made between each sample and infected WT cells. Data are the compilation of at least two separate 453 experiments and are expressed as means + SEM. D) Quantification of host cell death during R. parkeri 454 infection of BMDMs. LDH release was measured at 24 hpi upon R. parkeri infection at an MOI of 1. 455 Statistical comparisons were made between each sample and WT B6 cells. Data are the compilation of 456 at least three separate experiments and are expressed as means + SEM. E) Images of BMDMs infected 457 with *R. parkeri*. Cells were infected at an MOI of 1 and images were captured at 96 hpi. The scale bar 458 is 100 µm. F) Measurement of the number of infectious *R. parkeri* in BMDMs over time. BMDMs were 459 infected with *R. parkeri* at an MOI of 1 and PFU were determined every 24 hpi. Statistical comparisons 460 were made between each sample and WT B6 cells at each time point. Data are the compilation of at 461 least three separate experiments and are expressed as means + SEM. G) Measurement of the number

462 of infectious R. parkeri in BMDMs over time. Cells were infected with R. parkeri at an MOI of 0.2 and 463 PFU were measured over time. Statistical comparisons were made between each sample and WT B6 464 cells at each time point. Data are the compilation of at least three separate experiments and are expressed as means + SEM. "Supe" indicates supernatant collected at 24 hpi from Casp1^{-/-}Casp11^{-/-} 465 466 BMDMs infected with R. parkeri at an MOI of 1. H) Measurement of IFN-I in supernatants of cells 467 infected with R. parkeri. Supernatants from infected cells were harvested at 24 hpi and used to stimulate 468 a luciferase-expressing cell line and relative light units (RLU) were compared between each sample 469 and uninfected cells. Statistical comparisons were made between each sample and infected WT cells. 470 Data are the compilation of at least three separate experiments and are expressed as means + SEM. 471 I) Measurement of the number of infectious *R. parkeri* in BMDMs over time. Cells were infected at an 472 MOI of 1 and PFU were measured over time. The α -IFNAR antibody was added at T=0 to a final 473 concentration of 1 µg/ml. Statistical comparisons were made between each sample and WT B6 cells at 474 each time point. Statistical analyses for data in panels A, B, C, F, G, and I were performed using a two-475 tailed Student's T-test, where each sample at each time point was compared to the control. Statistical 476 analyses in panels D and H were performed using an ANOVA with multiple comparisons post-hoc test. 477 *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001, ns=not significant.

478

479 Figure 2: Pyroptosis masks cGAS-induced IFN-I

480 A) Quantification of host cell death during *R. parkeri* infection of BMDMs. LDH release was measured 481 at 24 hpi upon *R. parkeri* infection of the indicated BMDMs at an MOI of 1. Statistical comparisons were 482 made between each sample and WT B6 cells. Data are the compilation of at least three separate 483 experiments and are expressed as means + SEM. B) Measurement of IFN-I abundance in supernatants 484 of infected BMDMs. The indicated BMDMs were infected with *R. parkeri* at an MOI of 1. Supernatants 485 from infected cells were harvested at 24 hpi and used to stimulate a luciferase-expressing cell line and 486 compared to uninfected cells. Statistical comparisons were made between each sample and infected 487 WT cells. Data are a compilation of at least three separate experiments and are expressed as means 488 + SEM. C) Measurement of bacterial PFU in BMDMs over time. BMDMs were infected with R. parkeri

489 at an MOI of 1 and PFU were measured over time. Statistical comparisons were made between each 490 sample and WT B6 cells at each time point. Data are the compilation of at least three separate 491 experiments and are expressed as means <u>+</u> SEM. Statistical analyses for bacterial abundance in panel 492 C were performed using a two-tailed Student's T-test, where each sample at each time point was 493 compared to the control. Statistical analyses for LDH assays and IFN-I abundance in panels A and B 494 were performed using an ANOVA with multiple comparisons post-hoc test. *p<0.05, **p<0.01, 495 ***p<0.001, ****p<0.0001, ns=not significant.

