CD11a polymorphisms regulate TH2 cell homing and TH2-related disease

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CD11a polymorphisms regulate T<sub>H2</sub> cell homing and T<sub>H2</sub>-related disease

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Abstract

**Background**—T<sub>H2</sub>-dependent diseases vary in severity according to genotype, but relevant gene polymorphisms remain largely unknown. The integrin CD11a is a critical determinant of allergic responses, and allelic variants of this gene might influence allergic phenotypes.

**Objective**—We sought to determine major CD11a alleles in mice and human subjects and their importance to allergic disease expression.

**Methods**—We sequenced mouse CD11a alleles from C57BL/6 and BALB/c strains to identify major polymorphisms; human CD11a single nucleotide polymorphisms were compared with allergic disease phenotypes as part of the international HapMap project. Mice on a BALB/c or C57BL/6 background and congenic for the other strain's CD11a allele were created to determine the importance of mouse CD11a polymorphisms in vivo and in vitro.

**Results**—Compared with the C57BL/6 allele, the BALB/c CD11a allele contained a nonsynonymous change from asparagine to aspartic acid within the metal ion binding domain. In general, the BALB/c CD11a allele enhanced and the C57BL/6 CD11a allele suppressed T<sub>H2</sub> cell-dependent disease caused by the parasite Leishmania major and allergic lung disease caused by the fungus Aspergillus niger. Relative to the C57BL/6 CD11a allele, the BALB/c CD11a allele conferred both greater T-cell adhesion to CD54 <em>in vitro</em> and enhanced T<sub>H2</sub> cell homing to lungs <em>in vivo</em>. We further identified a human CD11a polymorphism that significantly associated with atopic disease and relevant allergic indices.

**Conclusions**—Polymorphisms in CD11a critically influence T<sub>H2</sub> cell homing and diverse T<sub>H2</sub>-dependent immunopathologic states in mice and potentially influence the expression of human allergic disease.
The mechanisms that control the recruitment, or homing, of T\(\text{H}_2\) cells collectively represent an essential immune regulatory checkpoint that crucially influences outcomes of diverse infectious and inflammatory processes.\(^1\) The homing of T\(\text{H}_2\) cells is specifically regulated through the expression and regulation of leukocyte function–associated antigen 1 (LFA-1), representing the heterodimeric association of CD18 and CD11a.\(^2\) In the absence of stimulation, LFA-1 is maintained in a low-affinity state that weakly binds the LFA-1 receptor CD54 (intercellular adhesion molecule 1) through the inserted (I) domain of CD11a.\(^3\) High concentrations of divalent cations stabilize and chemokines activate LFA-1 to a high-affinity state, allowing circulating T cells to tightly adhere to CD54 on endothelial surfaces and extravasate to sites of inflammation and infection.\(^5\)\(^,\)\(^6\)

In part, the importance of LFA-1 for T\(\text{H}_2\) cell homing is determined by successive rounds of cell division during which LFA-1 expression is maintained or enhanced while expression of alternate homing integrins is reduced.\(^9\) Blockade or genetic deletion of LFA-1 does not interfere with the induction of robust T\(\text{H}_2\) responses but precludes efficient T\(\text{H}_2\) cell homing to sites of infection and inflammation without interfering with the development and homing of other T\(\text{H}_1\) effector subsets.\(^9\) Therefore selective blockade of T\(\text{H}_2\) cell homing through LFA-1 antagonism results in diverse phenotypes in mice, including reduced expression of allergic lung disease and improved control of the intramacrophage parasite *Leishmania major* in C57BL6 and BALB/c mice.\(^9\) The effect of LFA-1 deletion in BALB/c mice is particularly striking, converting the normally progressive and lethal *L major* infection into a transient illness that more closely resembles the disease course of C57BL/6 mice.\(^9\)

