

Bystander immunotherapy as a strategy to control allergen-driven airway inflammation.

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1	Bystander immunotherapy as a strategy to control allergen-driven airway inflammation
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44 Abstract

45

Background: Allergic asthma is a chronic inflammatory disease that is characterized by airway hyper responsiveness (AHR), infiltration of Th2 cells in lungs and high levels of circulating IgE. Allergen-specific immunotherapy (SIT), in which patients are rendered tolerant by exposure to steadily increasing doses of the allergen, is the only curative treatment to date. Unfortunately, SIT is not suitable for treating multi-sensitized patients, and some allergens are too immunogenic to be used in desensitization protocols.

52

53 **Objective:** To investigate whether, and to understand how, regulatory $CD4^+$ T cells (T_{reg}) 54 specific for a third-party "drug" antigen could control allergic immune responses and lung 55 inflammation.

56

57 **Methods:** Mice were tolerized to ovalbumin (OVA), sensitized to ragweed, and eventually 58 challenged with aerosols of ragweed alone or ragweed and OVA together. Animals were then 59 monitored for cardinal features of allergic asthma including AHR and infiltration of Th2 cells in 60 lungs. In additional experiments aimed at elucidating the mechanisms of OVA-induced 61 suppression, OVA-tolerized mice were sensitized with the LACK model antigen, challenged with 62 LACK alone or LACK and OVA together, and LACK-specific T cells were visualized by flow 63 cytometry.

64

Results: In both the ragweed and the LACK model, allergen-induced airway inflammation and AHR were strongly reduced in mice challenged with both the allergen and OVA compared to mice challenged with the allergen alone. OVA-induced protection did not result from competition between OVA and the allergen, was mediated by OVA-specific CD25⁺ T_{reg} , required both CTLA-4 and ICOS signaling, and was partially dependent on IL-10. Bystander suppression was associated with reduced proliferation of allergen-specific Th2 cells and decreased numbers of airway DC migrating to the lungs.

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73 **Conclusion:** Our results demonstrate that T_{reg} specific for a third-party drug antigen could 74 control allergic immune responses and lung inflammation when re-stimulated *in vivo*. Clinical implications: This study paves the way for the development of a novel therapeutic
 strategy that could control allergen-specific Th2 responses in patients with allergic asthma.

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78 **Capsule summary**: This study provides the proof of concept that bystander suppression 79 mediated by T_{reg} specific for a third-party drug antigen could be used as an efficient strategy to 80 control allergen-specific Th2 cells and asthma symptoms in allergic individuals.

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Key words: asthma, immune tolerance, airway inflammation, Th2, T_{reg}, specific immunotherapy
 83

- 84 Abbreviations: Ag, antigen; APC, antigen-presenting cells; AHR, airway hyperresponsiveness;
- 85 Alum, aluminium hydroxide; BAL, bronchoalveolar lavages; BALF, bronchoalveolar lavage
- 86 fluids; **BSA**, bovine serum albumin; **CFSE**, carboxyfluorescence diacetate succinimidly ester;
- 87 **DCs**, dendritic cells; **ELISA**, enzyme-Linked immunosorbent assay; **FACS**, fluorescence
- 88 activated cell sorter; FITC, fluoresceinisothiocyanate; LACK, Leishmania homolog of receptors
- 89 for activated c kinase antigen; LN, lymph nodes; Ig, immunoglobulin; IFN, interferon; IL,
- 90 interleukin ; i.p., intraperitoneal ; i.n., intranasal; MCH, methacholine; MedLN, mediastinal LN;
- 91 **OVA**, ovalbumin; **PCR**, polymerase chain reaction; **PE**, phycoerythrin; **PBS**, phosphate-buffered
- saline; **PLN**, popliteal LN; **RNA**, ribonucleic acid; **SIT**, specific immunotherapy; **T**_{eff}, effector T
- 93 cells; **Tg**, transgenic; **Th**, T helper ; **T**_{reg}, regulatory T cells; **WT**, wild type
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95 Introduction

96 Allergic diseases affect up to 30% of the population and their prevalence has steadily 97 increased in recent decades probably due to numerous changes in the environment. Among 98 allergic diseases, asthma is a chronic inflammation of the lungs caused by an inappropriate 99 immune response to a single or multiple airborne allergens. This pathology has a substantial 100 economic burden for which the only curative and specific method of treatment to this day is 101 allergen-specific immunotherapy (SIT). SIT involves the administration by either subcutaneous 102 injection or mucosal application of increasing doses of the allergen to which the patient is allergic 103 to. Unfortunately, SIT is unsuitable for treating multi-sensitized patients, and some allergens are 104 too immunogenic to be used in desensitization protocols.