496

497 Figure 3: IRF5 is required for IFN-I-dependent restriction of *R. parkeri*

498 A) Measurement of *R. parkeri* PFU in BMDMs over time. Cells were infected at an MOI of 0.2, and 499 PFU were measured over time. 10,000 units of recombinant IFN- β (rIFN- β) was added at 0 hpi. 500 Statistical comparisons were made between each sample and untreated cells at each time point. Data 501 are the compilation at least three separate experiments and are expressed as means + SEM. B) 502 Measurement of bacterial PFU in BMDMs. Cells were infected at an MOI of 1, and PFU were measured 503 over time. Statistical comparisons were made between each sample and WT B6 cells at each time 504 point. Data are the compilation at least three separate experiments and are expressed as means + SEM. "Supe" indicates 200 µl of supernatant collected at 24 hpi from Casp1^{-/-}Casp11^{-/-} BMDMs infected 505 506 with R. parkeri at an MOI of 1. All statistical analyses were performed using a two-tailed Student's T-507 test. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001, ns=not significant. **C**) Transcript abundance of 508 antimicrobial genes upregulated by IFN-I. BMDMs were infected at an MOI of 2.3, treated with 10,000 509 U recombinant IFN- β , and RNA was harvested at 12 hpi. High-throughput RNA-sequencing was then 510 performed. The regulation of 7 candidate anti-rickettsial genes is shown. Genes are ordered from 511 highest to lowest in terms of upregulation by IFN- β in WT cells, with the exception of Nos2, which is 512 shown separately because it was more highly regulated by IRF1 than IRF5. Each data set was 513 normalized to *R. parkeri*-infected WT, IFN-β-untreated BMDMs.

21

515 Figure 4: The GBPs contribute to IFN-I-dependent and -independent inhibition of *R. parkeri* 516 growth

517 A) Measurement of R. parkeri PFU in BMDMs. BMDMs were infected with R. parkeri at an MOI of 0.2 518 and PFU were measured over time. "Supe" signifies 500 µl supernatant from infected Casp1^{-/-}Casp11⁻ 519 ¹ cells, which was added at 0 hpi. Statistical comparisons were made between each sample and 520 untreated cells at each time point. Data represent at least three separate experiments and are 521 expressed as means + SEM. B) Quantification of host cell death. LDH release was measured at 24 hpi 522 upon R. parkeri infection at an MOI of 1. Statistical comparisons were made between each sample and 523 WT B6 cells. Data are the compilation at least three separate experiments and are expressed as means 524 + SEM. C) Measurement of IFN-I abundance in supernatants of infected BMDMs. The indicated BMDMs 525 were infected with *R. parkeri* at an MOI of 1. Supernatants from infected cells were harvested at 24 hpi 526 and used to stimulate a luciferase-expressing cell line and compared to uninfected cells. Data are a 527 compilation of at least five separate experiments. **D,E,F**) Measurement of bacterial PFU in BMDMs. 528 Cells were infected at an MOI of 0.2, and PFU were measured over time. "Supe" signifies 500 µl 529 supernatant from infected Casp1^{-/-}Casp11^{-/-} cells, which was added at 0 hpi. L-NIL was resuspended in 530 water and added to a final concentration of 1 mM at T=0. IFN-γ was added at 0 hpi. Statistical 531 comparisons were made between each sample and untreated cells at each time point. Data are the 532 compilation of at least three separate experiments and are expressed as means + SEM. G) A 533 representative image of GBP2 localization to the surface of *R. parkeri*. WT BMDMs were infected at an 534 MOI of 1, and were imaged at 3 hpi using 100x confocal immunofluorescence microscopy. Cyan staining 535 is phalloidin; green staining is α -GBP2 (ProteinTech); red staining is α -*Rickettsia*. The red and green 536 channels of the indicated white box are increased in size and shown on the right and bottom left. The 537 scale bar is 5.6 µm. H) Quantification of GBP2 localization to the surface of *R. parkeri*. WT BMDMs 538 were infected at an MOI of 1, and were imaged at 3 hpi using 100x confocal immunofluorescence 539 microscopy. For IFN- β treated cells, BMDMs were treated overnight with 100 U recombinant IFN- β . 540 Each data point represents an individual experiment, and each experiment consists of at least 10 541 separate images, and each image contained approximately 20 bacteria. Statistical analyses for panels

A, C, D, E, F, and H were performed using a two-tailed Student's T-test, where each sample at each time point was compared to the control. Statistical analyses for LDH assays in panel B were performed using an ANOVA with multiple comparisons post-hoc test. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001, ns=not significant.

546

547 Figure 5: In mice, inflammasome activation antagonizes IFN-I production, leading to tissue-548 specific effects on *R. parkeri* burdens

