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CARD9+ Microglia Orchestrate Antifungal Immunity via IL-1β and CXCL1-mediated Neutrophil Recruitment

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1	CARD9 ⁺ Microglia Orchestrate Antifungal Immunity via IL-1 β and CXCL1-mediated
2	Neutrophil Recruitment
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4	CARD9 ⁺ Microglia Promote Antifungal Immunity via IL-1 eta and CXCL1-mediated
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35 Abstract

37	The C-type lectin receptor/Syk adaptor CARD9 facilitates protective antifungal immunity within
38	the central nervous system (CNS), as human CARD9-deficiency causes fungal-specific CNS-
39	targeted infection susceptibility. CARD9 is required for neutrophil recruitment to the fungal-
40	infected CNS, which mediates fungal clearance. Here, we investigated host and pathogen
41	factors that promote protective neutrophil recruitment during Candida albicans CNS invasion.
42	IL-1 β was essential for CNS antifungal immunity by driving CXCL1 production, which recruited
43	CXCR2-expressing neutrophils. Neutrophil-recruiting IL-1 β and CXCL1 production was
44	induced in microglia by the fungal-secreted toxin Candidalysin, in a p38-cFos-dependent
45	manner. Importantly, microglia relied on CARD9 for production of IL-1 β , via both <i>II1b</i>
46	transcriptional regulation and inflammasome activation, and of CXCL1 in the fungal-infected
47	CNS. Microglia-specific Card9 deletion impaired IL-1 β and CXCL1 production and neutrophil
48	recruitment, and increased CNS fungal proliferation. Taken together, an intricate network of
49	host-pathogen interactions promotes CNS antifungal immunity, which is impaired in human
50	CARD9-deficiency leading to CNS fungal disease.

51 Introduction

52

53	The CNS is invaded by microorganisms during systemic infections, yet the mechanisms of
54	CNS-specific anti-microbial immunity remain poorly-understood. This is particularly true for
55	CNS fungal infections, which present unmet diagnostic and treatment challenges, leading to
56	unacceptably high mortality (>50%) ¹ . Fungal CNS invasion is enhanced by fungal-specific risk
57	factors, including HIV infection, neutropenia, corticosteroid use, and Bruton's tyrosine kinase
58	inhibition ¹ . However, the most striking human risk factor for selective CNS fungal infection
59	susceptibility is inherited deficiency of the C-type lectin receptor (CLR)–Syk adaptor CARD9.
60	
61	CARD9 relays fungal-sensing signals downstream of the CLR superfamily of pattern
62	recognition receptors, including Dectin-1, Dectin-2, Dectin-3 and Mincle. Syk kinase is
63	recruited to phosphorylated ITAM sequences of CLRs or their signaling partner FcR γ to form
64	the CARD9-BCL10-MALT1 signalosome, which activates downstream effectors including
65	NFκB, NLRP3 inflammasome and MAPK signaling.
66	
67	CARD9-deficient patients manifest fungal-specific infection susceptibility, predominantly in the
68	CNS by Candida albicans ^{2, 3, 4} . We previously showed that CARD9-deficiency in humans and
69	mice confers a fungal- and brain-specific defect in neutrophil recruitment, which is detrimental
70	for control of CNS fungal invasion ⁵ . However, the CNS cellular and molecular cues that
71	promote protective neutrophil recruitment during C. albicans invasion and their dependence on

72 CARD9 *in vivo* remain unknown.

74 Herein, we systematically investigated host and pathogen factors that promote protective 75 neutrophil influx into the C. albicans-infected CNS to better understand the pathogenesis of 76 human CARD9-deficiency. We uncover an intricate pathway by which the C. albicans-secreted 77 toxin Candidalysin engages microglia to produce IL-1ß and CXCL1 for protective recruitment of 78 CXCR2-expressing neutrophils. Importantly, microglial IL-1 β and CXCL1 production depends 79 on CARD9 and specific deletion of microglial CARD9 impairs neutrophil recruitment to the C. 80 albicans-infected CNS. Collectively, our data unveil complex host-pathogen interactions that recruit protective neutrophils during fungal CNS invasion and reveal the mechanism that 81 82 underlies CNS fungal susceptibility in CARD9-deficiency.

83 **Results**

84

85 CLR Functional Redundancy during Fungal CNS Invasion

86 As previously shown, CARD9 is essential for protective CNS immunity against C. albicans, 87 principally through promoting early neutrophil recruitment (Fig. 1a). We first investigated the 88 relative contribution of CARD9-coupled CLRs, which are expressed by brain-resident microglia (Supplementary Fig. 1), in mediating this protective neutrophil recruitment. We infected mice 89 deficient in Dectin-1 (*Clec7a*^{-/-}), Dectin-2 (*Clec4n*^{-/-}), Dectin-3 (*Clec4d*^{-/-}) and Mincle (*Clec4e*^{-/-}) 90 and measured brain neutrophil accumulation at 24 h post-infection (Supplementary Fig. 2). 91 92 We chose this time-point since it is the peak of the neutrophil response in wild-type animals 93 and neutrophil depletion at this time-point increases susceptibility to brain fungal invasion⁵. 94

Animals individually deficient in CARD9-coupled CLRs recruited neutrophils to the infected
brain normally (Fig. 1b). Despite this, we observed increases in fungal brain burdens at 72 h
post-infection in mice deficient in Dectin-1 or Dectin-2 but not Dectin-3 or Mincle (Fig. 1c),
suggesting that Dectin-1 and Dectin-2 employ neutrophil recruitment-independent mechanisms
to protect against brain fungal proliferation. Indeed, brain-infiltrating neutrophils from Dectin-1
and Dectin-2-deficient animals exhibited reduced fungal phagocytosis (Supplementary Fig.
3), consistent with prior findings⁶.

102

To activate CARD9-dependent signaling, phosphorylation occurs on the ITAM sequence within
 the intracellular tail of Dectin-1 or FcRγ, which Dectin-2, Dectin-3 and Mincle associate with.

105 Therefore, we assessed whether deletion of all four CLRs affected the neutrophil response in

the infected brain. We used mice doubly-deficient in Dectin-1 and FcRγ (*Clec7a^{-/-}Fcerg1^{-/-}*)
and found that loss of both Dectin-1 and the FcRγ-coupled CLRs phenocopied Card9deficiency with significantly decreased neutrophil recruitment and corresponding increased
brain fungal burdens (**Fig. 1b,c**). Taken together, CARD9-coupled CLRs functionally
compensate to mediate neutrophil recruitment-dependent protection against *C. albicans* CNS
invasion.

112

113 MALT1 is Required for Defense against CNS Candidiasis

114 The CARD9-MALT1-BCL10 signalosome is necessary for transducing fungal-sensing 115 intracellular signals. Human deficiencies of MALT1 or BCL10 cause defective innate and 116 adaptive immune responses, and many of these patients die in childhood from bacterial and 117 viral infections². Human MALT1-deficiency additionally manifests with mucosal candidiasis, suggesting that antifungal immunity is impaired in these patients. However, whether MALT1-118 119 deficiency also predisposes to brain-targeted candidiasis is unknown. To test this, we infected 120 Malt1-deficient mice and assessed control of CNS C. albicans growth. Malt1-/- animals 121 recruited neutrophils to the brain similarly to wild-type, however these animals exhibited 122 uncontrolled brain fungal growth at 72 h post-infection (Fig. 1d). Therefore, MALT1 is critical 123 for protective CNS immunity against *C. albicans*; however, the MALT1-dependent protective 124 mechanisms operating in this tissue are independent of neutrophil recruitment, unlike CARD9. 125

126 Candida Drives CNS Neutrophil Influx via IL-1β and CXCL1

127 To determine the local cues that recruit protective neutrophils in the infected CNS, we 128 examined key cytokine and chemokine circuits using gene-deficient mice. We began by

129 infecting IL-1 receptor (IL-1R)-deficient mice, because production of IL-1 β by human peripheral blood mononuclear cells upon fungal stimulation depends on CARD9^{5, 7}, and because IL-1R 130 was previously shown to promote neutrophil recruitment to fungal-infected mucosal tissues^{8, 9}, 131 and to the bacterial-infected brain¹⁰. Upon *C. albicans* challenge, IL-1R-deficient mice 132 133 phenocopied Card9-deficient mice, with a loss of early brain neutrophil recruitment and accompanying increased fungal brain burdens (Fig. 2a,b). Consistent with this, loss of the IL-134 135 1R signaling adaptor MyD88 caused similar defects in recruiting neutrophils and controlling 136 fungal proliferation in the infected brain (Fig. 2a,b).