105 SIT induces a state of peripheral tolerance characterized mainly by the generation of 106 allergen-specific T_{reg}, suppressed effector cell proliferation and cytokine production against major allergens ^{1, 2}. Indeed, IL-10 and/or TGF- β producing T_{reg} are the key factors for specific 107 immunotherapy in humans, considered as a model of tolerance induction $^{3-5}$. It has been shown 108 109 that treating naive (non-sensitized) mice with OVA aerosols leads to IgE-unresponsiveness to OVA ⁶, and induces the development of OVA-specific T_{reg} which prevented the development of 110 asthma upon subsequent sensitization and challenge with OVA⁷. Moreover, lung draining lymph 111 112 node (LN) dendritic cells (DC) first encountering an inhaled antigen transiently produced IL-10⁷. These phenotypically mature DC induce the development of $CD4^+$ T_{reg} that also produce high 113 amounts of IL-10 $^{8}\!\!.$ TGF- β -expressing T_{reg} also play a role for tolerance induction to inhaled 114 antigens ^{5, 9}. Interestingly, it has been suggested that tolerance induced by SIT was not only 115 limited to the administered allergen but also conferred protection against other allergens ^{10, 11}. 116 117 This non-specific and beneficial action of SIT could be explained by the activation of bystander 118 T_{reg}.

Bystander immunosuppression was first described by Bullock *et al.* as a process in which antigen (Ag)-specific T_{reg} inhibit T effector cell responses directed to both the targeted Ag and to a co-localized third-party Ag ¹². Due to the requirement that the tolerogen and the Ag have to be physically linked, i.e. presented by the same antigen-presenting cell (APCs), in order for suppression to occur, the terms « linked suppression » or « linked recognition » were coined. In the field of oral tolerance, the term « bystander suppression » was introduced to describe an inhibition of a T cell memory response as a result of a regulatory response to an unrelated but

- 126 colocalized tolerogen $^{13, 14}$. In bystander suppression, the tolerogen and third-party Ag do not 127 need to be presented by the same APC. In this case, soluble mediators induce suppression of the 128 response directed to the third-party Ag $^{13, 15-17}$.
- 129 In the present study, we have sought to investigate whether T_{reg} specific for a third-party 130 "drug" antigen could control allergic immune responses and lung inflammation in mice. To this 131 aim, mice were tolerized to OVA, sensitized to ragweed, and eventually challenged with aerosols 132 of ragweed alone or ragweed and OVA together.
- 133

134Results

135

Exposure to OVA aerosols inhibits ragweed- and LACK-induced allergic asthma in OVA tolerized mice

138 To determine whether T_{reg} specific for a third-party antigen could control allergic airway 139 inflammation in mice, mice were tolerized to OVA through intranasal administrations, sensitized 140 to ragweed and further exposed to aerosols of ragweed alone or ragweed and OVA together (Fig 141 $\mathbf{1}, \mathbf{A}$). When compared to the group challenged with ragweed only, mice exposed to both ragweed 142 and OVA showed decreased AHR (Fig 1, B) and reduced numbers of total cells, eosinophils and 143 lymphocytes in broncho alveolar lavage fluids (BALF) (Fig 1, C). We further analyzed airway-144 infiltrating T cells for expression of the Th2- marker, T1/ST2 (IL-33Ra). Compared to the mice 145 challenged with ragweed alone, the number of $T1/ST2^+$ CD4⁺ T cells was reduced in the BALF 146 of mice challenged with ragweed and OVA aerosols (Fig 1, D). The amounts of IL-5 and IL-13 147 in the lungs were also reduced upon challenge with ragweed and OVA while IFN- γ levels 148 remained low and similar in both groups (Fig 1, E). This phenomenon was not observed in mice 149 than have not been tolerized to OVA prior to ragweed sensitization (Supplementary Figure 1) 150 further supporting a role for OVA-specific T_{reg} in this phenomenon and ruling out the possibility 151 that it could result from competition between OVA and ragweed for antigen presentation.

152 To generalize our findings and to further dissect the mechanisms involved, we switched to 153 another experimental model in which BALB/c mice were sensitized to the model antigen LACK. 154 tolerized to OVA through intranasal administration, and further exposed to aerosols of LACK 155 alone, or LACK and OVA (Fig 2, A). As observed with ragweed, mice exposed to both 156 LACK/OVA showed decreased AHR (Fig 2, B), and reduced total numbers of cells, eosinophils 157 and lymphocytes in BALF (Fig 2, C). Compared to control mice challenged with LACK alone, 158 both the frequency and the number of T1/ST2⁺ CD4⁺ T cells were reduced in the BALF of mice 159 challenged with LACK/OVA aerosols (Fig 2, D). The amounts of IL-4, and IL-13 in the BALF 160 were also reduced upon challenge with LACK/OVA aerosols while IFN-y levels remained low 161 and similar in both groups (Fig 2, E). The lungs of LACK/OVA-challenged mice contained less 162 IL-4-, and IL-5- secreting CD4⁺ T cells than those challenged with LACK only as demonstrated 163 by intracellular cytokine staining of lung cells upon LACK restimulation (Fig 2, F). In contrast, 164 the amounts of LACK-specific -IgE and -IgG1 were not affected by additional OVA exposure 165 (**Fig 2**, *G*). Altogether, our results suggested that exposure to OVA aerosols could inhibit allergic 166 airway inflammation and allergen-specific Th2 immune responses in mice that have been 167 tolerized with OVA prior to sensitization and challenge with either ragweed or LACK.

