The Binding Site of Human Adenosine Deaminase for CD26/Dipeptidyl Peptidase IV: The Arg142Gln Mutation Impairs Binding to CD26 but Does Not Cause Immune Deficiency

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Abstract

Human, but not murine, adenosine deaminase (ADA) forms a complex with the cell membrane protein CD26/dipeptidyl peptidase IV. CD26-bound ADA has been postulated to regulate extracellular adenosine levels and to modulate the costimulatory function of CD26 on T lymphocytes. Absence of ADA–CD26 binding has been implicated in causing severe combined immunodeficiency due to ADA deficiency. Using human–mouse ADA hybrids and ADA point mutants, we have localized the amino acids critical for CD26 binding to the helical segment 126–143. Arg142 in human ADA and Gln142 in mouse ADA largely determine the capacity to bind CD26. Recombinant human ADA bearing the R142Q mutation had normal catalytic activity per molecule, but markedly impaired binding to a CD26−/− ADA-deficient human T cell line. Reduced CD26 binding was also found with ADA from red cells and T cells of a healthy individual whose only expressed ADA has the R142Q mutation. Conversely, ADA with the E217K active site mutation, the only ADA expressed by a severely immunodeficient patient, showed normal CD26 binding. These findings argue that ADA binding to CD26 is not essential for immune function in humans.

Key words: adenosine deaminase deficiency • severe combined immunodeficiency • T lymphocyte • protein–protein interaction • adenosine deaminase complexing protein

Introduction

Adenosine deaminase (ADA)1 deficiency in humans causes profound lymphopenia, which is evident by midgestation and results in SCID during infancy (1, 2). Various effects of adenosine (Ado) and 2′-deoxyadenosine (dAdo) have been implicated in pathogenesis (3). Skeletal, neurologic, and hepatic abnormalities that occur in some patients may also be due to the metabolic disorder, but these are of less clinical relevance than the immunodeficiency. In contrast to the phenotype in humans, ADA knockout mice have normal lymphoid development at birth and die perinatally of hepatic and pulmonary injury (4, 5). Lymphopenia develops postnatally in strains genetically engineered to express only placental ADA, but these animals die at a few weeks of age from lung injury (4–6). The greater lymphoid selectivity in humans than mice may reflect differences in purine metabolism, tissue sensitivity to ADA substrates, or timing of development. In this context it is intriguing that human ADA, but not murine ADA, can form a complex with a multifunctional membrane protein that has a role in regulating thymocyte proliferation and T cell activation.

The ADA of human erythrocytes behaves as a soluble monomer of ~41 kD. Larger forms (≥200 kD) found in extracts of other tissues of humans, rabbits, and cattle are due to the binding of ADA to a homodimeric membrane glycoprotein with subunit Mr ~ 110 kD (7–9). This so-called “ADA complexing protein” (ADA-CP) was shown to bind two ADA monomers with Ks of 4–20 nM (8, 10). ADA-CP occurs on secretory or absorptive surfaces of epithelia of liver, gut, kidney, and exocrine glands (11). Mice lack “large” forms of ADA (12), but were found to possess a membrane protein with the tissue distribution and size of ADA-CP, which cross-reacted with Ab to human ADA-
CD45 tyrosine phosphatase, and the HIV-1 Tat protein
CD26 has been reported to bind collagen, fibronectin, the
dues 294 and 340–343 of the cysteine-rich segment are es-
552), and a COOH-terminal region that bears the serine
glycosylated “stalk”, a cysteine-rich segment (residues 290–
brane anchor, a large extracellular domain consisting of a
ficiencies about the nature of the CD26 binding site; neither do
binding of ADA mutations from individuals with ADA de-
the amino acid residues of human ADA that are essential
CD26, that such binding protects lymphocytes from ef-
thymocytes and blood T cells with a helper/memory pheno-
tion used for screening ADA–CD26 binding had DPPIV activity
Among lymphoid cells, CD26 is expressed on medullary
thymocytes, blood T cells with a helper/memory pheno-
type, activated B cells, and NK cells. On T cells, CD26 acts as a “co-stimulator” of antigen receptor–mediated activa-
tion (24). CD26 ligation also stimulates the proliferation of
thymocytes and other hematopoietic cells in the mouse and
rat (25, 26). It has been variously proposed that in humans,
binding of ADA is important to the costimulatory function of
CD26, that such binding protects lymphocytes from ef-
effects of extracellular Ado, and that immunodeficiency in
patients with ADA deficiency is due to the absence of
CD26–associated ADA (15, 23, 24, 27). We have defined the
amino acid residues of human ADA that are essential
for CD26 binding, and investigated the effects on CD26
binding of ADA mutations from individuals with ADA de-
ficiency. Our findings are not consistent with some predic-
cations about the nature of the CD26 binding site; neither do
they support the hypothesis that ADA–CD26 binding is es-
sential for immune function in humans.

