

Extrathymic expression of Aire controls the induction of effective TH17 cell-mediated immune response to Candida albicans

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Extended Data Fig. 1	Extended Data Fig. 1	EF1.pdf	a-c) GO enrichment analysis of upregulated differentially expressed genes from unstimulated vs. <i>C. albicans</i> - stimulated ILC3 subsets including cILC3s (related to Figure 2a) (a); MHCII ⁺ ILC3s (related to Figure 2b) (b); Aire ⁺ ILC3s (related to Figure 2c) (c). d) Heatmap of Pearson correlation according to gene expression values between individual samples analyzed in Fig. 2a-c.
Extended Data Fig. 2	Extended Data Fig. 2	EF2.pdf	a) Imaging flow cytometry analyzing the physical interaction between HKCA with DCs or with Aire ⁺ ILC3s. CD11c ⁺ DCs or Lineage negative cells and were isolated from mouse popliteal lymph nodes using MACS-beads depletion and then incubated with CPD-stained HKCA for 90 minutes. Samples were stained for Aire, MHCII, CD11c, Lamp1 (lysosomal marker) and DAPI and analyzed by Imaging flow cytometry. In both, Aire ⁺ ILC3 cells (Aire ⁺) and DCs (CD11c ⁺) HKCA associate with lysosomes (Lamp1). Representative images out of five independent repetitions of the experiment are shown. (b) Comparison of the capacity of different immune subsets (B cells, Ly6C ⁺ MNPs, CD11b ⁺ MNPs, CD11c ⁺ MNPs and Aire ⁺ ILC3s) to endocytose CPD-labeled HKCA in an invitro endocytosis assay. Subsets were FACS-sorted and incubated with HKCA-CPD for three hours. The frequency of cells that have successfully internalized HKCA-CPD was then assessed by flow cytometry (mean \pm SD; n=3). c-d) Comparison of the capacity of either wild-type or Galectin-3/Dectin-1-double knockout (dKO) Aire+ ILC3s (c) or DCs (d) to endocytose CPD-labeled HKCA at 37°C or at 4°C. Corresponding populations were isolated from WT or dKO mice and kept either at 37°C or at 4°C. The internalization of HKCA-CPD by the cells was measured by FACS (mean \pm SD; n=3). e) Representative gating

			strategy of APC populations related to Fig. 3c-g. f) FACS analysis of in vitro assay related to Fig. 3c. g) Graphical summary of an experimental setting relevant for data shown in h. Wild-type mice were intravenously stimulated HKCA in specified timepoints and APC population were isolated using gating strategy depicted in e. h) FACS analysis of in vitro assay related to Fig. 3d-f assessing the antigen presentation of the corresponding cell subsets.
Extended Data Fig. 3	Extended Data Fig. 3	EF3.pdf	a) FACS analysis validating that Aire protein expression in the entire LN- resident cell compartment is exclusively restricted to a CD45 positive, lineage negative (Lin 1: TCR- β , CD19, Gr-1, F4/80, CD11b, CD11c, NK1.1; Lin 2: CD3, B220), NKp46-negative, MHCII positive, Roryt positive cell subset – previously identified as Aire ⁺ ILC3s. b) Back-gating FACS analysis of Aire- expressing cells identified in a. c) Representative flow cytometry dot plots showing the frequencies of transferred CD45.1 ⁺ OT-II T cell (red gate) vs. CD45.1/CD45.2 double positive control T cell (blue gate) populations 2 days after the transfer. d) Statistical analysis of ratios related to b (n=5 per group, mean \pm SD, two-tailed Student's t-test) of OT-II vs. control T two days after the transfer. e-f) Survival curves of WT (Aire ^{+/+}) and knockout (Aire ^{-/-}) mice (n \geq 10 mice per group) on either on NOD (e) and C57Bl/1 (f) genetic background after systemic challenge with live <i>C. albicans</i> . Mice were <i>i.v.</i> injected every second day by heat-killed <i>C. albicans</i> (HKCA) for the duration of three weeks. Subsequently, mice were infected by alive <i>C. albicans</i> and monitored for survival. Long-rank (Mantel-Cox) test was used to calculate the indicated p-value. g-h) Quantitative PCR analysis assessing the presence of <i>C.</i> <i>albicans</i> -specific DNA in the liver (g) and small intestine (h) from <i>Rorc</i> -Cre ⁻ Aire ^{fl/fl} (WT) and <i>Rorc</i> -Cre ⁺ Aire ^{fl/fl} (ILC3 ^{ΔAire}) mice (n=6, mean \pm SD, two- tailed Student's t-test). i-j) ELISA assessing the amount of IL-17 (i) or (IL-22) (j) autoantibodies in the sera of 8-week old untreated Aire ^{+/+} vs. Aire ^{-/-} mice on NOD and C57Bl/6 (B6) genetic background (n=6 per group, two-tailed Student's t-test). Data are shown as mean of optical density \pm SD. Data are

			shown as mean of optical density ± SD. P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.001, * = p-value < 0.05, ns = not significant.
Extended Data Fig. 4	Extended Data Fig. 4	EF4.pdf	 a) Experimental outline relevant to data shown in Fig 6a, b: Wild-type mice were orally colonized by <i>C. albicans</i> and analyzed in indicated time points. b) Representative FACS gating strategy of Als1-tet⁺ T cells. c) Experimental outline relevant to data shown in (d) and (e): WT (Aire^{+/+}) and Aire^{-/-} were orally colonized by <i>C. albicans</i> and analyzed after two weeks. d) Representative FACS plot of Als1-tet⁺ T cells. Cells were isolated from pLNs and spleens of mice described in (c). Counts of Als1-tet⁺ cells are highlighted in red rectangles (left panel). Statistical analysis of the same representative experiment showing the total counts (mean ± SD, two-tailed Student's t-test, n=6). Representative experiment is shown. e) Quantitative PCR analysis assessing the presence of <i>C. albicans</i>-specific DNA in the ileal part of small intestine from WT and Aire-deficient mice (n=5, mean ± SD, two-tailed Student's t-test). f) Experimental outline relevant to data shown in (g) and (h). Bone-marrow (BM) chimeras restricting Aire expression either to hematopoietic (Aire^{+/+} BM → Aire^{-/-}) or stromal compartment (Aire^{-/-} BM → Aire^{+/+}) were generated by reciprocal BM transfer to recipient mice after 900 rad whole-body irradiation. Six weeks after the BM transfer, the mice were orally colonized by <i>C. albicans</i> and analyzed after two weeks. g) Representative FACS plot of Als1-tet⁺ T cells. Counts of Als1-tet⁺ cells are highlighted in red rectangles (left panel). Statistical analysis of the same representative experiment showing the total counts (n=6, mean ± SD, two-tailed Student's t-test). Representative experiment is shown. h) Quantitative PCR analysis assessing the presence of <i>C. albicans</i>-specific DNA in the ileal part of small intestine from reciprocal bone-marrow chimeras (n=6, mean ± SD, two-tailed Student's t-test). i) Experimental outline relevant to data shown in j and k. CD90-disperate chimeras were created by adoptive transfer of T cells and B-lymphocytes from CD90.1 mice to Rag1^{-/}

			control antibody prior the <i>C. albicans</i> oral colonization and then each third day and analyzed after two weeks. j) Representative FACS plot of Als1-tet ⁺ T cells. Counts of tetramer positive cells are highlighted in red rectangles (left panel). Statistical analysis of the same representative experiment showing the total counts (n=6, mean \pm SD, two-tailed Student's t-test). Representative experiment is shown. k) Quantitative PCR analysis assessing the presence of <i>C. albicans</i> -specific DNA in the ileal part of small intestine from CD90- disparate chimeras (n=6, mean \pm SD, two-tailed Student's t-test). P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.001, * = p-value < 0.05, ns = not significant.
Extended Data Fig. 5	Extended Data Fig. 5	EF5.pdf	a-e) Flow cytometry analysis of <i>C albicans</i> -specific T cells (using Als1-tet) after oral colonization. Aire whole-body knockout mice (Aire ^{-/-}), their wild-type littermates (Aire ^{+/+}), WT (<i>Rorc</i> -Cre ⁻ Aire ^{fl/fl}) and ILC3 ^{ΔAire} (<i>Rorc</i> -Cre ⁺ Aire ^{fl/fl}) were orally colonized by <i>C. albicans</i> and analyzed after 24 hours (left panel) or two weeks (right panel). a) Representative FACS plot of Als1-tet ⁺ T cells. Cells were isolated from pLNs and spleens (SLO), oral mucosa, esophagus and intestine of mice described above 24 hours (left panel) or 2 weeks (right panel) post <i>C. albicans</i> colonization. Counts of Als1-tet ⁺ cells are highlighted in red rectangles. b-e) Statistical analysis of the Als1-tetramer counts (n=6, mean ± SD, two-tailed Student's t-test in SLO (b), oral mucosa (c), esophagus (d) and small intestine (e) 24 hours (left panel) or 2 weeks (right panel) post <i>C. albicans</i> colonization (n=6, mean ± SD, two-tailed Student's t-test in SLO (b), oral mucosa (c), esophagus (d) and small intestine (e) 24 hours (left panel) or 2 weeks (right panel) post <i>C. albicans</i> colonization (n=6, mean ± SD, two-tailed Student's t-test). P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.05, ns = not significant.
Extended Data Fig. 6	Extended Data Fig. 6	EF6.pdf	a) Representative FACS gating strategy of Roryt ⁺ CD4 ⁺ TCR- β^+ T _H 17 cells in pLNs two weeks after <i>C. albicans</i> colonization of <i>Rorc</i> -Cre ⁻ Aire ^{fl/fl} (WT) and <i>Rorc</i> -Cre ⁺ Aire ^{fl/fl} (ILC3 ^{ΔAire}) mice. b) Statistical analysis of related to a). The plot is showing the frequency from parent gate (n=6, mean ± SD, two-tailed Student's t-test). c) Representative FACS gating strategy of Roryt ⁺ CD4 ⁺ TCR- β^+ T _H 17 cells in mesenteric lymph nodes (mLN) two weeks after <i>C. albicans</i>

			colonization of <i>Rorc</i> -Cre ⁻ Aire ^{fl/fl} (WT) and <i>Rorc</i> -Cre ⁺ Aire ^{fl/fl} (ILC3 ^{ΔAire}) mice. d) Statistical analysis of related to c). The plot is showing the frequency from parent gate (n=6, mean ± SD, two-tailed Student's t-test). e) Representative FACS gating strategy of Roryt ⁺ CD4 ⁺ TCR- β^+ T _H 17 cells in lamina propria two weeks after <i>C. albicans</i> colonization of <i>Rorc</i> -Cre ⁻ Aire ^{fl/fl} (WT) and <i>Rorc</i> -Cre ⁺ Aire ^{fl/fl} (ILC3 ^{ΔAire}) mice. f) Statistical analysis of related to e). The plot is showing the frequency from parent gate (n=6, mean ± SD, two-tailed Student's t-test). P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.001, * = p-value < 0.05, ns = not significant.
Extended Data Fig. 7	Extended Data Fig. 7	EF7.pdf	a-d) Colony forming units (CFU)-based assay determining the overgrowth of <i>C</i> albicans 1 day (left panel) and 14 days (right panel) after the oral colonization. The tissues were isolated from Aire whole-body knockout mice (Aire ^{-/-}), their wild-type littermates (Aire ^{+/+}), <i>Rorc</i> -Cre ⁻ Aire ^{fl/fl} (WT) and <i>Rorc</i> -Cre ⁺ Aire ^{fl/fl} (ILC3 ^{ΔAire}) mice. CFU was determined by plating the lysates obtained from the kidney (a), oral cavity (b), esophagus (c), small intestine (d) (n=6, mean ± SD, two-tailed Student's t-test). P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.05, ns = not significant.
Extended Data Fig. 8	Extended Data Fig. 8	EF8.pdf	a) Experimental outline relevant to data shown in (b-d). <i>Rorc</i> -Cre ⁻ Aire ^{fl/fl} (WT) and <i>Rorc</i> -Cre ⁺ Aire ^{fl/fl} (ILC3 ^{ΔAire}) were first primed by repeated injections of heat-killed <i>C. albicans</i> (HKCA) for two weeks, then exposed to prolonged protocol of oropharyngeal candidiasis and analyzed five days later. b) Representative FACS plot of Als1-tet ⁺ T cells . Cells were isolated from pLNs and spleens of mice described in a). Counts of Als1-tet ⁺ cells are highlighted in red rectangles (left panel). Statistical analysis of the same representative experiment showing the total counts of Als1-tet ⁺ cells (n=6, mean ± SD, two- tailed Student's t-test). Representative experiment is shown. c) Representative FACS plot of Als1-tet ⁺ T cells. Cells were isolated from tongue mucosae of mice described in a). Counts of Als1-tet ⁺ cells are highlighted in red rectangles (left panel). Statistical analysis of the same representative

			the total counts of Als1-tet ⁺ cells (n=6, mean \pm SD, two-tailed Student's t-test). Representative experiment is shown.
Extended Data Fig. 9	Extended Data Fig. 9	EF9.pdf	 a) Representative FACS-sorting strategy of Aire⁺ ILC3s from reporterAire-GFP positive mice. The plots are showing the frequency from parent gates. Lineage: CD3, CD19, CD11c, CD11b, N.K.1, F4/80, Gr1, B220. b) Back-gating of Aire-GFP⁺ cells.
Extended Data Fig. 10	Extended Data Fig. 10	EF10.pdf	a) Experimental outline relevant to data shown in (b-c). WT (<i>Rorc</i> -Cre ⁻ Aire ^{fl/fl}) and ILC3 ^{ΔAire} (<i>Rorc</i> -Cre ⁺ Aire ^{fl/fl}) mice were transferred with naïve OT-II CD4 ⁺ T cells and subsequently injected with HKCA-OVA four times during a single week. b) Statistical analysis of the frequencies of OT-II T-cells subtypes (n=4, mean ± SD, two-tailed Student's t-test). c) GO enrichment analysis of upregulated differentially expressed genes from <i>Rorc</i> -GFP ⁺ OT-II T cells vs Non-proliferating OT-II T cells comparison. d) Graphical model summarizing the role of Aire ⁺ ILC3s in the induction of Candida-specific T _H 17 response. While the immediate immune response to <i>C. albicans</i> infection is dominated by neutrophils, monocytes and macrophages, Aire ⁺ ILC3s become essential in the later phase, as they facilitate clonal expansion of the primed T _H 17 cell clones in the LN. The expanded candida-specific T _H 17 clones subsequently limit <i>C.</i> <i>albicans</i> overgrowth at mucosal surfaces and its dissemination into epithelial tissues; e) Scheme illustrating the putative mechanism through which Aire ⁺ ILC3 cells induce the expansion of candida-specific T _H 17 clones.