168

The inhibition of LACK-induced airway inflammation in OVA-tolerized mice is mediated by OVA-specific CD25⁺ T_{reg}

171 Previous experiments have shown that the intranasal administration of OVA to BALB/c mice induces OVA-specific T_{reg} that express CD25. To investigate whether these cells were 172 173 responsible for the inhibition of LACK-induced airway inflammation in OVA-tolerized LACK-174 sensitized mice, CD4⁺ T cells were purified from the spleen of mice that have been previously 175 tolerized to OVA or bovine serum albumine (BSA), or treated with PBS. Cells were injected into 176 LACK-sensitized recipients that were further challenged with both LACK and OVA (Fig 3, A). 177 Compared to mice injected with CD4⁺ T cells purified from BSA-tolerized or PBS-treated 178 donors, mice injected with CD4⁺ T cells from OVA-tolerized mice exhibited decreased AHR 179 (Fig 3, B), reduced numbers of total cells, eosinophils and lymphocytes (Fig 3, C) and $T1/ST2^+$ $CD4^+$ T cells in BALF (**Fig 3**, *D*), and reduced numbers of IL-4 and IL-5-secreting $CD4^+$ T cells 180 181 in lungs (Fig 3, E). In striking contrast, no inhibition of LACK-induced airway inflammation or LACK-specific Th2 immune responses were observed when CD4⁺ T cells from OVA-tolerized 182 183 mice depleted of CD25⁺ cells prior to injection into LACK-sensitized mice. Lastly, both airway 184 inflammation and LACK-specific Th2 immune responses were inhibited when recipient mice 185 were injected with CD25⁺ CD4⁺ T cells purified from the spleen of OVA-tolerized mice (Fig 3, 186 F-I). Therefore, the inhibition of LACK-induced airway inflammation that was observed in 187 OVA-tolerized mice upon challenge with OVA relied on the activation of OVA-specific CD25⁺ 188 T_{reg}.

189

190 CTLA-4, ICOS, and IL-10 dependency of OVA-specific T_{reg} in the control of LACK 191 induced asthma

192 T_{reg} mediate suppression through various mechanisms including the secretion of inhibitory 193 cytokines such as IL-10, the induction of cytolysis, metabolic disruption and the inhibition of 194 antigen presentation by dendritic cells (DCs) through a CTLA-4-dependent mechanism ¹⁸.

195 Furthermore, it was recently shown that ICOS mediated the generation and function of CD25⁺ CD4⁺ FoxP3⁺ T_{reg} conveying respiratory tolerance ¹⁹, and that ICOS expression defined a subset 196 of IL-10 secreting T_{reg}^{20} and was required for the production of IL-10 by these cells ²¹. 197 198 Therefore, we sought to elucidate the role of CTLA-4, ICOS and IL-10 in the inhibition of 199 LACK-induced airway inflammation induced by OVA aerosols. To this aim, LACK-sensitized 200 OVA-tolerized mice were challenged with both LACK and OVA and treated or not with blocking 201 mAbs directed to CTLA-4, ICOS or IL-10R (Fig 4, A). CTLA-4 blockade prevented the 202 inhibition of AHR induced by OVA aerosols, as well as the reduction in the number of total cells, 203 eosinophils, lymphocytes and T1/ST2⁺ Th2 cells in BALF, and IL-4- and IL-5-producing LACK-204 specific CD4⁺ in lungs (Fig 4, *B-D*). Likewise, blocking ICOS/ICOSL interactions abolished the 205 protection that was induced by OVA aerosols in OVA-tolerized LACK-sensitized mice but did 206 not have any detectable effect in mice that have not been tolerized to OVA (Fig 4, F, G). In 207 contrast to anti-CTLA-4 and anti-ICOS mAbs that restored both AHR and allergic airway 208 inflammation in OVA-tolerized mice exposed to OVA, anti-IL-10R mAbs restored AHR but not 209 allergic airway inflammation (Fig 4, F, G). Therefore, the inhibition of AHR and airway 210 inflammation that was induced by OVA aerosols in OVA-tolerized mice upon sensitization and 211 challenge with LACK was dependent on both CTLA-4, ICOS and partially on IL-10.

212

213 **OVA-specific** T_{reg} inhibited the proliferation of LACK-specific Th2 cells and the subsequent 214 airway inflammation