A) Measurement of IFN- β transcripts in infected mice. C57BL/6 mice were infected i.v. with 10⁷ R. 549 parkeri, and qPCR was used to analyze IFN-β and actin transcript abundance at 72 hpi in spleens. IFN-550 551 β was normalized to actin to determine relative copy number per mouse spleen. Data are the 552 combination of at least 3 separate experiments. B) Measurement of bacterial burdens in mouse spleens. 553 Mice were infected i.v. with 10⁷ *R. parkeri*, and bacterial burdens were determined in spleens at 72 hpi 554 via plague assay. Colors were arbitrarily assigned to distinguish between genotypes. Data are the 555 combination of at least 3 separate experiments. Statistical analyses were performed using Mann-556 Whitney U test. Bars denote medians. C) Measurement of bacterial burdens in mouse livers. Mice were 557 infected i.v. with 10⁷ R. parkeri and bacterial burdens were determined in livers at 72 hpi via plaque 558 assay. For α -IFN- γ (BioLegend), mice were injected i.v. with 300 µl at 30 minutes postinfection (mpi), 559 200 µl at 24 hpi, and 200 µl at 48 hpi, totaling 800 µl (0.8 µg antibody). Colors are used to distinguish 560 between genotypes. Data are the combination of at least 3 separate experiments, with the exception of 561 experiments using the α -IFN- γ antibody, which are a combination of 2 separate experiments. Statistical 562 analysis performed for gPCR experiments in panel A used a two-tailed Student's T Test. Statistical 563 analyses for *in vivo* experiments in panels B and C were performed using a Mann-Whitney U test. Bars 564 denote medians. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001, ns=not significant. **D**) Textual summary 565 of results from in vivo infections.

566

567 Figure 6: Mice mutated for both *Ifnar* and *Ifngr* succumb to *R. parkeri* infection

568 A) Survival of mice lacking interferon signaling. Mice in the C57BL/6 background were infected i.v. with 569 the indicated concentrations of *R. parkeri*, and survival was measured over time. Mice were euthanized 570 if their core body temperature fell below 90° F or if they exhibited symptoms of severe infection, such 571 as severe hunching, scruffing, or inability to move around the cage. Infections from at least 6 mice are 572 shown for each genotype. Data are the combination of at least three separate experiments. B) 573 Measurement of mouse weight during infection. Mice were infected with 10⁷ R. parkeri and weight was 574 measured every 24 hpi. Data are normalized to the weight starting at the initial time of infection. C) 575 Measurement of mouse core body temperature during infection. Mice were infected with 10⁷ R. parkeri 576 and body temperature was measured every 24 hpi, using a rectal thermometer. D) Survival of mice 577 lacking interferon signaling. Mice of the 129 genotype lacking IFN-I and IFN- γ receptors (AG129) were 578 infected i.v. with the indicated concentrations of *R. parkeri* and mouse health was measured over time. 579 Mice were euthanized if their core body temperature fell below 90° F or if they exhibited symptoms of 580 severe infection. At least 6 mice were used for each condition. Data are the combination of at least 3 581 separate experiments.

582

583 Figure 7: Inflammasome-mediated antagonism of IFN-I promotes intracellular growth of *R.* 584 *parkeri*

A) Model depicting intracellular growth of *R. parkeri* in WT cells (left) or in cells lacking inflammasome
 signaling (right). B) Visual representation of the antimicrobial effects of interferons on cytosolic
 pathogens. Red zones indicate approximate restrictive effects of interferons.

588

589 STAR Methods

590 Contact for reagent and resource sharing

591 Further information and requests for reagents may be directed to and will be fulfilled by the Lead 592 Contact Dr. Matthew D. Welch (welch@berkeley.edu).

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594 EXPERIMENTAL MODEL AND SUBJECT DETAILS

24

595 **Bacterial strains**

596 All R. parkeri used in this study were in the Portsmouth strain originally obtained from 597 Christopher Paddock (Centers for Disease Control and Prevention) and were authenticated by whole 598 genome sequencing (NCBI Trace and Short-Read Archive: Sequence Read Archive (SRA), accession 599 number SRX4401164). To prepare the bacteria for infections, confluent T175 flasks of female African 600 green monkey kidney epithelial Vero cells (authenticated by STR analysis) grown in DMEM (Gibco 601 11965-092) with glucose (4.5 g/L) supplemented with 2% fetal bovine serum (FBS, GemCell) were infected with 5 x 10⁶ R. parkeri per flask and rocked for 10 min at 37°C. Infected cells were scraped and 602 603 collected at 5-6 dpi when ~90% of cells were highly infected, as indicated by cell rounding. Scraped 604 cells were centrifuged at 12,000 x G for 20 min at 4°C. Pelleted cells were then resuspended in K-36 605 buffer (0.05 M KH₂PO₄, 0.05 M K₂HPO₄, 100 mM KCl, 15 mM NaCl, pH 7) and transferred to a cold 606 glass Dounce homogenizer. Bacteria were released from infected cells by repeated douncing (60 607 strokes). The dounced solution was then centrifuged at 200 x G for 5 min at 4°C to pellet host cell debris. 608 The supernatant containing *R. parkeri* was overlaid on a 30% MD-76R (Merry X-Ray) solution. 609 Gradients were then centrifuged at 18,000 rpm in an SW-28 ultracentrifuge swinging bucket rotor 610 (Beckman/Coulter) for 20 min at 4°C to separate remaining host cells debris from the bacteria. Bacterial 611 pellets were resuspended in brain heart infusion (BHI) media (BD, 237500) and stored at -80°C.