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Extrathymic expression of Ai	re contro	Is the induction of effective $T_{\rm H}17$ cell-m	ediated immune response to Candida albicans
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42 Patients with loss of function in the gene encoding the master regulator of central tolerance AIRE suffer from a devastating disorder called autoimmune polyendocrine syndrome type 1 (APS-1), 43 44 characterized by a spectrum of autoimmune diseases and severe mucocutaneous candidiasis. While 45 the key mechanisms underlying the development of autoimmunity in APS-1 patients are well-46 established, the underlying cause for the increased susceptibility to Candida albicans infection 47 remains less understood. Here we show that Aire⁺MHCII⁺ type-3 innate lymphoid cells (ILC3) could sense, internalize and present Candida albicans, and had a critical role in the induction of 48 49 candida-specific T_H17 cell clones. Extrathymic Rorc-Cre-mediated deletion of Aire resulted in 50 impaired generation of candida-specific $T_H 17$ cells and subsequent overgrowth of C. albicans in 51 the mucosal tissues. Collectively, our observations identify a previously uncharacterized regulatory 52 mechanism for effective defense responses against fungal infections.

The transcriptional regulator Aire has an essential role in the induction of self-tolerant T cells during thymic development by controlling the expression of thousands of self-antigen genes in medullary thymic epithelial cells (mTECs)^{1, 2}. Presentation of self-antigens by mTECs is essential for the deletion of self-reactive T cell clones³ or their conversion into regulatory T cells (T_{reg} cells) ^{4, 5}. *Aire* deficiency results in impaired T_{reg} cell generation and escape of self-reactive T cells into the periphery, leading to breakdown of immunological tolerance to various parenchymal tissues¹.

60 Patients with AIRE deficiency develop a rare genetic disorder called autoimmune polyendocrine 61 syndrome type-1 (APS-1; also known as autoimmune polyendocrinopathy candidiasis ectodermal 62 dystrophy – APECED, OMIM: 240300), which is characterized by autoimmune pathologies such 63 as hypoparathyroidism and primary adrenocortical insufficiency (Addison's disease), with additional autoimmune disorders such as hypothyroidism, type-1 diabetes, premature ovarian 64 65 failure, pernicious anemia, vitiligo, alopecia, keratitis or intestinal malabsorption occurring with lower frequency ^{6,7}. In addition, the vast majority (75-100%) of APS-1 patients develop chronic 66 67 mucocutaneous candidiasis (CMC), mainly characterized by C. albicans overgrowth in the oral cavity, esophagus and nails as early as 1 year of age (median 5 years)^{8,9}. Because candidiasis is 68 a common complication in people born with a loss of function mutation in various genes linked to 69 the CD4⁺ helper type 17 (T_H17 cell)-mediated response (e.g. $RORC^{10}$, $IL17F^{11}$, $STAT3^{12}$, 70 *CLEC7A*¹³, *CARD9*¹⁴ or *STAT1*¹⁵), T_H17 cells are assumed to have an indispensable role in long 71 term protection against C. albicans infection¹⁶. APS-1 patients have been reported to develop 72 autoantibodies against the T_H17 effector cytokines IL-17A, IL17-F and/or IL-22^{17, 18}, suggesting 73 74 that the increased susceptibility to C. albicans might also be caused by an autoimmune-mediated 75 mechanism. However, a considerable fraction of APS-1 patients with very low or no IL-17 or IL-22 autoantibodies still develop CMC^{17, 18}, indicating the correlation between the IL-17- and/or IL-76 77 22-specific autoantibodies and candidiasis is incomplete and that additional mechanism(s) may be 78 involved.

79 In addition to its well-established expression pattern in the thymus, Aire was reported to be expressed in a rare population of cells residing in the lymph nodes (LNs) and bearing the hallmarks 80 of a subset of MHCII⁺ type-3 innate lymphoid cells (ILC3)¹⁹. Considering that these Aire⁺ MHCII⁺ 81 82 ILC3 (hereafter Aire⁺ ILC3s) also express the molecular machinery for antigen presentation and T cell activation (MHCII, CD80, CD86, ICOSL)¹⁹, we investigated whether extrathymic expression 83 84 of Aire in this ILC3 subset may contribute to the adaptive immune response to C. albicans. Here 85 we show that Aire⁺ ILC3 cells sensed and internalized C. albicans, and effectively presented C. 86 albicans epitopes on their MHCII. Moreover, extrathymic (but not thymus-specific) ablation of 87 Aire impaired the expansion of the candida-specific T_H17 cell pool and resulted in overgrowth of 88 C. albicans at various mucosal surfaces.

89 **RESULTS**

90 Aire⁺ ILC3s express receptors involved in *C. albicans* sensing

91 To test whether peripheral expression of Aire in ILC3s was required for the induction or modulation 92 of an effective immune response to C. albicans infection, we determined whether ILC3s in general 93 and Aire⁺ ILC3s in particular expressed pattern recognition receptors (PRRs) for *C. albicans*. We 94 used Rorc-Cre⁺flox-STOP-flox-tdTomato reporter mice that were crossed with Aire-GFP⁺ transgenic reporter mice (hereafter Rorc^{Tomato} Aire^{GFP} mice), in which the tdTomato reporter is 95 expressed in all cells with a history of *Rorc* expression and the green fluorescent protein (GFP) is 96 97 expressed in cells with an active Aire locus. ILC3s isolated from the popliteal lymph node (pLN) 98 by flow cytometry-based sorting as lineage⁻(CD3, TCR-β, CD45RB, CD19, Gr1, F4/80, CD11b, 99 CD11c) Rorc-tdTomato⁺ cells were divided according to their MHCII and Aire-GFP expression 100 into MHCII⁻Aire-GFP⁻ conventional ILC3s (hereafter cILC3s), MHCII⁺Aire-GFP⁻ ILC3s (hereafter MHCII⁺ ILC3s) and MHCII⁺Aire-GFP⁺ ILC3s (hereafter Aire⁺ ILC3s) (Fig. 1a). In 101 102 addition, to compare the molecular characteristics of these three ILC3 subsets with conventional antigen presenting cells (APCs) that can sense C. albicans²⁰, we isolated CD11c⁺CD11b⁺ MHCII⁺ 103 dendritic cells (DCs) from the Rorc^{Tomato} Aire^{GFP} mice. All sorted populations were analysed by 104 105 bulk RNA sequencing. Clustering analysis highlighted relatively large transcriptional similarity 106 between all cell subsets expressing MHCII (i.e. MHCII⁺ ILC3, Aire⁺ ILC3 and DC), and in 107 particular between the Aire⁺ ILC3s and MHCII⁺ ILC3s (Fig. 1b). All three ILC3 subsets were, as 108 expected, characterised by low expression of genes specific for hematopoietic stem cells (Cd34, 109 Slamf1), T cells (Cd4, Cd8, Foxp3); B cells (Cd19, Cd79a); granulocytes (Ly6c1, Ly6g5b, Siglecf); 110 macrophages and monocytes (Itgam, Siglech); dendritic cells (Itgax, Csflr, Cd207) and had high 111 expression of genes associated with ILC3s (Rorc, Il1r1, Il7r, Kit, Ccr6) (Fig. 1c). Unlike cILC3s, 112 MHCII⁺ ILC3 and the Aire⁺ ILC3s had high expression of genes encoding MHCII (H2-Aa, H2-113 Ab1) (Fig. 1d), while Aire⁺ILC3s also had high expression of Aire and genes encoding

114 costimulatory molecules (Cd80, Cd86) at levels comparable to DCs (Fig. 1d). The Aire⁺ ILC3s also expressed transcripts for several key PRRs implicated in sensing C. albicans, such as $Tlr2^{21, 22}$, 115 dectin-1 (*Clec7a*)^{23, 24} or galectin-3 (*Lgals3*)^{25, 26} (Fig. 1e) and signaling molecules downstream of 116 117 these receptors (Myd88, Syk) (Fig. 1f). The protein expression of the C. albicans-sensing receptors 118 including dectin-1, galectin-3 and Tlr2 on Aire⁺ ILC3s was confirmed by flow cytometry in steadystate conditions and was comparable to the expression on these receptors on DCs (Fig. 1g). These 119 120 results collectively suggested that Aire⁺ ILC3s expressed genes implicated in antigen presentation, T cell activation¹⁹ and also C. *albicans* sensing. 121

122

123 C. albicans induces transcriptional changes in ILC3 subsets

124 To test whether C. albicans could induce the activation of ILC3s in general, and Aire⁺ ILC3s in 125 particular, we performed bulk RNA sequencing of Aire⁺ ILC3s, MHCII⁺ ILC3s and cILC3s sorted from pLN of *Rorc*^{Tomato} *Aire*^{GFP} mice that were intravenously injected at 1 day intervals for 3 days 126 127 with heat-killed Candida albicans (HKCA) or PBS and analyzed 24 hours after the last injection. All three ILC3 subsets, and particularly the Aire⁺ ILC3s subset isolated from HKCA-challenged 128 129 mice showed transcriptional changes compared to PBS-challenged controls (Fig. 2a-c). cILC3s 130 from HKCA-challenged mice significantly upregulated 149 genes (Fc ≥ 2.0 , adjusted p-value ≤ 0.05) 131 compared to PBS-treated controls (Fig. 2a), with genes encoding several PRR implicated in 132 C. albicans-sensing (Lgals3, Clec7a, Clec4d, Clec4e and Cd209b) or molecules involved in 133 antigen presentation (H2-Aa, H2-Ab1, H2-Eb1 and Cd74), co-stimulation (Cd86) or induction of 134 pro-inflammatory response (II1a, II1b and II18) among the most upregulated (Fig. 2a, Extended 135 Data Fig. 1a). This suggested that cILC3s may upregulate *Candida*-sensing receptors and/or MHCII 136 molecules in response to the inflammatory conditions engendered by C. albicans stimulation. 137 MHCII⁺ ILC3s from the HKCA-challenged mice significantly induced the expression of 641 genes, including *Aire*, which was the most significantly upregulated gene (p-value $\leq 1 \times 10^{-23}$; FC>7) 138

139 compared to their PBS-their counterparts, and further upregulated the expression of Clec7a (dectin-140 1), H2-Ab1 (MHCII) and Cd86, while downregulating the expression of several key ILC3-specific 141 genes, such as *II17a*, *II17f* and *II12rb1* (Fig. 2b, Extended data Fig. 1b). These observations 142 suggested that the HKCA challenge potentiated the antigen presentation capacity of the MHCII⁺ 143 ILC3 subset, while limiting their effector functions as IL-17-producing cells (Fig. 2b, Extended 144 Data Fig. 1b). Moreover, fold-change/fold-change analysis comparing MHCII⁺ ILC3s from 145 HKCA-stimulated vs. PBS-stimulated mice and Aire⁺ ILC3 vs. MHCII⁺ ILC3s from PBS-treated mice showed that MHCII⁺ ILC3s from HKCA-treated mice acquired a transcriptional signature 146 147 similar to that of Aire⁺ ILC3s (Fig. 2d). This suggested that exposure of MHCII⁺ ILC3s to HKCA 148 could likely initiate the acquisition of a transcriptional signature characteristic of Aire⁺ ILC3s (Fig. 149 2d, e, Extended Data Fig. 1d).