C57BL/6 mice control *L major* infection by mounting a dominant T\(\text{H}_1\) immune response, whereas BALB/c mice do not control the infection because of production of an aberrant T\(\text{H}_2\)-predominant antiparasite response.\(^10\)\(^,\)\(^11\) Although several genetic loci have been shown to contain genes that promote resistance to *L major*,\(^12\) the specific genetic causes for the dominant T\(\text{H}_2\) response and susceptibility in BALB/c mice have yet to be clearly defined.\(^13\)

Airway challenge of C57BL/6 and BALB/c mice with the same allergen also produces variable immune and physiologic responses, the latter assessed as airway hyperreactivity (AHR), although allergic eosinophil- and T\(\text{H}_2\)-predominant inflammation is consistently observed.\(^14\) The many genetic factors that underlie the mouse allergic lung inflammatory response overlap considerably with those influencing the mouse response to *L major* infection and include CD11a. BALB/c mice that are deficient in CD11a demonstrate impaired T\(\text{H}_2\) cell homing to the site of infection and arrested progression of *L major* infection, which is similar to the healer phenotype observed in the C57BL/6 mouse strain.\(^9\) The apparent increase in resistance in the absence of the BALB/c CD11a allele suggests that the BALB/c CD11a allele is an important factor in the pathogenesis of *L major* in the BALB/c mouse strain. As a gene located within a genomic region identified for carrying *L major* resistance in C57BL/6 mice, it is possible that polymorphisms in CD11a might in part determine the divergent immune and physiologic outcomes of BALB/c and C57BL/6 mice.

To investigate the existence of polymorphisms in CD11a, we sequenced and identified strain-specific polymorphisms in CD11a of BALB/c and C57BL/6 mice and created mice congenic for CD11a on the BALB/c and C57BL/6 backgrounds to test the effect of CD11a...
alleles in the T\textsubscript{H}1- and T\textsubscript{H}2-mediated disease models of \textit{L major} infection and allergic lung disease.

**Methods**

**Mice**

Animal studies were conducted under protocols approved by Baylor College of Medicine’s Institutional Animal Care and Use Committee; mice were all housed under specific pathogen-free conditions and were fed standard mouse chow \textit{ad libitum}. BALB/c, C57BL/6, and recombination-activating gene (Rag) 1\textsuperscript{−/−} mice were purchased from Jackson Laboratories (Bar Harbor, Me). Rag2\textsuperscript{−/−} mice were acquired from Taconic (Hudson, NY). CD11a\textsuperscript{−/−} mice on the BALB/c (10 generations backcrossed) and C57BL/6 (8 generations backcrossed) backgrounds were provided by Dr C. M. Ballantyne.\textsuperscript{16} CD11a-congenic mice on the BALB/c (Balb\textsuperscript{C57-CD11a}) and C57BL/6 (C57\textsuperscript{balb-CD11a}) backgrounds were created by crossing CD11a\textsuperscript{−/−} mice of one background with the wild-type (WT) strain of the other background. Progeny were then backcrossed to the parent CD11a\textsuperscript{−/−} strain, selecting for CD11a expression on CD4 T cells for 8 generations. Homozygous congenic or litter-mate CD11a\textsuperscript{−/−} mice were recreated by crossing eighth-generation heterozygous mice and selecting for complete loss of the targeting construct by using PCR or CD11a by using flow cytometry, respectively.

**\textit{L major} infection**

\textit{L major} (strain MRHO/SU/59/P/LV39) was cultured and mouse infection was established, as previously described.\textsuperscript{17}

**Macrophage phenotype**

Popliteal lymph nodes were harvested from 3-week \textit{L major}–infected mice and processed for total RNA by using Trizol (Invitrogen, Carlsbad, Calif), according to the manufacturer’s methods. Primer sets (Applied Biosystems, Foster City, Calif) for inducible nitric oxide synthase (iNOS; Mm00440502\_m1), Arg1 (Mm00475988\_m1), found in inflammatory zone 1 (Fizz1; Mm00445109\_m1), and 18s RNA (4319413E-0710034) were used to quantify mRNA with the TaqMan One-Step RT-PCR Master Mix (4309169, Applied Biosystems) using the 7500 Real-Time PCR System (Applied Biosystems). Signals were used to calculate relative expression levels in accordance with standard practice and expressed as the relative expression ratios of select genes.

