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DOI: 10.1101/2024.02.22.581301

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Document Version Other version

Citation for published version (Harvard): Onyishi, CU, Fejer, G, Mukhopadhyay, S, Gordon, S & May, RC 2024 'Loss of the scavenger receptor MARCO results in uncontrolled vomocytosis of fungi from macrophages' bioRxiv. https://doi.org/10.1101/2024.02.22.581301

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1 Loss of the scavenger receptor MARCO results in uncontrolled vomocytosis of fungi

2 from macrophages.

- 3
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- 6

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- 21

22 Abstract

23 Vomocytosis, also known as nonlytic exocytosis, is a process whereby fully phagocytosed 24 microbes are expelled from phagocytes without discernible damage to either the phagocyte or 25 microbe. Although this phenomenon was first described in the opportunistic fungal pathogen 26 Cryptococcus neoformans in 2006, to date, mechanistic studies have been hampered by an 27 inability to reliably stimulate or inhibit vomocytosis. Here we present the fortuitous discovery 28 that macrophages lacking the scavenger receptor MAcrophage Receptor with COllagenous 29 domain (MARCO), exhibit near-total vomocytosis of internalised cryptococci within a few 30 hours of infection. Our findings suggest that MARCO's role in modulating vomocytosis is 31 independent of its role as a phagocytic receptor and instead may be driven by variation in 32 cytoskeletal arrangement between wildtype and MARCO-deficient macrophages.

33

34 Introduction

35 *Cryptococcus neoformans* is an opportunist fungal pathogen that causes life-threatening

36 meningitis, mainly in immunocompromised individuals such as HIV/AIDS patients (1).

37 Infection is thought to begin with the inhalation of the fungi into the lungs where it

38 encounters macrophages of the innate immune system that serve as the first line of defence

39 against infection (1). The interaction between *C. neoformans* and macrophages can lead to a

40 range of outcomes including fungal survival and replication within macrophages (2,3), lateral

41 transfer of cryptococci between macrophages (4,5), lysis of macrophages (6) and

42 vomocytosis, also called nonlytic exocytosis (7,8).

43

44 Vomocytosis is a nonlytic expulsion mechanism where fully phagocytosed fungi are expelled

45 from the macrophage with no evidence of host cell damage (7–9). Vomocytosis occurs

46 through the fusion of *Cryptococcus*-containing phagosome with the plasma membrane in a

47 manner that is modulated by the actin cytoskeleton (10). It has also been shown to require

48 phagosome membrane permeabilization (10) and a failure to fully acidify the phagosome

49 (11,12). Previous studies have identified the mitogen-activated protein kinase ERK5 and the

- 50 phospholipid binding protein Annexin A2 as regulators of vomocytosis, with ERK5
- 51 inhibition and Annexin A2 deficiency leading to increased and decreased vomocytosis,

52 respectively (13,14). Moreover, stimulation of macrophages with type 1 interferons (IFNα

53 and IFNβ), mimicking viral coinfection, increased cryptococcal vomocytosis (15). Very little

54 else is known about host regulators of vomocytosis.

5	5
J	J

56 Here we present the chance observation that the scavenger receptor MAcrophage Receptor 57 with Collagenous structure (MARCO) is a key modulator of vomocytosis. Typically, 58 vomocytosis rates from wildtype macrophages are between 10-20%, but this number rises 59 close to 100% in *Marco^{-/-}* macrophages. Further investigation indicates that this impact on 60 vomocytosis is likely independent of MARCO's role in phagocytosis but may instead result 61 from the previously documented cytoskeletal dysfunction seen in *Marco^{-/-}* macrophages (16). 62 As well as providing a powerful experimental tool for the future investigations into this 63 phenomenon, this finding also has important implications for interpreting infection assays conducted in Marco-/- animals. 64 65

66 Results and Discussion

67 *Marco^{-/-}* macrophages show increased vomocytosis of non-opsonised *C. neoformans.*