150 Finally, Aire⁺ ILC3s from HKCA-stimulated mice significantly upregulated 777 genes compared 151 to PBS-treated mice, including cytokines or cytokine receptors (II6, Bmp2, Il7r, Il23r, Il2rb), 152 chemokines (Ccl2), C. albicans-sensing receptors (Clec7a), cell adhesion molecules (Vcam1, 153 Cadm1, co-stimulatory molecules (Cd86) and enzymes involved in proinflammatory response 154 (Ptgs2) (Fig 2c). Moreover, the HKCA challenge lead to a significant upregulation of 116, which 155 along with TGF- β , induces the polarization of naïve T cells to the T_H17 subset and their clonal expansion ^{27, 28, 29} (Fig. 2c, Extended Data Fig. 1c). These results indicated that HKCA induced 156 157 distinct transcriptional programs in the three ILC3 subsets analyzed, including the upregulation of 158 genes involved in antigen presentation, co-stimulation and candida sensing in cILC3s and MHCII⁺ 159 ILC3s, the upregulation of *Aire* in MHCII⁺ ILC3s and the upregulation of *Il6* in the Aire⁺ ILC3s, 160 suggesting that Aire⁺ ILC3s could play a role in the induction or regulation of the adaptive immune 161 response to C. albicans.

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163 Aire⁺ ILC3s internalize *C. albicans* for antigen presentation

164 Next, we investigated whether Aire⁺ ILC3s could uptake C. albicans by endocytosis. To this end, 165 we performed imaging flow cytometry (Imagestream) analysis of MACS-enriched lineage negative 166 (Lin^{neg}: CD3⁻, TCR-β⁻, CD45RB⁻, CD19⁻, Gr1⁻, F4/80⁻, CD11b⁻, CD11c⁻) cell fraction or 167 CD11c⁺DCs isolated from pLN and incubated ex-vivo with cell proliferation dye (CPD)-stained 168 HKCA (HKCA-CPD). After 30 minutes of co-incubation, most Aire⁺ ILC3s were physically 169 associated with HKCA-CPD (Fig. 3a), whether through cell-cell interactions or internalization of 170 the HKCA-CPD into their cytoplasm. After 90 minutes of co-incubation, most Aire⁺ ILC3s had intracellular CPD signals (Fig. 3b), suggesting ingestion of HKCA-CPD. Vesicles containing 171 172 internalized fungi fused with the lysosomes in both Aire⁺ILC3s and DCs (Extended Data Fig. 2a), 173 suggesting that Aire⁺ ILC3s had endocytic capacity for *C. albicans*. Moreover, while the capacity of Aire⁺ ILC3s to endocytose HKCA-CPD was comparable to CD11c⁺ or CD11b⁺ mononuclear 174 175 phagocytes (MNPs) (Extended Data Fig. 2b), the endocytic capacity of both MHCII⁺ ILC3s and 176 cILC3s was substantially lower than that of Aire⁺ ILC3s (Extended Data Fig. 2b). Moreover, similarly to DCs, the endocytic capacity of Aire⁺ ILC3s cells was partially dependent on the 177 178 expression of dectin-1 and galectin-3 and was reduced at 4°C (Extended Data Figure 2c, d). 179 To assess the capacity of Aire⁺ ILC3s to present candida-derived antigens under *in vitro* conditions, 180 we FACS-sorted pLN-resident Aire⁺ ILC3s and other types of APCs (B cells, Ly6C⁺ MNPs, 181 CD11b⁺ MNPs, CD11c⁺ MNPs) from Balb/c wild-type mice using a gating strategy described in

187 presentation capacity (Fig 3c). The Aire⁺ ILC3s showed substantial capacity to present HKCA-

Extended Data Fig. 2e), stimulated them with HKCA or transgenic HKCA expressing OVA

(HKCA-OVA) and incubated them for 17 hours with an NFAT-GFP DO11.10 TCR reporter cell

line⁴, which turn on GFP expression in response to TCR stimulation by OVA³²³⁻³³⁹ peptide

presented in the context MHCII molecules. Based on this *in vitro* assay, CD11b⁺MNPs were the

most efficient APCs (Fig. 3c), while CD19⁺ B cells and Ly6C⁺ MNPs showed negligible antigen

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derived antigens, comparable to that of CD11c⁺ MNPs and ~5-fold lower than that of CD11b⁺
MNPs (Fig. 3c; Extended Data Fig. 2f).

190 To assess the capacity of Aire⁺ ILC3s to present candida-derived antigens under more physiological 191 conditions, we sorted Aire⁺ ILC3s and other types of APCs (B cells, Ly6C⁺ MNPs, CD11b⁺ MNPs, 192 CD11c⁺ MNPs) from the pLNs of Balb/c wild type mice injected with either HKCA-OVA or 193 HKCA at 12, 24 or 72 hours before sorting (Extended Data Fig. 2g). The sorted cells were then co-194 incubated with the NFAT-GFP DO11.10 TCR reporter cells line for 17 hours and their antigen 195 presentation capacity was determined by the frequency of cells expressing the NFAT-GFP reporter. 196 CD11b⁺MNPs isolated 12 hours after the HKCA challenge showed the highest antigen presentation capacity, as measured by the frequency of NFAT-GFP⁺ cells ($\sim 1.7\%$) (Fig. 3d, Extended Data Fig. 197 2h). Their capacity to induce the NFAT-GFP signal declined at 24 hours (Fig. 3e, Extended Data 198 199 Fig. 3d) and largely diminished at 72 hours after the HKCA challenge (Fig. 3f, Extended Data Fig. 200 2h). Both, Aire⁺ ILC3s and CD11c⁺ MNPs induced the highest frequency of NFAT-GFP⁺ cells 201 (1.2% and 0.7% respectively) at 72 hours after the HKCA challenge (Fig. 3f, Extended Data Fig. 202 2h). These data suggested that while $CD11b^+$ MNPs dominated the early antigen-presentation 203 phase, Aire⁺ ILC3s were more efficient at presenting candida-derived antigens at later timepoints. 204 To examine the capacity of Aire⁺ ILC3s to present C. albicans-derived antigens to T cells in the 205 pLNs, we performed *ex vivo* two-photon imaging in *Aire*-GFP reporter mice that were adoptively 206 transferred with OT-II T cells bearing OVA-specific TCR and endogenously expressing tdTomato fluorescence protein (OT-II^{tdTomato}). The mice were intravenously stimulated either with HKCA-207 208 OVA or HKCA as control, and two-photon excitation microscopy on explanted pLNs was used to visualize the interactions between Aire-GFP⁺ cells and OVA-specific OT-II^{tdTomato} T cells (Fig. 3g). 209 While we did not observe any physical interaction between *Aire*-GFP⁺ cells and OT-II^{tdTomato} T cells 210 211 in pLNs isolated from mice stimulated with HKCA, we observed numerous such interactions in

212 pLNs isolated from mice stimulated by HKCA-OVA (Fig. 3g). These observations suggested that

Aire⁺ILC3s might present candida-derived antigens to T cells and that they could exert this function
at later timepoints compared to conventional APCs.

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216 Aire⁺ ILC3s are required for induction of candida-specific T_H17

217 To determine whether Aire⁺ ILC3s, and, in particular, the expression of *Aire* in these cells regulated 218 the adaptive immune response to C. albicans, we aimed to conditionally inactivate Aire in Aire⁺ ILC3s. To this end, we generated *Rorc*-Cre⁺Aire^{fl/fl} (hereafter ILC3^{Δ Aire}) ^{30, 31}, in which Aire is 219 220 ablated in all cells that either actively express *Rorc* or have *Rorc* expression history. Because the expression of Aire in the pLNs of wild-type mice was restricted to Lin^{neg}Roryt⁺MHCII⁺ cells¹⁹ 221 222 (Extended Data Fig. 3a, b), this strategy allowed us to specifically inactivate Aire expression in Aire⁺ ILC3s. As controls, we generated *Foxn1*-Cre *Aire*^{fl/fl} mice (hereafter TEC^{Δ Aire}), in which Aire 223 224 is deleted in thymic epithelial cells (TEC). Using flow cytometry, we could not detect any Aire protein expression in samples prepared from the pLN of ILC3^{Δ Aire} mice, while Aire protein was 225 detected in the ILC3^{Δ Aire} thymus samples (Fig. 4a). Conversely, Aire protein was detected in the 226 pLNs, but not the thymus, of $\text{TEC}^{\Delta Aire}$ mice (Fig. 4a). We next intravenously transferred naïve OT-227 II T cells from CD45.1 expressing GFP fluorescent protein under the Rorc-promoter (OT-II-228 229 Rorc-GFP) equally mixed with control polyclonal T cells from CD45.1/CD45.2 wild-type mice into CD45.2 positive wild-type or ILC3^{Δ Aire} mice, followed by stimulation of recipient mice with 230 HKCA-OVA or HKCA as control in consecutive 2 days intervals and analyzed 2 or 14 days after 231 232 the first injection. While no significant differences in OT-II-Rorc-GFP T cell cellularity and/or 233 proliferation were observed at day 2 post-transfer between the HKCA-OVA or HKCA-stimulated wild-type vs. ILC3^{ΔAire} mice (Extended Data Fig. 3 c,d), the OT-II-Rorc-GFP T cells transferred in 234 235 the HKCA-OVA-stimulated wild-type mice showed a ~2.5 fold proliferative expansion at day 14 post-transfer compared to the HKCA-OVA-stimulated ILC3^{Δ Aire} mice (Fig. 4b, c). The frequency 236

of CD45.1⁺ OT-II T cells in the HKCA-OVA-treated ILC3^{Δ Aire} mice was decreased by > 8-fold 237 238 compared to HKCA-OVA-treated wild-type controls (Fig. 4b, c). Moreover, approximately 10% 239 of the CD45.1⁺ OT-II T cells transferred in the HKCA-OVA-stimulated wild-type mice 240 differentiated into Rorc-GFP⁺ T_H17 cells (Fig. 4d). In contrast, the frequency (Fig. 4d) and number (Fig. 4e) of CD45.1⁺ Rorc-GFP⁺ OT-II T_H17 cells in HKCA-OVA-treated ILC3^{△Aire} was reduced 241 242 by ~20 fold and ~100-fold, respectively, compared to HKCA-OVA-treated wild-type controls at 243 day 14 post-transfer, suggesting that Aire expression in ILC3s was required for the induction of 244 effector $T_H 17$ cells in response to *C. albicans* stimulation.

245 Next, we performed MHCII tetramer staining for C. albicans-specific epitopes derived from the 246 agglutinin-like protein 1 (Als1). Specifically, pooled spleen or pLN-resident CD4⁺ T cells isolated from ILC3^{Δ Aire}, TEC^{Δ Aire} or their wild-type littermates that had been stimulated with HKCA for two 247 248 weeks prior to isolation were stained with Als1-specific tetramers (Als1-Tet) and analyzed by flow 249 cytometry. We observed a significant reduction in the number of Als1-Tet⁺CD44⁺CD4⁺ activated T cells isolated from the spleen and pLNs of ILC3^{Δ Aire} mice compared to wild-type or TEC^{Δ Aire} 250 251 mice (Fig 4f, g), suggesting that extrathymic, but not thymic expression of Aire was critical for the 252 induction of candida-specific T cells.

To investigate whether a similar defect of T cell response to candida was found in APS-1 patients, we stimulated the peripheral blood mononuclear cell (PBMC) fraction from APS-1 patients or healthy individuals with HKCA and monitored T cell proliferation four days later. We observed a significant decrease (~3-fold) in both the frequency of HKCA-induced Ror γ T⁺ T_H17 cells (Fig. 4h, i), as well as in the amount of IL-17A released in supernatants from PBMCs isolated from APS-1 vs. healthy controls (Fig. 4j). Collectively, these data suggest that expression of Aire in ILC3s was required for the expansion of candida-specific CD4⁺ Ror γ t⁺ T cell clones.

261 Extrathymic Aire is critical for effective response to C. albicans

262 Next, we sought to determine whether loss of Aire expression in ILC3s could impair the clearance 263 of live C. albicans in vivo. Because C. albicans is generally completely absent in mice housed under specific pathogen free (SPF) conditions³², implying that SPF mice would have limited 264 adaptive immune response to this pathogen, we pretreated wild-type and ILC3^{Δ Aire} mice for 3 weeks 265 with intravenous (*i.v.*) injections of HKCA, before *i.v.* administration of live C. albicans. ILC3^{Δ Aire} 266 267 mice were significantly more susceptible to C. albicans challenge, with only ~10% survival at 14 268 days post-challenge, compared to wild-type littermates, which had a ~50% survival rate at this time point (Fig. 5a). Moreover, the ILC3^{Δ Aire} mice showed a significantly higher burden of C. 269 270 albicans in their kidneys in comparison to wild-type littermate controls (Fig. 5b, c and Extended Data Fig. 3g, h). Similarly, *Aire^{-/-}* mice on either B6 or NOD genetic backgrounds had significantly 271 272 lower survival rate following *i.v.* injection of live C. albicans (with 20% and 0% survival respectively) compared to their $Aire^{+/+}$ littermates (with 60% and 45% survival respectively) 273 (Extended Data Fig. 3e-f). While the Aire^{-/-} mice on the B6 background had significantly poorer 274 275 survival rate than their wild-type littermates, they did not develop detectable levels of IL-17- and/or 276 IL-22-specific autoantibodies (Extended Data Fig. 3i-j), further suggesting that the impaired anti-277 candida response was primarily T cell dependent.