**Dynamic adhesion assay**

Adhesion of CD4\textsuperscript{+} cells was quantitated by means of video microscopy under continuous flow conditions \textit{in vitro}.\textsuperscript{18,19} For further information, see the Methods section in this article’s Online Repository at www.jacionline.org.

**Flow cytometry**

Lymphocytes were stained with CD4-Pacific Blue, CD3–allophycocyanin, and CD11a–phycoerythrin (PE), CD11b-PE, or CD11c-PE (all from BD PharMingen, San Jose, Calif) by using standard staining methods.

**Adoptive transfer of ovalbumin-specific CD4\textsuperscript{+} T cells**

Mice were challenged with ovalbumin (OVA)–alum weekly for at least 3 weeks, spleens were harvested, and CD4+ cells were collected, selected under T\textsubscript{H}2 responses, differentially labeled fluorescently, and injected into strainspecific Rag-deficient mice for comparison of
T\textsubscript{H}2 cell homing efficiency. For further information, see the Methods section in this article's Online Repository.

**Allergic airway disease**

Mice were challenged with a clinical isolate of \(4 \times 10^5\) *Aspergillus niger* conidia every 2 days for 8 challenges and assessed for allergic airway disease, as previously described.\(^\text{20}\)

**Human single nucleotide polymorphism analysis and correlation**

Human studies were conducted under Institutional Review Board–approved protocols at Harvard Medical School and Brigham and Women's Hospital.

**Genotyping and data management**

Of the 1041 participating parent-child trios in the Childhood Asthma Management Program (CAMP) trial,\(^\text{21}\) 92.9% are included in this analysis. Using data from European Americans (CEU) in the International HapMap project,\(^\text{22}\) we applied a linkage disequilibrium (LD)–tagging algorithm (minor allele frequency, ≥10%; \(r^2 > 0.8\)) to capture common variation in CD11a and its 10-kb flanks.\(^\text{23}\) For this study, additional single nucleotide polymorphisms (SNPs) were genotyped to evaluate reported functional variation (\(r_s\)) within the genomic region of CD11a. All SNPs were genotyped with the SEQUENOM iPLEX platform (Sequenom, San Diego, Calif).\(^\text{24}\) The 8 polymorphic SNPs successfully genotyped capture of greater than 84% of the HapMap SNPs with a minor allele frequency of 10% or greater in CD11a and its 10-kb flanks in CEU trios at an \(r^2\) value of greater than 0.8.

**Statistical analysis**

Mouse statistics were performed, as described previously.\(^\text{25}\) For human data, the family-based association analyses in CAMP were performed with the family-based association test statistic implemented in GoldenHelix PBAT version 3.6.\(^\text{26}\) In family-based samples Hardy-Weinberg equilibrium was tested in parental data.

**Results**

**Identification of polymorphisms in CD11a**

To determine whether CD11a polymorphisms distinguish the divergent responses of BALB/c and C57BL/6 mice to *L major* infection, we first sequenced the full CD11a mRNA from C57BL/6 and BALB/c mice (GenBank no. JN986841.1, Fig 1). Of 3 variances from the canonical C57BL/6 sequence, 2 of the polymorphisms were found in the metal ion binding domain (MIBD) of CD11a, with the C57BL/6 allele encoding asparagine at position 598 and proline at position 603 and the BALB/c allele encoding aspartic acid and alanine at the same positions, respectively. The BALB/c allele further contained an additional glutamine at position 1007 (Fig 1).