68 While investigating the role of scavenger receptors in the phagocytosis of C. neoformans

69 using a non-transformed GM-CSF dependent alveolar-like macrophage cell line derived from

70 wildtype and *Marco^{-/-}* C57BL/6 mice (17), we observed that, in LPS-stimulated macrophages,

71 MARCO-deficiency led to decreased non-opsonic phagocytosis of *C. neoformans* (Figure

1A). Using live cell imaging to observe the interaction between *C. neoformans* and

73 macrophages, we noted a dramatic increase in the vomocytosis of *C. neoformans* from

74 *Marco^{-/-}* macrophages with 80% to 100% of infected macrophages experiencing at least one

vomocytosis event (Figure 1B; Supplementary Video 1). No other host or pathogen factor has

76 been found to increase the rate of vomocytosis to such an extent. Not only did Marco^{-/-}

77 macrophages show elevated vomocytosis, but 75% of nonlytic expulsion events occurred

78 within the first 1 h 40 mins of the initiation of the timelapse video, with the median time to a

vomocytosis event being 0.92 h (55 mins) (Figure 1C; Supplementary Video 2). In contrast,

80 vomocytosis in wildtype macrophages occurred over a wider range of time with the median

81 time to vomocytosis being 10.6 h (10 h:35 mins) (Figure 1C). Figure 1D provides a

82 representative image of non-opsonised *C. neoformans* being expelled from *Marco^{-/-}*

83 macrophages.

85 MARCO-deficiency leads to elevated vomocytosis of 18B7 antibody-opsonised C.

86 *neoformans* and yeast-locked *Candida albicans*, but not heat killed *C. neoformans* or

87 latex beads.

To investigate the generality of this phenomena, we infected macrophages with heat-killed C. 88 89 neoformans, 7 µm diameter latex beads, anti-GXM 18B7 antibody-opsonised C. neoformans, and a yeast-locked TetOn-NRG1 C. albicans strain that constitutively expresses Nrg1 90 91 transcription factor, thereby preventing yeast to hypha formation (18). In line with previous 92 data showing that inert particles do not undergo vomocytosis (7,8), we observed no vomocytosis of heat-killed C. neoformans by either wildtype or Marco^{-/-} macrophages out of 93 55 infected macrophages observed (Table 1), and only a single event (amongst 239 infected 94 95 cells) when macrophages were "infected" with latex beads (Table 2). Notably, Marco-/macrophages showed decreased phagocytosis of heat-killed cryptococci (Supplementary 96 97 Figure 1A) and latex beads (Supplementary Figure 1B) compared to wildtype cells. 98 99 Next, macrophages were infected with antibody-opsonised fungi to drive uptake via FcyRs. As expected, there was no difference in antibody-opsonised phagocytosis between wildtype 100 and Marco^{-/-} macrophages (Figure 2A). However, vomocytosis was elevated in Marco^{-/-} 101 102 macrophages compared to wildtype cells (Figure 2B). Finally, macrophages were infected 103 with a veast-locked C. albicans strain that fail to undergo filamentation and observed over a 6 104 h period. As expected, since phagocytosis of *Candida* is predominantly driven by Dectin-1 (19), there was no difference in phagocytosis between wildtype and *Marco^{-/-}* macrophages 105 106 (Figure 2C). Surprisingly, we observed increased vomocytosis of yeast-locked Candida from 107 MARCO-deficient macrophages (Figure 2D and E; Supplementary Video 3). The percentage 108 of *Marco^{-/-}* macrophages that experienced at least one vomocytosis event was not as dramatic 109 as that observed with *Cryptococcus*; however, vomocytosis of wildtype *C. albicans* is rare, happening at a rate of <1% over a 6-hour period (20). Therefore, a rate of 30% over 6 hours 110 in *Marco^{-/-}* cells is significant for this fungal pathogen. Given that elevated vomocytosis was 111 112 observed when phagocytosis was mediated by non-opsonic receptors (Figure 1B), FcyR 113 (Figure 2B) and Dectin-1 (Figure 2D), it seems likely that the role of MARCO in 114 vomocytosis is independent of the mechanism of uptake. 115

- 116

117 Treatment of wildtype MPI cells with inhibitors of MARCO does not phenocopy

118 increased vomocytosis seen in *Marco^{-/-}* cells.