Next, we assessed the adaptive immune response to live C. albicans in a mucosal model of 278 infection. We used a modified version of an existing protocol³³, which is based on the oral 279 280 administration of live C. albicans to mice in 3 consecutive times with 1 day intervals, resulting in 281 a long-lasting colonization of their gastro-intestinal tract. First, we assessed whether oral candida 282 colonization of wild-type mice resulted in the expansion of candida-specific T cells in the secondary 283 lymphoid organs (SLO) (Fig. 6a, b; Extended Data Fig. 4 a, b). Flow cytometry indicated that 284 while wild-type mice not treated with oral C. albicans harbored relatively low numbers of Als1-Tet⁺ T cells (3-5 naïve CD44⁻ T cells/SLO in each mouse analyzed), the number of Als1-Tet⁺ T 285

286 cells increased by ~50-100-fold at day 14 post C. albicans colonization (Fig. 6a, b), and virtually 287 all of them had an activated memory phenotype, as evidenced by high expression of CD44 (Fig. 6a). In contrast, *C. albicans* colonization of ILC3^{ΔAire} mice (Fig. 6c, d), Aire^{-/-} mice (Extended Data 288 Fig. 4c-e) or bone-marrow chimeric mice in which Aire deficiency occurred in the hematopoietic. 289 290 but not in the stromal compartment (Extended Data Fig. 4f-h), resulted in significantly decreased 291 numbers of Als1-Tet⁺ T cells in the SLO than did C. albicans colonization of their corresponding 292 wild-type littermate controls. Moreover, low numbers of Als1-Tet⁺ T cells were also observed in CD90.2 Rag1^{-/-} mice that were adoptively transferred with CD90.1⁺ T cells and B cells and treated 293 294 with a CD90.2-specific antibody to deplete the endogenous CD90.2 lymphoid compartment, thus rendering them ILC-deficient ³⁴ compared to isotype control-treated mice (Extended Data Fig. 4j-295 296 k).

297 We next investigated the presence of candida-specific T cells at different mucosal sites of the 298 gastrointestinal tract at day 1 or day 14 post C. albicans administration. While Als1-Tet⁺ T cells could not be detected in the oral cavity, esophagus or small intestine mucosa from wild-type, Aire 299 ^{*/-*} or ILC3^{Δ Aire} mice 24 hours post *C. albicans*-challenge (Extended Data Fig. 5), they were present 300 301 in these locations in wild-type mice at day 14 post-challenge (Fig. 6g, h; Extended Data Fig. 5). In contrast, ILC3^{Δ Aire} or *Aire*^{-/-} mice had markedly reduced numbers of Als1-Tet⁺ T cells at these sites 302 303 along the gastrointestinal tract, in particular in the lamina propria, at day 14 post-challenge 304 compared to their corresponding wild-type controls (Fig. 6g, h; Extended Data Fig. 5). In addition, 305 the number of the bulk Roryt⁺ T_H17 population in pLN and mLN isolated was higher in wild-type mice compared to ILC3^{Δ Aire} mice at day 14 post *C. albicans* colonization (Extended Data Fig. 6). 306 307 In contrast the number of $Roryt^+$ T_H17 cells within lamina propria were similar in wild-type mice and ILC3^{Δ Aire} mice at this time point (Fig. 6i,k), suggesting that while ILC3-specific Aire deficiency 308 309 impaired the expansion of candida-specific T_H17 cells in the lymphoid organs and lamina propria, 310 the effect on the general $T_{\rm H}17$ population was smaller and more variable. In line with the decreased number of candida-specific T cells at mucosal surfaces, the $Aire^{-/-}$ and $ILC3^{\Delta Aire}$ mice had a significantly higher burden of *C. albicans* in their gastrointestinal mucosal sites compared to their wild-type littermates at day 14 post-challenge (Fig. 6k, 1; Extended Data Fig. 7).

Finally, we also utilized a more conventional model for oropharyngeal candidiasis (OPC)³⁵, in 314 which the oral mucosal tissue in mice was exposed to live C. albicans for 90 minutes. In this setting, 315 we pre-treated wild-type and ILC3^{Δ Aire} mice with *i.v.* injection of HKCA every 2 days for 14 days 316 317 prior to the OPC challenge with live C. albicans, and the oral mucosa was analyzed at day 5 after the OPC challenge (Extended Data Fig. 8a). At this timepoint, we observed significantly less Als1-318 319 Tet⁺ T cells (Extended data Fig. 8b,c) and a higher burden C. albicans (Fig. 6m) in the oral mucosa of ILC3^{ΔAire} compared to wild-type mice (Extended Data Fig. 7b,c). Collectively, these data 320 321 demonstrated that Aire expression in Aire⁺ ILC3s was required for the induction of adaptive 322 immune responses to *C. albicans* at the mucosal surfaces.

323

324 Aire⁺ ILC3 induce survival of candida-specific T_H17 clones

325 To explore how expression of Aire in Aire⁺ ILC3s promoted the expansion of *C. albicans* specific 326 T cells, we assessed the impact of Aire deficiency on the gene expression profiles of Aire⁺ILC3s and candida-specific T cells in response to C. albicans challenge. For this purpose, we crossed Aire⁻ 327 ^{/-} mice with Aire-GFP reporter mice, to generate Aire-GFP-Aire^{-/-} mice. Aire⁺ ILC3s sorted (based 328 on Aire-GFP expression) from Aire-GFP-Aire^{-/-} or Aire-GFP-Aire^{+/+} littermates that had received 329 330 HKCA or PBS *i.v.* every day for 3 days (Extended Data Fig. 9) were analyzed by bulk RNAseq 331 analysis on day 4. Analysis of these transcriptomes indicated that Aire regulated the expression of 332 hundreds of genes, which strongly overlapped with the genes upregulated in response to HKCA (Fig. 7a, b). Specifically, the Aire⁺ ILC3s isolated from HKCA-stimulated Aire-GFP-Aire^{-/-} mice 333 334 had impaired induction of genes encoding cytokines (116, 1118, Bmp2), C. albicans-sensing receptors (*Clec7a*,), cell adhesion molecules (*Vcam1*, *Cadm1*), co-stimulatory molecules (*Cd86*) and enzymes involved in proinflammatory response (*Ptgs2*) compared to their HKCA-stimulated *Aire-GFP-Aire*^{+/+} counterparts (Fig 7a, b). These data suggested that Aire regulated the expression of several key molecules that may be critical for T_H17 differentiation (*Il6*), T cell clonal expansion (*Cd80/Cd86*; *Vcam1*) and pro-inflammatory signaling (*Il18*, *Ptgs2*).

340 To investigate how Aire deficiency impacted the transcriptional program of C. albicans-specific T cells, we adoptively transferred *Rorc*-GFP⁺ OT-II T cells into wild-type or ILC3^{△Aire} mice that were 341 342 stimulated with HKCA-OVA or HKCA every second day. Rorc-GFP⁺ OT-II T cells were then 343 isolated from pLN on day 7 post-transfer and analyzed by bulk RNA sequencing (Extended Data Fig. 10a). At this time point, the frequencies of *Rorc*-GFP⁺ OT-II T cells in ILC3^{Δ Aire} mice were 344 decreased by ~3-fold compared to wild-type littermates (Extended Data Fig. 10b). We observed 345 346 only mild increase in the frequency of CPD⁺ proliferating *Rorc*-GFP⁻ OT-II T cells in wild-type vs ILC3^{Δ Aire} mice (Extended Data Fig. 10b), suggesting that Aire was not involved in the regulation 347 348 of *Rorc*-GFP⁻ T cell priming.

349 The transcriptional analysis indicated that the *Rorc*-GFP⁺ OT-II T cells had substantially different transcriptomes compared to the non-proliferating CPD⁻ OT-II T cells. Specifically, the Rorc-GFP⁺ 350 351 OT-II T cells isolated from wild-type mice upregulated 551 genes, including Rorc, Cd44, chemokine receptor Ccr5 and proliferation marker Mki67 and downregulated 220 genes, including 352 353 Sell and Cd69, compared to the non-proliferating CPD⁻ OT-II T cells from wild-type mice (Fig. 7C 354 and Extended Data Fig. 10b), highlighting their activated/memory phenotype and readiness to exit 355 the lymph node and move to the effector site. We detected more than a thousand differentially 356 expressed genes between the *Rorc*-GFP⁺ OT-II T cells derived from the wild-type compared to the 357 ILC3^{AAire} mice (Fig 7c, and Extended Data Fig. 10b). Specifically, *Rorc*-GFP⁺ OT-II T cells isolated from ILC3^{ΔAire} mice had significantly reduced expression of anti-apoptotic factors such as survivin 358 359 (*Birc5*) or $Bcl2l12^{36}$ and a less activated/memory phenotype, as suggested by the decreased expression of the chemokine receptor *Cxcr3*, *Il2rb* (encoding the IL-2 receptor beta subunit), *Igf2r* ³⁷, *Il12rb1* (the key subunit of IL-23 receptor; Fig 7d, e) and *Cd28*, encoding the CD80/86 receptor essential for T cell co-stimulation (Fig. 7e). The GO annotation of cell processes indicated that the differentially regulated genes we highly enriched for factors involved in the regulation of cell cycle and/or mitosis (Fig. 7f). Collectively, these data suggested that Aire controlled the expression of genes in Aire⁺ ILC3s (e.g. *Il6*, *Cd86*, *Ptgs2*) that were critical for the subsequent induction of a prosurvival transcriptional program in T_H17 cells (Extended Data Fig. 10d, e).

367

368 DISCUSSION

369 Here we show that extrathymic expression of Aire in Aire⁺ ILC3s was required for the expansion 370 of candida-specific CD4⁺ T cells, in particular $T_{\rm H}17$ clones, in response to C. albicans infection 371 and for consequently limiting the pathogenicity of this opportunistic pathogen at mucosal tissues. 372 We found that Aire⁺ ILC3s expressed receptors implicated in *C. albicans* detection (e.g. dectin-1, 373 galectin-3) at levels comparable to DCs and effectively endocytosed C. albicans. Moreover, upon 374 C. albicans uptake, Aire⁺ ILC3s presented candida-derived antigens to CD4⁺ T cells through 375 MHCII. Therefore, our study suggests that effective response to C. albicans infection involves 376 division of labor between different types of phagocytes and APCs, with Aire⁺ ILC3s playing a non-377 redundant role in promoting the survival and subsequent expansion of candida-specific T cell 378 clones in the LN.

While ILCs are generally viewed as the innate analogs of T cells, a growing body of evidence suggests that some ILC subsets are, similarly to Aire⁺ ILC3, equipped with potent endocytic and/or antigen presentation capacity^{38, 39, 40, 41, 42, 43}. For instance, spleen-derived NCR⁻CCR6⁺MHCII⁺ ILC3s, were reported to internalize latex beads and present model protein antigen to CD4⁺ T cells *in vitro*⁴³. Similarly, MHCII⁺ ILC2 can endocytose and present OVA protein and induce antigen-

specific T cell proliferation⁴², while intestinal MHCII⁺ ILC3s were reported to regulate T cell 384 responses to bacterial antigens in an antigen dependent manner^{38, 40, 41}, arguing that some ILC3 385 subsets may act as potent APCs, with the capacity to modulate antigen-specific T cell responses in 386 387 different contexts and in different anatomical niches³⁸. Moreover, our data also suggested that LN-388 resident MHCII- ILC3s could upregulate the expression of MHCII and CD86 coding genes in response to C. albicans challenge. This is in line with reports showing that the pro-inflammatory 389 390 cytokine IL-1ß promotes the expression of MHCII and costimulatory molecules on spleen-, but not gut-derived MHCII⁻ ILC3s⁴³. 391

392 Moreover, our data indicated that the impaired expansion of candida-specific CD4⁺ T cells in SLO of ILC3^{ΔAire} mice was associated with their decreased accumulation at different mucosal sites, 393 including the oral cavity, esophagus and intestine, and with increased burden of C. albicans at these 394 395 mucosal tissues. It is likely that the impaired surveillance of the mucosal tissues due to a loss of 396 candida-specific CD4 T cells resulted in reduced ability to control the fungal burden at the mucosal 397 surfaces and loss of barrier integrity. In Aire-deficient patients, in which both the extrathymic and the thymic expression of Aire are defective, additional mechanisms (e.g. autoantibodies against IL-398 17, IL-22^{17, 18, 44}, defensins⁴⁵ and mucins⁴⁶ or loss of Paneth cells⁴⁵) may further impair the integrity 399 of mucocutaneous surfaces and thereby enhance the invasiveness of C. albicans. Therefore, the 400 401 increased susceptibility to C. albicans in Aire-deficiency may combine aspects of the role of Aire 402 in central tolerance with its role in shaping the C. *albicans*-specific $T_{\rm H}17$ response in the periphery.