Of particular interest were the position 598 and 603 mutations. The change from asparagine (C57BL/6) to aspartic acid (BALB/c) occurs at a site within the MIBD that is predicted to directly bind divalent cations, as required for optimal CD11a function (UniProt). The MIBD is known specifically to regulate the stability and affinity state of LFA-1 and the homing efficiency of T cells.\(^\text{5}\) The polymorphisms occurring in the BALB/c CD11a allele effectively increase the net negative charge within the MIBD, as well as potentially reducing steric hindrance to cation binding. Together, we hypothesized that this combination of polymorphisms would enhance the binding of divalent cations by the BALB/c CD11a allele, leading to an enhanced affinity for CD54 and potentially augmented T\textsubscript{H}2 cell homing.
BALB/c CD11a congenic mice are resistant to *L major* infection

T<br__/>H cells from BALB/c mice express more robust T<br__/>H2 cell development under neutral and biased in vitro activation conditions, a finding that parallels the enhanced T<br__/>H2 cell–dependent susceptibility to *L major* infection of BALB/c mice.13,27,28 To determine whether such enhanced T<br__/>H2 responses ascribed to the BALB/c mouse are in part attributable to CD11a polymorphisms, we created mice congenic for CD11a on the BALB/c and C57BL/6 backgrounds, termed Balb<sup>C57</sup>-CD11a and C57<sup>Balb</sup>-CD11a, respectively, and assessed their ability to control infection with *L major*. As expected,9,13,29 wild type C57BL/6 mice challenged with *L major* controlled the infection, as indicated by an increase followed by a reduction in footpad swelling, whereas BALB/c mice were unable to control the infection, as assessed by progressive footpad swelling (Fig 2, A). C57<sup>balb-CD11a</sup> mice also controlled infection as efficiently as WT C57BL/6 mice,13,30 as did BALB/c CD11a<sup>−/−</sup> mice, as previously reported.9 In contrast, relative to WT BALB/c mice, congenic Balb<sup>C57-CD11a</sup> mice showed a marked attenuation in disease progression (Fig 2, A). BALB/c mice further showed significantly higher parasite burdens in the footpads than C57BL/6 mice at both 3 and 6 weeks after *L major* infection (Fig 2, B and C). In contrast, both Balb<sup>C57-CD11a</sup> and BALB/c<sup>-CD11a−/−</sup> mice had lower footpad parasite burdens compared with WT BALB/c mice at the 3-week time point. Dissemination of infection from the footpad to the spleen was observed at 3 weeks in all mice. However, WT BALB/c mice had significantly higher parasite burdens in the spleen than all groups at both time points (Fig 2, D and E). Moreover, BALB/c<sup>-CD11a−/−</sup> and WT C57BL/6 mice resolved splenic infection by 6 weeks, and although Balb<sup>C57-CD11a</sup> mice retained spleen parasites at 6 weeks, the burdens were approximately 10 times lower relative to WT BALB/c mice (Fig 2, E). For all other parameters, C57<sup>balb-CD11a</sup> mice did not differ from WT C57BL/6 mice.

The intermediate phenotype of Balb<sup>C57-CD11a</sup> mice, as assessed based on footpad swelling caused by *L major* (Fig 2, A), was reflected in 2 additional immune parameters. First, relative to WT BALB/c mice, the relative abundance of IL-4–secreting cells, expressed as the ratio of IL-4–secreting to IFN-γ–secreting cells from popliteal lymph nodes, was lower in Balb<sup>C57-CD11a</sup> mice, reflecting more predominant IFN-γ secretion. Second, lymph node cells of Balb<sup>C57-CD11a</sup> mice showed significantly lower expression of a marker of alternatively (ie, T<br__/>H2 cell) activated macrophages (found in inflammatory zone 1 [Fizz1]) relative to the classically (ie, T<br__/>H1 cell) activated macrophage marker inducible nitric oxide synthase relative to WT BALB/c mice (Fig 2, G). Together, these findings demonstrate that possessing the C57BL/6 variant of CD11a in BALB/c mice can influence the course of *L major* infection, favoring weaker T<br__/>H2 responses and improved disease control.

