Title
Altered ligands reveal limited plasticity in the T cell response to a pathogenic epitope.

Permalink
https://escholarship.org/uc/item/2pb2b2mh

Journal
The Journal of experimental medicine, 189(7)

ISSN
0022-1007

Authors
Pingel, S
Launois, P
Fowell, DJ
et al.

Publication Date
1999-04-01

DOI
10.1084/jem.189.7.1111

Peer reviewed
Altered Ligands Reveal Limited Plasticity in the T Cell Response to a Pathogenic Epitope

By Sabine Pingel,§ Pascal Launois,§ Deborah J. Fowell,* Christoph W. Turck,* Scott Southwood,‡ Alessandro Sette,¶ Nicolas Gläichenhaus,§ Jacques A. Louis,§ and Richard M. Locksley*‡

From the *Division of Medical Sciences, Stanford University School of Medicine, Stanford, California 94305, and the §Department of Microbiology and Immunology, University of California, San Francisco, California 94143; the ¶Medical Institute, University of Lausanne, Epalinges CH-1066, Switzerland; and the ‡Institute of Molecular and Cellular Pharmacology, University of Nice, Valbonne 06560, France.

Summary

Experimental leishmaniasis offers a well characterized model of T helper type 1 cell (Th1)-mediated control of infection by an intracellular organism. Susceptible BALB/c mice aberrantly develop Th2 cells in response to infection and are unable to control parasite dissemination. The early CD4+ T cell response in these mice is oligoclonal and reflects the expansion of Vß4/Va8-bearing T cells in response to a single epitope from the parasite homologue of mammalian RACK1 (LACK) antigen. Interleukin 4 (IL-4) generated by these cells is believed to direct the subsequent Th2 response. We used T cells from T cell receptor–transgenic mice expressing such a Vß4/Va8 receptor to characterize altered peptide ligands with similar affinity for I-Ab. Such altered ligands failed to activate IL-4 production from transgenic LACK-specific T cells or following injection into BALB/c mice. Pretreatment of susceptible mice with altered peptide ligands substantially altered the course of subsequent infection. The ability to confer a Th1 phenotype on otherwise susceptible mice using altered peptides that differed by a single amino acid suggests limited diversity in the endogenous T cell repertoire recognizing this antigen.

Key words: Leishmania major • CD4+ T cell subsets • altered peptide ligands • LACK antigen

Leishmania major is a complex intracellular protozoan parasite of macrophages in the vertebrate host that requires a robust Th1 response for control of infection (reviewed in 1). Requirements for MHC class II molecules, T cells, IFN-γ, IL-12, and NO 2 have delineated a pathway by which CD4+ T cells differentiate into Th1 effector cells that activate macrophages and confer resistance to the disease. Among mice with I-Aß class II molecules, the CD4+ T cell immune response is focused dominantly on a single epitope from the parasite LACK² antigen, which results in oligoclonal expansion of cells bearing Vß4/Va8 heterodimeric TCR in the regional lymph nodes (2, 3). The dominant nature of this antigen was demonstrated using TCR-transgenic mice that expressed a monoclonal αβ T cell repertoire to this epitope; such mice displayed remarkable control over the organism, despite the absence of T cells that recognize any parasite-specific antigens other than the single epitope from LACK (4).

Unlike resistant strains of mice, BALB/c mice develop aberrant Th2 responses after challenge with L. major and sustain fatal disseminated disease. The early CD4+ T cell response is also dominated by expansion of LACK-specific T cells that express Vß4/Va8 TCR. However, activation of these cells results in the rapid production of IL-4 that drives the subsequent Th2 response (5, 6). Deletion of LACK-reactive cells, either through thymic expression of LACK as a transgene (7) or superantigen-mediated deletion of Vß4-expressing CD4+ T cells (5), attenuated the early production of IL-4 and promoted the differentiation of Th1 effector cells that controlled disease in infected BALB/c mice. These and other data have suggested a model whereby susceptibility to L. major in BALB strain mice is driven by a confluence of factors involving the inherent propensity to produce IL-4 after activation in the setting of an appropriately sized precursor pool of LACK-reactive CD4+ T cells (8).
The L. major model provides an opportunity to test the plasticity of the wild-type CD4+ T cell repertoire in susceptible mice. The conserved charge and length of the peptide-interacting complementarity determining region 3 (CDR3) domain from LACK-reactive T cell hybridomas suggested the observation that a single parasite antigen served as the focus of their recognition (2). The constraints on the entire LACK-reactive repertoire in the intact animal remain unknown, however. We decided to indirectly test such constraints by attempting to abrogate activation of these cells in vivo using altered ligands. We demonstrate substantial protection against disease in this system, suggesting that identification of dominant antigens from organisms can be used to target pathogenic T cells that mediate progressive disease in a highly specific manner.