- 119 To explore whether the vomocytosis phenotype seen in *Marco^{-/-}* can be induced in wildtype
- 120 macrophages, we exposed wildtype cells to polyguanylic acid potassium salt (polyG), a
- 121 MARCO ligand and inhibitor (21–23), and quantified vomocytosis. Although polyG pre-
- 122 treatment decreased the phagocytosis of non-opsonised C. neoformans (Figure 3A), unlike
- 123 genetic knockout of *MARCO*, the inhibition of MARCO using a ligand did not result in an
- 124 increase in vomocytosis (Figure 3B). Since polyG functions as a competitive inhibitor and
- 125 likely does not block MARCO-mediated downstream signalling, this suggests that the impact
- 126 of MARCO on vomocytosis can be mechanistically separated from its ligand-binding
- 127 activity.
- 128
- 129 Next, MARCO receptor on wildtype macrophages was blocked using increasing
- 130 concentrations of an anti-MARCO ED31 antibody. Anti-MARCO antibody reduced
- 131 MARCO-mediated phagocytosis in a dose-dependent manner (Figure 3C), without impacting
- 132 the rate of vomocytosis in wildtype macrophages (Figure 3D). According to the
- 133 manufacturers, the anti-MARCO ED31 antibody recognises the ligand binding domain of
- 134 MARCO receptors and can therefore compete for receptor binding with *C. neoformans*
- 135 without impacting intracellular MARCO signalling (24). Taken together, the role of MARCO
- in vomocytosis is most likely independent of its role in uptake, hence the inability of
- 137 inhibitors that act on the ligand binding site to phenocopy the genetic knock out of MARCO
- 138 receptor.
- 139

140 There is a noticeable difference in actin morphology wildtype and *Marco^{-/-}*

141 macrophages.

Granucci et al. (16) identified a role for MARCO in cytoskeletal remodelling of microglial and dendritic cells. Moreover, repeated actin polymerization and depolymerisation around phagosomes containing cryptococci leading to the formation of transient actin 'cages' has been shown to prevent vomocytosis (10). We therefore wondered whether the actin

146 cytoskeleton may be perturbed in MARCO-deficient macrophages.

- 147
- 148

149 Rhodamine-conjugated phalloidin staining of uninfected macrophages revealed wildtype 150 macrophages to be more compact than MARCO-deficient macrophages, which were larger 151 and with expansive ruffle-like structures (Figure 4A, white arrows; Supplementary Figure 2). 152 Similarly, in C. neoformans infected macrophages, wildtype macrophages appeared more 153 rounded and had well-formed filopodial protrusions (Figure 4B; yellow arrows). Though 154 *Marco^{-/-}* cells also had instances of filopodial protrusions from the cell periphery (yellow 155 arrows), these macrophages appeared larger, were less organised and had extensive ruffles 156 (Figure 4B, white arrows). Taken together, there is a clear difference in actin organisation between wildtype and Marco^{-/-} cells. It is thus possible that MARCO's role in actin 157 158 remodelling and organisation is one reason for the elevated vomocytosis seen in Marco^{-/-} 159 macrophages. 160 161 Our observation raises a number of questions for future investigation. Firstly, given that loss

162 of MARCO leads to elevated vomocytosis, then one role for MARCO may be to sense

163 phagosomal content and prevent premature expulsion, potentially by regulating the formation

164 of actin 'cages' the have been show to block phagosome fusion with the plasma membrane

165 (10). It is also possible that MARCO activity is linked to the MAPK ERK5, since ERK5