137

138 We next assessed which IL-1R ligands were important for driving CNS protection by infecting mice deficient in IL-1 α , IL-1 β or both. Mice lacking IL-1 α had a small reduction in neutrophil 139 140 numbers and a slight increase in fungal brain burden at 24 h post-infection (Fig. 2a,b). 141 However, the lack of IL-1 α appeared to be compensated by IL-1 β , since *II*1 $a^{-/-}$ animals 142 recovered and controlled fungal brain infection similar to wild-type by 72 h post-infection (Fig. 143 **2b**). In keeping with the critical contribution of IL-1 β , mice deficient in IL-1 β , or both IL-1 α /IL-1β, exhibited significantly reduced neutrophil accumulation and were highly susceptible for 144 145 fungal brain invasion (**Fig. 2a,b**). Therefore, IL-1 β is a critical mediator of neutrophil 146 recruitment to promote control of C. albicans brain infection.

147

Downstream of IL-1R, the production of local chemotactic mediators recruits immune cells to infected tissues. Previously, we showed that the CNS-neutropenia observed in mouse and human CARD9-deficiency is not caused by neutrophil-intrinsic chemotaxis defects⁵, but rather by insufficient local production of soluble chemotactic mediators. However, which among the

several chemoattractants and their receptors recruit(s) protective neutrophils to the *C. albicans*-infected CNS is unknown.

154

155 CARD9 was shown to drive production of the CXCR2 ligands CXCL1 and CXCL2 during 156 inflammatory arthritis¹¹ and murine subcutaneous phaeohyphomycosis¹². During systemic C. 157 albicans infection, CCR1 drives renal neutrophil accumulation and immune-related kidney 158 destruction¹³, the leukotriene B4 (LTB₄) receptor LTB4R1 promotes detrimental pulmonary neutrophil accumulation¹⁴, and CXCR1 mediates neutrophil-dependent fungal killing in the 159 kidney¹⁵. However, the role of these receptors in CNS anti-*Candida* immunity is unknown, 160 161 while CXCR2 and fMet-Leu-Phe (fMLP) receptor FPR1 have not been examined in anti-Candida defense. 162

163

To test the relative dependence on these major neutrophil-targeted chemoattractant receptors 164 in protecting the fungal-infected brain, we infected mice deficient in CCR1, CXCR1, CXCR2, 165 166 LTB4R1 or FPR1 and measured neutrophil recruitment and fungal brain burdens. We found no involvement of the CCL3–CCR1, CXCL5–CXCR1, LTB4–LTB4R1 or fMLP–FPR1 axes in 167 168 controlling fungal brain infection, in line with normal early neutrophil recruitment in infected 169 *Ccr1*^{-/-}, *Cxcr1*^{-/-}, *Ltb4r1*^{-/-} and *Fpr1*^{-/-} animals (Fig. 2c,d and Supplementary Fig. 4). In 170 contrast, CXCR2-deficient mice had significantly reduced neutrophil accumulation and 171 corresponding significantly increased fungal brain growth (Fig. 2c,d). These data demonstrate the importance of the CXCR2 axis in neutrophil-mediated protection against C. albicans brain 172 173 infection.

175 Next, we wondered which CXCR2 ligand may recruit protective neutrophils to the fungal-176 infected brain. We infected Cxcl1-/- mice that lack expression of the potent neutrophil 177 chemoattractant CXCL1. Notably, these animals had decreased neutrophil recruitment to the 178 brain post-infection and exhibited a similar CNS invasion susceptibility phenotype to the Cxcr2-179 ⁻ mice (**Fig. 2c,d**). Therefore, the CXCL1/CXCR2 chemokine axis is critical for protection against *C. albicans* brain invasion by recruiting protective neutrophils. Importantly, this data 180 181 indicates that the reduced CXCL1 in the human CARD9-deficient C. albicans-infected CSF is 182 biologically relevant and significant⁵.

183

184 IL-1β Activates CXCL1 in the Fungal-Infected Brain

Since both IL-1ß and CXCL1 were required for protection, we investigated whether their 185 activation in the infected brain was simultaneous or sequential. We measured IL-1ß and 186 187 CXCL1 in brain homogenates at 24 h post-infection in animals lacking these inflammatory 188 mediators using ELISA. We found no defect in IL-1 β levels in CXCL1-deficient infected brains; 189 however, we discovered a significant defect in CXCL1 production in the absence of IL-1 β (Fig. 190 **3a**). To define the IL-1^β-dependent brain cellular sources of CXCL1, we infected wild-type and 191 IL-1β-deficient mice and used intracellular flow cytometry. CXCL1 and pro-IL-1β were produced by multiple myeloid phagocytes in the fungal-infected brain, including resident 192 193 microglia, the most numerous immune cells in the brain, recruited Ly6C^{hi} monocytes which have been implicated in controlling *C. albicans* CNS invasion¹⁶, and neutrophils themselves 194 195 (Fig. 3b). Interestingly, *II1b^{-/-}* microglia recovered from *C albicans*-infected brains had a profound defect in CXCL1 production, exhibiting significant reductions under every ex vivo 196 restimulation condition tested (Fig. 3c). Ly6C^{hi} monocytes isolated from *II1b^{-/-} C. albicans*-197

infected brains produced less CXCL1 when restimulated *ex vivo* with LPS, with no differences detected under non-stimulated or zymosan-stimulated conditions. Neutrophil production of CXCL1 did not differ between the two mouse groups (**Fig. 3c**). Therefore, IL-1 β is required for subsequent CXCL1 production from resident microglia and recruited monocytes, which in turn recruits CXCR2-expressing neutrophils to the fungal-infected brain (**Fig. 2c**).

203

204 Candidalysin is a Fungal Avirulence Factor in the Brain

205 Use of genetically-deficient mice allowed us to map the host pathway promoting protection

against *C. albicans* brain infection, in which IL-1 β –IL-1R–MyD88 signaling activates CXCL1

207 production by resident microglia and recruited monocytes to mobilize neutrophils into the CNS.

208 To identify the pathogen-associated factors that induce this protective host pathway, we

209 infected animals with *C. albicans* strains lacking known virulence factors and assessed

210 neutrophil recruitment and IL-1 β and CXCL1 production in the infected brain.

211

C. albicans hyphae are the predominant CNS-invasive morphology of *C. albicans*⁵ and hyphal 212 213 formation is associated with important virulence traits such as toxin and protease production, 214 adhesion, invasion, and immune system activation¹⁷. Thus, we first asked whether neutrophil 215 recruitment was impaired during infection with the $hgc1\Delta/\Delta$ C. albicans strain which cannot 216 filament *in vivo*¹⁸. Indeed, infection with hypha-deficient $hgc1\Delta/\Delta$ C. albicans significantly 217 impaired neutrophil recruitment and enhanced fungal CNS tissue invasion relative to the isogenic wild-type *C. albicans* strain (Fig. 4a). Thus, filamentation is strikingly not required for 218 219 C. albicans invasion of brain tissue, in contrast to other organs such as the kidney¹⁸.

220

221 Candidalysin is a recently-described peptide toxin encoded by ECE1 and expressed 222 exclusively by *C. albicans* hyphae¹⁹. Candidalysin was shown to mediate epithelial cell 223 damage via pore formation in the plasma cell membrane resulting in IL-1 α release and pro-224 inflammatory cytokine production. Hence, Candidalysin-null mutants were highly attenuated in murine oropharyngeal and vulvovaginal candidiasis models^{19, 20, 21}. Instead, we found that lack 225 of Candidalysin promoted brain infection, and that this phenotype was specific to the 226 227 Candidalysin peptide since mutant strains deficient in the entire gene (*ece1* Δ/Δ) or specifically 228 in the Candidalysin-encoding portion of the gene ($ece1\Delta/\Delta + ECE1_{\Delta 184-279}$) were both hyper-229 virulent for C. albicans brain invasion (Fig. 4b).