Materials and Methods

Reagents. The LACK (156-173) Peptide (ICFSPSLEHIPIVSGSWD), LACK-N164 and LACK-K164 mutant peptides with the designated alterations of H at position 164, and the OVA (designated IR/173) were passaged and maintained as described (4, 5). Groups of 4–10 mice were infected in the hind footpads with purified metacyclic (4 × 10^6) or stationary phase (2 × 10^6) promastigotes as previously described (4, 5, 11). After inoculation, disease progression was monitored using a metric caliper to quantitate footpad size. Animals were killed at the designated times, and the popliteal lymph nodes were harvested for the evaluation of cell types and cytokine production as described (5). Serum was collected terminally for quantitation of IgE by ELISA as described (11), and the footpads and spleens were used to quantitate the parasite burden by limiting dilution (12).

MHC Binding A flitches. I-A^d molecules were affinity purified from cell lysates of A20 lymphoma cells using anti-I-A^-mAb M 69-K (American Type Culture Collection [ATCC]). Peptides were tested for binding to I-A^d as measured by their capacity to inhibit the binding of [125]I-radiolabeled OVA 323-336 as previously described (13).

Stimulation of ABLE T Cells In Vivo. Spleen and lymph node cells were harvested from BALB/c ABLE TCR-C^d mice and used to produce single-cell suspensions after disruption through a 0.75-mNylon mesh filter. Cells were washed, and the resulting populations of ABLE T cells and APC were distributed to duplicate wells of round-bottom microtiter plates (10^5 cells/well) in cell culture medium (RPMI 1640, 5% FCS, 2% Hyclone, 2 mM pyruvate, 50 μM b-2-ME, and 100 U/ml each of penicillin and streptomycin) with the indicated concentrations of synthetic peptides in a final volume of 0.2 ml. Supernatants were harvested at 48 h for cytokine analysis by ELISA. The wells were pulsed at 48 h with 1 μCi [3H]thymidine, and cell proliferation was assessed 18 h later.

TCR Antagonism Assay. CD4^+ T cells from BALB/c ABLE TCR-C^d mice were enriched from spleen and lymph node cell suspensions by antibody- and complement-mediated lysis of B cells. MHC class II-, and CD8^-expressing cells using mAbs 11d1, BP107, and 3155 (ATCC), respectively, and low-toxicity rabbit and guinea pig complement (Cedarlane Labs., Ltd.). The resulting populations were 80% Vbeta^+ cells, of which 35–40% were CD4^- and the remainder CD4^-CD8^- as previously described (14). Irradiated spleen cell populations from BALB/c ABLE TCR-C^d mice were used as APC. Antagonism was quantitated using the method of DeMigistis et al. (13), with slight modifications. APC (10^6 cells/ml) were prepulsed with suboptimal concentrations of the wild-type LACK peptide (0.008–0.2 μM as established in preliminary experiments) in culture medium for 2 h at 37°C. The APC were washed, irradiated, and distributed to 96-well round-bottom microtiter plates (2 × 10^5 cells/well) and further incubated with varying concentrations of the designated peptides (0.01–100 μM) for 2 h at 37°C. The plates were washed, and enriched ABLE T cells were added using 1.6 × 10^5 T cells/well in 0.2 ml medium. After 48 h, the supernatants were collected and analyzed for cytokines by ELISA. Proliferation was assessed at hour 66, and the final 18 h with 1 μCi [3H]thymidine/well.

Immune Response in Mice Injected with rLACK Proteins. Groups of BALB/c mice were injected in the hind footpads with 5, 25, or 50 μg of the designated rLACK protein or 25 μg chicken egg OVA in 50 μl buffer. At various time points, the popliteal lymph nodes were harvested and mRNA purified for analysis of IL-4 transcripts using a semiquantitative reverse transcriptase (RT)-PCR assay as previously described (15). Groups of these treated control or adult thymectomized mice were challenged either 24 h or 10,
BALB/c ABLE-Cα mice were immunized in both footpads with 25 μg of the purified rLACK proteins or chicken egg OVA in 50 μl buffer and, 24 h later, the popliteal lymph nodes were collected and single-cell suspensions prepared. Cells (2 × 10^6) were analyzed by flow cytometry (FACSVantage™: Becton Dickinson) after incubation with a combination of fluorescein isothiocyanate–conjugated anti-Vβ4, PE-conjugated anti-CD4, and biotinylated anti-CD69 mAbs followed by streptavidin-tricolor (all from Caltag Labs).

Immunization with Altered LACK Proteins. Nonthymectomized or adult thymectomized BALB/c mice were immunized in the hind footpad with 25 μg purified rLACK proteins or chicken egg OVA in 50 μl of 50 mM Tris/100 mM NaCl, pH 8.0. Mice were infected 24 h later with the designated strains of L. major promastigotes in the left footpad and the course of infection monitored as described above.