166 activity has been implicated in disruptions in actin cytoskeleton during oncogenic

167 transformation (25,26) and is known to modulate vomocytosis (13). Additionally, MARCO

168 may be upstream of Annexin A2, another host signalling molecule found to modulate

169 vomocytosis (14). Annexin A2 plays a significant role in a range of cellular processes

170 including exocytosis and binding to actin to modulate cytoskeleton arrangement (27,28),

171 processes that have been linked to nonlytic expulsion. Finally, we note that this hitherto

172 undocumented impact of MARCO loss on pathogen expulsion will be important for

- investigators to consider when using *Marco^{-/-}* cells or animals for a range of other infection
 assays.
- 175

176 Conclusion

177 Here we present a novel role for MARCO in modulating the vomocytosis of *C. neoformans*.

178 The increase in vomocytosis observed in *Marco^{-/-}* macrophages is the most dramatic change

in vomocytosis rate observed to date. Increased vomocytosis in *Marco^{-/-}* macrophage was also

180 observed when macrophages were infected with a yeast-locked *C. albicans* strain, suggesting

181 that MARCO's modulation of vomocytosis is a broadly relevant phenomenon. Given that

182 MARCO-deficiency still resulted in elevated vomocytosis of antibody-opsonised *C*.

183 *neoformans* and *C. albicans*, we propose that MARCO's role in vomocytosis is independent

184 of the mode of uptake, and instead that MARCO may modulate vomocytosis through its role

185 in actin remodelling. We hope this finding will inspire new research aimed at understanding

186 the mechanism and clinical consequence of vomocytosis during host-pathogen interaction.

- 187
- 188

189 Materials & Methods

190 Max Plank Institute (MPI) Cell Culture

- 191 Max Plank Institute (MPI) cells are a non-transformed, granulocyte-macrophage colony-
- 192 stimulating factor (GM-CSF)-dependent murine macrophage cell line that is functionally
- 193 similar to alveolar macrophages (17,29). MPI cell lines isolated from wildtype and
- 194 MAcrophage Receptor with COllagenous structure knockout (*Marco^{-/-}*) mice were cultured in
- 195 Roswell Park Memorial Institute (RPMI) 1640 medium [ThermoFisher] supplemented with
- 196 10% heat inactivated FBS [Sigma-Aldrich], 2 mM L-glutamine [Sigma-Aldrich], and 1%
- 197 Penicillin and Streptomycin solution [Sigma-Aldrich] at 37°C and 5% CO₂. Each flask was
- 198 further supplemented with 1% vol/vol GM-CSF conditioned RPMI media prepared using a
- 199 X-63-GMCSF cell line.
- 200

201 Phagocytosis Assay

Twenty-four hours before the start of the phagocytosis assay, MPI cells were seeded onto 24well plates at a density of $2x10^5$ cells/mL in complete culture media supplemented with 1%

- 204 vol/vol GM-CSF. The cells were then incubated overnight at 37°C and 5% CO₂. The
- 205 following day, macrophages were stimulated with 10 ng/mL lipopolysaccharide (LPS) from
- 206 Escherichia coli [Sigma-Aldrich; Cat#: L6529] and 1% vol/vol GM-CSF for 24 h. At the
- same time, an overnight culture of *Cryptococcus neoformans* var. *grubii* KN99α strain, that
- 208 had previously been biolistically transformed to express green fluorescent protein (GFP)(30),
- 209 was set up by picking a fungal colony from YPD agar plates (50 g/L YPD broth powder
- 210 [Sigma-Aldrich], 2% Agar [MP Biomedical]) and resuspending in 3 mL liquid YPD broth
- 211 (50 g/L YPD broth powder [Sigma-Aldrich]). The culture was then incubated at 25°C
- 212 overnight under constant rotation (20rpm).
- 213
- 214 After overnight LPS stimulation, macrophages were infected with non-opsonised C.
- 215 neoformans. To prepare C. neoformans for infection, an overnight C. neoformans culture was