230

231 The increased ability of the Candidalysin-null mutants to proliferate within the brain directly 232 correlated with the degree of neutrophil recruitment. We found a near absence of neutrophils in 233 the brains of wild-type animals infected with Candidalysin-null strains and observed hyphal 234 forms growing in the brain parenchyma with no neutrophilic reaction (**Fig. 4c**). In contrast, the 235 Candidalysin-producing parental strain and the re-integrant control strain promoted neutrophil 236 recruitment at 24 h post-infection, and these neutrophils clustered around invading hyphae 237 (Fig. 4c). In line with the absence of neutrophils in the brains of mice infected with 238 Candidalysin-null strains, IL-1 β and CXCL1 were significantly reduced in brain homogenates from animals infected with these strains (Fig. 4d). Therefore, Candidalysin is a key fungal 239 240 factor that activates the IL-1β–CXCL1 protective pathway *in vivo*. Notably, in contrast to its role 241 in the mucosa, Candidalysin acts as an avirulence factor in the brain by instigating protective 242 host CNS immunity, underscoring the tissue-specific opposing roles that a microbial factor may 243 play during infection with the same pathogen¹⁷.

245 We next wondered whether other C. albicans hyphae-associated secreted proteins also 246 activate protective neutrophil responses in the brain. Secreted aspartyl proteases (Saps) are 247 enzymes with extracellular proteolytic activity and are linked to virulence²². C. albicans Saps promote neutrophil recruitment during vulvovaginal candidiasis in mice^{23, 24}. Expression of the 248 249 SAP4-6 subfamily is coordinately regulated with hyphal formation²², therefore we tested 250 whether these hyphal-associated Saps contributed towards virulence during brain invasion. 251 Wild-type animals infected with the triple-deficient strain $sap4/5/6\Delta/\Delta$ had comparable brain 252 fungal burdens to animals infected with the complemented control strain (Fig. 4e). In line with 253 this, we saw no difference in CNS neutrophil recruitment in these animals, indicating that C. 254 albicans Saps exhibit tissue-specific roles in promoting neutrophil recruitment during infection^{23, 24} (Fig. 4e). Therefore, protective CNS neutrophil recruitment is activated by 255 256 Candidalysin, and not by other *C. albicans* hyphae-secreted enzymes.

257

258 Candidalysin Drives Microglial IL-1β and CXCL1 in vivo

259 Since Candidalysin activates host CNS immunity, we sought to define the Candidalysin-

responsive CNS immune cells. We infected wild-type mice with either the parental strain of *C*.

albicans (BWP17) or the Candidalysin-null strain (*ece1* Δ / Δ), and analyzed IL-1 β and CXCL1

production using intracellular flow cytometry at 24 h post-infection (**Fig. 5**).

Although all brain phagocytes produced both IL-1 β and CXCL1, microglia were the only

population to exhibit dependence on Candidalysin, since microglia isolated from $ece1\Delta/\Delta$ -

infected brains produced significantly less IL-1 β and CXCL1 *ex vivo* (**Fig. 5a,b**). In contrast,

266 Ly6C^{hi} monocytes and neutrophils did not depend on Candidalysin for IL-1β and CXCL1

production, suggesting that other as-yet unidentified fungal factors activate this pathway in
these phagocytes. Together, our data show that Candidalysin acts on microglia to stimulate IL1β release, which in turn drives CXCL1 production that is required for protective neutrophil
CNS recruitment.

271

272 Candidalysin Drives Differing Glial IL-1β–CXCL1 ex vivo

273 To gain mechanistic insights into how microglia respond to Candidalysin, we cultured the microglia cell line BV-2²⁵ in the presence of synthetic Candidalysin and measured IL-1β and 274 275 CXCL1 in the supernatants by ELISA. In line with our in vivo work, we found time- and dose-276 dependent IL-1ß production by BV-2 cells in response to Candidalysin (Fig. 6a). However, we 277 did not detect CXCL1 from BV-2 cells stimulated under these conditions. We first considered 278 that this could be due to Candidalysin-induced damage that may prevent BV-2 cells from 279 producing CXCL1 after IL-1β secretion. Indeed, as shown for epithelial cells¹⁹, Candidalysin 280 mediated dose-dependent cell damage to BV-2 microglia (Fig. 6b). Alternatively, additional 281 signals beyond IL-1β, derived from non-microglial CNS cells, might be required for microglial 282 CXCL1 induction, acting in trans. To test this hypothesis, we co-cultured BV-2 cells with 283 immortalized C8-D1A astrocytes in the presence of Candidalysin and measured IL-1ß and 284 CXCL1 in the supernatants. We chose astrocytes since they are known to respond to IL-1 β to produce inflammatory mediators, including CXCL1, in other models of CNS inflammation^{26, 27,} 285 286 ²⁸. We found that astrocytes responded to Candidalysin to produce CXCL1 (Fig. 6c), but not 287 IL-1 β (data not shown), and that CXCL1 production significantly increased when astrocytes 288 and microglia were co-cultured (Fig. 6c). To confirm that microglia are a relevant cellular 289 source of CXCL1 detected during microglia-astrocyte co-culture, we performed intracellular

staining for CXCL1 and found that BV-2 microglia are significant producers of CXCL1 in
response to Candidalysin, but only when astrocytes were present (Fig. 6d). Therefore,
astrocytes provide additional signals to microglia that are needed for CXCL1 production in
response to Candidalysin.

294

295 We next investigated the pathway activated by Candidalysin in BV-2 microglia to produce IL-296 1 β . Candidalysin was previously shown in epithelial cells to activate c-Fos, in a p38-dependent 297 manner, and the phosphatase MKP-1²⁰. We thus asked whether the same pathways are 298 activated by Candidalysin in BV-2 microglia. We found that Candidalysin sequentially and 299 dose-dependently activated MKP-1 and c-Fos (Fig. 6e), and chemical inhibition of p38 or c-300 Fos significantly reduced IL-1 β release by Candidalysin-stimulated BV-2 cells (**Fig. 6f**). 301 Therefore, microglia produce IL-1 β in response to Candidalysin via activation of p38 and c-302 Fos.

303

304 The Microglia IL-1 β -CXCL1 Response Requires CARD9

CARD9-deficiency is the only known risk factor that uniquely predisposes to CNS candidiasis 305 306 in the absence of iatrogenic intervention^{2, 5}. We first examined whether CARD9-deficiency 307 causes developmental defects in resident microglia, but found no defects in abundance or activation markers at steady state in *Card9^{-/-}* microglia, which accumulated in similar numbers 308 309 as wild-type microglia after fungal infection (**Supplementary Fig. 5**). Since *C. albicans* 310 activates the microglial IL-1 β -CXCL1 axis to regulate protective neutrophil CNS recruitment, 311 we next analyzed the dependence on CARD9 for induction of this pathway in microglia post-312 infection *in vivo*. We hypothesized that CARD9 is required for these functions, as microglia

highly express CARD9 and we previously found reduced transcription of CXC chemokines by *Card9-/-* microglia harvested from the *C. albicans*-infected brain⁵. We infected *Card9+/+* and *Card9-/-* animals with wild-type Candidalysin-expressing *C. albicans*, isolated phagocytes from the brain and measured pro-IL-1 β and CXCL1 production following *ex vivo* restimulation. We found significantly decreased frequencies of CXCL1+ and pro-IL-1 β + cells in the fungal-infected *Card9-/-* brain, and these decreases mapped to microglia (**Fig. 7a,b**).