**CD11a polymorphisms influence allergic lung disease**

To confirm these findings in a distinct disease model, we next evaluated the effect of CD11a polymorphisms on the development of infectious allergic lung disease, a disease model in which CD11a–dependent T<br__/>H2 cells exert a more dominant role compared with the *L major* model.9 WT BALB/c and C57BL/6 mice were challenged intranasally with the conidia of the fungus *A niger* and assessed for the development of AHR, an index of airway obstruction, in response to acetylcholine challenge, airway eosinophilia, and lung and airway cytokine secretion (Fig 3).20 Based on a much greater increase in respiratory system resistance induced by acetylcholine relative to naive mice, both BALB/c and C57BL/6 mice had AHR, although as expected, this IL-13–dependent response was more pronounced in BALB/c mice (Fig 3, A and B).31,32 However, whereas C57<sup>Balb-CD11a</sup> mice showed significantly enhanced AHR, BALB/c<sup>C57-CD11a</sup> mice showed significantly attenuated AHR relative to that seen in genotypematched WT control animals. Similarly, conidium-challenged Balb<sup>C57-CD11a</sup> mice showed a 2-fold lower recruitment and C57<sup>Balb-CD11a</sup> mice showed a 2-
fold higher recruitment of IL-4–secreted cells to the lung when compared with genotypematched WT control animals (Fig 3, B and D). There were no significant differences in either bronchialalveolar lavage fluid cellularity, especially eosinophils, or airway glycoprotein secretion between the 2 congenic strains (see Figs E1 and E2 in this article’s Online Repository at www.jacionline.org). Differences in cellular proliferation could not account for the differences noted between WT and congenic T cells because Th2 cells from both genotypes proliferated identically in vitro (see Fig E3 in this article’s Online Repository at www.jacionline.org).

Analysis of culture supernatants of lung homogenates from fungus-challenged mice revealed that BalbC57-CD11a mice secreted less IL-5 and IL-13 relative to WT BALB/c mice, whereas no significant difference was observed between congenic and WT mice on the C57BL/6 background (Fig 3, E and F). Concentrations of other inflammatory chemokines and cytokines (CCL11 [eotaxin], IL-17A, and CCL17 [thymus and activation-regulated chemokine]) did not differ from lung supernatants of the same mice (see Fig E4 in this article's Online Repository at www.jacionline.org). Thus the BALB/c allele confers increased susceptibility to the development of allergic lung disease and more efficient recruitment of IL-4–producing cells to the lungs regardless of the mouse’s genetic background. Conversely, having the C57BL/6 CD11a allele correlated with improved resistance to allergic lung disease in either strain.

**CD11a polymorphisms influence CD54 adhesion and Th2 cell homing in vivo**

The previous results are potentially explained by several functions ascribed to LFA-1, including modulation of T-cell signaling and development, in addition to homing.33-36 Our prior analysis of CD18 and CD11a function did not reveal an essential role for LFA-1 for either T-cell proliferation or differentiation but did demonstrate its critical importance for Th2 cell homing.2,9 To determine whether allelic variants of CD11a similarly affect T-cell homing, we next sought to demonstrate the effect of polymorphisms of CD11a on adherence to the LFA-1 receptor CD54 under dynamic flow conditions.9 Th2 cells derived from WT BALB/c mice showed the greatest time-dependent adherence to CD54 while rolling when compared with WT C56BL/6 T cells (Fig 4, A). In contrast, BalbC57-CD11a T cells showed a marked reduction in the number of rolling cells when compared with WT BALB/c T cells and were not significantly different from WT C57BL/6 T cells. Conversely, C57Balb-CD11a T cells showed significantly increased adhesion when compared with both WT C57BL/6 and BalbC57-CD11a T cells. We also assessed the average velocity of rolling T cells but observed no strain-specific differences (see Fig E5 in this article's Online Repository at www.jacionline.org). Differences in CD11a expression or additional integrins known to bind CD54 (CD11b and CD11c) could explain these results but were not detected (see Fig E6 in this article's Online Repository at www.jacionline.org). Thus similar to the observed in vivo phenotypes, allelic variants of CD11a substantially influence the ability of LFA-1 to bind CD54, with the BALB/c CD11a allele conferring increased adherence and the C57BL/6 CD11a allele conferring reduced adherence under dynamic conditions.