612 Titers were determined for *R. parkeri* stocks via plaque assays by serially diluting the bacteria 613 in 6-well plates containing confluent Vero cells, which were plated ~24 h prior. Plates were then spun 614 for 5 min at 300 x G in an Eppendorf 5810R centrifuge. At 24 hpi, the media from each well was 615 aspirated and the wells were overlaid with 4 ml/well DMEM with a final concentration of 5% FBS and 616 0.7% ultrapure agarose (Invitrogen, 16500-500). When plagues were visible at 5-6 dpi, an overlay of 617 0.7% agarose in DMEM containing 2.5% neutral red solution (Sigma, N6264) was added and plates 618 incubated overnight until plagues were clearly visible. Plagues were then counted to determine bacterial 619 concentrations.

620 *L. monocytogenes* strain 10403S was originally obtained from Dr. Daniel Portnoy (UC Berkeley).

25

622 Animal experiments

623 Animal research using mice was conducted under a protocol approved by the UC Berkeley 624 Institutional Animal Care and Use Committee (IACUC) in compliance with the Animal Welfare Act and 625 other federal statutes relating to animals and experiments using animals. The UC Berkeley IACUC is 626 fully accredited by the Association for the Assessment and Accreditation of Laboratory Animal Care 627 International and adheres to the principles of the Guide for the Care and use of Laboratory Animals. 628 Infectious disease studies were performed in a biosafety level 2 facility. All animals were maintained at 629 the UC Berkeley campus and all infections were performed in accordance with the approved Welch lab 630 Animal Use Protocol. Mice were age matched between 8 and 18 weeks old. Mice were selected for 631 experiments based on their availability, regardless of sex. PFU data for each gender is reported in 632 Figure S5 at 48 and 72 hpi. All mice were healthy at the time of infection and were housed in 633 microisolator cages and provided chow and water. Experimental groups were littermates of the same 634 sex that were randomly assigned to experimental groups. All mice were phenotypically healthy at the 635 time of the experiment, as determined by their weight, temperature, and movement around the cage. 636 For experiments with mice mutated for *Ifnar* and *Ifngr*, as described in Figure 6, mice were immediately 637 euthanized if they exhibited severe degree of infection, as defined by a core body temperature dropping 638 below 90° F or lethargy that prevented the mouse from moving around the cage normally.

639

640 Mouse genotyping

641 Casp1^{-/-} (Rauch et al., 2017), Sting^{gt/gt} (Sauer et al., 2011), Cgas^{-/-} (Marcus et al., 2018), Gsdmd⁻ ^{/-} (Rauch et al., 2017), *Irf5*^{-/-} (Purtha et al., 2012), *Ifit1*^{-/-} (Szretter et al., 2012), and *Ifit2*^{-/-} (Fensterl et al., 642 2012), *lipG^{-/-}* (Ishida et al., 2003), *Gbp^{chr3-/-}* (Yamamoto et al., 2012) mice were previously described. 643 Casp11^{-/-} (Wang et al., 1998), Irf1^{-/-} (Matsuyama et al., 1993), Ifnar^{-/-} (Müller et al., 1994), Ifngr^{-/-} (Huang 644 645 et al., 1993), *Ifnar^{-/-}Ifngr^{-/-}*, and WT mice were previously described and originally obtained from Jackson 646 Laboratories. For genotyping, ear clips were boiled for 15 min in 60 µl of 25 mM NaOH, quenched with 647 10 µl tris-HCl pH 5.5, and 2 µl of lysate was used for PCR using SapphireAMP (Takara, RR350) and primers specific for each gene. Mutant mice were genotyped using the following primers: Ifnar forward 648

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649 (F): CAACATACTACAACGACCAAGTGTG; Ifnar WT reverse (R): AACAAACCCCCAAACCCCAG; 650 Ifnar mutant R: ATCTGGACGAAGAGCATCAGG; WT Casp1/11 F: CATGCCTGAATAATGATCACC; 651 WT Casp1/11 R: GAAGAGATGTTACAGAAGCC; Casp1/11 mutant F: GCGCCTCCCCTACCCGG; 652 Casp1/11 mutant R: CTGTGGTGACTAACCGATAA; Cgas F: ACTGGGAATCCAGCTTTTCACT; Cgas 653 R: TGGGGTCAGAGGAAATCAGC; WT tlr4 F: CACCTGATACTTAATGCTGGCTGTAAAAAG; WT tlr4 654 R: GGTTTAGGCCCCAGAGTTTTGTTCTTCTCA; tlr4 mutant F: TGTTGCCCTTCAGTCACAGAGACTCTG; tlr4 mutant R: TGTTGGGTCGTTTGTTCGGATCCGTCG; 655 656 Sting F: GATCCGAATGTTCAATCAGC; Sting R: CGATTCTTGATGCCAGCAC; Gsdmd F: 657 ATAGAACCCGTGGAGTCCCA; and Gsdmd R: GGCTTCCCTCATTCAGTGCT.