403 Although our data are in line with the interpretation that the increased susceptibility to candida 404 infection in both humans and mice is due to defects in the $T_H 17$ response ^{12, 14, 15, 47, 48}, it was also 405 suggested that in the oral mucosa, Aire-deficient mice and humans have intact $T_H 17$ responses to 406 *C. albicans* and that these phenotypes are due to overproduction of interferon- γ (IFN- γ) in 407 CD4⁺ and CD8⁺ T cells at the epithelial barriers⁴⁹. In this specific study, however, the $T_H 17$ 408 response was measured only 24 hours post *C. albicans* challenge ⁴⁹, which might be too early a timepoint to assess an adaptive immune response, because *C. albicans* is virtually absent in mice housed under SPF conditions^{32, 33}. Indeed, we found that the induction of candida-specific T cells in SPF-housed mice peaked only two weeks post *C. albicans* colonization. In addition, the report assessed bulk T_H17 cell responses⁴⁹, and not candida-specific T cell clones as it was done in this study.

414 The molecular features of Aire⁺ ILC3s described here seem to largely overlap with a subset of Lin^{neg}Aire⁺Rorc⁺MHCII⁺ Janus cells that was recently identified using single-cell RNA profiling of 415 mouse LN⁵⁰, suggesting they might represent the same cell population. Although the study 416 suggested that the Lin^{neg}Aire⁺Rorc⁺MHCII⁺ Janus cells are a subset of tolerogenic DCs⁵⁰, they 417 lacked expression of key DC canonical markers such as Cd11c, Cd11b, Dec205, Clec9a or 418 $Cd4/Cd8^{50}$, as well as of other key myeloid or lymphoid markers. Moreover, similarly to Aire⁺ 419 420 ILC3s, the Janus cells highly expressed ILC3 markers, including Rorc, Il1r1, c-Kit, il7r, Id2, Ccr6, 421 Ccr7, Il18r1, as well as genes linked to antigen presentation and co-stimulation. Therefore, 422 although both Aire⁺ ILC3 and Janus cells lacked the expression of key ILC3 effector molecules 423 such as *Il17a*, *Il17f* or *Il22*, they bore molecular features more characteristic of MHCII⁺ innate 424 lymphoid cells, rather than of myeloid APCs. This notion is also supported by a recent study 425 showing that extrathymic $Lin^{neg}Aire^+Rorc^+$ MHCII⁺ cells have an interconverting potential with ILC3s in a fate mapping analysis based on Aire-Cre reporter mice⁵¹. Irrespectively of nomenclature, 426 427 our study provides experimental evidence that the expression of Aire in LN-resident Lin^{neg}Aire⁺Rorc⁺ MHCII⁺ cells has a critical and a non-redundant role in the induction of candida-428 429 specific T cells and the control of C. albicans colonization at mucosal tissues.

431 Collectively, our data not only help shedding more light on the mechanisms underlying chronic432 mucocutaneous candidiasis in Aire-deficient individuals, but also help identifying an additional

- 433 functional role for Aire, beyond its well-established role in central tolerance induction in the
- 434 thymus.
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450

451 AUTHORS CONTRIBUTION

JA and JD conceived the project, designed the experiments and wrote the manuscript; JD performed
the experiments and analyzed the data; JA supervised the study; AB, YG, OB-N, NK, YG, TG, IZ,
HB, KK, EV, BEO and ESH assisted at different aspects of the study, including data or sample
acquisition and analysis. DF contributed by the essential reagent. LSB and ZS performed the twophoton microscopy and analyzed the data.

457

458 **CFI statement :** The authors declare that they have no conflict of interest

460 FIGURE LEGENDS

461 Figure 1| Aire⁺ ILC3s express *C. albicans* sensing receptors.

462 a) FACS sorting strategy of cILC3s, MHCII⁺ ILC3s and Aire⁺ ILC3s subsets isolated from the popliteal lymph nodes of *Rorc*^{tdTomato} Aire^{GFP} reporter mice. Representative plots are shown. **b**) 463 464 Heatmap of Pearson correlation according to gene expression values between individual samples as in a). c-f) RNA-seq based heatmap of expression of lineage specific genes (c), genes associated 465 with MHCII-presentation or co-stimulation (d), genes encoding receptors implicated in sensing 466 467 and/or phagocytosis of C. albicans (e) and genes encoding signaling molecules downstream of C. 468 albicans sensing receptors (f) in cILC3s, MHCII⁺ ILC3s, Aire⁺ ILC3s and MHCII⁺ CD11c⁺ DCs isolated from popliteal lymph nodes of Rorc^{tdTomato} Aire^{GFP} at steady-state. g) Flow cytometry of 469 Roryt⁺ cILC3s, Roryt⁺ MHCII⁺ ILC3s, Roryt⁺ MHCII⁺Aire⁺ ILC3s and MHCII⁺ CD11c⁺ DCs 470 471 isolated from popliteal lymph nodes of wild-type animals under steady-state conditions. Grey filled 472 histogram indicates staining in control dectin-1 or galectin-3-deficient mice or isotype control for 473 CD80, CD86, TLR2. Representative figure, $n \ge 3$.

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475 Figure 2 C. albicans induces transcriptional changes in ILC3 subsets

476 a-c) Volcano plots of bulk RNA-seq analyses showing differential gene expression in nonstimulated vs. HKCA stimulated cILC3 (a), MHCII⁺ ILC3 (b) and Aire⁺ ILC3 (c) subpopulations 477 isolated from the popliteal lymph nodes of *Rorc*^{tdTomato} *Aire*^{GFP} mice *i.v.* injected by HKCA or PBS 478 479 for three consecutive days, using gating strategy as in Fig. 1a. Dashed lines indicate the cutoff for fold-change (Fc)=2.0 and p-value=0.05. Wald test was used to calculate the p-value. Based on their 480 481 function, the selected genes were highlighted in blue (antigen presentation, co-stimulation), red 482 (cytokine and chemokine signaling), green (candida sensing receptors), purple (Aire); d) Fold 483 change:fold change (Fc:Fc) graph of RNA-seq data showing differential gene expression in Aire⁺ 484 ILC3s in contrast to stimulated or non-stimulated MHCII⁺ ILC3s. Comparison of fold-change of 485 Aire⁺ ILC3s vs. MHCII⁺ ILC3s (x-axis) and non-stimulated vs. HKCA stimulated MHCII⁺ ILC3s 486 (y-axis) isolated by FACS using a gating strategy as in Fig. 1a. Dashed lines indicate the Fc=2.0487 cutoff. e) RNA-seq-based heatmap showing the level of expression of selected effector genes in 488 ILC3s, MHCII⁺ ILC3s and Aire⁺ ILC3s with or without HKCA stimulation. Data are plotted as the 489 z-score calculated for particular row. Data are derived from three independent biological replicates 490 for each population (a-e).

492 Figure 3 | Aire⁺ ILC3s internalize *C. albicans* for antigen presentation

493 **a-b**) Imaging flow cytometry showing the physical interaction between HKCA and Aire⁺ ILC3s 494 isolated from popliteal lymph nodes of wild-type mice and incubated ex-vivo with CPD-stained 495 HKCA for 30 (a) or 90 minutes (b). Samples were stained for Aire and DAPI (a-b) and MHCII 496 (b). Shown are representative images of one out of five independent experiments. Scale bar: 8µm. 497 **c-f)** In vitro reporter assay measuring the capacity of B cells, $Ly6C^+MNPs$, $CD11b^+MNPs$, $CD11c^+$ 498 MNPs and Aire⁺ ILC3s to endocytose HKCA transgenically expressing OVA (HKCA-OVA) either 499 in vitro (c) or in vivo (d-f) and subsequently to present HKCA-derived OVA antigens on their 500 MHCII molecules to DO11.10-TCR NFAT-GFP reporter cell line that was co-incubated with the 501 sorted populations for 17 hours. The corresponding APC populations were FACS-sorted according 502 to gating strategy shown in Extended Data Fig. 3a, b. Antigen presentation capacity was measured 503 as a frequency of NFAT-GFP⁺ reporters in the specific sample. Representative data set out of three 504 independent experiments are shown (n=5 for each experiment), mean \pm SD depicted. (d-f) Dot plot 505 of B cells, Ly6C⁺ MNPs, CD11b⁺ MNPs, CD11c⁺ MNPs and Aire⁺ ILC3s depicting APC capacity 506 at 12 hours (d), 24 hours (e) or 72 hours (f) after HKCA-OVA i.v. injection of HKCA-OVA into 507 wild-type mice; g) Two-photon microcopy of explanted popliteal lymph nodes showing interaction between Aire-GFP⁺ cells and candida-specific T cells from Aire-GFP reporter mice adoptively 508 509 transferred with OT-II^{tdTomato} T cells stimulated by either HKCA or HKCA-OVA. Representative 510 figures from two experiments are shown.

511

512 Figure 4 Aire⁺ ILC3s are essential for the generation of *C. albicans*-specific T_H17 response

a) Flow cytometry analysis showing disruption of Aire-expression Rorc-Cre⁻Aire^{fl/fl} (WT) and 513 *Rorc*-Cre⁺Aire^{fl/fl} (ILC3^{Δ Aire}) mice. Shown are representative FACS plots (n>6) of intracellular 514 Aire-staining of thymic stroma (upper panel) or lineage negative cells from pLNs (lower panel); **b**-515 e) Flow cytometric analysis of OT-II T cell proliferation and differentiation in WT or ILC3^{△Aire} 516 517 mice transferred with naïve OT-II and control CD4⁺T cells in 1:1 ratio and subsequently injected 518 with HKCA or HKCA-OVA every second day for 2 weeks showing representative flow cytometry 519 dot plot show the frequencies of transferred CD45.1⁺ OT-II T cell vs. CD45.1/CD45.2 double 520 positive control T cell populations on day 14 post-transfer (b), the corresponding statistical analysis 521 showing ratios (mean \pm SD, two-tailed Student's t-test) of OT-II vs. control T cells (n=5 per group) 522 (c), the frequencies of OT-II Rorc-GFP⁺ cells (d) and the corresponding statistical analysis of the

523 experiment showing the total counts (mean \pm SD, two-tailed Student's t-test) of HKCA-induced 524 OT-II *Rorc*-GFP⁺ cells (e); f-g, Flow cytometric analysis assessing counts of Als1-tetramer positive CD4 T cells in WT, ILC3^{Δ Aire} and TEC^{Δ Aire} mice that were injected intravenously with HKCA 525 526 every second day for two weeks showing representative FACS plot (f) and the corresponding 527 statistical analysis depicting the total counts (mean \pm SD, two-tailed Student's t-test) in all mice 528 (n=5) (g); h-i) Flow cytometric analysis assessing the proliferation of human Ror γT^+ T_H17 cells in 529 PBMCs isolated from APS-1 patients or healthy controls showing representative dot plots of the 530 frequency of proliferating Ror γT^+ T_H17 cells measured by CPD dilution (**h**) and the corresponding statistical analysis of the experiment depicting average frequency (mean \pm SD, two-tailed Student's 531 532 t-test, n=10 per group) (i); j) ELISA assessing amounts of IL-17A in the PBMC supernatants from 533 proliferation assay described in (h). Data are shown as mean of IL-17A concentration \pm SD, n=10 534 for each group, two-tailed Student's t-test. P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.001, * = p-value < 0.05, ns = not significant. 535

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Figure 5| Mice lacking extrathymic expression of Aire have reduced survival after systemic challenge with live *C. albicans*

a) Survival curves of WT (Cre⁻Aire^{fl/fl}) and ILC3^{Δ Aire} (*Rorc*-Cre⁺Aire^{fl/fl}) mice (n \geq 10 per genotype 539 540 group) injected *i.v.* with HKCA every two days for three weeks prior the infection by live C. albicans. Long-rank (Mantel-Cox) test was used to calculate the p-value: 0.0414; b) Quantitative 541 PCR analysis of C. albicans-specific DNA in the kidney from WT and ILC3^{Δ Aire} mice (n=6) 542 infected with C. albicans as in a (mean \pm SD, two-tailed Student's t-test); c) Colony forming units 543 544 (CFU)-based assay determining the overgrowth of *C albicans* in kidneys of mice described in (a), (n=6, mean ± SD, two-tailed Student's t-test). P-value indicators: *** = p-value < 0.0001, ** = p-545 546 value < 0.001, * = p-value < 0.05, ns = not significant.