Materials and Methods

Materials

DEAE-Sepharose fast flow, Superose 12, and CNBr-activated
Sepharose 4B were obtained from Amersham Pharmacia Biotech.
ADA-Sepharose was prepared by coupling calf mucosal ADA
(Sigma–Aldrich and Boehringer) to CNBr-activated Sepharose
4B according to the manufacturer’s instructions. Plasmid pZC11
and Echericha coli SO3834 were provided by Dr. Rod Kellems
(University of Texas, Houston, TX). Goat anti–human ADA an-
tiserum was provided by Dr. Dan Wiginton (University of Cin-
cinnati, Cincinnati, OH). The mouse mAb 1C5 was prepared
against ADA purified from human T cell leukemia cells (Hersh-
field, M.S., unpublished details).

Lymphoid Cells and Cell Lines

Activated T lymphocytes were prepared by culturing blood
mononuclear cells with phytohemagglutinin and IL-2 as de-
scribed (28), except that the cells were grown in AIM-V serum-
free medium (Life Technologies). The HTLV-1–transformed
AlNle cell line derived from an ADA-deficient patient (provided
by Dr. Ken Weinberg, Los Angeles Children’s Hospital, Los An-
geles, CA) was also maintained in AIM-V medium.

Enzyme Assays

DPPIV activity was determined using Gly-Pro-p-nitroanilide
tosylate (Sigma–Aldrich) as substrate, monitoring A405 at 37°C
(29). ADA activity was determined by monitoring the decrease in
A263 at 37°C using as substrate 150 μM Ado in 50 mM Tris-
HCl, pH 7.4. Both the CD26 and ADA assays were performed in
96-well microwell plates (Costar UV plate; Corning Inc.) in reac-
tion volumes of 0.2 μl, using a SpectraMax plus spectrophotome-
ter (Molecular Devices). ADA was also assayed by a radiochem-
tical-TLC method (28). Protein was determined using the
bicinchoninic acid (BCA) method (Pierce Chemical Co.) with
BSA as standard.

Rabbit Kidney CD26/DPPIV

CD26/DPPIV was purified from New Zealand white rabbit
kidney homogenates by a published procedure involving DEAE-
Sepharose and ADA-Sepharose 4B chromatography (30). In
some preparations, the ADA-Sepharose column was eluted with
6 M urea in 0.01 M KPO4, pH 7.4, 0.1% Triton X-100, as de-
scribed (31). Fractions with DPPIV activity were pooled, dial-
ed against 0.01 M KPO4, pH 7.4, 0.1% Triton X-100, and
stored at –20°C. The purified preparation had a DPPIV–specific
activity of 7,883 μmol min⁻¹ mg⁻¹. A partially purified prepara-
tion used for screening ADA–CD26 binding had DPPIV activity
of 6.5–7.2 μmol min⁻¹ mg⁻¹. ADA activity of these prepara-
tions was <0.003 μmol min⁻¹ mg⁻¹.