319

320 Since production and secretion of mature IL-1 β depends on pro-IL-1 β expression and 321 consecutive inflammasome-dependent processing, we asked whether microglia depend on 322 CARD9 for pro-IL-1β transcription and/or inflammasome activation. We FACS-sorted microglia 323 from wild-type and Card9^{-/-} infected brains, and examined *II1b* transcription by qRT-PCR, and levels of pro-IL-1 β and cleaved and pro-caspase-1 by immunoblot. We found significantly 324 325 decreased *II1b* transcription in *Card9*^{-/-} microglia, which we confirmed at the protein level (Fig. 326 **7c,d**). These data are in line with the reported CARD9-dependent pro-IL-1 β transcription in 327 bone marrow-derived dendritic cells post-viral infection²⁹. We also found significantly reduced 328 cleaved caspase-1 in Card9^{-/-} microglia (Fig. 7d), indicating that Card9 also operates at the 329 level of inflammasome activation for IL-1 β production. Given that c-Fos mediated 330 Candidalysin-induced IL-1 β production by BV-2 cells (**Fig. 6e,f**), we measured c-Fos 331 expression in WT and Card9^{-/-} microglia by immunoblot and found significantly decreased c-332 Fos expression in *Card9*^{-/-} microglia (**Fig. 7d**). 333

We next examined the NLRP3 inflammasome in FACS-sorted WT and *Card9^{-/-}* microglia. We focused on NLRP3 because CARD9 was reported to negatively regulate NLRP3 activation

336	during macrophage Salmonella infection ³⁰ , and we recently showed that Candidalysin
337	activates NLRP3 in bone marrow-derived macrophages ³¹ . We found significantly decreased
338	NLRP3 protein expression in <i>Card9^{-/-}</i> microglia (Fig. 7e). Of interest, <i>Nlrp3^{-/-}</i> animals had
339	significantly decreased neutrophil accumulation to the C. albicans-infected brain and increased
340	fungal load post-infection, consistent with a potential role of Card9-dependent NLRP3-
341	inflammasome activation for protective neutrophil influx in the fungal-infected CNS (Fig. 7f).
342	Together, microglia require CARD9 for c-Fos activation and for production of mature IL-1 β via
343	<i>II1b</i> transcriptional regulation and inflammasome activation, to activate the IL-1 β –CXCL1 axis
344	in response to fungal invasion.
345	
346	Microglial CARD9 Deletion Causes CNS Fungal Invasion
347	We next directly examined the impact of genetic Card9 deletion specifically within microglia by
348	utilizing mice expressing tamoxifen-inducible Cre recombinase under the Cx3cr1 promoter
349	(<i>Cx3cr1</i> ^{CreER}) ³² . These mice has been used to genetically manipulate long-lived CX3CR1 ⁺
350	microglia while leaving short-lived CX3CR1 ⁺ monocytes and monocyte-derived macrophages

- unaffected. We bred $Cx3cr1^{CreER}$ animals to Card9-floxed mice¹¹, tamoxifen-pulsed the
- 352 progeny to activate Cre expression and waited 4-6 weeks to allow replenishment of short-lived
- 353 non-microglia CX3CR1⁺ cells from the bone marrow, while long-lived microglia remained
- 354 Card9-deficient (**Supplementary Fig. S6**). *C. albicans* infection of *Card9*^{fl/fl}*Cx3cr1*^{CreER+/-}
- animals revealed a significant dependence on *Card9* expression by the long-lived CX3CR1⁺
- cellular compartment for control of fungal brain growth (Fig. 8a), while fungal control in the
- kidney was unaffected in microglia-specific conditional *Card9*^{-/-} mice (**Fig. 8a**).
- 358

To analyze whether the susceptibility to brain infection in Card9^{fl/fl}Cx3cr1^{CreER+/-} mice was 359 360 related to a neutrophil recruitment defect, we quantified neutrophils within the infected brains of *Card9*^{fl/fl}*Cx3cr1*^{CreER+/-} mice and their Cre-negative littermates. We found that microglial 361 362 deletion of Card9 significantly reduced the protective early influx of neutrophils into the fungal-363 infected brain (Fig. 8b), which correlated with significantly decreased expression of microglial 364 pro-IL1 β and CXCL1 in the conditional *Card9*^{-/-} mice (**Fig. 8c**). Together, our data shows that 365 CARD9-expressing microglia orchestrate control of fungal brain invasion, in part by responding 366 to fungus-secreted Candidalysin, to produce IL-1β-induced CXCL1, which recruits CXCR2expressing neutrophils that are required for CNS fungal clearance (Supplementary Fig. S7). 367

368 **Discussion**

370 Herein, we demonstrate the critical contribution of CARD9-mediated IL-1β and CXCL1 in 371 recruiting protective neutrophils to the fungal-infected CNS. We identify microglia as major producers of CARD9-dependent IL-1β and CXCL1 during *C. albicans* CNS invasion and the 372 373 fungal-secreted toxin Candidalysin as a critical pathogen-derived factor activating this 374 pathway. Our study offers novel insights into the network of host and fungal factors that protect 375 against CNS fungal invasion and unveil the mechanism of CNS fungal susceptibility in 376 inherited CARD9-deficiency. 377 378 Systemic candidiasis is a leading cause of nosocomial bloodstream infection with mortality 379 >50% despite therapy³³. Neutropenia is the major predisposing factor for systemic candidiasis and *Candida* CNS invasion in particular^{5, 34}. Moreover, CNS invasion is prevalent during 380 381 systemic candidiasis in low-birth weight neonates and also occurs as an iatrogenic 382 complication post-neurosurgical procedures^{35, 36}. Strikingly, CARD9-deficiency is a primary 383 immunodeficiency disorder (PID) characterized by heightened susceptibility to fungal infections 384 of which CNS candidiasis is a hallmark^{2, 3}. CARD9-deficiency is the only known PID that 385 causes fungal-specific infection susceptibility without other infectious or non-infectious 386 manifestations, and the only PID that causes fungal disease in which CNS is a primary target 387 tissue¹⁷. We previously demonstrated that *Candida* CNS disease in CARD9-deficiency is 388 caused by a fungal- and brain-specific defect in neutrophil recruitment⁵. CNS neutropenia is 389 now confirmed in several CARD9-deficient patients with CNS candidiasis^{7, 37, 38}. Nonetheless,

how CARD9 mediates protective neutrophil trafficking into the fungal-infected CNS remainedunclear.

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393 Our analysis of mice deficient in several CLR, cytokine and chemokine circuits uncovered a) 394 the functional redundancy among CLRs, which may suggest the presence of yet-undiscovered 395 CARD9-coupled receptors for tissue-specific protection against fungal invasion; b) the 396 indispensable role of the CARD9 partner MALT1 in controlling CNS fungal invasion independent of neutrophil recruitment, which implies that MALT1^{-/-} patients may be at risk for 397 398 CNS fungal disease; and c) the critical contribution of IL-1 β -CXCL1-mediated neutrophil 399 recruitment for control of CNS fungal invasion. CXCR2 was known to mediate neutrophil trafficking during viral infection, parasitic meningitis³⁹ and fungal pneumonia⁴⁰, and herein we 400 401 reveal its importance for recruiting neutrophils during C. albicans CNS infection, principally 402 through binding CXCL1. In contrast, CCR1, CXCR1 and LTB4R1 are dispensable despite 403 them regulating neutrophil recruitment and function in other *C. albicans*-infected tissues^{13, 14, 15}. 404 These studies further underscore the organ- and context-specific dependence on chemotactic 405 molecules for protective host immunity.

406

We showed that IL-1 β is required for CXCL1 induction, in line with earlier work which showed IL-1 β -induced CXCL1 production controlling neutrophil accumulation during bacterial peritonitis and autoimmune, traumatic or bacterial neuroinflammation^{10, 41, 42}. Importantly, microglia are the primary myeloid cellular source of IL-1 β -dependent CXCL1 production *in vivo*. During oral candidiasis, IL-1R is also required for neutrophil accumulation to the oral mucosa⁸, as we have shown for systemic infection in the brain. However, further attesting to

the presence of tissue-specific anti-*Candida* immune response cues, the IL-1R-dependent
response in the oral epithelium is largely controlled by IL-1α released by damaged
keratinocytes⁸, whereas we found that IL-1α plays a modest role in the control of brain fungal
invasion. In fact, IL-1α release by epithelial cells, in both the oral and vaginal mucosal barriers,
is driven by exposure to the fungal secreted toxin Candidalysin^{19, 20, 21}.

418

419 Candidalysin enables the establishment of C. albicans mucosal infections, since Candidalysin-420 deficient strains are avirulent in these models^{19, 21}. In contrast, we found that Candidalysindeficient strains are hyper-virulent for the brain, associated with decreased IL-1^β and CXCL1 421 422 production and impaired neutrophil recruitment. These results indicate that Candidalysin is not 423 only a classical virulence factor, but also an immune modulator, which exerts context-specific 424 effects on the immune system. We propose that this dual function of Candidalysin is the result 425 of a co-evolutionary event; the fungus developed an efficient toxin to damage host 426 membranes, and, in response, the host evolved a sensitive Candidalysin detection system to 427 defend against this common mucosal pathogen. Whether Candidalysin is recognized by a 428 specific microglial innate receptor to mediate the protective IL-1 β -CXCL1 axis is unclear, since 429 the toxin mediates cellular damage which could also activate glial cells. Therefore, identifying 430 how host epithelial and immune cells recognize Candidalysin merits investigation.