Cytokine Analysis and Serum IgE Determination. IL-4 and IFN-γ were measured using sandwich ELISA with mAbs 11B11 and biotinylated BVD6 for IL-4 detection and R46A2 and biotinylated XMG1.2 for IFN-γ detection as described (11). Samples were normalized to standard recombinant controls. The limits of detection in these assays were 50 pg/ml for IL-4 and 1 ng/ml for IFN-γ. Cytokine production by individual lymphocytes was assessed by ELISPOT assay as described (11). Total serum IgE was measured using ELISA with mAbs B1.4 and 6B3 and biotinylated XMG1.2 for IFN-γ detection as described (11). Cytokine mRNA transcript abundance was quantitated using RT-PCR with the competitor plasmid pPQRS as described (5, 15). In brief, cDNA samples were first normalized for expression of a constitutively expressed gene, hypoxanthine phosphoribosyltransferase (HPRT), and then quantitated for expression of IL-4 and IFN-γ as compared with competitor pseudogene transcripts amplified within the same reaction. The ratio of the authentic and competitive amplicons was quantitated using densitometry.

Results

Construction of Analogue Peptides of the LACK Antigenic Determinant. Using overlapping synthetic peptides and a panel of T cell hybridomas generated from BALB/c mice immunized with the recombinant protein, a single I-A^d–restricted epitope in LACK was localized to amino acids 156–173, comprising the sequence ICFSPSLEHPIVVSGSWD (data not shown). Almost all hybridomas reactive to LACK expressed a Vβ4/Vα8 heterodimeric TCR, although considerable junctional diversity was apparent. The putative CDR3 peptide contact domain, however, was generally conserved in length and charge, with a negatively charged QE or QD motif in the TCR β chain of each of the LACK-reactive hybridomas (3). Similarly, hybridomas established from the lymph node cells of infected BALB/c mice that expressed the Vβ4 TCR contained the QE motif in the CDR3; one had a charged WD motif at the same position (2). Such features suggested that a positively charged amino acid within the LACK antigenic determinant represented a critical TCR contact residue. Based on the use of histidine and other charged residues at TCR contact points among peptides binding to I-A^d (16), we mutated the histidine at position 164 in the wild-type peptide (LACK) to asparagine or lysine, thus creating peptides LACK-N164 and LACK-K164, respectively.

Binding Affinities of LACK Analogue Peptides to I-A^d. The relative affinities for MHC class II molecules by LACK and the LACK analogues were tested by assaying their capacities to compete with an I-A^d ligand of known affinity, chicken egg OVA peptide323–336. By this assay, each of the LACK–derived peptides displayed binding affinities for I-A^d in the same nanomolar range as the OVA323–336 reference peptide; if anything, they showed slightly stronger affinities (Table I). Substitution of H164 in the wild-type LACK determinant by N or K did not, therefore, affect its binding affinity for MH C class II molecules.

A analogue LACK Peptides Do Not Aivate LACK-reactive T Cells from TCR-transgenic Mice. ABLE mice express a transgenic TCR derived from a LACK-reactive Vβ4/Vα8 T cell clone that is activated by the LACK156–173 peptide in the context of I-A^d (4). These mice have been crossed to BALB/c TCR-Cα0 mice, thus creating BALB/c ABLE TCR-Cα0 mice. These mice express a monochonal αβ T cell repertoire consisting exclusively of the LACK-reactive TCR transgene and were used as a source of T cells, designated ABLE T cells. ABLE T cells proliferated in response to the LACK wild-type peptide at low concentrations (7 nM) but not after stimulation with the LACK-N164 or LACK-K164 analogues, even at concentrations up to 4 μM (Fig. 1).

Although ABLE T cells generated both IFN-γ and IL-4 in culture supernatants after incubation with the LACK peptide, neither cytokine was detected after incubation with the two analogue peptides nor with the irrelevant OVA peptide that also binds I-A^d (Fig. 1). Thus, a single amino acid substitution at position 164 in the LACK T cell epitope substantially altered reactivity of the transgenic T cells, indicating that this amino acid position is likely to be a critical TCR contact residue.