215 To further dissect the molecular mechanisms by which OVA-specific T_{reg} inhibited LACK-216 induced allergic asthma, CD4⁺ T cells from Thy1.1⁺ LACK-specific WT15 TCR transgenic mice 217 were differentiated in vitro under Th2 polarizing conditions, labeled with CFSE, and injected into 218 OVA- or BSA- tolerized recipients before being challenged with LACK/OVA (Fig 5, A). As 219 compared to BSA-tolerized mice, OVA-tolerized animals injected with LACK-specific Th2 cells 220 exhibited decreased AHR (Fig 5, B), and reduced numbers of eosinophils and lymphocytes in the 221 BALF (Fig 5, C). In addition, both the frequency and the number of donor Th2 cells were 222 reduced in the BALF, lung and MLN, but not in the blood of OVA-tolerized mice compared to 223 BSA-tolerized mice (Fig 5, D). Furthermore, LACK-specific Th2 cells had undergone more 224 divisions in BSA-tolerized mice than in OVA-tolerized mice (Fig 5, E) further suggesting that 225 OVA aerosols inhibited the proliferation of LACK-specific Th2 cells possibly at the level of 226 antigen presentation. To directly test this latter hypothesis, LACK-specific Th2 cells were 227 injected into either OVA- or BSA-tolerized mice, and further challenged with both LACK and 228 OVA. Cells were then purified from the mediastinal LN (MedLN) and distal popliteal LN (PLN) 229 and incubated with CFSE-labeled LACK-specific WT15 CD4⁺ T cells to assess their ability to 230 induce proliferation. As expected, PLN cells did not induce T cell proliferation (Fig 5, F).. 231 Furthermore, LACK-specific T cells proliferated more vigorously when incubated with MedLN 232 cells from BSA-tolerized mice than when incubated with MedLN cells from OVA-tolerized mice 233 (Fig 5, F). To investigate whether this phenomenon resulted from quantitative (i.e. different 234 number of LACK-loaded DCs in OVA-tolerized and BSA-tolerized mice) or qualitative 235 differences (i.e. similar number of LACK-loaded DCs in OVA-tolerized and BSA-tolerized mice 236 but different ability to induce T cell proliferation), we administered FITC-labeled latex beads to 237 OVA- and BSA-tolerized mice at the time of the challenge with LACK and OVA, and we measure the frequency of bead⁺ DCs in MedLN 16 hrs later. While bead⁺ DCs were readily 238 239 detected in the MedLN of both OVA- and BSA-tolerized mice and expressed similar surface 240 levels of CD80, CD86, OX40L and MHC class II molecules (not shown), the frequency of bead⁺ 241 DCs was decreased from 4.7 \pm 0.7 % in BSA-tolerized mice to 1.4 \pm 0.3 % in OVA-tolerized animals (Fig 5, H). Altogether, our data suggested that OVA-specific T_{reg} inhibited the 242 243 proliferation of LACK-specific Th2 cells by preventing the migration of airway DCs to MedLN. 244

245 **Discussion**

246 In his paper, we have demonstrated that OVA-specific T_{reg} could inhibit allergic airway 247 inflammation induced by sensitization and challenge with ragweed. OVA-specific T_{reg} were also 248 efficient to protect mice from allergic airway inflammation induced by the immunodominant 249 LACK antigen further validating our finding to second allergen. In addition, BSA-tolerized mice 250 were protected from LACK-induced allergic asthma when challenged with both LACK and BSA 251 further suggesting that the phenomenon that we have observed was not restricted to OVA-252 specific T_{reg} (data not shown). Interestingly, the phenomenon that we have described in this paper 253 may explain the protective effects of SIT against unrelated allergens other than the one primarily targeted as reported in several epidemiological studies ^{10, 11, 22}. 254

255 In the LACK model, we have found that the number of allergen-specific Th2 cells in 256 BALF, the frequency and number of IL-4 and IL-5 producing CD4⁺ T cells as well as the IL-4, 257 IL-5, and IL-13 amounts in lungs were decreased in mice challenged with both LACK and OVA 258 compared to mice challenged with LACK alone. OVA-induced protection did not result from a 259 competition between OVA and LACK for antigen presentation because it was only observed in 260 mice that have been previously tolerized with OVA. In contrast, experiments in which CD4⁺ T 261 cells were purified from the spleen of PBS-treated, BSA- or OVA-tolerized mice and injected 262 into LACK-sensitized recipient mice showed that protection was antigen-specific and mediated 263 by T_{reg}.

264 Further experiments showed that CTLA-4 was absolutely required for OVA-induced 265 suppression of LACK-mediated asthma. While we did not elucidate the mechanisms by which 266 CTLA-4 acted, CTLA-4 was shown to be essential for the function of naturally occurring T_{reg} which constitutively express this molecule. Indeed, CTLA-4 deficiency in T_{reg} impaired both 267 268 their *in vivo* and *in vitro* suppressive function. In addition, CTLA-4 prevented CD28 signaling in 269 effector T cells (T_{eff}) by competing with CD80 and CD86, and/or induced the synthesis of the 270 tryptophan catabolizing enzyme indoleamine 2,3-dioxygenase (IDO) by the APCs leading to T-271 cell suppression by both local depletion of tryptophan and induction of apoptosis via tryptophan catabolites ²³. While we were unable to detect increased death of LACK-specific T cells in OVA-272 273 tolerized mice challenged with both OVA and LACK (not shown), we cannot rule out the 274 possibility that this phenomenon was partially responsible for OVA-induced protection.