We next examined the role of CD11a polymorphisms on the homing of Th2 cells in vivo. OVA-specific Th2 cells from WT and congenic mice were fluorescently labeled, combined in a 1:1 ratio, and adoptively transferred into genotype-appropriate Rag-deficient mice. Recipient mice were then intranasally challenged with OVA and A niger conidia for 3 days, and lungs were assessed for the presence of labeled donor T cells (Fig 4, B-D). Although equivalent donor cells were transferred to each mouse (Fig 4, B, inset), a greater number of WT BALB/c T cells when compared with BalbC57-CD11a T cells were observed in the lungs after A niger and OVA challenge (Fig 4, B). The total numbers of lung T cells (see Fig E7 in this article's Online Repository at www.jacionline.org) and relative percentage of recruited T
cells (Fig 4, C) were determined in multiple mice and consistently showed that T cells containing the BALB/c CD11a allele were more efficient at homing to the lungs, regardless of the genetic background of the donor mouse (Fig 4, C and D). In contrast, homing of CD4+ TH1 cells was not affected by CD11a polymorphisms in the in vivo homing model (Fig 4, E). These experiments show that the BALB/c CD11a allele significantly enhances the homing efficiency of TH2 cells when compared with the C57BL/6 CD11a allele.

**Human CD11a polymorphism correlates to allergic disease**

The unexpected importance of allelic variants of CD11a to the expression of mouse allergic disease suggested that human CD11a allelic variants might have genetic importance. To begin to determine this, we first identified 16 SNPs in the region of CD11a on chromosome 16 (Fig 5, A) from participants in the CAMP study. Two SNPs (rs1064524 and rs2230433) were found to cause missense mutations and 1 (rs2230434) synonymous mutation in exons of CD11a. LD analysis showed 3 LD blocks across the CD11a SNPs (Fig 5, B).

Genotyping was performed in 6 SNPs, including the 2 missense mutations. The missense mutation rs1064524 represents a change from a positive polar (Arg) to a hydrophobic (Trp) amino acid in the inserted domain of CD11a. Analysis of the transmission of rs1064524 from parents to asthmatic probands in relation to allergic disease phenotypes was analyzed by using a dominant model of inheritance (Fig 5, C). The frequency of rs1064524 in the CAMP cohort significantly correlated with the occurrence of multiple allergic disease phenotypes and TH2 cell activity, including atopic dermatitis (self-report), peripheral blood eosinophil counts, serum IgE levels, AHR (DRSR), and allergic rhinitis (hay fever; self-report).

**Discussion**

The divergent but critical roles of CD11a in regulating the mouse immune and physiologic responses in models of *L major* infection and allergic lung disease suggested a broader role for this adhesion molecule in immune regulation and disease. We have now identified distinct alleles for CD11a occurring within BALB/c and C57BL/6 mice that explain in part the distinct immune responses of these mouse strains. Functionally, the BALB/c CD11a allele conferred greater adhesion to CD54, more efficient homing of TH2 cells in vivo, and more robust expression of TH2 cell–dependent disease, including more progressive disease caused by *L major* infection and enhanced allergic lung disease. We further identified a unique allelic variant of CD11a that segregates with multiple indices of human allergic disease. Together, these findings confirm the importance of CD11a to the expression of allergic disease through the regulation of TH2 cell homing and provide the first evidence that polymorphic variants of CD11a are significant disease determinants in human allergic disease.