216 washed two times in 1X PBS, counted using a haemocytometer, and fungi incubated with 217 macrophages at a multiplicity of infection (MOI) of 10:1. The infection was allowed to 218 proceed for 2 h at 37° C and 5% CO₂. Where applicable, macrophages were pre-treated with 219 400 ug/mL polyguanylic acid potassium salt (polyG) [Sigma-Aldrich; Cat#: P4404], rat anti-220 mouse MARCO ED31 clone monoclonal antibody [BioRad; Cat#: MCA1849], or anti-rat 221 IgG1 isotype control [Invitrogen; Ca#: 14430182] for 30 mins at 37°C prior to infected with 222 non-opsonised C. neoformans. In these cases, infection was carried out still in the presence of 223 polyG ligand or antibodies.

224

For infection with antibody-opsonised C. *neoformans*, 1×10^6 yeast cells in 100 µL PBS were

226 opsonized for 1 h using 10 µg/mL anti-capsular 18B7 antibody (a kind gift from Arturo

227 Casadevall, Albert Einstein College of Medicine, New York, NY, USA). For infection with

heat-killed C. neoformans, fungi were killed by heating at 56°C for 30 mins. After 2 h

229 infection at 37°C, macrophages were washed 4 times with PBS to remove as much

230 extracellular *C. neoformans* as possible.

231

To explore vomocytosis of *Candida albicans*, a yeast-locked *C. albicans* strain was used (a kind gift from Hung-Ji Tsai, University of Birmingham, Birmingham, United Kingdom). The yeast-locked TetOn-NRG1 *C. albicans* strain constitutively express the Nrg1 transcription factor, thereby preventing yeast to hypha transition (18). A colony was suspended in 10 mL YPD broth and incubated overnight at 30°C and 180 rpm. Prior to their use in infection, a *C. albicans* overnight culture was diluted 1:100 in fresh YPD broth, then incubated at 30°C and

238 180 rpm for 3 h till cells were in exponential phase. Cells in exponential phase were washed

with PBS, counted and macrophages were infected at MOI 2:1.

240

241 Time-lapse Imaging

After infection, extracellular fungi were removed and fresh media containing 1% vol/vol

243 GM-CSF (with or without relevant inhibitor) was added back into the wells. For infection

244 with *C. albicans*, imaging began immediately following infection. Live-cell imaging was

245 performed using a Zeiss Axio Observer [Zeiss Microscopy] or Nikon Eclipse Ti [Nikon] at

246 20X magnification. Images were acquired every 5 mins for 16 hours at 37°C and 5% CO₂.

- 247 The resulting videos were analysed using Fiji [ImageJ], at least 200 macrophages were
- 248 observed, and vomocytosis was scored according to the following guideline:

249	1.	One vomocytosis event is the expulsion of internalized cryptococci from an infected
250		macrophage, regardless of the number of cryptococci expelled if they do so
251		simultaneously.

- 252 2. Vomocytosis events are scored as independent phenomena if they occur in different253 frames or from different macrophages.
- Vomocytosis events are discounted if the host macrophage subsequently undergoes
 lysis or apoptosis within 30 min.
- 256

257 F-Actin Staining and Confocal Microscopy

258 Macrophages were seeded on 13 mm cover slips placed into 24-well plates. Staining was 259 performed on macrophages fixed with 4% paraformaldehyde for 10 mins at room 260 temperature and permeabilised with 0.1% Triton X-100 diluted in PBS for 10 mins at room 261 temperature. F-actin filaments were stained using 2 units of rhodamine-conjugated phalloidin 262 stain [Invitrogen; Cat#: R415] diluted in 400 µl 1% BSA in 1X PBS and incubated for 20 263 mins at room temperature. Cells were washed with PBS, then counter stained with 0.5 µg/mL 264 DAPI for 5 mins at room temperature to visualize the nucleus. After PBS washes, glass slides 265 were mounted using Fluoromount mounting medium [Sigma; Cat#: F4680]. Z-stack images 266 were acquired using the Zeiss LSM880 Confocal with Airyscan2, laser lines 405, 488, 561 267 and 640 nm, and at 63X oil magnification. Image acquisition was performed using the ZEN Black software [Zeiss Microscopy] and the resulting images were analysed using the Fiji 268 269 image processing software [ImageJ].