275 Blocking ICOS-ICOSL interactions resulted in the same effect as CTLA-4 inhibition, a result in agreement with a previous study showing that ICOS^{-/-} T_{reg} did not confer protection upon 276 transfer to asthmatic mice demonstrating a crucial role of ICOS in their suppressive function ¹⁹. 277 278 As ICOS was shown to define a subset of IL-10-producing T_{reg} , we next investigated whether 279 OVA-induced suppression of the disease required IL-10. IL-10R blockade prevented OVA-280 induced suppression of AHR, but did not affect LACK-induced inflammation in our model. It 281 remains to be determined whether IL-10 is produced by the $CD25^+$ T_{reg} or by other cells as previously described Kearley et al.²⁴. Taken together, these data underline the multiple and 282 283 complex effects of $CD25^+$ T_{reg}. While these cells have been proposed to act via cell contact-284 dependent mechanisms in vitro, these cells have been proposed to work through various 285 mechanisms including inhibitory cytokines and non-cytokine-dependent mechanisms in vivo, depending on the experimental conditions²⁵. Our findings also suggest that inflammation and 286 287 AHR can be uncoupled and are in agreement with previous studies that demonstrate that effects on inflammation are not always predictive of AHR changes ²⁶⁻²⁹. Indeed, this might also hold true 288 289 in human asthma, in which anti-IL-5 mAb treatment reduced blood and lung eosinophilia but did not affect lung function ³⁰. 290

291 Adoptive transfer of CFSE-labeled LACK-specific Th2 cells into OVA-tolerized recipient 292 mice demonstrated that OVA-specific T_{reg} significantly reduced the number of LACK-specific 293 Th2 cells in the BALF, lung and MLN by affecting their proliferation. Imaging data in mice have 294 showed that T_{reg} do not directly interact with T_{eff} but rather with DC, altering the latter and diminishing subsequent DC-T_{eff} cells conjugate formation *in vivo*^{31, 32} a phenomenon that could 295 296 explain our results. In addition, Derks and colleagues have envisioned two hypotheses of APC function in bystander suppression ³³: a passive APC model, in which the APCs would present 297 298 MHC-peptide to the T_{reg}, stimulating them to produce immunosuppressive cytokines that would 299 further binds their cognate receptors on the third-party T_{eff}, or an active APC model, in which the 300 APCs would propagate regulatory effects from the T_{reg} to the T_{eff} through various APC products. 301 These two hypotheses remain to be tested in our model.

302 Our results demonstrate that T_{reg} specific for a third-party drug antigen could control 303 allergic immune responses and lung inflammation when re-stimulated *in vivo*. This study paves 304 the way for the development of a novel therapeutic strategy that could control allergen-specific 307

308 FIGURE LEGENDS

309 Figure 1. AHR, airway inflammation and cytokine levels in ragweed-sensitized, OVA-310 tolerized mice upon ragweed challenge. (A) Experimental protocol. Mice were treated with 311 three i.n. injections of OVA, and four i.n. injections of ragweed. Mice were then either 312 challenged with ragweed or ragweed/OVA, and analyzed one and two days after the last i.n. 313 injection. (B) AHR. Whole body plethysmography in mice exposed to ragweed (filled squares), 314 ragweed/OVA (empty circles), or PBS (crosses). (C) Number and phenotype of BALF cells. 315 BALF cells were analyzed by FACS in mice exposed to ragweed only (black bars), to both 316 ragweed and OVA aerosols (empty bars) or to PBS (grey bars). Eosinophils, E; neutrophils, N; 317 lymphocytes, L; macrophages, M. (D) Frequency and number of Th2 cells in the airways. BALF 318 cells were stained with CD4 and T1/ST2 mAbs and analyzed by FACS. (E) Cytokine levels in 319 lung cells. IL-4, IL-5 and IFN- γ levels were assessed by ELISA after *in vitro* stimulation with 320 LACK protein (0.1mg/ml), anti-CD28 (1µg/ml) and brefeldin A (5µg/ml). Data are expressed as mean \pm s.e.m. of 2 experiments with n=8 mice per group. n.s., non significant; * P<0.05; ** 321 322 *P*<0.01.

323 Figure 2. AHR, airway inflammation and cytokine levels in OVA-tolerized mice exposed to 324 both OVA and LACK aerosols. (A) Experimental protocol. Mice were sensitized with two 325 intra-peritoneal injections of LACK in Alum, treated with three i.n. injections of OVA, and 326 challenged daily for 5 days with LACK aerosols or LACK/OVA aerosols. Mice were analyzed one and two days after the last aerosol. (B) AHR. Whole body plethysmography (right), and 327 328 dynamic lung resistance and compliance (left) were monitored in mice exposed to LACK 329 aerosols (filled squares), LACK/OVA aerosols (empty circles) or PBS (crosses, dashed line). (C) 330 Number and phenotype of BALF cells. BALF cells were counted and analyzed by FACS in mice 331 exposed to PBS (dashed bars), LACK (black bars) or LACK/OVA aerosols (empty bars). Data 332 show the number of eosinophils (E), neutrophils (N), lymphocytes (L), macrophages (M). (D) 333 Frequency and number of Th2 cells in the airways. BALF cells were stained with anti-CD4, CD3 334 and T1/ST2 mAbs and analyzed by FACS. Data show representative FACS profiles, numbers 335 indicate the mean frequency \pm s.e.m and histograms show the absolute numbers of T1ST2⁺ CD4⁺ 336 T cells for the indicated groups. (E) Cytokine levels in BALF. Mice were analyzed for IL-4, IL-5, 337 IL-13 and IFN- γ by cytometric bead array (CBA). (F) Cytokine secretion by lung CD4⁺ T cells.