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Figure 6| Extrathymic expression of Aire is critical for an effective T_H17 response to *C*. *albicans* at mucosal sites

a-b) Flow cytometry analysis assessing frequencies of *C. albicans*-specific Als1-Tet⁺, CD4⁺T cells in pLNs of WT mice orally colonized by *C. albicans* and analyzed at different time points post

552 colonization. Representative FACS plot showing counts of Als1-tet⁺, *C. albicans*-specific CD4⁺T

- cells (highlighted in red rectangles in the upper panel and red dots in the lower panel showing T
- cell activation markers CD44 vs CD69) (a) with a corresponding statistical analysis of the same

555 experiment showing the total counts (mean +SD, two-tailed Student's t-test (b); c-i) Flow 556 cytometry analysis assessing frequencies of C. albicans-specific Als1-Tet⁺, CD4⁺T cells in pLNs (c-f) or intestinal lamina propria (g-j) of WT vs. ILC3^{Δ Aire} mice orally colonized by C. albicans and 557 analyzed two weeks post colonization. Representative FACS dot plots showing counts of tetramer 558 559 positive CD4 T cells (c, g) and Ror γ T⁺CD4 T cells (e, i) are highlighted in black and red rectangles 560 respectively. Statistical analyses showing the total counts (mean +SD, two-tailed Student's t-test, 561 n=6 per group) corresponding to data shown in (c, e, g, i) are shown in (d, f, h, j) respectively. 562 Representative experiment out of three independent biological replicates is shown. (k) Quantitative PCR analysis assessing the presence of C. albicans-specific DNA in the ileal part of small intestine 563 from WT and ILC3^{Δ Aire} mice (mean +SD, two-tailed Student's t-test, n=6 per group). I) Colony 564 565 forming units (CFU)-based assay determining the overgrowth of C albicans in small intestinal 566 tissues 14 days after oral colonization (mean +SD, two-tailed Student's t-test, n=6 per group). m) 567 CFU-based assay determining the overgrowth of *C albicans* in oral mucosa 6 days after OPC 568 challenge (mean +SD, two-tailed Student's t-test, n=6 per group). P-value indicators: *** = p-value < 0.0001, ** = p-value < 0.001, * = p-value < 0.05, ns = not significant. 569

570

571 Figure 7 Aire⁺ ILC3s induce pro-survival program in candida-specific T_H17 clones

572 a) Volcano plot of bulk RNA-seq analysis showing differential gene expression in Aire⁺ ILC3s that were isolated from pLN of HKCA-stimulated Aire^{GFP}Aire^{+/+} vs Aire^{GFP}Aire^{-/-} mice. Dashed lines 573 indicate the Fc=2.0 and p-value=0.05 cutoff. Selected genes are highlighted in blue (antigen cell 574 575 adhesion, co-stimulation), red (cytokine and chemokine signaling), green (candida sensing 576 receptors). b) Fold change: fold change (Fc:Fc) graph of RNA-seq data showing differential gene 577 expression in Aire⁺ ILC3s promoted by HKCA stimulation. Comparison of fold-change of HKCAstimulated vs. non-stimulated Aire⁺ ILC3s isolated from Aire^{GFP}Aire^{+/+} vs Aire^{GFP}Aire^{-/-} (x-axis) 578 579 and Aire⁺ ILC3s from stimulated $Aire^{GFP}Aire^{+/+}$ vs stimulated $Aire^{GFP}Aire^{-/-}$ (y-axis). c) Volcano 580 plot of RNA-seq data showing differential gene expression of *Rorc*-GFP⁺ versus non-proliferating OT-II T cells derived from WT (*Rorc*-Cre⁻Aire^{fl/fl}) versus ILC3^{ΔAire} (*Rorc*-Cre⁺Aire^{fl/fl}) mice. The 581 mice were transferred with naïve OT-II CD4⁺T cells and subsequently injected with HKCA-OVA 582 583 four times during a single week; d) Volcano plot of RNA-seq data showing differential gene expression of *Rorc*-GFP⁺ OT-II T cells derived from WT versus ILC3^{Δ Aire} mice treated as in (c). 584 Dashed lines indicate the Fc=2.0 and p-value=0.05 cutoff. Data are derived from three independent 585 replicates. (e) RNA-Seq based heatmap showing the level of expression of selected genes in *Rorc*-586 GFP⁺ OT-II T cells isolated from WT and ILC3^{ΔAire} mice. Data are plotted as the z-score calculated 587

- 588 for particular row. (f) GO enrichment for upregulated differentially expressed genes from Rorc-
- 589 GFP⁺ OT-II T cells derived from WT versus ILC3^{Δ Aire} mice.

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766 MATERIAL AND METHODS

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768 Mice

The Aire^{fl/fl} (Jax: 031409;³⁰), Aire^{-/-} (004743 C57BL/6J and 006360 NOD genetic background;¹ 769 and⁵²), Aire-GFP (Aire-IGRP-GFP;⁵³) a kind gift of Dr. M. S. Anderson (University of California. 770 San Francisco, USA), CD45.1 congenic strain (002014), CD90.1 congenic strain (000406), Clec7a-771 ^{/-} (012337,⁵⁴), Foxn1-Cre (018448,⁵⁵), Lgals3^{-/-} (006338,⁵⁶), OT-II (004194;⁵⁷), Rag1^{-/-} (002216;⁵⁸) 772 *Rorc*-Cre (022791;³¹), *Rorc*-GFP (007572;³¹), *Rosa*-tdTomato (007914;⁵⁹) strains were used in the 773 774 study. Unless indicated otherwise, all mouse strains were of C57Bl6/J genetic background and were 775 purchased from Jackson laboratories if not indicated otherwise. Mice were housed in the premises 776 of Weizmann Institute of Science in SPF conditions. All experiments were approved by local 777 ethical committee (IACUC) under numbers: 39661117-2, 01420218-2, 04690718-2, 14850619-2. 778 Usually, 6-8 weeks old mice were used for the experiments with the exception of bone marrow 779 chimeras, where mice were 12-15 weeks old. For the generation of bone marrow chimeras, recipients were irradiated by single dose of 900 rad and transplanted by 1.10^7 bone marrow cells. 780 781 Only mice with reconstitution level higher than 95% were used for experiments. For the generation of CD90-disperate chimeras the protocol described elsewhere was used³⁴. Briefly, 6 weeks old 782 Rag1^{-/-} mice were intravenously adoptively transferred by 8.10⁷ MACS-enriched T cells and B-cells 783 784 from CD90.1 mice. These were let to homeostatically proliferate for two months. After this period, 785 mice were intraperitoneally injected by 250µg of CD90.2 depleting antibody (BioXcell) every 3 786 days. Whenever possible, littermates were used as the controls.

787

788 Human samples

Patients were included from Norwegian National Registry of Organ Specific Autoimmune Diseases
and fulfilled the APS-1 diagnostic criteria. All APS-1 patients suffer from *C. albicans* infection.
Gender matched controls were recruited from the local blood bank at Haukeland University

Hospital. All participants gave informed and written consent, and the study was approved by The Regional Committee for Medical and Health Research Ethics for Western Norway. PBMC were obtained from the whole blood by centrifugation in Ficoll-Pague (GE Healthcare), freeze and stored in liquid nitrogen. The proliferation assay was done for all samples together; whole PBMC fraction (5.10⁶ cells) was stained by Cell proliferation dye and stimulated by 1.10⁵ HKCA particles (Invivogen). Proliferative response was measured 4 and 6 days later.

798

799 Material

All material and reagents used in this study are described in particular relevant section and in detailspecified in Suppl. Table 1 and 2.

802

803 Infection by live *C. albicans*

Experimental mice were intra-venously injected every two days by 10^6 particles of heat-killed *C*. *albicans* in PBS for the duration of three weeks. After this period, mice were infected by single dose of 10^5 particles of live *C. albicans*. Mice were then monitored daily for their well-being and were sacrificed when they lose 20% of their initial weight of show signs of distress.

808

809 Mucosal colonization by C. albicans

In order to establish the gastrointestinal colonization by *C. albicans*, mice were supplemented *ad libitum* by 1mg/ml of ampicillin in drinking water and were kept on it during the experiment. After two days, mice were colonized by 50 ul of 10^6 particles *C. albicans* in PBS. The inoculation was performed dropwise into the mouth of mice. Experimental mice were monitored daily for their well-being.

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816 ELISA

For the detection of autoantibodies, the ELISA microtiter plate was pre-coated by 5µl/ml of recombinant IL-17 or IL-22 in bicarbonate buffer overnight in 4°C. The plate was washed and blocked by 5% milk. Detection of autoantibodies was performed using anti-IgG specific antibody conjugated to HRP (Jackson Immunoresearch). For the detection of human IL-17A cytokine the Human IL-17A ELISA kit (Biolegend) was used according to manufactures instructions.

822

823 C. albicans strains and preparation of HKCA

The wild-type *C. albicans* strain used in the study is of SC5314 origin⁶⁰. GFP and OVA coding sequence were inserted in the coding frame after the c-terminal end of the Eno1 gene resulting in generation of OVA-expressing strain derived from the wild-type⁶¹. Both strains were a kind gift of Dr. Judith Berman (Tel Aviv University, Israel). *C. albicans* was grown in 30°C using YPD agar. HKCA variant was prepared by heat-inactivation of the yeast in 60°C for one hour in thermoshaker. The heat-inactivation was tested by seeding the HKCA on YPD plates.

830

831 Cell isolation for flow cytometry and cell sorting

Aire-ILC3, MHCII-ILC3, ILC3s and DCs were isolated as described previously¹⁹. Although all 832 systemic LNs were found to contain Aire-ILC3 like cells¹⁹, for consistency most of the experiments 833 834 was done by analysis of cells and cellular responses in the popliteal LNs, unless stated otherwise. Briefly, lymph nodes were collected and subjected to several rounds of enzymatic digestion by 835 836 Dispase I. (Roche). Single cell suspension was depleted of lineage positive cells using LS-column 837 based MACS enrichment by cocktail of biotinylated antibodies (TCR-β, CD3, CD19, B220, 838 CD11b, F4/80, Gr1, CD11c, Biolegend) and anti-biotin microbeads (Miltenyi Biotec). T cells were 839 isolated by meshing the skin-draining lymph nodes and spleens through 40 µm nylon mesh. For 840 surface staining, cells were incubated with antibodies for 25 minutes on ice. DAPI (Sigma) of

841 viability dye eF506 (eBioscience) were used for live/dead cells discrimination. For intracellular 842 staining fixation the Foxp3 / Transcription factor staining buffer set (eBioscience) was used 843 according to manufacturer's recommendation. Subsequently, intracellular targets were stained by 844 antibodies for one hour in room temperature. Cells from the small intestinal lamina propria, oral 845 cavity or esophagus were collected by enzymatic digestion. Briefly, small intestinal tissue was 846 subjected to two rounds of epithelial cells removal by incubation with 2mM EDTA in HBSS for 20 847 minutes in 37°C. All the tissues were digested in 1mg/ml of Collagenase D and (Roche) for 1 hour 848 and immune cells were enriched by Percoll gradient (Sigma Aldrich). For details concerning 849 antibodies please refer to Suppl. Table 2. Flow cytometry analysis and cell sorting were performed 850 using BD CantoII, LSRII and AriaIII machines (BD). FlowJO (V10; Tristar) software was used for 851 flow cytometry data analysis.

852

853 **RNA sequencing**

Single cell suspensions were directly FACS sorted to Lysis/Binding buffer (Invitrogen) and frozen
on dry ice. RNA was isolated using Dynabeads (Invitrogen) according to manufactures protocol.
The MARS-seq protocol described elsewhere was followed to generate the sequencing libraries⁶².
The sequencing of the library was performed using the NextSeq high output kit and NextSeq 500
sequencer (Ilumina). Obtained data were analyzed for differential gene expression using the UTAP
pipeline⁶³.

860

861 Imagestream analysis of endocytosis

Lineage negative cell population or enriched DCs were co-incubated in-test described time period together with Cell proliferation dye eF660 (Thermo) stained 10⁵/ml HKCA particles in 37°C. Cells were fixed by Foxp3 / Transcription factor staining buffer set (eBioscience) and stained immediately after the end of incubation period and subjected to Imagestream analysis (Amnis). Data were analyzed using Ideas (v6.2) software (Amnis).

867 Image acquisition by TPLSM

MACS-isolated tdTomato⁺ cells were adoptively transferred to *Aire*-GFP hosts and stimulated by 868 869 heat-killed HKCA or HKCA-OVA. Zeiss LSM 880 upright microscope fitted with Coherent 870 Chameleon Vision laser was used for lymph node imaging experiments. Images were acquired with 871 a femtosecond-pulsed two-photon laser tuned to 930 nm. The microscope was fitted with a filter 872 cube containing 565 LPXR to split the emission to a PMT detector (with a 579-631 nm filter for 873 tdTomato fluorescence) and to an additional 505 LPXR mirror to further split the emission to 2 874 GaAsp detectors (with a 500-550nm filter for GFP fluorescence). Pictures were acquired at $512 \times$ 875 512 x-y resolution and the zoom was set to 1.5.

876

877 Analysis of endocytosis and antigen presentation capacity by FACS

Experimental mice were intravenously injected by 10^6 Cell proliferation dye eF660 (Thermo) stained HKCA. Mice were analyzed in indicated described time periods. For intravascular staining, mice were injected five minutes prior the analysis intravenously by 5 µg of anti-mouse CD45 BV605 monoclonal antibody (30-F11, Biolegend). For the antigen presentation assays experimental mice were intravenously injected by 10^6 HKCA or HKCA-OVA particles. Cells with antigen presentation capacity were isolated using FACS-sort and incubated with DO11.10 TCR NFAT-GFP cell line⁴ for 17 hours in ratio 1:5. GFP-fluorescence was measured using FACS.