The unusual susceptibility of BALB/c mice to *L major* infection has been studied intensively for more than 40 years and is known to be both T cell based and polygenic in nature, involving at least 6 genetic loci. Among the many T cell–specific factors linked to resistance to *L major* infection, our data indicate that the importance of chromosome 7 to *L major* resistance is in part due to CD11a, the gene for which (*Itgal*) lies within this chromosome. Replacement of the cognate CD11a allele in BALB/c mice with the C57BL/6 allele improved resistance to *L major* in mice on the BALB/c background, as assessed based on reduced, although still progressive, footpad swelling; reduced parasite burdens at multiple time points; and reduced alternative macrophage activation. Nonetheless, as expected, the effect was not as pronounced as deletion of CD11a entirely (Fig 2 and Lee et al). In contrast, congenic C57BL/6 mice were indistinguishable from WT C57BL/6 mice...
with regard to control of *L. major* infection (Fig 2 and data not shown). Thus CD11a is a major determinant of resistance to *L. major* infection only in BALB/c mice; other factors are more important in determining resistance on the C5BL/6 genetic background. These observations are consistent with the downregulation of LFA-1 expression and CD54-independent homing that has previously been shown for T<sub>H</sub>1 cells, which predominate in *L. major*–infected C57BL/6 mice.⁹,³⁷

A more consistent role for CD11a allelic variants was evident in experimental allergic lung disease, a model in which T<sub>H</sub>2 cells are clearly linked to disease expression across genotypes.¹⁴ Regardless of the mouse genetic background, possession of the BALB/c CD11a allele resulted in greater AHR, an important disease marker that is strongly dependent on the recruitment of T<sub>H</sub>2 cells to the lung.²,⁹ The BALB/c CD11a allele further enhanced recruitment to the lung of IL-4–secreting cells and the secretion of allergy-related cytokines from the lungs of BALB/c mice (Fig 3). A similar effect of the BALB/c allele was seen in C57BL/6 mice, although enhanced secretion of IL-5 and IL-13 from the lung was not seen. Thus, as with *L. major* infection, although CD11a is important for disease expression in both C57BL/6 and BALB/c mice, additional factors in C57BL/6 mice modify disease expression independently of LFA-1.

Our findings demonstrate that distinct CD11a alleles influence the efficiency of T<sub>H</sub>2 cell homing. These observations are entirely consistent with the enhanced adhesion of the BALB/c CD11a allele to CD54 *in vitro* (Fig 4, A) and our prior demonstration of the general importance of CD11a for T<sub>H</sub>2 cell homing.⁹ Thus independent of the absolute level of gene expression, mouse CD11a exists in functionally hypomorphic (ie, the C57BL/6 allele) or hypermorphic (ie, the BALB/c allele) variants, as assessed by adhesion to CD54, which critically influence the expression of T<sub>H</sub>2-dependent diseases, and our preliminary analyses indicate that the same paradigm is also true in human subjects. Although additional study is required, we propose that CD11a might confer enhanced or reduced susceptibility to diverse allergic diseases based on the expressed alleles.

Our findings leave unresolved which amino acid changes or combinations thereof within the BALB/c CD11a allele determine the enhanced adhesion to CD54. We speculate that the greater negative charge within the MIBD of the BALB/c allele promotes more robust interaction with divalent cations, which in turn promotes a more consistently activated conformation that more efficiently engages CD54.⁴,³⁸ Similarly, we suspect that the Arg to Trp change within the inserted domain of the rs1064524 allele of human CD11a also enhances binding to CD54, although through a mechanism that might not involve cation binding. Obviously, this allele is not present in the mouse, and hence there is no direct polymorphism link between our human genetics and the mouse genetics described here. Therefore much further work is required to resolve the atomic and molecular basis of the distinct CD11a phenotypes. Clarification of these mechanisms and the improved understanding of CD11a function that this will bring might provide new insight into the importance of CD11a as a disease marker and how LFA-1 can be safely manipulated to improve outcomes for a variety of diseases involving T<sub>H</sub>2 cells.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