338 IL-4, and IL-5-secreting CD4⁺ T cells were assessed by FACS after *in vitro* stimulation with 339 LACK protein (0.1mg/ml), anti-CD28 (1µg/ml) and brefeldin A (5µg/ml). Data show 340 representative FACS profiles, numbers indicate the mean frequency \pm s.e.m and histograms show 341 the absolute numbers for the indicated groups. (G) Immunoglobulins. Levels of serum LACK-342 specific -IgE and -IgG1 were assessed in mice upon challenge with LACK or LACK/OVA 343 aerosols. All data show either individual mice with bar indicating the mean, with n = 6 mice per group pooled from four different experiments. n.s., non significant, * P<0.05, ** P<0.01, *** 344 345 *P*<0.0001.

346 Figure 3. AHR, airway inflammation and cytokine levels in mice injected with CD4⁺ T cells 347 from OVA-tolerized mice. (A) Experimental protocol. Mice were sensitized with two i.p. injections of LACK in Alum, and injected 9 d later with 4 X 10⁶ CD4⁺ T cells (A-E), or 4 X 10⁶ 348 $CD25^{-}CD4^{+}$ T cells (**F-I**) or 1.5 X 10⁶ CD25⁺CD4⁺ T cells (**F-I**) prepared from the spleen of mice 349 350 exposed to OVA, BSA, or PBS. Sensitized mice were then challenged with LACK/OVA aerosols 351 for five days and analyzed one and two days after the last aerosol. (**B**, **F**) AHR. Whole body 352 plethysmography was monitored in the indicated mice challenged to LACK/OVA aerosols in 353 response to increased doses of inhaled methacholine. Control mice (vehicle) were sensitized with 354 LACK, non-transferred and challenged with PBS. (C, G) Number and phenotype of BALF cells. 355 BALF cells were analyzed by FACS for the number of eosinophils (E), neutrophils (N), 356 lymphocytes (L), macrophages (M). (D, H) Frequency and number of Th2 cells in the airways. 357 BALF cells were stained with anti-CD3, -CD4 and T1/ST2 mAbs and analyzed by FACS. (E, I) 358 Cytokine secretion by lung CD4⁺ T cells. IL-4, and IL-5-secreting CD4⁺ T cells were assessed by 359 FACS after in vitro stimulation with LACK protein (0.1mg/ml), anti-CD28 (1µg/ml) and 360 brefeldin A (5µg/ml). Data show numbers of IL-4 and IL-5-secreting CD4⁺ T cells in lungs. All data show either individual mice or are expressed as mean \pm s.e.m, with n = 5 mice per group 361 362 pooled from two experiments. P values have been calculated by comparing OVA-tolerized mice to BSA-tolerized mice, ns, non significant, * P < 0.05, ** P < 0.01, *** P < 0.001. 363

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Figure 4: AHR and airway inflammation in mice treated with anti-CTLA-4, anti-IL-10R, or anti-ICOS. (A) Experimental protocol. Mice were sensitized with two i.p. injections of LACK in 367 Alum, treated with three i.n. injections of OVA, treated or not with anti- CTLA-4 mAb, -IL-10R, 368 -ICOS, or IgG1 isotype mAb at the indicated time, and challenged daily for 5 days with LACK or 369 LACK/OVA aerosols. Mice were analyzed one and two days after the last aerosol. (B) and (F) 370 AHR. Whole body plethysmography in response to increasing doses of inhaled methacholine in 371 the indicated groups of mice. (C) and (G) Number and phenotype of BALF cells. BALF cells 372 were analyzed by FACS in the indicated groups of mice. Eosinophils, E; neutrophils, N; 373 lymphocytes, L; macrophages, M. (**D**). Number of Th2 cells in the airways. BALF cells were 374 stained with anti -CD3, -CD4 and T1/ST2 mAbs and analyzed by FACS. (E). IL-4, and IL-5-375 secreting CD4⁺ T cells were assessed by FACS after in vitro stimulation with LACK protein 376 (0.1 mg/ml), anti-CD28 (1 ug/ml) and brefeldin A (5 ug/ml). Data are expressed as mean \pm s.e.m 377 of 3 experiments with n=6 mice per group. n.s., non significant; * P<0.05; ** P<0.01; *** 378 *P*<0.0001.