Bovine ADA cDNA

Standard methods were used to amplify, clone, and sequence
cDNA (32, 33). Degenerate reverse transcription PCR primers
were used to clone the ADA coding region from bovine thymus
RNA: (+) 5’ATCGAAGCTTCCATGGCCCAGACRCCCGC-
MTC, (−) 5’GGCCATGGAGAGGTAGCCACGACACCTT-
CACAGACA. Both strands were sequenced by the dideoxy
method (sequence data are available from EMBL/GenBank/DDBJ
under accession no. AF280603). The expressed product of the
bovine ADA cDNA was enzymatically active (Kelly, S.J., and M.S.
Hershfield, unpublished data).

Construction of Human–Mouse ADA cDNA Hybrids and Point
Mutants, and Expression of ADA for CD26 Binding Studies

cDNAs consisting of segments from both the human (34) and
mouse (35) ADA coding regions were made by overlap extension
PCR (36; Tables I and II). The P126Q, R142Q, R149Q,
A215T, and E217K ADA mutants have been identified in human
subjects (37–40); their expression in E. coli strain SO3834 has
been reported (41). Other mutations (see Results) were intro-
duced into the wild-type human or mouse ADA cDNAs by
PCR mutagenesis essentially as described (37, 41). All final ADA
cDNA PCR products were cloned into pBluescript II KS or
pBluescript SK and fully sequenced using the ABI 377 PRISM
DNA Sequencing Instrument (Applied Biosystems).
ADA cDNAs were ligated into the NcoI site of pZ (derived from pZC11 from which wild-type human ADA cDNA had been excised [42]). pZ/ADA plasmids were used to transform E. coli SO3834, which has a deletion of the bacterial ADA gene (42). Single transformant colonies were grown at 37°C in Luria broth/ carb/tet medium to constitutively express ADA, as described (41). Under these conditions, the yield of wild-type mouse ADA activity, aliquots of these lysates (20-µg protein) were analyzed by Western blotting using goat anti–human ADA antisera, as described (41). Western blotting was also performed with the 1C5 mouse mAb to human ADA (1:2,000 dilution of ammonium sulfate–concentrated mouse ascites; a goat anti–mouse IgG–horse radish peroxidase conjugate (Santa Cruz Biotechnology, Inc.) was used as a second Ab, and was detected with the Enhanced Chemiluminescence System (Amersham Pharmacia Biotech).

**ADA-CD26 Binding**

In the standard assay, dialyzed (0.01 M KPO₄, pH 7.4) lysates of pZ/ADA cDNA-transformed E. coli SO3834 cells containing 100–250 nmol/min of ADA activity (diluted as necessary with dialyzed lysate of untransformed SO3834 cells) were combined with 10–30 nmol/min (DPPIV activity) of rabbit CD26 in 0.01 M KPO₄, pH 7.4, 0.1% Triton X-100, total volume 150 μL. After incubating for 2 h at 37°C, the entire mixture was injected onto a 1.5 × 30 cm Superose 12 column equilibrated with 50 mM Tris-HCl, pH 8.4 (fast protein liquid chromatography [FPLC] system; Amersham Pharmacia Biotech). The column was eluted with this buffer at room temperature, flow rate 0.5 ml/min. Fractions (0.5 ml) were assayed for ADA and DPPIV activities, and the percentage of total ADA activity associated with the peak of DPPIV activity (CD26) was calculated. In some experiments the column was equilibrated and eluted with 50 mM Tris-HCl, pH 8.4, containing 150 mM NaCl. This shifted the peak of free (41 kD) ADA activity from fraction 25 to fraction 27, but neither the DPPIV activity peak nor the amount of ADA activity associated with the DPPIV peak were altered.