431

We found that Candidalysin selectively activates microglia for IL-1β and CXCL1 production, a
 self-renewing macrophage population that contributes towards neuroinflammation in
 neurodegenerative disorders and promotes pathogen and dead cell clearance within the
 CNS⁴³. Interestingly, downstream of Candidalysin-induced microglial IL-1β secretion, which

occurs via c-Fos activation, we show that additional signals derived from astrocytes acting *in trans* are required for microglia to secrete CXCL1. Whether direct microglia-astrocyte contact
is required or astrocyte-derived soluble factors acting on microglia are needed remains
unknown. Indeed, microglia are known to interact with astrocytes to drive or suppress
inflammation^{44, 45}. Therefore, the molecular factors that drive microglia-astrocyte cross-talk
within the fungal-infected brain warrant further investigation.

442

443 Lastly, we examined the dependence on CARD9 for the microglia-mediated, IL-1 β -CXCL1-444 dependent pathway that recruits protective neutrophils, using fully Card9-deficient and 445 conditional microglia-specific Card9^{-/-} mice. We show that CARD9 is critical for c-Fos 446 activation and for the production of both IL-1ß and CXCL1 by microglia in the infected CNS operating at the levels of both transcriptional pro-IL-1ß regulation and inflammasome activation 447 448 for IL-1 β generation, with NLRP3 being at least partly involved, as shown with *Microsporum* 449 infection⁴⁶. Together, these data shed light into the pathogenesis of inherited CARD9-450 deficiency by outlining a pathway of CARD9-dependent microglial production of sequential IL-451 1β and CXCL1 that recruits protective neutrophils into the fungal-infected CNS.

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Future studies should examine how Card9 promotes microglial innate functions beyond
orchestrating neutrophil recruitment such as fungal uptake and killing. Of note, the phenotype
of the microglia-specific conditional knockout mice is less severe than that of *Card9^{-/-}* mice,
which may reflect the important role of astrocytes, which express Card9 post-*Candida*infection¹³, in priming microglial CXCL1 production. Future work should also examine the
potentially differential tissue-specific dependence on Card9 for neutrophil recruitment by non-

CNS tissue-resident macrophages, such as Kupffer cells, as previously shown for macrophages and dendritic cells⁴⁷, which will help further understand the CNS-specificity of fungal disease in CARD9-deficiency. Beyond understanding the pathogenesis of inherited CARD9-deficiency, our findings have important implications for recognizing the potential fungal infection risk in patients who are increasingly receiving Syk inhibitors for the treatment of autoimmune and malignant diseases^{48, 49}. Surveillance of Syk inhibitor-treated patients and research in conditional Syk-deficient mice will help determine their CNS fungal disease risk.

In summary, we present evidence of an intricate host immune pathway that protects the CNS
from invading fungi. This work uncovers the complex interactions occurring between the host
and the most common human fungal pathogen within the CNS, and sheds novel mechanistic
light into the pathogenesis of inherited CARD9-deficiency.

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481	
482	Authors' Contributions

R.A.D, B.H, J.R.N, T.M.H and M.S.L designed the study. R.A.D, M.S, V.O, B.Z, and I.M.D
performed the experiments. B.C.S, A.C.B, K.D.M-B, S.A.L, Y.I, S.G.F, G.D.B, B.H, J.R.N and
T.M.H provided key reagents/mouse lines and intellectual input into the experimental design
regarding their use. R.A.D. and M.S.L assembled and wrote the manuscript.

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488 Data Availability

The data that support the findings of this study are available from the corresponding authorsupon request.

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681 Methods

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683 **Mice**

Animals (males and females) were used at 8-12 weeks of age and were maintained in 684 individually ventilated cages under specific pathogen-free conditions at the 14BS facility at the 685 686 National Institutes of Health (Bethesda, MD, USA), the Memorial Sloan Kettering Cancer Center Comparative Medicine Shared Resources (New York, NY, USA), or the Medical 687 688 Research Facility at the University of Aberdeen (UK). The following strains (and their 689 respective WT controls/littermates) were obtained from the NIAID Taconic contract; Cxcr2^{-/-}, *II1r^{-/-}, Ltb4r1^{-/-}, Fpr1^{-/-}*. All other strains and their respective controls/littermates were bred in-690 691 house at the NIH (Clec7a-/-, Clec4n-/-, Clec4e-/-, Myd88-/-, Ccr1-/-, Cxcr1-/-, Cxcl1-/-, II1a-/-, II1b^{-/-}, II1a^{-/-}II1b^{-/-}, NIrp3^{-/-}, Card9^{fl/fl}Cx3CR1^{CreER+/-}), Memorial Sloan-Kettering Cancer Center 692 693 (*Clec7a^{-/-}Fcer1g^{-/-}*), University of Aberdeen (*Clec4d^{-/-}*), or USUHS (*Malt1^{-/-}*)⁵¹. Mice homozygous for the *Card9*^{tm1a} allele were purchased from the Wellcome Trust Sanger Institute 694 695 (EUCOMM Project No. 44813), and these animals were bred with the FLPer deleter strain 696 (Jackson Laboratories) to remove the FRT-flanked knock-out first cassette, generating *Card9*^{tm1c} homozygous mice (referred to as *Card9*^{fl/fl} in this manuscript)^{52,53}. Homozygous 697 *Card9*^{fl/fl} animals were bred with heterozygous *Cx3cr1^{CreER}* transgenic animals (Jackson 698 Laboratories) to generate Card9^{fl/fl}Cx3cr1^{CreER+/-} mice and littermate controls. Soon after 699 weaning (~5-6 weeks old), Card9^{fl/fl}Cx3cr1^{CreER+/-} mice and their controls were treated with two 700 701 10mg doses of tamoxifen (Sigma) administered in corn oil by oral gavage, given 48 h apart. 702 After 4-6 weeks, these animals were infected and analyzed as outlined in the Figure legends.

All experimentation conformed to conditions approved by the Animal Care and Use Committeeof the National Institute of Allergy and Infectious Diseases.

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706 Candidiasis Model and Fungal Burden Determination

707 *Candida albicans* strains used in this study were SC5314, BWP17, $ece1\Delta/\Delta$, $ece1\Delta/\Delta+ECE1$, 708 $ece1\Delta/\Delta + ECE1_{\Delta 184-279}$ ¹⁹, CAI4+Clp10 and $sap4/5/6\Delta/\Delta$, and $hgc1\Delta/\Delta$ and $hgc1\Delta/\Delta + HGC1^{18}$. 709 Yeast was serially passaged three times in YPD broth, grown at 30°C with shaking for 18-24 h 710 at each passage. Yeast cells were washed in PBS, counted, and injected intravenously via the lateral tail vein. Animals were infected with 1.3 × 10⁵ colony forming units (CFU) for analysis at 711 24 h post-infection, or 7 × 10⁴ CFU for analysis at 72 h post-infection, unless otherwise stated 712 in the corresponding Figure legends. For analysis of brain fungal burdens, animals were 713 714 euthanized and brains weighed, homogenized in PBS, and serially diluted before plating onto YPD agar supplemented with Penicillin/Streptomycin (Invitrogen). Colonies were counted after 715 716 incubation at 37°C for 24-48 h.

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718 Analysis of Brain Neutrophil Recruitment by FACS

Leukocytes were isolated from brains using previously described methods⁵⁴, resuspended in PBS and stained with Live/Dead fluorescent dye (Invitrogen) for 10 min on ice. Cells were then stained with fluorophore-conjugated antibodies in the presence of anti-CD16/32 and 0.5% BSA for 30 min on ice. Samples were washed in PBS/0.5% BSA/0.01% sodium azide and acquired using the BD Fortessa instrument equipped with BD FACS Diva software (BD Biosciences). FlowJo (TreeStar) was used for the final analysis. Anti-mouse antibodies used in this study were: CD45 (30-F11), CD11b (M1/70), both from eBiosciences, and Ly6G (1A8), Ly6C (AL21), both from BD Biosciences.

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728 Histology

Brains were removed from infected mice at the indicated time points and fixed in 10% formalin
for 24 h before embedding in paraffin wax. Tissue sections were stained with periodic acidSchiff (PAS).