379 Figure 5. AHR, airway inflammation, cytokine levels and T cell proliferation in OVA-380 tolerized mice injected with LACK-specific Th2 cells and exposed to LACK/OVA. (A) 381 Experimental protocol. Mice were treated with three i.n. injections of OVA or BSA, injected with 382 1.5 X 10⁶ CFSE-labeled Thy1.1⁺ LACK-specific Th2 cells. Mice were injected intranasally 2 d 383 later with LACK/OVA and analyzed 4 and 5 days later. (B) AHR. Whole body plethysmography 384 was monitored in mice tolerized to OVA (empty circles) or BSA (filled squares, and crosses) and 385 exposed to aerosols of LACK/OVA (full lines) or to PBS (dashed lines). Data are expressed as 386 mean \pm s.e.m. (C) Number and phenotype of BALF cells. BALF cells were analyzed by FACS in 387 the indicated groups of mice. Eosinophils, E; neutrophils, N; lymphocytes, L; macrophages, M. 388 Data are expressed as mean \pm SEM of 3 experiments with n=6-8 mice per group. (**D**) Numbers of Thy1.1⁺ LACK-specific Th2 cells in BALF, lung, and medLN and frequency in the blood of 389 390 BSA- (filled bars) or OVA- (empty bars) tolerized mice challenged with LACK/OVA aerosols. 391 (E) Representative plots of CFSE (left panels) and MFI of CFSE (right panels) of the indicated 392 mice. (F) In vitro antigen presentation assay. Whole cell suspensions prepared from the MedLN 393 and PLN of OVA- or BSA-exposed mice were incubated for 3 days with CFSE-labeled LACK-394 specific Th2 cells. Data show representative CFSE plots for the indicated mice with the 395 frequency of divided cells as mean \pm s.e.m of 2 experiments with n=6 mice per group. (G) Mice 396 underwent the same protocol as shown in panel A, but also received fluorescent latex beads

- 397 during LACK/OVA challenge. Beads⁺CD11c⁺ migratory DCs were analyzed by FACS in the 398 MedLN of BSA- (filled bars) or OVA- (empty bars) tolerized mice challenged with LACK/OVA 399 aerosols. n.s., non significant; * P<0.05; ** P<0.01; *** P<0.001.
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401 Supplementary Figure 1. AHR and airway inflammation in LACK-sensitized mice 402 challenged with LACK alone or LACK and OVA aerosols. (A) Experimental protocol. Mice 403 were sensitized with two i.p. injections of LACK in Alum, and challenged daily for 5 days with 404 LACK or LACK/OVA aerosols, or PBS. Mice were analyzed one and two days after the last 405 aerosol. (B) AHR. Whole body plethysmography in mice exposed to aerosols of LACK (filled 406 squares), LACK/OVA (empty circles) or PBS (crosses). (C) Number and phenotype of BALF 407 cells. BALF cells were analyzed by FACS in mice exposed to LACK only (black bars), to both 408 LACK and OVA aerosols (empty bars) or to PBS (grey bars). Eosinophils, E; neutrophils, N; 409 lymphocytes, L; macrophages, M. Data are expressed as mean \pm s.e.m. of 2 experiments with 410 n=8 mice per group. n.s., non significant.

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412 **METHODS**

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Mice. 6-week old BALB/c mice were purchased from The Centre d'Elevage Janvier (France) and housed under SPF conditions. LACK TCR transgenic mice (WT15 RAG-1 KO) on the BALB/c background as previously described ³⁴ were bred in our animal facility at the Institut de Pharmacologie Moleculaire et Cellulaire (Valbonne, France). In this study, WT15 transgenic mice were further crossed onto RAG-1^{-/-} Thy1.1^{+/+} BALB/c mice. All experimental protocols were approved by the local animal ethic committee.

Reagents. LACK recombinant protein was produced in *E. coli*, purified as described ³⁵, and
detoxified using an Endotrap column (Profos). Lipopolysaccharide contents in LACK protein
were below 5 ng/mg as determined using Limulus Amoebocyte Lysate (LAL) assay (Pierce).
LACK₁₅₆₋₁₇₃ peptide was purchased from Mimotopes. T1/ST2 mAbs were purchased from MD
Biosciences. Monoclonal antibodies to CD3, CD4, CD25, Thy1.1, CD11c, IA/IE, CD80, CD86,
IL-4 and IL-5 were purchased from BD Biosciences.

426 Induction of allergic asthma and tolerization to OVA. Sensitization was performed by 2 427 intraperitojneal (i.p.) injections of 10 µg of LACK in 2 mg of Aluminium hydroxide (Alum) (Pierce) at day 0 and 7. On days 12, 13 and 14, mice were tolerized to OVA by injecting i.n. 100 428 µg of LPS-free OVA (Profos) as described ⁷. From day 23 to day 27, mice were either exposed to 429 430 LACK (0.15%) or to LACK plus OVA (0.2%) aerosols (administered 8 hour apart) for 20 min using an ultrasonic nebulizer (Ultramed, Medicalia). Mice were analyzed on day 28 and 29 for 431 432 AHR and airway inflammation, respectively. When indicated, mice were injected with either 0.5 433 mg of anti-CTLA-4 mAb (9H10), anti-ICOS (17G9), or anti-IL-10R (1B1.3A) every other day 434 over the challenge period starting one day before the first aerosol. For ragweed-induced asthma, 435 mice were first tolerized to OVA by receiving i.n. injections of OVA on days 0, 1, and 2 and further sensitized to ragweed via i.n. administrations of 25 µg ragweed (Greer laboratories) on 436 437 days 11, 15, 19, and 23. Mice received a last challenge of ragweed on day 27 or ragweed and 438 OVA on days 26 and 27. Mice were analyzed on day 28 for AHR and on day 29 for airway 439 inflammation.