885

886 Adoptive T cell transfer and stimulation of mice by HKCA

Naïve ovalbumin specific TCR⁺ OT-II cells from CD45.1⁺ mice and wild-type derived CD4⁺ T cells (CD45.1/CD45.2) were isolated using Naïve CD4⁺ T cells isolation kit (Miltenyi Biotec), mixed in 1:1 ratio, stained by Cell proliferation dye eF660 (Thermo) and transfer via tail vain to recipient mice. Once in two days, mice were injected by 10^6 HKCA or HKCA-OVA particles via tail vein.

892

893 Tetramer staining of *C. albicans* specific T cells

Als1 tetramers conjugated with PE and APC were used to detect *C. albicans* specific T cells from HKCA stimulated mice or mice after *C. albicans* colonization. The staining by tetramers and pulldown of tetramer positive cells by anti-PE and anti-APC conjugated microbeads (Miltenyi Biotec) was performed as described elsewhere⁶⁴. Each batch of tetramer reagent was titrated to determine the optimal staining concentration. We thank the NIH Tetramer Core Facility for providing tetramer reagents.

900

901 Isolation of DNA from tissue and intestinal content and quantification of *C. albicans* burden

902 Approximately 3mm of the ileum or kidney or liver tissue were surgically resected including its 903 content. DNA was extracted using Quick-DNA kit (Zymo research) according to manufactures instructions. 10ng of isolated DNA was used for downstream quantitative PCR reaction using Syber 904 905 -green (Roche) and following set of primers for detection of C. albicans (pF: TTTATCAACTTGTCACACCAGA , pR:ATCCCGCCTTACCACTACCG) and bacterial 906 907 16S ribosomal subunit (pF:ACTCCTACGGGAGGCAGCAGT, 908 pR: ATTACCGCGGCTGCTGGC) as the calibrator. The relative C. albicans DNA content in the samples was calculated using method described elsewhere⁶⁵. 909

910

911 Isolation of RNA from tissues and quantification of gene expression

Approximately 0.1g of the tongue tissue, esophagus, ileal part of small intestine and kidney was
collected and RNA was extracted using Nucleospin RNA Mini kit (Macherey Nagel) according to
manufactures instructions. Isolated RNA was subjected to reverse transcription reaction using
RevertAid RT Reverse Transcription Kit (Thermo). Quantitative PCR reaction using Syber -green
(Roche) and following set of primers was used; *Il17a* (pF: TGACCCCTAAGAAACCCCCA, pR:
TCATTGTGGAGGGCAGACAA), *Il17f* (pF: GAAGGCTGGGAACTGTCCTC , pR:

918 CGGAGTTCATGGTGCTGTCT), Il22 (pF:TTGACACTTGTGCGATCTCTGA, pR: AAAGGTGCGGTTGACGATGT), 919 Casc3 housekeeping and as gene (pF: TTCGAGGTGTGCCTAACCA, pR: GCTTAGCTCGACCACTCTGG). The relative gene 920 921 expression was calculated using method described elsewhere⁶⁵.

922

923 Mouse model of Oropharyngeal candidiasis (OPC)

924 Mice were first primed by repeated injection of 10^6 HKCA particles every second day for the 925 duration of two weeks. Then, previously established protocol was followed³⁵. Briefly, mice were 926 sedated and exposed to *C. albicans* orally for 1.5 hours using cotton swabs soaked with *C. albicans* 927 diluted in PBS (10^7 particles/ml). Mice were analyzed five days post oral inoculation.

928

929 Determination of C. albicans CFU

0.2g of the tongue, esophagus, ileal part of small intestine and kidney tissue was mechanically
disrupted in PBS and plated on Sabouraud Dextrose Agar (Merck) in two dilutions. Number of
colonies was calculated after 24 and 48 hours. The CFU were recalculated per g of original tissue.

933

934 Statistical analysis

935 Unless indicated otherwise, statistical significance was assessed using two-tailed Student's t test

936 calculated in GraphPad Prism program. For summarizing the p-value, following marks were used:

937 *** = p-value < 0.0001, ** = p-value < 0.001, * = p-value < 0.05, ns = not significant.

938

939 Access codes for all transcriptomics data

940 All RNA-seq data are available at GEO under accession number: GSE203158

941 Methods references

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Figure 1

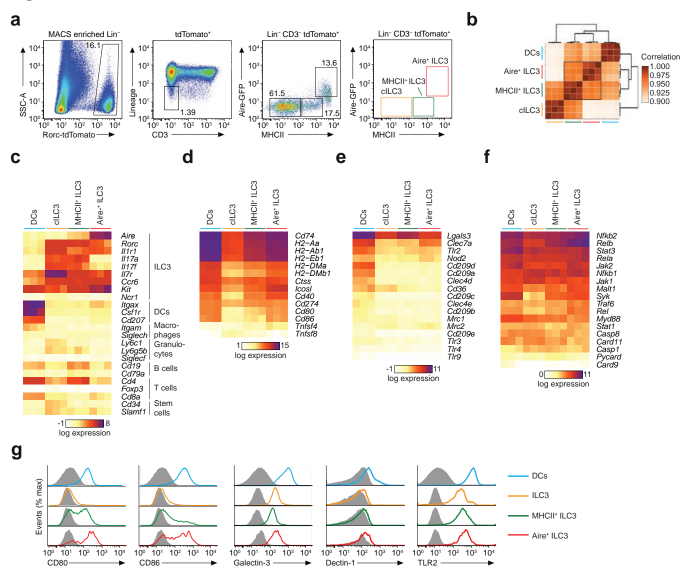
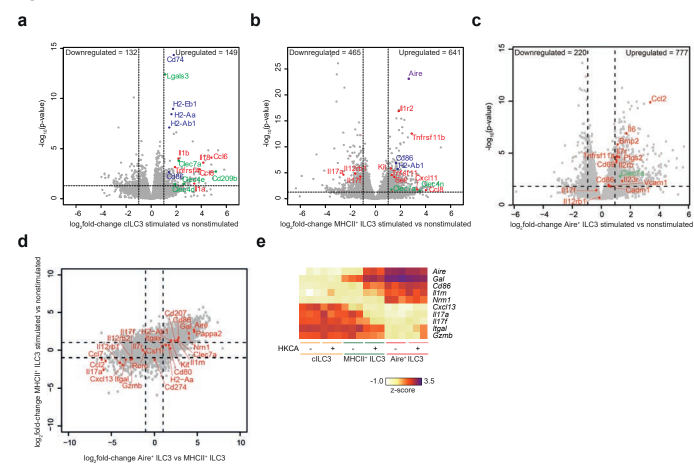
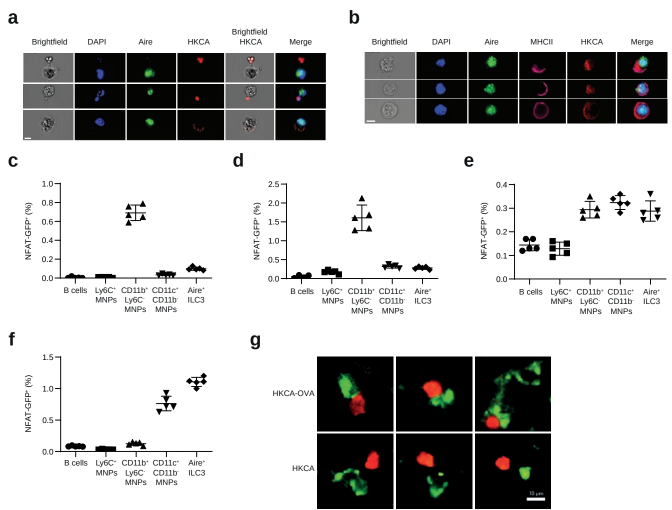


Figure 2



6

Figure 3



Aire-GFP OTII tdTomato

Figure 4

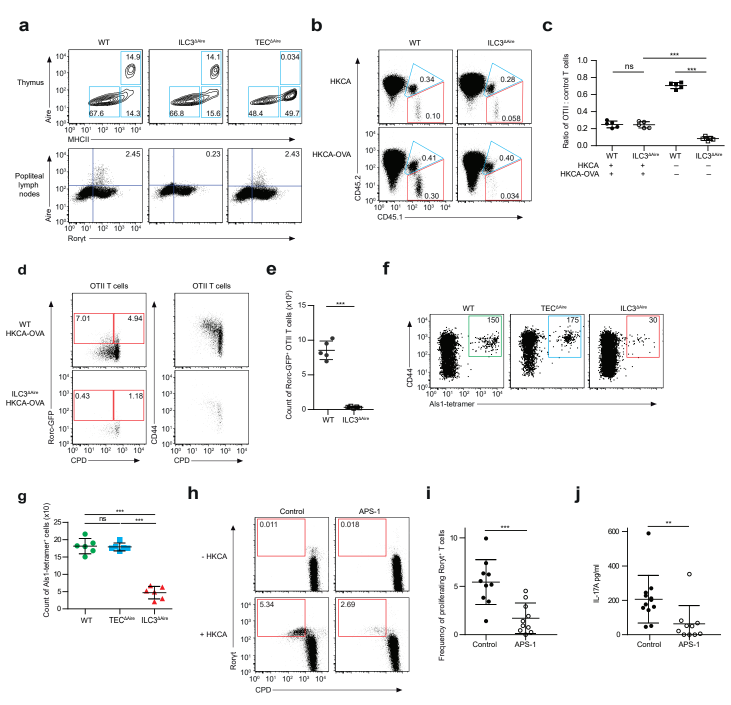


Figure 5

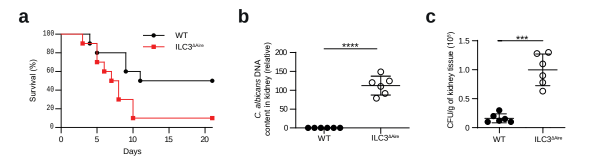


Figure 6

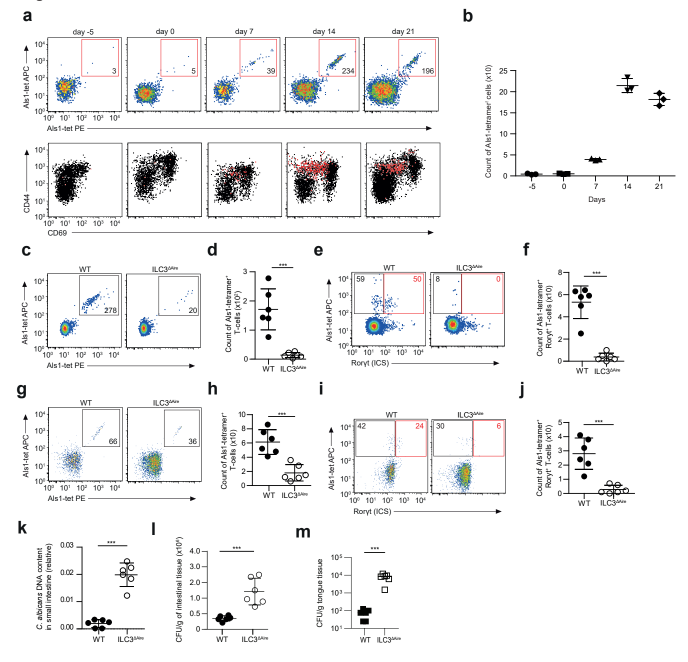
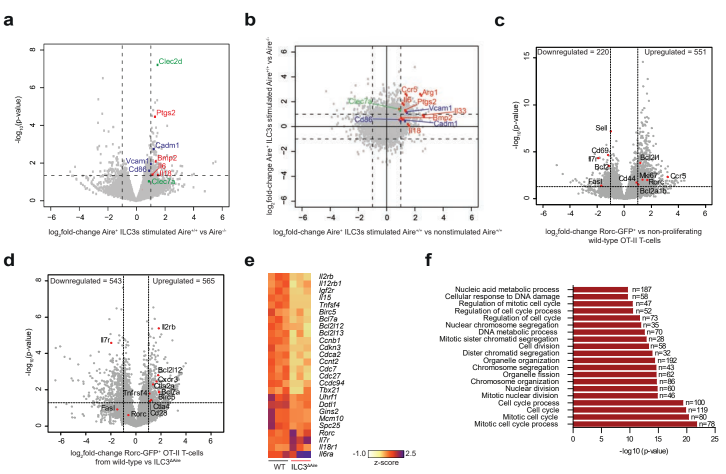
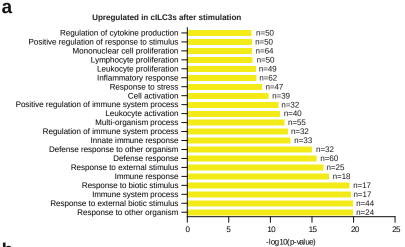