658

659 **Deriving bone marrow macrophages**

660 To obtain bone marrow, male or female mice were euthanized and femurs, tibias, and fibulas 661 were excised. Visible muscle and connective tissue were removed from the bones and the bones were 662 sterilized with 70% ethanol. Bones were washed with BMDM media (20% HyClone FBS, 1% sodium 663 pyruvate, 0.1% β-mercaptoethanol, 10% conditioned supernatant from 3T3 fibroblasts, in Gibco DMEM 664 containing glucose and 100 U/ml penicillin and 100 ug/ml streptomycin) and ground using a mortar and 665 pestle. Bone homogenate was passed through a 70 µm nylon Corning Falcon cell strainer (Thermo 666 Fisher Scientific, 08-771-2) to remove particulates. Filtrates were centrifuged in an Eppendorf 5810R at 667 1,200 RPM (290 x G) for 8 min, supernatant was aspirated, and the remaining pellet was resuspended 668 in BMDM media. Cells were then plated in non-TC-treated 15 cm petri dishes (at a ratio of 10 dishes 669 per 2 femurs) in a volume of 30 ml BMDM media and incubated at 37° C. An additional 30 ml BMDM 670 media was added 3 d later. At 7 d the media was aspirated and cells were incubated at 4°C with 15 ml 671 cold PBS (Gibco, 1x pH 7.4, no ions) for 10 min. BMDMs were then scraped from the plate, collected 672 in a 50 ml conical tube, and centrifuged at 1,200 RPM (290 x G) for 5 min. The PBS was then aspirated, 673 cells were resuspended in BMDM media, and cell numbers were counted using trypan blue and a 674 hemocytometer. For freezing, the pellet was resuspended in BMDM media with 30% FBS and 10%

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675 DMSO at 10⁷ cells/ml. 1 ml aliquots were frozen in a Styrofoam box at -80°C for one day and then 676 moved to liquid nitrogen.

677 **METHOD DETAILS**

678 Mouse infections

For mouse infections, R. parkeri were prepared by diluting 30%-prep bacteria to 1 ml cold sterile 679 680 PBS, centrifuging the bacteria at 12,000 x G for 1 min (Eppendorf 5430 centrifuge), and resuspended in cold sterile PBS to the desired concentration (either 5x10⁷ PFU/ml, 2.5x10⁷ PFU/ml, 5x10⁶ PFU/ml, 681 682 or 5x10⁵ PFU/mI). Bacterial suspensions were kept on ice for the duration of the injections. For i.v. 683 injections, mice were exposed to a heat lamp while in their cages for approximately 5 minutes and then 684 each mouse was moved to a mouse restrainer (Braintree, TB-150 STD). The tail was sterilized with 685 70% ethanol and then 200 µl of bacterial suspensions were injected using 30.5 gauge needles into the 686 lateral tail vein. For monitoring body temperatures of infected mice, mice were placed in a mouse 687 restrainer, and a rodent rectal thermometer (BrainTree Scientific, RET-3) was used to measure 688 temperature. For delivering the anti-IFN- γ antibody (BioLegend, 505847), mice were injected i.v. with 689 300 µl at 30 min p.i., 200 µl at 24 hpi, and 200 µl at 48 hpi, totaling 800 µl (0.8 µg antibody). Mice were 690 monitored daily for clinical signs of disease, such as hunched posture, lethargy, or scruffed fur. Only 691 mice lacking both interferon receptors exhibited such symptoms of disease, and if this occurred, mice 692 were monitored daily for these symptoms, as well as for loss in body weight and temperature. If a mouse 693 displayed severe symptoms of infection, as defined by a reduction in body temperature below 90°F or 694 an inability to move around the cage normally, the animal was immediately and humanely euthanized 695 using CO₂ followed by cervical dislocation, according to IACUC-approved procedures.