270

271 Statistics

272 GraphPad Prism Version 9 for Mac (GraphPad Software, San Diego, CA) was used to

273 generate graphical representations of experimental data. Violin plots were generated using R

274 programming. Inferential statistical tests were performed using Prism. The data sets were

- assumed to be normally distributed based on results of Shapiro-Wilk test for normality.
- 276 Consequently, to compare the means between treatments, the following parametric tests were
- 277 performed: unpaired two sided t-test, one-way ANOVA followed by Tukey's post-hoc test.
- 278 When data failed the normality test, Mann-Whitney U nonparametric test was used. Variation

between treatments was considered statistically significant if p-value < 0.05.

280

281 Acknowledgments

- 282 We thank Hung-Ji Tsai for the provision of yeast-locked C. albicans strain. C.U.O is
- supported by a PhD studentship from the Darwin Trust of Edinburgh. R.C.M. gratefully
- acknowledges support from the BBSRC and European Research Council Consolidator
- Award.
- 286

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373 Figures and Tables



374

375 Figure 1: Macrophages were stimulated overnight with LPS then infected with non-opsonised 376 Cryptococcus neoformans. (A) Phagocytosis was quantified as the number of internalised cryptococci 377 per 100 macrophages. (B) Vomocytosis was quantified over a 16 h period and presented as the 378 percentage of infected macrophages that experienced one or more vomocytosis events. At least 200 379 macrophages were observed. Data is presented as mean ± SEM; ***p<0.001, ****p<0.0001 in an 380 unpaired two sided t-test. (C) The time at which individual vomocytosis events took place was quantified and expressed as decimals. Wildtype (n=32); Marco^{-/-} (n=56). A violin plot with an 381 382 overlapping box plot was created using the ggplot2 package on R; ****p<0.0001 in a Mann-Whitney 383 test. (D) Representative image showing vomocytosis of GFP-expressing C. neoformans from Marco^{-/-} 384 macrophages. Time is presented in hh:mm:ss; white and red arrows follow the course of expulsion 385 events. Data is representative of three independent experiments. 386 387 388

390 Table 1: Quantification of the vomocytosis of heat-killed *C. neoformans* in wildtype and

391 *Marco^{-/-}* macrophages.

	# infected macrophages counted	# vomocytosis events observed
Wildtype	39	0
Marco ^{-/-}	16	0

392

393

394 Table 2: Quantification of the vomocytosis of latex beads in wildtype and Marco-/-

395 macrophages.

	# infected macrophages counted	# vomocytosis events observed
Wildtype	203	1
Marco ^{-/-}	36	0

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398

399 Figure 2: (A, B) Wildtype and Marco^{-/-} macrophages were stimulated with 10 ng/mL LPS overnight, 400 then infected with anti-GXM 18B7 antibody opsonised C. neoformans. After 2 h infection, images 401 were acquired every 5 mins for 16 h. (C, D) LPS stimulated macrophages were infected with a yeast-402 locked TetO-NRG1 C. albicans strain that constitutively expresses Nrg1 transcription factor, thereby 403 preventing yeast to hyphae transition. Images were acquired every 5 mins for 6 h. (A, C) Phagocytic 404 index (%) represents the percentage of macrophages that phagocytosed one or more fungal cells. (B, 405 **D**) Vomocytosis (%) is the percentage of infected macrophages that experienced at least one 406 expulsion events. At least 200 macrophages were observed per condition. Data is representative of 407 two independent experiments. Data shown as mean \pm SEM; ns, not significant; *p<0.05; 408 ****p<0.0001 in a t-test. (E) Representative image showing vomocytosis of C. albicans from Marco ^{/-} MPI cells. Time is presented in hh:mm:ss; red arrows follow the course of a vomocytosis event; 409 410 scale bar = $10 \mu m$.