Figure 7

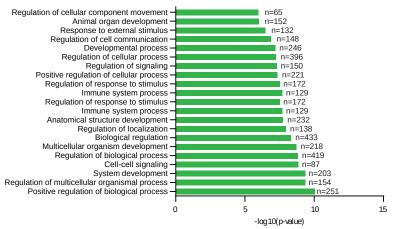


Extended Data Figure 1 (related to Figure 2A-C)



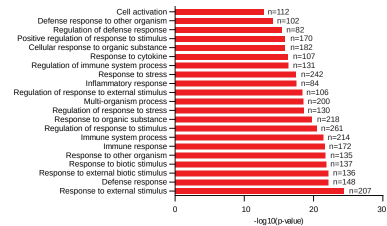
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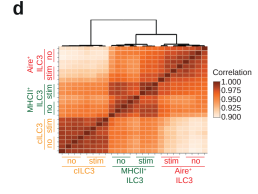
Upregulated in MHCII+ ILC3s after stimulation



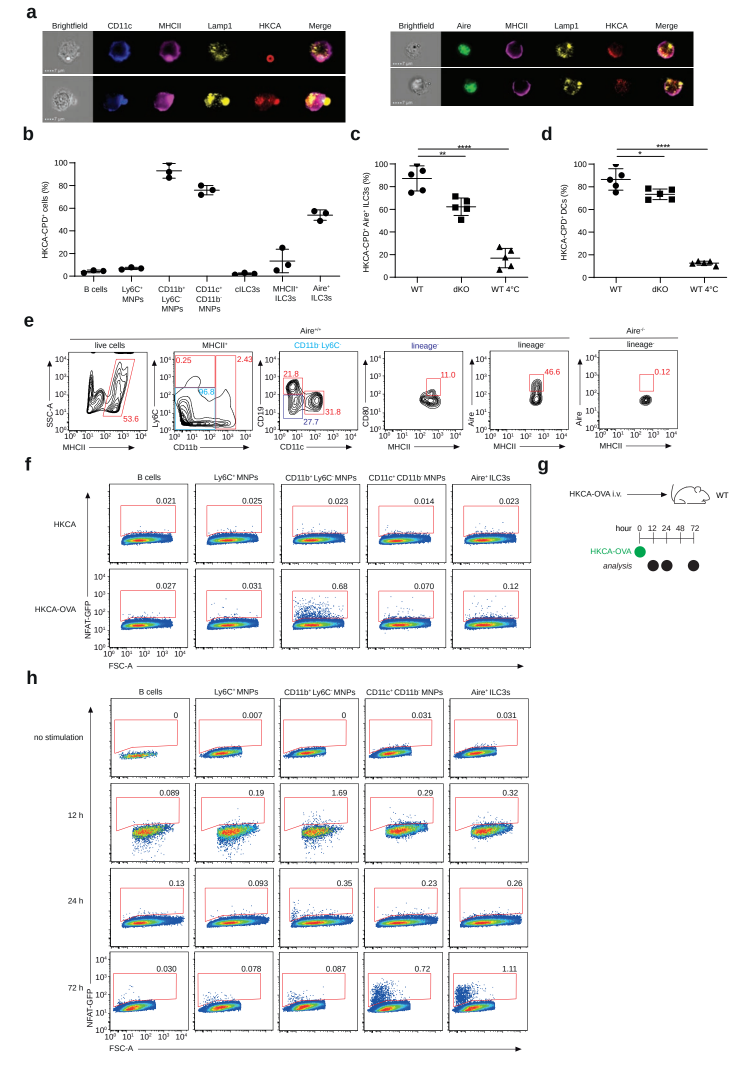
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Upregulated in Aire+ ILC3s after stimulation

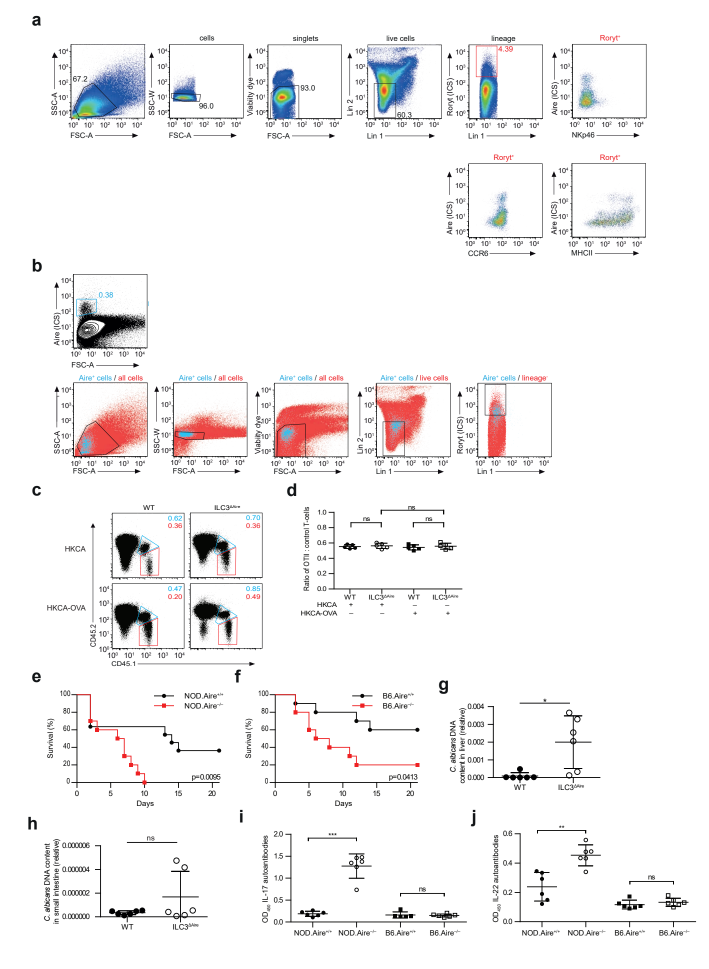




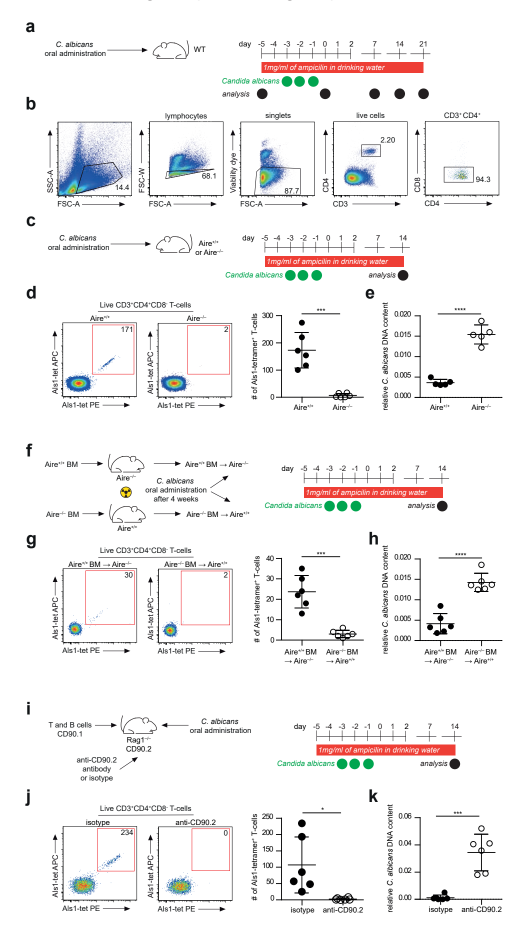
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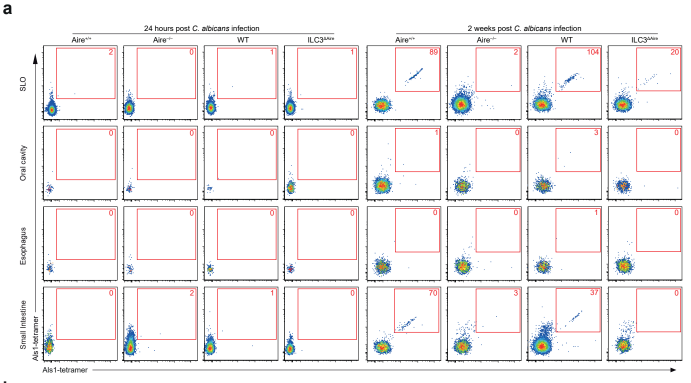
Extended Data Figure 3 (related to Figure 4 and 5)



Extended Data Figure 4 (related to Figure 6)



Extended Data Figure 5 (related to Figure 6)

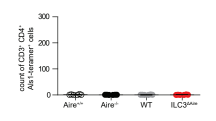




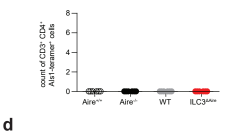
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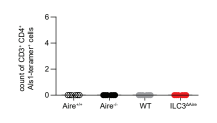




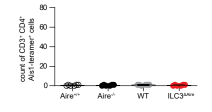
Oral cavity 24 hours post C.albicans colonization



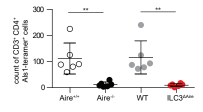
Esophagus 24 hours post C. albicans colonization



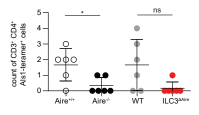
Small intestine 24 hours post C. albicans colonization



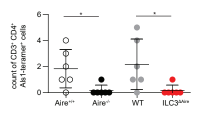
SLO 14 days post C.albicans colonization



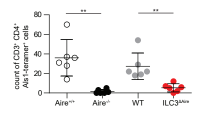
Oral cavity 14 days post C. albicans colonization



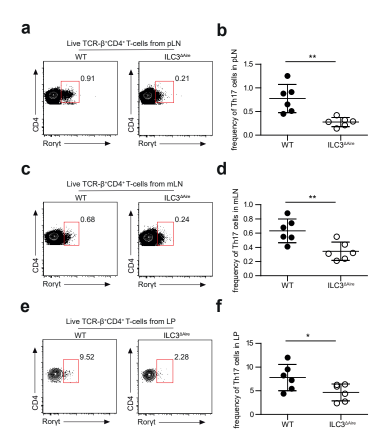
Esophagus 14 days post C. albicans colonization



Small intestine 14 days post C. albicans colonization



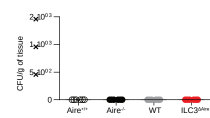
Extended Data Figure 6 (related to Figure 6)



Extended Data Figure 7 (related to Figure 6)

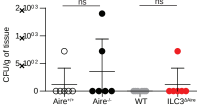
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Kidney 24 hours post C.albicans colonization



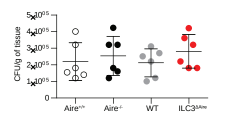


Kidney 14 days post C.albicans colonization

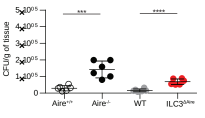


b

Oral cavity 24 hours post C.albicans colonization

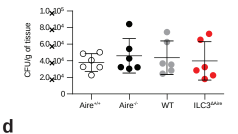


Oral cavity 14 days post C. albicans colonization

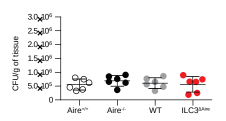


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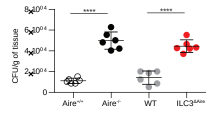
Esophagus 24 hours post C. albicans colonization



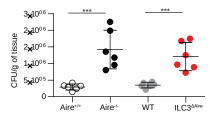
Small intestine 24 hours post C. albicans colonization



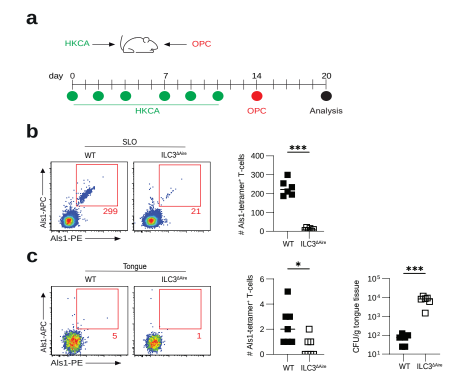




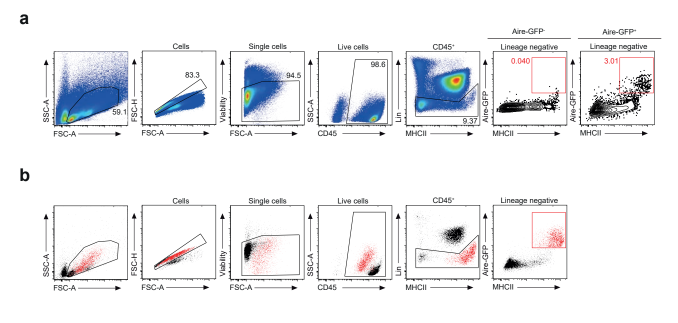
Small intestine 14 days post C. albicans colonization



Extended Data Figure 8



Extended Data Figure 9 (related to Figure 7A-B)



Extended Data Figure 10 (related to Figure 7C, D)