For harvesting organs, mice were euthanized at the indicated pre-determined times and doused with ethanol. Mouse organs were extracted in a biosafety cabinet and deposited into 50 ml conical tubes (Falcon) containing 4 ml sterile cold PBS (Gibco 10010-023, no ions) for the spleen, and containing 8 ml cold sterile PBS for the liver. Organs were kept on ice and were each homogenized for approximately 10 s using an immersion homogenizer (Fisher, Polytron PT 2500E) at 22,000 RPM. Organ

701 homogenates were spun at 290 x G for 5 min to pellet cell debris (Eppendorf 5810R centrifuge). 20 µl 702 of organ homogenates were plated into confluent Vero cells (plated 48 h prior) in 12-well plates, and 703 then serial diluted. The plates were then spun at 260 x G for 5 min at room temperature (Eppendorf 704 5810R centrifuge) and incubated at 33°C overnight. To reduce the possibility of contamination, organ 705 homogenates were plated in duplicate and the second replicate was treated with 50 µg/ml carbenicillin 706 (Sigma) and 200 µg/ml amphotericin B (Gibco). The next day, at approximately 16 hpi, the cells were 707 gently washed by replacing the existing media with 1 ml DMEM containing 2% FBS. The media was 708 then aspirated and replaced with 2 ml/well of DMEM containing 0.7% agarose, 5% FBS, and 200 µg/ml 709 amphotericin B. When plaques were visible, at approximately 6 dpi, 1 ml of DMEM containing 0.7% 710 agarose, 1% FBS, and 2.5% neutral red (Sigma) was added to each well. Plaques were counted 24 h 711 later.

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713 Infections in vitro

714 Aliguots of BMDMs were thawed on ice, diluted into 9 ml of DMEM, centrifuged in an Eppendorf 715 5810R at 1,200 RPM (290 x G) for 5 minutes, and the pellet was resuspended in 10 ml BMDM media 716 without antibiotics. The number of cells was counted using Trypan blue (Sigma, T8154) and a 717 hemocytometer (Bright-Line), and 5 x 10⁵ cells were plated into 24-well plates. Approximately 16 h later, 718 30% prep R. parkeri were thawed on ice and diluted into fresh BMDM media to the desired concentration (either 10⁶ PFU/ml or 2x10⁵ PFU/ml). Media was then aspirated from the BMDMs, replaced with 500 µl 719 720 media containing R. parkeri, and plates were spun at 300 G for 5 min in an Eppendorf 5810R. Infected 721 cells were then incubated in a humidified CEDCO 1600 incubator set to 33°C and 5% CO₂ for the 722 duration of the experiment.

For measuring PFU, supernatants were aspirated from individual wells, and each well was gently washed twice with 500 μ l sterile milli-Q-grade water. 1 ml of sterile milli-Q water was then added to each well and repeatedly pipetted up and down to lyse the host cells. Serial dilutions of lysates were added to confluent Vero cells in 12 well plates that were plated 24 or 48 h prior. Plates were then spun at 300 x G using an Eppendorf 5810R centrifuge for 5 min at room temperature and incubated at 33°C

overnight. At ~16 hpi, media was aspirated and replaced with 2 ml/well of DMEM containing 0.7% agarose and 5% FBS. When plaques were visible, at approximately 6 dpi, 1 ml of DMEM containing 0.7% agarose, 1% FBS, 200 μ g/ml amphotericin B (Invitrogen, 15290-018), and 2.5% neutral red (Sigma) was added to each well. Plaques were counted 24 h later. For neutralizing IFN-I signaling, an ultra-LEAF-purified α -IFNAR-1 antibody (BioLegend, 127323) was added to a final concentration of 1 μ g/ml at 0 hpi. For experiments with recombinant TNF- α , 200 ng was added to each well in a 24-well plate, and products from two different vendors were tested (BioLegend 575202 and Thermo PMC3014).

For collecting supernatant from $Casp1^{-/-}Casp11^{-/-}$ cells, $5x10^5$ BMDMs in 24-well plates were infected with *R. parkeri* at an MOI of 1 and at 24 hpi supernatants were pooled and stored at -80°C. For adding the supernatant to infected BMDMs, either 200 or 500 µl of supernatant was removed at 20 mpi from the untreated cells and replaced with the supernatant from $Casp1^{-/-}Casp11^{-/-}$ cells.