**Gel Mobility Shift Assay.** [35S]Met-labeled ADA was generated in vitro from ADA cDNA constructs in pBluescript, using the TNT Coupled Wheat Germ Extract System (Promega) according to the manufacturer’s instructions (this system was used in preference to rabbit reticulocyte extracts because it possessed very low endogenous ADA-like activity). Translation products were analyzed by SDS-PAGE and fluorography, and were also electrophoresed on cellulose acetate and stained for ADA activity in situ, as described (28). Aliquots of the translation mixtures containing equal amounts of [35S]-labeled products were then incubated (2 h, 37°C) with 0.7–2.2 nmol/min of rabbit CD26/DPPIV in 0.01 M KPO₄, pH 7.4, 0.1% Triton X-100, total volume 20 μl. These reaction mixtures were then electrophoresed on a 3–15% gradient nondenaturing PAGE gel (pH 8.8, 30% acrylamide, 0.8% bisacrylamide; running conditions 60 V/cm, 18 h, 4°C [43]). [35S]ADA was located by fluorography.

**Recombinant ADA Binding to AlNe T Cells**

Aliquots (25 μl) of pZ/ADA cDNA-transformed SO3834 lysates containing 400 nmol/min of ADA activity were added to 2 × 10⁶ AlNe T cells in 1 ml of serum-free growth medium and incubated 2 h at 37°C. The cells were then washed three times with PBS, pH 7.4, and lysed by freezing and thawing (or by sonication, which gave identical results) in 150 μl of 25 mM Tris-HCl, pH 7.5, 1 mM EDTA, 1 mM DTT, 0.1% Triton X-100. After centrifugation, the supernatant was passed through 0.2 μm filters and dialyzed at 4°C against 0.01 M KPO₄, pH 7.4. In addition to assaying ADA activity, aliquots of these lysates (20-µg protein) were analyzed by Western blotting using goat anti–human ADA antisera, as described (41). Western blotting was also performed with the 1C5 mouse mAb to human ADA (1:2,000 dilution of ammonium sulfate–concentrated mouse ascites; a goat anti–mouse IgG–horse radish peroxidase conjugate (Santa Cruz Biotechnology, Inc.) was used as a second Ab, and was detected with the Enhanced Chemiluminescence System (Amersham Pharmacia Biotech).
cubated for 60 min at 37°C in 1 ml of medium containing aliquots of untransformed or ADA-expressing SØ3834 lysates. The cells were then washed with cold PBS containing 1% BSA and 0.1% sodium azide (PBSW) and processed for flow cytometry.

**Flow Cytometry**

Surface antigens and bound ADA on AlNe T cells were detected with the following reagents: unconjugated anti-ADA mAb 1C5 and control mAb P3; and fluorochrome-conjugated Ta1-RD1 (CD26), B1-RD1 (CD20), T11-RD1 (CD2), mouse IgG1-RD1, and either fluorescein- or PE-conjugated goat anti–mouse IgG1 provided by Beckman Coulter (Fullerton, CA). All Ab incubations were performed in PBSW at 4°C. After washing with PBSW, cells were fixed with 0.4% paraformaldehyde, and analyzed using an Epics Elite XL flow cytometer (Beckman Coulter). Data were processed using the FACSConvert™ and CellQuest™ software programs (Becton Dickinson).

**Results**

**Binding of Recombinant ADAs to Rabbit CD26.** Like the enzymes isolated from tissues (10), human and mouse ADA expressed in *E. coli* SO3834 differ markedly in ability to...