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References


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**Abbreviations**

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<tr>
<td>AHR</td>
<td>Airway hyperreactivity</td>
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<tr>
<td>CAMP</td>
<td>Childhood Asthma Management Program</td>
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<tr>
<td>LD</td>
<td>Linkage disequilibrium</td>
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<tr>
<td>LFA-1</td>
<td>Leukocyte function–associated antigen 1</td>
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<tr>
<td>MIBD</td>
<td>Metal ion binding domain, OVA, Ovalbumin</td>
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<td>PE</td>
<td>Phycoerythrin</td>
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<tr>
<td>Rag</td>
<td>Recombination-activating gene</td>
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<td>SNP</td>
<td>Single nucleotide polymorphism</td>
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Key messages

- Polymorphisms in the integrin CD11a in part explain differences in susceptibility of diverse mouse strains to T_{H2}-dependent disease.
- A distinct SNP in human CD11a is associated with several allergic disease indices.
- CD11a might represent a useful susceptibility biomarker and therapeutic target in allergic disease.
Fig 1.
Mouse CD11a exists in distinct allelic forms. The diagram schematically depicts major functional domains of the extracellular domain of CD11a and the amino acid differences between the BALB/c and C57BL/6 mouse strains. TM, Trans-membrane domain.
Fig 2.
Effect of CD11a polymorphisms on *L. major* infection. 

A, Footpad swelling of WT compared with congenic mice infected with *L. major* promastigotes. *P < .001 (n = 5 per group).

B-E, Parasite burdens from footpads and spleens at 3 (Fig 2, B and D) and 6 (Fig 2, C and E) weeks. *P < .05.

F, Ratio of IL-4– and IFN-γ–producing cells from popliteal lymph nodes at 3 and 6 weeks. *P < .05.

G, Ratio of *Fizz1* inducible nitric oxide synthase (*iNOS*), as assessed by using real-time quantitative PCR. *P < .05.
Fig 3.
CD11a polymorphisms influence allergic airway disease. **A**, Respiratory system resistance ($R_{RS}$) values comparing BALB/c and BALB/cC57 congenic mice challenged with PBS or *A. niger* conidia (AN). *P < .001 (n = 3 for BALB/c mice; n = 5 for all other groups).** B, Total lung IFN-γ– and IL-4–producing cells. *P < .001. C, Respiratory system resistance values comparing C57BL/6 and C57Balb congenic mice. *P < .05, (n = 5). D, Total lung IFN-γ– and IL-4–producing cells. *P < .001. E and F, IL-5 and IL-13 levels from lung homogenates. *P < .05 (n = 5). Ach, Acetylcholine; ND, not detectable.
Fig 4. CD11a polymorphisms influence $\text{T}_{\text{H}}2$ cell homing. A, $\text{T}$-cell adhesion to CD54 under dynamic flow. $^*P < .05$ (n = 3). B-E, Rag-deficient mice were reconstituted with labeled $\text{T}_{\text{H}}2$ cells from WT and congenic mice, respectively (Fig 4, B, inset) and intranasally challenged with $\text{A niger/OVA}$ to induce $\text{T}$-cell recruitment, after which lung-derived single-cell suspensions were analyzed for donor $\text{T}_{\text{H}}2$ (Fig 4, B [n = 1]; C, and D) and $\text{T}_{\text{H}}1$ (Fig 4, E) cells. $^*P < .05$. Data from one of 3 comparable experiments analyzing $1 \times 10^6$ cells per lung.
Fig 5.
Association of human CD11a SNPs to allergic disease. **A**, Location of SNPs within the human CD11a locus; exons are indicated in gray. **B**, Pairwise $r^2$ values for correlations between identified SNPs. Black squares, $r^2 = 1$; white squares, $r^2 = 0$; gray squares, $0 < r^2 < 1$ with intensity proportional to $r^2$. **C**, Significant correlations between SNPs and the indicated allergy phenotypes. DRSS, Log of dose-response slope to methacholine challenge; ns, not significant; HAYF, hay fever.