739 For infections with L. monocytogenes, cultures of L. monocytogenes strain 10403S were grown 740 in 2 ml sterile-filtered BHI shaking at 37° until stationary phase (~16 h). Cultures were centrifuged at 741 20,000 x G (Eppendorf 5430), the pellet was resuspended in sterile PBS (Gibco 10010-023), and diluted 742 100-fold in PBS. 10 µl of the diluted bacteria were then added to each well of a 24-well plate of BMDMs 743 that were plated ~16 h prior to infections at 5×10^5 cells/well. Bacteria were also plated out onto Luria 744 Broth agarose plates to determine the titer, which was determined to be $\sim 5 \times 10^5$ bacteria / 10 µl, for an 745 MOI of 1 (based on the ratio of bacteria in culture to number of BMDMs). Infected cells were incubated 746 in a humidified 37° incubator with 5% CO₂. 25 µg of gentamicin (Gibco 15710-064) was added to each 747 well (final concentration 50 µg/ml) at 1 hpi. At 30 mpi, 2, 5, and 8 hpi, the supernatant was aspirated 748 from infected cells, and cells were washed twice with sterile milli-Q water. Infected BMDMs were then 749 lysed with 1 ml sterile water by repeated pipetting and scraping of the well. Lysates were then serially 750 diluted and plated on LB agar plates, incubated at 37° overnight, and CFU were counted at ~20 h later.

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752 High-throughput DNA sequencing

For high-throughput DNA sequencing, 5×10^5 BMDMs were plated in 24-well plates and infected 16 h later with *R. parkeri* and treated with 10,000 activity units of recombinant IFN- β (PBL Cat # 12405-

755 1). To determine the percentage of cells that were successfully infected, we analyzed the infected cells 756 using immunofluorescence microscopy and observed that the multiplicity of infection (the average 757 number of bacteria per host cell) was 2.3, and that 71% of cells were infected. At 12 hpi, cells were 758 lysed and RNA was purified using an RNeasy purification kit (Qiagen). RNA guality was assessed using 759 an Agilent 2100 Bioanalyzer, and all samples had RIN values above 8.0. Transcripts were selected 760 using polyA selection (using Dynabeads mRNA Purification Kit, Invitrogen) and enzymatically 761 fragmented as part of the Apollo library prep kits (Wafergen PrepX RNA library prep for Illumina). 762 Libraries were constructed by using Apollo 324 (IntegenX Inc.), PCR-amplified, and multiplexed at the 763 Functional Genomics Lab at the University of California, Berkeley (http://gb3.berkeley.edu/gb3/fgl/). The 764 resulting libraries were sequenced at the Vincent J. Coates Genomics Sequencing Facility at the 765 University of California, Berkeley using single-end reads, 50 base length, with the Hiseg 2000 Illumina 766 platform. Sequence data were aligned to the Mus musculus C57BL/6 reference genome (reference 767 assembly GCA 000001635.8) using CLC Genomics Workbench (Qiagen). Fold regulation of genes for 768 each genotype was determined by referencing the sample uninfected, untreated BMDMs. Comparisons 769 were then made between the sequencing results from the infected, IFN-I-treated WT and the Irf mutant 770 genes. Each data set was composed of at least 55 million reads and 98.3% of the reads aligned with 771 the reference. Genes with low abundance of reads (Reads Per Kilobase of transcript, per Million 772 mapped reads, RPKM, of <10) in the infected WT BMDMs treated with IFN-I were excluded from the 773 analysis. Sequencing data were uploaded to GEO, accession number GSE128211.

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775 Microscopy

For brightfield microscopy, images were captured using an IX71 Olympus microscope with a UCPLFLN 20x 0.7 NA objective, OptiMOS sCMOS camera (QImaging), and Micro-Manager software (Edelstein et al., 2014). For immunofluorescence microscopy, 2.5-5 x 10⁵ BMDMs were plated overnight in BMDM media in 24-well plates with sterile 12 mm coverslips (Thermo Fisher Scientific, 12-545-80). Infections were performed as described above. At the indicated times post-infection, coverslips were washed once with PBS and fixed in 4% paraformaldehyde (Ted Pella Inc., 18505, diluted in 1 x PBS)

782 for 10 min at room temperature. Coverslips were then washed 3 times in PBS. Coverslips were washed 783 once in blocking buffer (1 x PBS with 2% BSA) and permeabilized with 0.5% triton X-100 for 10 min. 784 Coverslips were incubated with primary and secondary antibodies diluted in 2% BSA in PBS, each for 785 30 min at room temperature. R. parkeri were detected using mouse anti-Rickettsia 14-13 (originally 786 from Dr. Ted Hackstadt, NIH/NIAID Rocky Mountain Laboratories). GBP2 was detected with anti-GBP2 787 (ProteinTech, 11854-1-AP; Research Resource Identifier AB 2109336). Nuclei were stained with DAPI, 788 and actin was stained with Alexa-568 phalloidin (Life Technologies, A12380). Secondary antibodies 789 were Alexa-405 goat anti-mouse (A31553) and Alexa-488 goat anti-rabbit (A11008). Coverslips were 790 mounted in Prolong mounting media (Invitrogen). Samples were imaged with the Nikon Ti Eclipse 791 microscope with a Yokogawa CSU-XI spinning disc confocal with 60X and 100X (1.4 NA) Plan Apo 792 objectives, and a Clara Interline CCD Camera (Andor Technology) using MetaMorph software 793 (Molecular Devices). Rendered Z-stacks were used for quantifications. Images were processed using 794 FIJI (Schindelin et al., 2012) and brightness and contrast adjustments were applied to entire images. 795 Images were assembled using Adobe Illustrator. Representative images are a single optical section, in 796 which most or all bacteria were in the focal plane.