Table I. Affinity of LACK Analogue Peptides for I-A^d MHC Class II Molecules

| Peptide       | Amino acid sequence     | I-A^d |
|---------------|-------------------------|-------|
| LACK156–173   | ICFSPSLEHPIVVSGSWD      | 63    |
| LACK-K164     | ICFSPSLEKPIVVSGSWD      | 68    |
| LACK-N164     | ICFSPSLENPIVVSGSWD      | 100   |
| OVA323–336    | ISQAVHAAHAEINE          | 150   |

Pingel et al.
peptide before ABLE T cells were added and assessed for their capacity to proliferate and produce cytokines. LACK-K164 showed dose-dependent inhibition of proliferation to the wild-type peptide; 50% inhibition occurred at a concentration of 10 μM of the analogue peptide (Fig. 2). At similar concentrations, the peptide also inhibited IL-4 and IFN-γ production. LACK-N164 inhibited the proliferation of ABLE T cells only at very high concentrations (>100 μM). The production of IL-4 was inhibited comparably to the LACK-K164 peptide, but IFN-γ production was inhibited consistently less by LACK-N164 in multiple assays. The unrelated OVA323–336 peptide displayed no inhibitory activity. Thus, in the presence of otherwise stimulatory amounts of the wild-type LACK peptide, the two analogue peptides behaved as TCR antagonists. Of the TCR-mediated functions tested, LACK-N164 preferentially inhibited IL-4 production by ABLE T cells, whereas the capacity to proliferate and produce IFN-γ was less affected; LACK-K164 was more global in its inhibitory capacities.

LACK Analogues Fail to Activate LACK-specific T Cells In Vivo. The same amino acid substitutions were introduced into the full length rLACK protein by site-directed mutagenesis, creating rLACK-N164 and rLACK-K164 altered proteins. When tested in vitro for its capacity to stimulate ABLE T cells, the rLACK protein stimulated proliferation and IL-4 and IFN-γ production at molar concentrations comparable to those of the wild-type LACK peptide. In contrast, the altered rLACK proteins, as their peptide counterparts, did not stimulate proliferation or measurable cytokine production over a wide range of concentrations (data not shown).

To assess the activity of the rLACK proteins in vivo, ABLE-Cα0 mice were injected in the hind footpads with 25 μg of purified rLACK, rLACK-N164, rLACK-K164, or OVA. After 24 h, the popliteal lymph node cells were recovered and analyzed using flow cytometry for activation, as assessed by expression of CD69 and enlargement by light-scattering characteristics. Inoculation of rLACK effectively targeted the transgenic T cells: 80% of Vβ1 cells expressed CD69 (Fig. 3) and forward/side scattering increased significantly (data not shown). The total number of transgenic T cells actually decreased in the draining lymph nodes of mice injected with either the wild-type or the altered LACK peptides.
Lymph nodes (from 2.9 × 10^5 after OVA to 1.3 × 10^5 after rLACK), consistent with antigen-mediated deletion as previously described in other TCR-transgenic mice (17, 18). In contrast, Vβ4^+ T cells collected from animals injected with the rLACK analogues showed CD69 induction and forward/side scattering indices that were only modestly greater than those from cells collected from animals injected with the control protein, OVA (Fig. 3). Furthermore, the total number of transgenic T cells in these mice was not significantly different from the number of transgenic T cells in mice that received OVA (data not shown). Thus, as assessed by these criteria, the rLACK analogues did not activate ABLE T cells in vivo in a manner comparable to the cognate LACK protein containing the wild-type epitope.

**LACK Analogue Proteins Block Activation of the LACK-specific Response in Wild-type Mice.** Prior experiments demonstrated that the LACK antigen induced prominent IL-4 expression in Vβ4/Vα8 CD4^+ T cells after injection into BALB/c mice that reached levels 30–100-fold greater than after injection of a construct with the I-A^d epitope deleted (5). Over a 5–50 μg range of rLACK, IL-4 mRNA was induced 100-fold, whereas no IL-4 mRNA was induced by either the 41-amino acid LACK deletion mutant or rLACK-K164 (Fig. 4 A and data not shown). Although a 10-fold induction of IL-4 mRNA was seen after injection of 5 μg rLACK-N164, no IL-4 mRNA was induced after injection of 25- or 50-μg doses. None of the LACK derivatives caused induction of IFN-γ mRNA under the conditions used. After immunization with CFA, however, each of the constructs, rLACK, rLACK-K164, and rLACK-N164, was capable of inducing a proliferative response from subsequently isolated popliteal lymph node CD4^+ T cells in response to their respective LACK156–173 epitopes (stimulation indices increased 10–18-fold; data not shown). No proliferation was induced by any of the LACK epitopes after immunization with the LACK deletion mutant or OVA. The 25-μg protein dose was selected for use in subsequent experiments.

To assess the capacity to alter the response of the endogenous LACK-reactive repertoire, mice were first injected with 25 μg of the LACK analogue proteins and then, 24 h later, with either the authentic LACK protein or viable L. major promastigotes (challenge). After 16 h, the popliteal lymph node cells were collected, RNA was isolated, and the relative IL-4 mRNA levels were determined using RT-PCR as described in Materials and Methods. Results depicted fold increases in IL-4 transcripts as compared with mice immunized with the same peptides but not challenged in the secondary experiment and are representative of three comparable experiments.
L. major promastigotes, both shown previously to activate a brisk IL-4 mRNA response in BALB/c CD4+ T cells (5). In three separate experiments, prior injection of either analogue protein substantially decreased the subsequent activation of the IL-4 response, consistent with an alteration of the normal V84/Vα8 CD4+ T cell response (Fig. 4 B). Prior injection of the LACK construct with the deleted I-Aa epitope or an unrelated I-Aa epitope (OVA) did not affect the subsequent IL-4 response. Kinetic analysis, in which the second injection of recombinant LACK was delayed 10, 20, or 30 d after the initial immunization, revealed that IL-4 nonresponsiveness was maintained for 10–20 d in mice that had been injected with the mutated LACK analogues but then subsequently recovered. Recovery of IL-4 responsiveness to LACK or L. major injection was ablated by prior thymectomy (data not shown).