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733 Measurement of Cytokines and Chemokines in Brain Homogenates

⁷³⁴ Infected brains were isolated at 24 h post-infection and homogenized in 1 mL PBS

supplemented with 0.05% Tween20 and protease inhibitor cocktail (Roche). Homogenized

brains were centrifuged twice to remove debris and resulting supernatants snap-frozen on dry

ice and stored at -80 °C prior to analysis. IL-1 β and CXCL1 concentrations in the

homogenates was determined by ELISA (R&D Systems), following the manufacturers'

instructions.

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741 Ex Vivo Restimulations and Intracellular FACS Analysis

Animals were infected with 2 × 10⁵ CFU of the indicated *C. albicans* strains intravenously, and brain leukocytes isolated 24 h later. For these experiments, brains were first digested in RPMI supplemented with 0.8 mg/mL Dispase (Gibco), 0.2mg/mL Collagenase Type 4 (Worthington), and 0.1 mg/mL DNAse (Roche) at 37 °C for 30 min, then pipetted vigorously to create a homogenous suspension. These suspensions were centrifuged (1500 rpm, 5 min, 4 °C), pellets resuspended in 5 mL 40% Percoll (GE Healthcare) and centrifuged again at 1700 rpm 748 for 20 min at 4 °C to remove myelin. Cell pellets were washed in RPMI supplemented with 749 10% heat-inactivated fetal bovine serum and Penicillin/Streptomycin (Invitrogen), and added to 750 FACS tubes for stimulations. Cells were incubated for 4 h at 37 °C in the presence of 5 µg/mL 751 Brefeldin A (Sigma) and, where indicated, 1 µg/mL LPS (Sigma) or 62.5 µg/mL depleted 752 zymosan (Sigma). After stimulation, cells were washed in PBS and stained for surface markers 753 as above. Fixation/permeabilization was performed with the eBioscience Foxp3 staining kit, 754 and staining for CXCL1 (IC4532R, from R&D Systems) or pro-IL-1 β (NJTEN3; from 755 eBioscience) performed overnight at 4 °C. Samples were washed once in PBS/0.5% 756 BSA/0.01% sodium azide prior to acquisition using the BD Fortessa instrument equipped with 757 BD FACS Diva software (BD Biosciences). FlowJo (TreeStar) was used for the final analysis. 758 CXCL1⁺ and pro-IL-1 β ⁺ cells were determined by employing similar staining and gating in animals deficient in these mediators ($Cxc/1^{-/-}$, $ll1b^{-/-}$) as negative controls. 759

760

761 Cell Culture and Candidalysin Stimulations

762 BV-2 cells were kindly provided by F. Crews (University of North Carolina School of Medicine). 763 C8-D1A astrocytes were kindly provided by D. McGavern (NINDS, NIH). BV-2 were 764 maintained at 37 °C, 5% CO₂ in RPMI supplemented with L-glutamine and HEPES (pH 7.0 – 765 7.4; Corning), 10% heat-inactivated fetal bovine serum and Penicillin/Streptomycin 766 (Invitrogen). DMEM media was used as the base media for C8-D1A culture, with the same 767 supplements as listed above and cultured as for BV-2. For BV-2 single culture experiments, 768 cells were lifted using cell scrapers and seeded into 24 well-plates at 5×10^5 cells/well (BV-2) 769 and left to adhere for 2 h at 37°C with either: 50 ng/mL LPS (Sigma), T-5224 (APExBIO) 770 and/or SB203580 (Adipogen); see Figure legends for details of each experiment. After 2 h.

recombinant Candidalysin peptide (Peptide Protein Research) was added to the cells at the indicated concentrations and incubation at 37 °C continued. For co-culture experiments, 3 × 10^5 C8-D1A were added to each well of a 24 well-plate and incubated overnight at 37 °C. BV-2 cells and Candidalysin were added as described above. In both types of experiments, supernatants or cells were collected at the indicated time points after Candidalysin addition and analyzed for IL-1 β and CXCL1 by ELISA (R&D Systems), CXCL1 staining by intracellular flow cytometry, or by immunoblot.

778

779 Immunoblot Analysis

Whole cell lysates were suspended in RIPA buffer containing protease and phosphatase
inhibitors (Thermo Scientific). Lysates were separated in SDS-PAGE and transferred to a
nitrocellulose membrane, 0.2 μm (Bio-Rad Laboratories). The membrane was incubated with
the following primary antibodies: phospho-MPK1/MPK2 polyclonal [Ser296, Ser318] (Thermo
Scientific) and c-Fos (Cell Signaling), IL-1β [3A6] (Cell Signaling), Caspase-1 p20 [Casper-1]
(Adipogen Life Sciences) (Thermo Scientific), NLRP3 [D4D8T] (Cell Signaling). Normalization
was performed by probing the membrane with β-Actin antibody (Cell Signaling).

Chemiluminescence detection was performed with Clarity[™] Western ECL Substrate (Bio-Rad
Laboratories), using the ChemiDoc[™] MP Imaging System (Bio-Rad).

789

790 FACS/MACS Sorting of Microglia

791 Wild-type animals were infected with 1.3×10^5 CFU SC5314 and euthanized at 24 h post-

⁷⁹² infection. Brains were digested as above and leukocytes stained with sterile antibodies⁵⁵.

793 Ly6C^{hi} monocytes (CD45^{hi} CD11b⁺ Ly6C^{hi} Ly6G⁻) and microglia (CD45^{lo} CD11b⁺ Ly6G⁻ Ly6C⁻)

794 were FACS-sorted into sterile sorting buffer (HBSS supplemented with 2 mM EDTA, 10 % 795 FCS, 100 U/mL penicillin, 100 µg/mL streptomycin) using a FACS Aria instrument for downstream gRT-PCR and immunoblot analyses. Purity of cells were greater than >95%, on 796 797 average. In some experiments (qRT-PCR of CLRs in brain-resident microglia; **Supplementary** 798 Fig. 1), microglia were instead sorted by magnetic separation using anti-CD11b microbeads (Miltenyi). Cells were then centrifuged (1500 rpm 5 min, 4 °C) and resuspended in Trizol for 799 800 RNA purification or RIPA buffer for downstream immunoblot analysis. Depending on the 801 experiment, up to 5 animals were pooled for individual sorts, or individual mice were analyzed 802 separately (see Figure legends for details).

803

804 Generation of cDNA and qRT-PCR

805 RNA was extracted from sorted brain myeloid cells (defined using the gating strategy shown in 806 Supplementary Fig. 2) using Trizol (Invitrogen) and the RNeasy kit (Qiagen) per the 807 manufacturer's protocol. Purified RNA was used as a template for cDNA generation using the 808 qScript cDNA SuperMix kit (Quanta Biosciences) with oligodT and random primers. Quantitative PCR was performed by TagMan detection (PerfeCTa qPCR FastMix ROX; 809 810 Quanta BioSciences) with the 7900HT Fast Real-Time PCR System (Applied Biosystems). All 811 gPCR assays were performed in duplicate and the relative gene expression of each gene was determined after normalization with GAPDH transcript levels using the $\Delta\Delta$ CT method. TagMan 812 primers/probes (Clec7a, Clec4n, Clec4d, Clec4e, II1b, Card9, Gapdh) were predesigned by 813 814 Applied Biosystems.