### Table II. PCR Primers Used in Constructing Human–Mouse ADA Hybrids

| Primer sequence* | Location in ADA cDNA‡ |
|------------------|-----------------------|
| 1                | (+) CGCGCGGATTCGGGACGAGCTGA | M1–21 (EcoRI/NcoI) |
| 2                | (-) CATCTCTCAAAACTCATAGGGATCATCCTTGATGGCTCTCTGCA | Reverse of primer 3 |
| 3                | (+) TCCAAGAGAGCCATCAAGAGAGATGCTGCTCTTGATGGCTCTCTGCA | M223–243/H244–267 |
| 4                | (-) CGCGCGGATTCGGGACGAGCTGA | H1073–1092 (HindIII/NcoI) |
| 5                | (+) CGCGCGGATTCGGGACGAGCTGA | H1–24 (EcoRI/NcoI) |
| 6                | (-) CTCTCTCCTCCTCCTCAGCTGCTGCTCTTGAGGCTCTCTGAGG | Reverse of primer 7 |
| 7                | (+) CCCTGGAACCGAGACTGAAAGGGAGCGCTGAGGAAGTGGTGA | M346–375/H376–429 |
| 8                | (-) TGCTCTCCTCCTCCTCAGCTGCTGCTCTTGAGGCTCTCTGAGG | H346–375/M376–429 |
| 9                | (+) CTCTGATACGGTTGATAGCCTGCTTCTCTCCTGAGGAGGAGGA | M376–429/H346–375 |
| 10               | (-) GCACAGAGGACGGAGCATTCGGAGACTGCTA

*Cloning sites are underlined. The bold italic nucleotides are silent polymorphisms found in human ADA cDNA.

‡The numbers indicate nucleotide position in human (H) or mouse (M) ADA cDNA (adenine of the initiation codon 5’).
bind rabbit kidney CD26/DPPIV. When equal amounts of ADA activity were compared, 50% of human and 1% of mouse ADA activity coeluted with DPPIV activity from a Superose 12 FPLC column (Fig. 1, A and B). The difference in CD26 binding could also be demonstrated using nondenaturing PAGE to assess the effect of rabbit CD26 on the migration of 35S-labeled human and mouse ADA in vitro translation products (Fig. 1 C). Addition of unlabeled human ADA expressed in E. coli SO3834 (100 nmol/min) blocked the CD26-induced shift in migration of the human 35SADA translation product, whereas the same amount of recombinant mouse ADA had no effect (not shown).

Based on these results, we constructed human–mouse ADA cDNA hybrids (Tables I–III) and used the above assays to locate residues necessary for CD26 binding (analogous to human–rat CD26 “swap mutants” used to identify CD26 residues necessary for ADA binding [22]). To refine our search, we also used point mutants found in ADA-deficient individuals, which we have previously expressed in SO3834 (41). ADA mutants with sufficient catalytic activity were screened by FPLC, although in some cases using less ADA activity than in the “standard” assay (see Materials and Methods). The PAGE assay was used for ADA mutants with very low catalytic activity. Some mutants could not be tested because their activity was too low for the FPLC assay, and their in vitro translation products did not enter the nondenaturing gel (results not shown).

Initial hybrid screening placed the CD26 binding site between residues 82 and 248 (Fig. 2, A and B). This interval was narrowed by results with the R142Q point mutant, which expressed 28% of wild-type human ADA activity in SO3834 (41): in four standard FPLC binding experiments, 10.4 ± 2.9% (mean ± SD) of R142Q ADA activity coeluted with DPPIV activity (Fig. 2 C), resulting in 0.9 ± 0.5 ADA units bound per unit DPPIV. By comparison, in five standard experiments with wild-type human ADA, 48.7 ± 5.1% of ADA activity coeluted with DPPIV activity, resulting in the binding of 5.9 ± 1.4 units of ADA per unit of DPPIV (Fig. 2, A and B). We also tested the P126Q, R149Q, and A215T mutants, which expressed 0.3, 6.2, and 4.8% of wild-type ADA activity, respectively (41). Experiments with these mutants were performed with less ADA activity than under standard conditions. However, 75% to >90% of the added ADA activity coeluted with DPPIV activity (Fig. 2 C), suggesting that these mutations did not impair CD26 binding. Association of 35S-labeled R142Q in vitro translation product with CD26 was undetectable by PAGE, whereas 35S-labeled E217K, an active site mutant with <0.005% of wild-type ADA activity (41, 44), showed normal binding (Fig. 2 D). Reduced CD26 binding by the R142Q mutant was of particular interest because (a) glutamine is the “wild-type” amino acid at position 142 in mouse ADA, and (b) R142Q was identified in a healthy adult whose other ADA allele was not expressed (38).