411

412 **Figure 3:** Wildtype macrophages were stimulated overnight with 10 ng/mL LPS. The following day,

413 cells were pre-treated with polyguanylic acid (polyG) (**A**, **B**) or anti-MARCO ED31 monoclonal

414 antibody (mAb) for 30 mins (**C**, **D**) then infected with *C. neoformans* still in the presence of polyG or

anti-MARCO mAb. Images were acquired every 5 mins for 16 h. (A, C) The number of internalised
fungi at the beginning of the timelapse video was quantified. (B, D) Vomocytosis (%) is the

417 percentage of infected macrophages that experienced at least one vomocytosis events. At least 200

418 macrophages were quantified per condition. Data is representative of two independent experiments.

419 Data shown as mean \pm SEM; ns, not significant; *, p<0.5; **p<0.01 in an unpaired two-sided t-test

420 (**A**, **B**) and a one-way ANOVA followed by Tukey's post-hoc test (**C**, **D**).



422

423 Figure 4: Wildtype and Marco^{-/-} macrophages were stimulated with 10 ng/mL LPS overnight, left 424 uninfected (A) or infected with GFP expressing C. neoformans (B). Prior to confocal microscopy 425 imaging, macrophages were fixed, permeabilised and F-actin was stained with rhodamine-conjugated 426 phalloidin. Cells were counter-stained with DAPI to visualize the nucleus, then mounted onto glass 427 slides using Fluoromount mounting medium. Z-stack images were acquired using the Zeiss LSM880 428 using 63X Oil magnification. Z-stack maximum intensity projection was applied onto the images. Red 429 = F-actin (Phalloidin); Blue = Nucleus; Green with white dashed circle = C. neoformans. White 430 arrows show examples of macrophages with ruffle-like structures; Yellow arrows show examples of 431 filopodial protrusions. Scale bar = $10 \ \mu m$

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- 435 Supplementary Figure 1: Wildtype and *Marco^{-/-}* macrophages were infected with (A) *C. neoformans*
- 436 killed by heating at 56°C for 30 mins or (B) 7 µm latex beads. After 2 h infection, images were
- 437 acquired every 5 mins for 16 h. The number of internalised cryptococci or latex bead per 100
- 438 macrophages was quantified. At least 200 macrophages were observed per condition. Data shown as
- 439 mean \pm SEM; *p<0.05; **p<0.01 in a t-test. Data is representative of two independent experiments.
- 440



441

442 **Supplementary Figure 2:** Wildtype and *Marco^{-/-}* macrophages were stimulated with 10 ng/mL LPS

443 overnight, then stained with rhodamine-conjugated phalloidin to visualize F-actin distribution. Z-stack

444 images were acquired on the Zeiss LSM880 using 100X Oil magnification. Z-stack maximum

- intensity projection was applied onto the images. Red = F-actin (Phalloidin); Blue = Nucleus. Scale
- 446 bar = $10 \ \mu m$

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448	Supplementary Video 1: Representative video showing vomocytosis of C. neoformans from Marco ^{-/-}
449	macrophages. Video corresponds to Figure 1D. Time shown as: hh:mm:ss

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- 451 Supplementary Video 2: Example video showing rapid time-to-vomocytosis in Marco^{-/-}

452 macrophages. Time shown as: hh:mm:ss

- 454 Supplementary Video 3: Representative video showing vomocytosis of C. albicans from Marco^{-/-}
- 455 macrophages. Video corresponds to Figure 2E. Time shown as: hh:mm:ss
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