815

816 Statistics

817	Statistical analyses were performed using GraphPad Prism 7.0 software. Details of individual
818	tests are included in the figure legends. In general, data was tested for normal distribution by
819	Kolmogorov-Smirnov normality test and analyzed accordingly by unpaired t-tests or Mann
820	Whitney U-test. In cases where multiple data sets were analyzed, two-way ANOVA was used
821	with Bonferroni correction. In all cases, <i>P</i> values <0.05 were considered significant.
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843 Figure Legends

845 Fig. 1: CARD9-coupled C-type lectin receptors functionally compensate for one another for protective neutrophil recruitment to the fungal-infected brain. a. Card9-/- mice (n=4 846 847 animals) and their wild-type controls (n=4 animals) were intravenously infected with C. albicans 848 SC5314 and analyzed for neutrophil counts by flow cytometry at 24 h post-infection (left; dose: 1.3×10^5 CFU) and fungal growth within the brain at 72 h post-infection (right; dose: 7 × 10⁴ 849 850 CFU). **b.** Animals of the indicated genotype (WT n = 9 animals, $Clec7a^{-l-}$ n = 6 animals; WT n 851 = 12 animals, $Clec4n^{-/-}$ n = 10 animals; WT n = 6 animals, $Clec4d^{-/-}$ n = 6 animals; WT n = 6 852 animals, $Clec4e^{-/-}$ n = 8 animals; WT n = 10 animals, $Clec7a^{-/-}Fcer1g^{-/-}$ n = 9 animals) were intravenously infected with C. albicans SC5314 (2 × 10⁵ CFU for Clec4d^{-/-} and Clec7a^{-/-} 853 *Fcer1g*^{-/-} and their controls; 1.3×10^5 all others) and analyzed for neutrophil counts by flow 854 cytometry at 24 h post-infection and **c**, fungal burdens in the brain at 24 and 72 h post-infection 855 (WT n = 9 animals, $Clec7a^{-/-}$ n = 6 animals; WT n = 10 animals, $Clec4n^{-/-}$ n = 10 animals; WT 856 n = 10 animals, *Clec4d*^{-/-} n = 10 animals; WT n = 6 animals, *Clec4e*^{-/-} n = 4 animals; WT n = 857 858 10 animals, Clec7a^{-/-}Fcer1g^{-/-} n = 9 animals). **d**, Malt1^{-/-} mice and their littermate controls were 859 infected as above and analyzed for fungal burdens in the brain (right; WT n = 7 animals, *Malt1*- $^{-}$ n = 5 animals) and neutrophil recruitment to the brain at 24 h post-infection (left; WT n = 5 860 861 animals, *Malt1^{-/-}* n = 5 animals). In all cases, 'wild type' refers to appropriate matched control 862 animals for each knock-out line for gender, age and genetic background. Individual points 863 represent different mice. Data is pooled from 2 independent experiments and is shown as 864 mean +/- SEM, and analyzed by unpaired two-tailed t-test (panel a [left], b) or two-tailed Mann Whitney U-test (panel **a** [right], **c**, **d**). **P*<0.05, ***P*<0.01, ****P*<0.005, *****P*<0.001. 865

867	Fig. 2: IL-1 β and CXCL1 are critical for protective neutrophil recruitment to the fungal-
868	infected brain. a,b, Animals deficient in elements of the IL-1R signaling pathway or c,d,
869	chemokine receptors and their ligands, were infected and analyzed for neutrophil recruitment
870	at 24 h post-infection (a,c) and control of fungal brain infection (b,d) as in Fig. 1. 'Wild type'
871	refers to appropriate matched control animals for each knock-out line for gender, age and
872	genetic background. Individual points represent different mice; (a) WT n = 8 animals, <i>ll1r</i> ^{-/-} n =
873	8 animals; WT n = 7 animals, <i>Myd88^{-/-}</i> n = 6 animals; WT n = 12 animals, <i>II1a^{-/-}</i> n = 12
874	animals; WT n = 6 animals, $II1b^{-/-}$ n = 6 animals; WT n = 8 animals, $II1a^{-/-}II1b^{-/-}$ n = 6 animals.
875	(b) WT n = 8 animals, $II1r^{-}$ n = 7 animals; WT n = 7 animals, $Myd88^{-}$ n = 6 animals; WT n =
876	3-8 animals, $II1a^{-/-}$ n = 4-8 animals; WT n = 6 animals, $II1b^{-/-}$ n = 6 animals; WT n = 6 animals,
877	<i>ll1a⁻/⁻ll1b⁻/</i> ⁻ n = 6 animals. (c) WT n = 6 animals, <i>Ccr1⁻/</i> ⁻ n = 6 animals; WT n = 11 animals,
878	<i>Cxcr1</i> -/- n = 11 animals; WT n = 14 animals, <i>Cxcr2</i> -/- n = 16 animals; WT n = 8 animals, <i>Cxcl1</i> -
879	^{/-} n = 7 animals. (d) WT n = 6-7 animals, <i>Ccr1</i> ^{-/-} n = 6-7 animals; WT n = 7-10 animals, <i>Cxcr1</i> ^{-/-}
880	n = 7-10 animals; WT n = 8-10 animals, $Cxcr2^{-/-}$ n = 7-10 animals; WT n = 6-8 animals, $Cxcl1^{-}$
881	$^{-}$ n = 5-7 animals. Data is pooled from 2-3 independent experiments and shown as mean +/-
882	SEM, analyzed by unpaired two-tailed t-test (panel a, c) or two-tailed Mann Whitney U-test
883	(panel b, d). * <i>P</i> <0.05, ** <i>P</i> <0.01, *** <i>P</i> <0.005, **** <i>P</i> <0.001.

884

Fig. 3: Production of CXCL1 is dependent on IL-1β in the fungal-infected brain. a, Wild

type (n = 6/7 animals), $Cxcl1^{-/-}$ (n = 5 animals) and $ll1b^{-/-}$ (n = 6 animals) animals were infected

as in Fig. 1 and brains isolated at 24 h post-infection and analyzed for CXCL1 or IL-1 β

production by ELISA. Data is pooled from 2 independent experiments and analyzed by

889 unpaired two-tailed t-test. b, The relative proportions of myeloid cell populations (gated within 890 live CD45⁺ singlets) in the uninfected (n = 6 animals) and 24 h infected WT (n = 6 animals) 891 brain (left), and the relative proportion of myeloid cell populations producing CXCL1 (n = 3 892 animals) or pro-IL-1 β (n = 9 animals) in the 24 h infected brain (right). For the latter, total 893 CD45⁺CXCL1(or IL-1 β)⁺ cells were first gated and then cell types defined within this initial gate 894 using lineage markers (see below), using samples from the unstimulated condition. Data is 895 shown as the mean +/- SEM. c, Wild type (n = 3 animals) and $II1b^{-/-}$ mice (n = 4 animals) were 896 infected with 2 × 10⁵ C. albicans and brain cells analyzed for CXCL1 production by intracellular 897 flow cytometry 24 h later. Brain cells were restimulated ex vivo with 62.5 µg/mL depleted zymosan or 1 µg/mL LPS for 4 h in the presence of 5 µg/mL Brefeldin A. Representative plots 898 from the LPS-stimulated condition are gated on microglia (top; CD45^{int} Ly6G⁻ CD11b⁺), Ly6C^{hi} 899 900 monocytes (middle; CD45^{hi} Ly6C^{hi} Ly6G⁻ CD11b⁺) and neutrophils (bottom; CD45^{hi} Ly6C^{int} Ly6G^{hi} CD11b⁺), showing corresponding $Cxc/1^{-/-}$ cells as gating controls. In all panels, 'wild 901 902 type' refers to appropriate matched control animals for each knock-out line for gender, age and 903 genetic background. Individual points represent different mice. Data shown as mean +/- SEM, 904 and analyzed by unpaired two-tailed t-test. *P<0.05, **P<0.01.

905

906 Fig. 4: Candidalysin is a specific hyphal-secreted factor promoting neutrophil

907 **recruitment and control of fungal growth in the brain. a,b,c,e,** Animals were infected with 2

- $\times 10^5$ CFU of the indicated *C. albicans* strains (parental strains, closed symbols; deficient
- mutants, open symbols) and analyzed as in Fig. 1 for fungal burdens ($hgc1\Delta/\Delta$ n = 8 animals,

910 $hgc1\Delta/\Delta + HGC1 n = 7$ animals; BWP17 n = 10 animals, $ece1\Delta/\Delta n = 11$ animals, $ece1\Delta/\Delta + 10$

911 ECE1 n = 10 animals, $ece1\Delta/\Delta + ECE1_{\Delta 184-279}$ n = 11 animals; CAI4 + Clp10 n = 6 animals,

912 $sap4/5/6\Delta/\Delta$ + Clp10 n = 6 animals) and neutrophil recruitment (hqc1 Δ/Δ n = 10 animals). 913 $hgc1\Delta/\Delta + HGC1$ n = 10 animals; BWP17 n = 7 animals, $ece1\Delta/\Delta$ n = 7 animals, $ece1\Delta/\Delta +$ 914 ECE1 n = 7 animals, $ece1\Delta/\Delta + ECE1_{\Delta 184-279}$ n = 11 animals; CAI4 + Clp10 n = 6 animals, 915 $sap4/5/6\Delta/\Delta$ + Clp10 n = 7 animals). Histology shown in (c) is from 24 h post-infection, stained 916 with PAS. Scale bar is 50 µm. **d**, Whole brain homogenates from animals infected with 917 indicated strains were isolated at 24 h post-infection and analyzed for IL-1ß and CXCL1 using 918 ELISA (BWP17 n = 8-10 animals, $ece1\Delta/\Delta$ n = 8-10 animals, $ece1\Delta/\Delta$ + ECE1 n = 6-11 919 animals, $ece_1\Delta/\Delta + ECE_{1\Delta 184-279}$ n = 7-11 animals). Individual points represent different mice. 920 Data is pooled from 2-4 independent experiments and shown as mean +/- SEM, analyzed by 921 unpaired two-tailed t-test, or two-tailed Mann Whitney U-test (panel **a**, left). *P<0.05, **P<0.01, 922 ***P<0.005; ns = not significant.