Protection of Susceptible BALB/c Mice by Immunization with LACK Analogue Proteins. Based on the capacity of LACK analogue proteins to abrogate the early IL-4 response in BALB/c mice, we assessed the ability of immunization to render these mice resistant to progressive disease. Cohorts of mice were immunized once with 25 μg of the various recombinant proteins in the footpad and challenged 24 h later with a lethal infectious dose of wild-type L. major promastigotes of either the IR/173 or LV39 strain.

In three separate experiments with the IR/173 parasite, the course of disease was significantly attenuated in animals that received rLACK-N164; no attenuation was seen in animals that received rLACK-K164 or any of the control proteins, including the wild-type LACK protein (Fig. 5). Whereas animals in all of the other groups had to be killed by week 8, mice that received rLACK-N164 controlled disease up to 12 wk after inoculation, when the experiment was terminated. Parastolog control was confirmed by limiting dilution of tissues that demonstrated a 2–4-log reduction in parasite numbers. Immunologic analysis revealed a threefold reduction in the number of IL-4-producing cells in the draining lymph nodes and in serum IgE levels, whereas the number of IFN-γ-producing cells was similar in all groups.

Protection was more dramatic using the LV39 strain. Similar to the results using the IR/173 strain, rLACK-N164 but not rLACK-K164 provided lasting protection in a subgroup of nonthymectomized mice (Fig. 6 A). The LACK protein itself provided protection in approximately half of both wild-type and thymectomized BALB/c mice (Fig. 6, A and B). Strikingly, either of the altered LACK proteins, in contrast to the LACK deletion mutant or OVA controls, provided a complete protection in thymectomized mice that was sustained over 100 d (Fig. 6 B). When studied at the conclusion of these experiments, the cure phenotype was associated with attenuation of IL-4 production and control of parasite growth in the footpads that was entirely concordant with the lesion phenotype (data not shown).

Discussion

L. major includes a heterogeneous group of protozoa strains that express some 10,000 proteins from a 35.5-megabase genome (19). Despite this complexity, the early immune response is highly focused on a single epitope from the parasite LACK antigen in mice that express I-Aa MHC class II molecules. As shown here, targeting T cells that recognize this epitope using ligands that differed by a single amino acid from the natural epitope was capable of redirecting an otherwise ineffective immune response with a fatal outcome to a completely protective response with long-term cure. The specificity of the immune intervention suggests limited plasticity in the innate LACK-reactive repertoire in I-Aa-bearing mice, as well as limited ability of Leishmania parasites to mediate progressive infection in such mice in the absence of exuberant LACK recognition.

The mechanisms underlying the dominant recognition of the LACK epitope remain unclear. Recognition does not seem related simply to the abundance of the LACK protein. As compared with other parasite molecules like the major surface protease gp 63 or the major surface glycolipid LPG, which are present in ~5 × 10^5 and 3–5 × 10^6 molecules per organism (20, 21), respectively, LACK was less abundantly expressed. Quantitation against recombinant standards showed that LACK comprised only ~0.03% of total cellular protein or ~30,000 molecules per organism (Pingel, S., and R. Locksley, unpublished data). The LACK epitope displayed in vitro affinity for I-Aa that was comparable with endogenously eluted I-Aa peptides (22) and contained a centrally disposed histidine residue, creating a charged element that has been noted in other peptides that bind this MHC molecule. Presumably, the dominance of the epitope must result from some confluence of stability and processing of the peptide, the efficiency of targeting to MHC class II molecules, and/or the size of the responding T cell repertoire (23). Equally perplexing is the dominant nature of the Vβ4/Vα8 TCR response to the I-Aa/LACK peptide complex. The convergence of the immune response on the LACK epitope through use of a dominant Vα/Vβ-paired TCR has been reported using other immunogenic peptides (24), suggesting that other antigens or adjuvant-like molecules from live parasites do not affect this clonal affinity maturation process.
Earlier studies reported the ability to vaccinate susceptible BALB/c mice against *L. major* using LACK antigen administered in a manner such that LACK-specific Th1 cells were generated (3, 25). Indeed, LACK-specific Th1 cells alone were sufficient to establish substantial control over infection with *L. major*, demonstrating that immunoreactive LACK peptide is expressed in vivo at physiologically important levels (4). Despite the capacity of LACK-specific T cells to control infection with the parasite in vivo, such T cells are not required. Thus, BALB/c mice rendered deficient in CD4 T cells that recognize this dominant epitope, either through thymic expression and central deletion or by superantigen-mediated deletion of all Vβ4+CD4+ T cells, were capable of containing *L. major* infection (3, 5). These experiments suggested that LACK recognition was required for establishing the susceptible state of BALB/c mice, although neither method directly targeted epitope-specific T cells. Thus, overexpression of LACK antigen in the thymus might affect the T cell repertoire in ways other than deletion of LACK-reactive T cells. Similarly, deletion of all Vβ4+CD4+ T cells targets cells of additional specificities but unknown contributions to defense against *Leishmania*.