797

798 In vitro assays

799 For LDH assays, 60 µl of supernatant from wells containing BMDMs was collected into 96-well 800 plates. 60 µl of LDH buffer was then added. LDH buffer contained: 3 µl of "INT" solution containing 2 801 mg/ml tetrazolium salt (Sigma I8377) in PBS; 3 µl of "DIA" solution containing 13.5 units/ml diaphorase 802 (Sigma, D5540), 3 mg/ml β-nicotinaminde adenine dinucleotide hydrate (Sigma, N3014), 0.03% BSA, 803 and 1.2% sucrose; 34 µl PBS with 0.5% BSA; and 20 µl solution containing 36 mg/ml lithium lactate in 804 10 mM Tris HCl pH 8.5 (Sigma L2250). Supernatant from uninfected cells and from cells completely 805 lysed with 1% triton X-100 (final concentration) were used as controls. Reactions were incubated at 806 room temperature for 20 min prior to reading at 490 nm absorbance using an Infinite F200 Pro plate 807 reader (Tecan). Values for uninfected cells were subtracted from the experimental values, divided by

808 the difference of triton-lysed and uninfected cells, and multiplied by 100 to obtain percent lysis. Each 809 experiment was performed and averaged between technical duplicates and biological duplicates.

For the IFN-I bioassay experiments, 5 x 10^4 3T3 cells containing an interferon-sensitive 810 811 response element (ISRE) fused to luciferase (Jiang et al., 2005; McWhirter et al., 2009) were plated per 812 well into 96-well white-bottom plates (Greneir 655083) in DMEM containing 10% FBS, 100 U/ml 813 penicillin and 100 µg/ml streptomycin. 24 h later, media was replaced and confluent cells were treated 814 with 2 µl of supernatant harvested from BMDM experiments. After 4 h, media was removed and cells 815 were lysed with 40 µl TNT lysis buffer (20 mM Tris, pH 8, 200 mM NaCl, 1% triton-100). Lysates were 816 then injected with 40 µl firefly luciferin substrate (Biosynth) and luminescence was measured using a 817 SpectraMax L plate reader (Molecular Devices).

- 818
- 819 **qPCR**

820 For qPCR experiments using mouse tissue, 50 µl of organ homogenates was added to 600 µl 821 of RLT buffer (Qiagen) containing 1% β -mercaptoethanol and frozen at -80°C. Frozen lysates were later 822 thawed and RNA was purified and treated with DNAse, according to the manufacturer's protocol using 823 an RNeasy kit (Qiagen). RNA abundance was guantified using a NanoDrop ND-1000 and 200 ng RNA 824 was in vitro transcribed (ProtoScript II, NEB, M0368S) and diluted 10x in sterile nuclease-free water. 825 Real-time PCR was performed using SYBR Green (Thermo, A25742), 2 ul of cDNA, and 1 uM each of 826 the following oligonucleotides: F: GGCTGTATTCCCCTCCATCG; actin R: actin 827 GTCACCCACATAGGAGTCCTTC; IFN- β F: AGCTCCAAGAAAGGACGAACAT; and IFN- β R: 828 CCCTGTAGGTGAGGTTGATCTT. For normalization, values for IFN- β were divided by values for actin. 829 Measurements were acquired with a QuantStudio 5 real-time qPCR machine (Applied Biosystems).

830

831 QUANTIFICATION AND STATISTICAL ANALYSIS

832 Statistical parameters and significance are reported in the figures and the figure legends. Data 833 are determined to be statistically significant when p<0.05 by an unpaired two-tailed Student's T-Test an

834 ANOVA with multiple comparisons post-hoc test. For *in vivo* PFU data, data are determined to be

835	statistically significant when p<0.05 by a Mann-Whitney U test. Asterisks denote statistical significance
836	as: *, p<0.05; **, p<0.01; ***, p<0.001; ****, p<0.0001, compared to indicated controls. For animal
837	experiments, bars denote medians. Error bars indicate standard error (SE). All other graphical
838	representations are described in the figure legends. Statistical analyses were performed using
839	GraphPad PRISM.
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842	Table S1. Related to Figure 3. RNA-seq analysis of genes regulated by IRFs during <i>R. parkeri</i>
843	infection and IFN-I treatment.
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