In the crystal structure of murine ADA, Gln142 lies in the 18-residue segment 126–143, which forms the peripheral α2 helix (44; Fig. 3). A hybrid consisting of human ADA substituted with mouse residues 126–143 (referred to as “m126–143”) expressed 36% of wild-type human ADA activity in SO3834 (Table III), but it showed minimal binding to CD26 by FPLC (Fig. 2, A and B) and none by PAGE (not shown). Binding was also reduced markedly in other hybrids that possessed the murine α2 helix.
residues (Fig. 2, A and B). Hybrids in which all other regions of human ADA, except for residues 126–143, were replaced with their murine counterparts had substantial ADA activity (Table III) and showed essentially normal CD26 binding (Fig. 2, A and B). Modifying mouse ADA by substituting human residues 126–143 for the corresponding mouse residues (“h126–143”), or by making the Q142R point mutation, conferred some ability to bind CD26, which could be appreciated by FPLC (Fig. 2, A–C). Together, these results show that the amino acids critical for CD26 binding lie within segment 126–143 of human ADA.

As bovine ADA also binds CD26, we amplified cDNA carrying the ADA coding region from bovine thymus RNA. The predicted amino acid sequences of residues 115–150 of human, bovine, and murine ADA are compared in Fig. 3. Within segment 126–143, human and bovine ADA differ at only 2 of 18 positions, whereas human and murine ADA differ at 5, and murine and bovine ADA at 4 positions. The most notable differences are (a) residues 141–143 of human and bovine ADA are all charged, Glu-Arg-Asp, whereas only Glu141 is charged in the mouse sequence, Glu-Gln-Ala; and (b) residue 131 in mouse ADA is Asp, whereas Ala and Ser are found in human and bovine ADA, respectively. We prepared mutants of human ADA in which α2 helix residues from murine ADA replaced the corresponding human residues (“h126–143”), or by making the Q142R point mutation, conferred some ability to bind CD26, which could be appreciated by FPLC (Fig. 2, A–C). Together, these results show that the amino acids critical for CD26 binding lie within segment 126–143 of human ADA.

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Figure 3. Location of the α2 helix (CD26 binding site) in the murine ADA crystal structure. The RasMol model is based on coordinates, as reported (reference 44). The α2 helix (residues 126–143) is shown in orange, with space-filling display of residues D131, Q142, and A143. For orientation, the COOH-terminal helix (residues 337–351) is shown in blue. Bound inhibitor (rose) and the zinc ion (yellow) are space-filling structures at the active site. The panel below the model compares the partial amino acid sequences (residues 115–150) of human (HUM), murine (MUS), and bovine (BOV) ADA. Amino acids 126–143 of human ADA, green letters; other amino acids of bovine and murine ADA that differ from the human sequence, blue letters.

shown). AlNe cells express a substantial level of CD26 (see below). We compared the binding of several recombinant ADAs to CD26 on AlNe cells (see Figs. 4–6).

Virtually no ADA activity was found in Superose 12–fractionated extracts of AlNe cells that had been incubated with a high concentration (400 nmol/min) of recombinant murine ADA (Fig. 4 A), or with the m126–143 human–mouse hybrid (Fig. 4 B). After incubation with wild-type human ADA, 91% of cell-associated ADA activity coeluted with DPPIV, resulting in a ratio of ADA to DPPIV activity in this peak of 3.7 (Fig. 4 A). With the R142Q mutant, ~40% as much total ADA activity became cell associated as was found with wild-type human ADA. However, only 9% of the bound R142Q ADA activity coeluted with DPPIV, resulting in a ratio of R142Q ADA to DPPIV activity of 0.1 (Fig. 4 B). Virtually identical results were obtained for wild-type human ADA and the R142Q mutant in a second experiment using the same protocol. The R142Q ADA activity eluting as a 41-kD monomer may have resulted from unstable binding of mutant ADA to CD26 on AlNe cells, with subsequent dissociation in the extract or during FPLC. This transient binding probably reflects the high concentration of exogenous ADA activity used in these experiments (see below).