The ability of peptide ligands that differed at a single amino acid residue to affect the subsequent course of disease was evaluated. Figure 6A shows the course of *L. major* LV39 infection in BALB/c mice immunized with altered LACK ligands. (A) Groups of eight nonthymectomized BALB/c mice were immunized in the hind footpads with either vehicle control or 25 µg recombinant LACK, LACK-K164, LACK-N164, LACK ΔA41, or OVA. Mice were infected in one footpad the following day with *L. major* LV39 stationary phase promastigotes and the size of the local lesion quantitated as infected minus noninfected footpads using a metric caliper. A cohort of age-matched, resistant C57BL/6 mice was included as shown (△). Subgroups of resistant and susceptible mice occurred after immunization with LACK and LACK-N164. Results were comparable in two independent infections. (B) Groups of 7-8 BALB/c mice were thymectomized (△) at 8 wk of age and immunized and infected with *L. major* LV39 promastigotes as above. Control adult thymectomized BALB/c mice (BALB/c ATx) remained susceptible as shown in the first panel. Subgroups of resistant and susceptible mice occurred after immunization with LACK. Results were comparable in two independent infections.
ease in susceptible BALB/c mice argues strongly for highly conserved specificity to the Th2-mediating repertoire. Altered ligands are presumed to anergize or functionally alter discrete populations of T cells by their ability to establish incomplete signaling through the TCR complex (reviewed in 26). Modulation of cytokine patterns by altered ligands has been previously demonstrated (27), although application of this technology to an acute infectious process has been infrequently examined. Redistribution of Th subset differentiation, or immune deviation, has been reported by either variations in antigen dose (28, 29) or through use of altered peptide ligands (30). For a variety of reasons, we consider it unlikely that immune deviation can account for the protection mediated by the altered LACK proteins. First, over a wide dose range, the LACK peptide caused no shift in the production of IL-4 and IFN-γ relative to each other by the TCR-transgenic ABLE T cells. Second, the altered LACK peptides induced neither proliferation nor cytokine production by ABLE T cells. Third, the massive activation of ABLE T cells after injection of LACK was absent after injection of the altered LACK antigens. Finally, injections of lower doses of LACK or the altered LACK proteins into BALB/c mice did not induce early production of IFN-γ, rather than IL-4, mRNAs. We could, therefore, find no evidence for the establishment of a LACK-specific Th1 response that could account for the protection mediated by the altered LACK analogues.

More likely, protection of susceptible mice was accomplished through tolerance or deletion of LACK-reactive T cells, a mechanism consistent with previous experimental findings (5, 7). Both altered LACK proteins antagonized IL-4 production by transgenic T cells in response to LACK. When used to immunize BALB/c mice, they abrogated the early IL-4 response to L. major parasites. Treated mice were able to control parasite multiplication of the LV39 strain up to 5 wk; over prolonged periods, and with both the LV39 and IR/173 strains, mice pretreated with the rLACK-N164 protein demonstrated persistent immunity. Thymectomized BALB/c mice immunized with either rLACK-N164 or rLACK-K164 were completely protected, a finding consistent with the ability of the altered LACK proteins to abrogate IL-4 production by Vβ4/Vα8 CD4+ T cells in these mice. Presumably, the delayed yet progressive disease in non-thymectomized LV39-infected mice was dependent on new thymic emigrants. Immunization with the LACK protein itself conferred protection to some mice. As demonstrated using the TCR-transgenic mice, this presumably relates to the capacity of the cognate ligand to mediate peripheral deletion of high-affinity LACK-specific T cells. The observed differences in the overall grade of protection between the two strains of L. major were surprising, but such differences in virulence have been previously described (31, 32), as have subset responses within cohorts of identically treated mice, as seen here among rLACK- and rLACK-N164-treated mice (33–35). Boosting or otherwise optimizing the immunization schedule might have enhanced protection against the IR/173 strain. The rLACK-N164 ligand, however, conferred protection against both L. major strains, suggesting that identification of dominant antigens from pathogens can be used to target disease-producing T cells in a highly sequence-specific manner.