440 **Th2 cell transfers**. In some experiments, mice were first tolerized to OVA, BSA or PBS and 441 injected i.v on day 11 with LACK-specific CD4⁺ Th2 cells. Mice were challenged 24 hours later 442 with a single i.n. injection of LACK ($30 \mu g$) and OVA ($100 \mu g$), and assessed for AHR, and 443 immunological parameters 3 and 4 days later, respectively.

444 AHR. For non-invasive measurements, mice were analyzed one day after the last aerosol challenge using whole body plethysmography as described ³⁶. Invasive measurements of dynamic 445 lung resistance and compliance were performed one day after the last aerosol challenge using a 446 Flexivent apparatus (SCIREO, Emka Technologies) as previously described ³⁷. Briefly, mice 447 were anesthetized (5 ml/kg Dormitor 10 % (Medetomidine, Pfizer) - Imalgene 10% (Ketamine, 448 449 Merial) tracheotomized, paralyzed (5 ml/kg Pavulon 1% (Pancuronium bromide, Organon) and 450 immediately intubated with an 18-G catheter, followed by mechanical ventilation. Respiratory 451 frequency was set at 150 breaths/min with a tidal volume of 0.2 ml, and a positive-end expiratory 452 pressure of 2 ml H₂O was applied. Increasing concentrations of methacholine (0-24 mg/ml) were 453 administered at the rate of 20 puffs per 10 seconds, with each puff of aerosol delivery lasting 10 454 ms, via a nebulizer aerosol system with a 2.5-4 μ m aerosol particle size generated by a nebulizer 455 head (Aeroneb, Aerogen). Baseline resistance was restored before administering the subsequent 456 doses of methacholine.

457 Analysis of BALF cells. Mice were bled and a canula was inserted into the trachea. Lungs were 458 washed 3 times with 1 ml of warmed PBS. For differential BALF cell counts, cells were stained 459 with mAb anti-CCR3 (R&D), anti-Gr1, anti-CD3 and anti-CD19 mAbs (Becton Dickinson, BD) 460 and analyzed by FACS using a FACScalibur flow cytometer and Cellquest software. Eosinophils 461 were defined as CCR3⁺ CD3⁻CD19⁻, neutrophils as Gr-1^{high} CD3⁻CD19⁻, lymphocytes as 462 CD3⁺CD19⁺ and alveolar macrophages as large autofluorescent cells.

463 Serum antibody measurements. Serum LACK-specific IgG1 and IgE were measured by 464 ELISA. For IgG1 quantification, antigen-coated Maxisorp plates (Nunc) were incubated with 465 serial dilution of sera and biotinylated anti-IgG1 mAb (BD). For antigen-specific IgE, plates were 466 first coated with the respective capture mAb (BD), and incubated with serum dilutions. 467 Biotinylated-LACK antigen was then added. HRP-conjugated streptavidin (BD) and TMB (KPL) 468 were used for detection.

Tissue processing. Lungs, LN or spleens were cut to small pieces in HBSS containing 400 U
type I collagenase and 1 mg/ml DNAse I and digested for 30 min at 37°C. Cells were strained
through a 70 µm cell strainer. Erythrocytes were lysed with ACK lysis buffer.

472 **Cytokine assays.** Lung samples were homogenized in C^{2+} and Mg²⁺ free HBSS. BAL and lung 473 supernatants were used. Multiplex IL-4, IL-5, IL-13 and IFN-γ analysis were performed with 474 CBA using FACS array (BD Bisociences). For intracellular staining, cells were incubated with 475 100 µg/ml LACK and 1 µg/ml of anti-CD28 (BD) for 6 h. Brefeldin A (5 µg/ml, Sigma) was 476 added during the last 4 h. Cells were then stained with anti-CD4 mAb, fixed, permeabilized using 477 cytofix/cytoperm reagent (BD), stained with anti-IL-4, or IL-5 (BD) and analyzed by FACS.

478 **CD4⁺ T cell transfer.** Donor mice were tolerized to OVA, BSA or PBS as described above. 479 Cells were prepared from spleens 21 days later, and CD4⁺ T cells were enriched by negative 480 depletion using CD4 isolation kit (Dynal) and further sorted using a high-speed sorter 481 VANTAGE SETLO⁺ flow cytometer (BD) after staining with anti-CD3 and anti-CD4 mAbs. 482 CD4 purity was > 95%. In some experiments, enriched CD4⁺ T cells were stained with antibodies to CD25, CD4 and CD3, and CD25⁻ and CD25⁺ CD4⁺ T cell populations were sorted by FACS. 483 Sorted cells were then injected i.v. into sensitized mice (4 X 10^6 or 1.5 X 10^6 cells per mouse. 484 485 respectively).

486 Statistic analysis. ANOVA for repeated measures was used to determine the levels of difference 487 between groups of mice for plethysmography measurements. Comparisons for all pairs were 488 performed by Mann-Whitney U test. Significance levels were set at a *P* value of 0.05.

489

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494 **COMPETING INTEREST STATEMENT**

- 495 The authors declare no competing financial interests.
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Figure No.