923

924 Fig. 5: Microglia produce IL-1β and CXCL1 in a Candidalysin-dependent manner.

925 Animals were infected with wild-type *C. albicans* (BWP17; closed bars) or a Candidalysin-null

strain (*ece1* Δ / Δ ; open bars), and brain cells isolated 24 h later. Brain leukocytes were

927 restimulated as in Fig. 3, and intracellular staining for **a**, IL-1 β (unstimulated, n = 8 animals;

228 zymosan, n = 4 animals; LPS n = 6 animals) and **b**, CXCL1 (unstimulated, n = 12 animals;

29 zymosan, n = 6 animals; LPS n = 9 animals) was analyzed by flow cytometry. Box-and-whisker

930 plots show the minimum/maximum values (whiskers), the 25th/75th percentiles and the median.

931 Data is pooled from 2-4 independent experiments and analyzed by unpaired two-tailed t-tests.

P*<0.05, *P*<0.01. Representative staining is shown for LPS-stimulated microglia (gated as in

Fig. 3) from wild-type mice infected with indicated strains, or BWP17-infected cytokine-

934 deficient mutants as control.

936	Fig. 6: Candidalysin activates IL-1 β production from microglia via p38-cFos signaling
937	and promotes CXCL1 production from microglia through interactions with astrocytes.
938	BV-2 microglia were seeded into 24-well plates at 5 \times 10 ⁵ per well and left to adhere for 2 h in
939	the presence of 50 n g/mL LPS (for priming) before the addition of purified Candidalysin at the
940	indicated concentrations. Cell culture supernatants were analyzed for a , IL-1 β production or b ,
941	LDH release after 24 h of stimulation. Data is shown with the mean +/- SEM, individual points
942	represent individual culture wells (n = 4). c-d, In some experiments, BV-2 cells were co-
943	cultured with 3 × 10^5 C8-D1A astrocytes, and CXCL1 production analyzed in the supernatant
944	by ELISA (n = 10 individual culture wells) or by intracellular flow cytometry (n = 3-5 individual
945	culture wells; data shown with the mean). In d , microglia and astrocytes were distinguished by
946	CD45 staining, and CXCL1 production assessed within CD45 ⁺ (microglia) and CD45 ⁻
947	(astrocyte) gates. Histogram is gated on CD45 ⁺ microglia. e, To measure cFos and pMKP1/2
948	activation, BV-2 cells were stimulated with the indicated Candidalysin concentrations for 30 or
949	120 min and BV-2 cells then lysed and analyzed for cFos and pMKP1/2 by immunoblot,
950	normalizing to β -actin. Immunoblots shown are representative of 2 independent experiments. f ,
951	BV-2 cells were cultured in the presence of the indicated cFos and p38 inhibitors for 2 h prior
952	to stimulating with 20 μM Candidalysin, and IL-1 β measured in the supernatant by ELISA after
953	24 h (data shown as mean +/- SEM; n = 6 individual culture wells). All data is pooled from 2
954	independent experiments and analyzed by one-way ANOVA. * <i>P</i> <0.05, ** <i>P</i> <0.01, *** <i>P</i> <0.005,
955	**** <i>P</i> <0.0001.

957	Fig. 7: CARD9 is required for the production of IL-1 β , via regulation of pro-IL-1 β
958	transcription and inflammasome activation, and of CXCL1 in the fungal-infected brain.
959	a,b, <i>Card9</i> ^{+/+} (n = 13 animals) and <i>Card9</i> ^{-/-} (n = 13 animals) animals were infected with 2 ×
960	10 ⁵ CFU wild-type <i>C. albicans</i> (BWP17), and brain cells isolated 24 h later. Brain leukocytes
961	were restimulated as in Fig. 4, and intracellular staining for pro-IL-1 β and CXCL1 analyzed by
962	flow cytometry in total CD45 ⁺ cells (LPS-stimulated condition shown) (a) or microglia alone,
963	normalized to <i>Card9</i> ^{+/+} results (b). Panels a,b show pooled data from 4 independent
964	experiments, analyzed with two-tailed unpaired t-test. Data shown as mean +/- SEM (a) or with
965	minimum/maximum values (whiskers), the 25 th /75 th percentiles and the median (b). c,
966	Microglia were FACS-sorted from pooled <i>Card9</i> ^{+/+} (n = 4 animals) and <i>Card9</i> ^{-/-} animals (n = 4
967	animals) at 24 h post-infection and analyzed by unpaired two-tailed t-test for <i>II1b</i> expression by
968	qRT-PCR, or d,e, the indicated proteins by immunoblot (Caspase and IL-1 β blots; WT n = 6
969	animals, <i>Card9</i> ^{-/-} n = 7 animals; cFos blot; WT n = 10 animals, <i>Card9</i> ^{-/-} n = 10 animals; NLRP3
970	blot; WT n = 8 animals, <i>Card9^{-/-}</i> n = 8 animals;). Graphs in (d,e) represent the band pixel
971	density normalized to the wild type control, and are shown with mean +/- SEM and analyzed by
972	unpaired two-tailed student t-tests. Example blots are representative of 3 independent FACS
973	sorts/experiments; pooled data is shown in the graphs above. f , <i>NIrp3</i> ^{-/-} animals and their wild-
974	type controls were infected with 1.3×10^5 CFU <i>C. albicans</i> and analyzed by unpaired two-
975	tailed t-tests for neutrophil recruitment to the brain 24 h later (left; WT n = 9 animals, $NIrp3^{-/-}$ n
976	= 8 animals) and by two-tailed Mann-Whitney U-test for fungal brain burdens at 72 h post-
977	infection (right; WT n = 14 animals, <i>NIrp3^{-/-}</i> n = 14 animals), as described in Fig. 1. * <i>P</i> <0.05,
978	** <i>P</i> <0.01, *** <i>P</i> <0.005.

- 980 Fig. 8: CARD9 is required specifically in microglia for neutrophil recruitment and control
- 981 of fungal invasion in the CNS. a, *Card*9^{fl/fl}*Cx3cr1*^{CreER-/-} and *Card*9^{fl/fl}*Cx3cr1*^{CreER+/-} littermates
- 982 (n=8-13) were tamoxifen-pulsed at 4-5 weeks of age, left to rest for 4-6 weeks and then
- 983 infected with 1.3 × 10⁵ CFU *C. albicans* (SC5314) intravenously and analyzed for brain and
- kidney fungal burdens (*Card9*^{fl/fl}*Cx3cr1*^{CreER_/-} n = 8-10 animals; *Card9*^{fl/fl}*Cx3cr1*^{CreER+/-} n = 8
- animals), **b**, neutrophil recruitment to the brain at 24 h post-infection (*Card9*^{fl/fl}*Cx3cr1*^{CreER-/-} n =
- 986 13 animals; $Card9^{fl/fl}Cx3cr1^{CreER+/-}$ n = 9 animals), and **c**, intracellular staining for pro-IL-1 β and
- 987 CXCL1, as described in Fig. 3 (*Card9*^{fl/fl}*Cx3cr1*^{CreER_/-} n = 8 animals; *Card9*^{fl/fl}*Cx3cr1*^{CreER+/-} n =
- 888 8 animals). Data is pooled from 2-4 independent experiments and is shown as mean +/- SEM,
- analyzed by two-tailed Mann-Whitney U-tests (panel **a**) or two-tailed unpaired t-tests (panel **b**,
- 990 **c**). **P*<0.05, ***P*<0.01.