Previous studies have documented that different T cell clonotypes can respond to the same antigen within a given T cell repertoire. Furthermore, considerable cross-reactivity is an essential feature of the T cell receptor, which assures that pathogenic peptides are efficiently recognized (36). Our results suggest that the endogenous LACK-specific repertoire is highly constrained in its CDR3 recognition domain. Limited plasticity of the endogenous T cell repertoire has been previously noted with certain peptide antigens (37), suggesting that infectious diseases may have contributed to the evolutionary divergence of V region genes. Expression of dominant epitopes by parasites, together with the diversity and size of the responding host T cell repertoire, might greatly affect the outcome of such infections and thus contribute to the highly diverse clinical manifestations of leishmaniasis in human populations.

L. major contains two LACK genes expressed in tandem from the same chromosome. Apart from what is known regarding their mammalian homologues, little is known regarding the biochemical action of these proteins. Aside from the potential vaccine use of this antigen (3, 25), additional study promises to shed much light on the coevolution of host and parasite within the context of the immune system, MHC recognition, and the T cell repertoire. Such studies may have great implications for our understanding of the basis for susceptibility and resistance to infectious diseases.

The authors thank Z.-E. Wang, L. Stowring, C. McArthur, and E. Weider for technical assistance.

This work was supported by grants from the National Institutes of Health (AI26918), the Howard Hughes Medical Institute, the Swiss National Science Foundation, the World Health Organization, and the European Union. D.J. Fowell was supported by a Juvenile Diabetes Foundation International Fellowship.

Address correspondence to Richard M. Locksley, UCSF, Box 0654, C-443, 521 Parnassus Ave., San Francisco, CA 94143. Phone: 415-476-5859; Fax: 415-476-9364; E-mail: locksley@medicine.ucsf.edu or Jacques A. Louis, WHO Research and Training Center, University of Lausanne, Epalinges, Switzerland. Phone: 41-21-692-5703; Fax: 41-21-692-5705; E-mail: jacques.louis@ib.unil.ch

Received for publication 23 October 1998 and in revised form 12 February 1999.

1118 Altered LACK Peptides Modulate Experimental Leishmaniasis
References

1. Reiner, S.L., and R.M. Locksley. 1995. The regulation of immunity to Leishmania major. Annu. Rev. Immunol. 13: 151–177.

2. Reiner, S.L., Z.-E. Wang, F. Hatam, P. Scott, and R.M. Locksley. 1993. Th1 and Th2 cell antigen receptors in experimental leishmaniasis. Science. 259:1457–1460.

3. Mougneau, E., F. Altare, A.E. Wakil, S. Zheng, T. Coppola, Z.-E. Wang, R. Waldmann, R.M. Locksley, and N. Glächenhaus. 1995. Expression cloning of a protective Leishmania antigen. Science. 268:56–56.

4. Reiner, S.L., D.J. Fowell, N.H. Moskowitz, K. Swier, D.R. Brown, C.R. Brown, C.W. Turck, P.A. Scott, N. Killeen, and R.M. Locksley. 1998. Control of Leishmania major by a monoclonal αβ T cell repertoire. J. Immunol. 160:88–89.

5. Launois, P., I. Maillard, S. Pingel, K.G. Swihart, I. Xenarios, H. Acha-O’Rea, H. Diggelmann, R.M. Locksley, H.R. MacDonal, and J.A. Louis. 1997. IL-4 rapidly produced by Vg4 Vox CD4+ T cells instructs Th2 development and susceptibility to Leishmania major in BALB/c mice. Immunity. 6:541–549.

6. Launois, P., K.G. Swihart, G. Milon, and J.A. Louis. 1997. Early production of IL-4 in susceptible mice infected with Leishmania major rapidly induces IL-12 unresponsiveness. J. Immunol. 158:3317–3324.

7. Julia, V., M. Assoulaezegad, and N. Glächenhaus. 1996. Resistance to Leishmania major induced by tolerance to a single antigen. Science. 274:421–423.

8. Fowell, D.J., and R.M. Locksley. 1999. Leishmania major infection of inbred mice: unmasking genetic determinants of susceptibility to Leishmania major in BALB/c mice. Immunity. 38:57–67.

9. Bouvier, J., R.J. Etges, and C. Bordier. 1985. Identification and purification of membrane and soluble forms of the major surface protein of Leishmania promastigotes. J. Biol. Chem. 260:15504–15509.

10. Hunt, D.F., H. Michel, T.A. Dickinson, J. Shabanowitz, A.L. Cox, K. Sakaguchi, E. Appella, H.M. Grey, and A. Sette. 1997. Peptides presented to the immune system by the murine class II major histocompatibility complex molecule I-A
d. Science. 256:1817–1820.

11. Seo, S., A. Casrouge, J.D. Altman, M. Haury, J. Kanellopoulos, J.-P. Abastado, and P. Kourilsky. 1998. Individual variations in the murine T cell response to a specific peptide reflect variability in naïve repertoires. Immunity. 9:169–178.

12. Bouso, P., A. Casrouge, J.D. Altman, M. Haury, J. Kanellopoulos, J.-P. Abastado, and P. Kourilsky. 1998. Individual variations in the murine T cell response to a specific peptide reflect variability in naïve repertoires. Immunity. 9:169–178.

13. Boucher, W., E. Mangeat, and P. M. Davis. 1995. Antigen-specific development of primary and memory T cells in vivo. Science. 268:106–111.

14. Gurunathan, S., D.L. Sacks, D.R. Brown, S.L. Reiner, H. Charest, N. Glächenhaus, and R.A. Seder. 1997. Vaccination with DNA encoding the immunodominant LACK parasite antigen confers protective immunity to mice infected with Leishmania major. J. Exp. Med. 186:1137–1147.

15. Kersh, J.G., and P.M. Allen. 1996. Essential flexibility in the T-cell recognition of antigen. Nature. 380:495–498.

16. Windhagen, A., C. Scholz, P. Hollsberg, H. Fukaura, A. Sette, and D.A. Hafler. 1995. Modulation of cytokine patterns of human autoreactive T cell clones by a single amino acid substitution of their peptide ligand. Immunity. 2:373–380.

17. Honak, N.A., K. Shibuya, A.W. Heath, K.M. Murphy, and A. O’Garra. 1995. Modulation of cytokine patterns of human autoreactive T cell clones by a single amino acid substitution of their peptide ligand. Immunity. 2:373–380.

18. DeMagistris, M.T., J. Alexander, M. Coggeshall, A. Altman, F.C. Gaeta, H.M. Grey, and A. Sette. 1992. Antigen analog-major histocompatibility complexes act as agonists of the T cell receptor. Cell. 7:545–555.

19. Fowell, D.J., J. Magram, C.W. Turck, N. Killeen, and R.M. Locksley. 1997. Impaired Th2 subset development in the absence of CD4. Immunity. 6:559–569.

20. Rainer, S.L., S. Zheng, Z.-E. Wang, L. Stowers, and R.M. Locksley. 1994. Leishmania promastigotes evade interleukin 12 (IL-12) production by macrophages and stimulate a broad range of cytokines from CD4+ T cells during initiation of infection. J. Exp. Med. 179:447–456.

21. Sette, A., S. Buus, S. Colon, J.A. Smith, C. Miles, and H.M. Grey. 1987. Structural characteristics of an antigen required for its interaction with M and recognition by T cells. Nature. 328:395–399.

22. Murphy, K.E., A.B. Hemberger, and D.Y. Loh. 1990. Induction of antigen-specific apoptosis of CD4+CD8+ TCR + thymocytes in vivo. Science. 250:1720–1723.

23. Liblau, R.S., R. Tisch, K. Shokat, X.-D. Yang, N. Dumont, C.C. Goodnow, and H.O. M. McDevitt. 1996. Intravenous injection of soluble antigen induces thymic and peripheral T cell apoptosis. Proc. Natl. Acad. Sci. USA. 93:3031–3036.

24. Ravel, C., P. Debussay, J.M. Blackwell, A.C. Ivens, and P. Bastien. 1998. The complete chromosomal organization of the reference strain of the Leishmania genome project, L. major ‘Friedlin’. Parasitol. Today. 14:301–303.

25. M.C. O’Garra, M.J., and A. Bacic. 1990. The glycoinositolphospholipid profiles of two Leishmania major strains that differ in lipophosphoglycan expression. Mol. Biochem. Parasitol. 38:57–67.
tance in reconstituted nude mice and several F1 hybrids infected with Leishmania tropica major. J. Immunogenet. 10:395-412.
34. Mitchell, G.F., E. Handman, and T.W. Spithill. 1985. Examination of variables in the vaccination of mice against cutaneous leishmaniaiasis using living avirulent cloned lines and killed promastigotes of Leishmania major. Int. J. Parasitol. 15:677-684.
35. Varkila, K., R. Chatelain, L.M. Leal, and R.L. Coffman. 1993. Reconstitution of C.B-17 scid mice with BALB/c T cells initiates a T helper type-1 response and renders them capable of healing Leishmania major infection. Eur. J. Immunol. 23:262-268.
36. Mason, D. 1998. A very high level of cross-reactivity is an essential feature of the T cell receptor. Immunol. Today. 19:395-403.
37. Nanda, N.K., R. Apple, and E. Sercarz. 1991. Limitations in plasticity of the T-cell receptor repertoire. Proc. Natl. Acad. Sci. U.S.A. 88:9503-9507.