We next examined binding of recombinant ADAs to the surface of AlNe cells by flow cytometry, using mouse mAb 1C5 to human ADA. Western blot experiments indicated that 1C5 recognizes a COOH-terminal epitope present in human but not in mouse ADA, and it reacts approximately as well with the R142Q and E217K mutants as with wild-type human ADA (Fig. 5). In these experiments (and in flow cytometry studies shown below in Fig. 6 D), equal units of R142Q and wild-type ADA activity were compared (Fig. 5 A), whereas equal amounts of total SO3834 lysate protein were used to compare E217K and wild-type ADA (Fig. 5 B). Taken together with previous studies of expression (38, 41), these results indicate that the R142Q protein has the same activity per molecule as wild-type ADA, but is somewhat less stable in cells, whereas the E217K protein is about as stable as the wild-type enzyme, but is catalytically inactive. The stability of E217K is unusual because most ADA missense mutations identified in SCID patients result in unstable proteins that are difficult to detect with antisera to ADA, both in patient-derived cell lines and when expressed in E. coli SO3834 (41, 46).

In flow cytometry experiments, AlNe cells expressed high levels of CD26 (Fig. 6 A). They did not bind anti-ADA mAb 1C5 (Fig. 6 B). Incubating AlNe cells with recombinant wild-type human ADA (followed by washing) induced 1C5 binding (Fig. 6, C and D), as did the R142Q and E217K mutants (Fig. 6 D). 1C5 binding increased with the amount of ADA to which the cells were exposed (Fig. 6 D). Based on mean fluorescence values, at the lowest concentrations tested, E217K induced about half the 1C5 binding induced by wild-type ADA, but equivalent binding occurred at the higher end of the range (because expression in SO3834 is somewhat variable [41], the amounts of wild-type and E217K ADA protein may not have been identical). 1C5 binding was detected at all concentrations of wild-type and E217K proteins tested, but was not measurable with R142Q ADA at 0.3 and 1.7 nmol/min per ml medium. Wild-type ADA induced 7.5-fold more 1C5 binding than R142Q at ADA activities of 8.3 and 42 nmol/min, and fourfold greater binding at 209 nmol/min (very similar results were found in a second experiment not shown). ADA bound to CD26 is derived from an extracellular source (47). Serum ADA1 activity (the isozyme derived from the ADA gene) for 320 normal humans was reported to be 4.2 ± 1.5 nmol/min per ml (48). Thus, under physiological conditions, binding of circulating R142Q ADA to T cells via CD26 may be 10-fold lower than normal.

Binding of Naturally Occurring R142Q ADA to Human CD26. Human ADA expressed in E. coli might interact differently with CD26 than would ADA produced in human cells. To address this possibility, we studied the binding of ADA in extracts of erythrocytes and cultured T cells from the individual in whom the R142Q allele was identified, whose second ADA allele carries a nonsense mutation in codon 3 (38). A limited quantity of R142Q cells was
available for these studies, but the results support those obtained with the recombinant enzyme.

As a control for the erythrocyte studies, we used red cells from an ADA heterozygote whose nonwild-type allele also has a nonsense mutation. ADA activities of the control and R142Q hemolysates was 18.8 and 5.5 nmol/h per mg protein (normal: 63 ± 41 nmol/h per mg), respectively. As erythrocytes lack CD26, we examined CD26 binding by incubating aliquots of hemolyzed red cells with AlNe T cells (ADA-deficient). After washing, Triton X-100 extracts of the AlNe cells were analyzed by FPLC (Fig. 7 A). About 2.6-fold more control (wild-type) than R142Q ADA activity eluted as monomer, proportional to the amounts of ADA activity in the incubations. However, ~14-fold more control (wild-type) than R142Q ADA activity coeluted with DPPIV activity.

For the T cell studies we assessed the interaction of endogenous ADA with endogenous CD26 (Fig. 7 B). The control T cells were obtained from an individual presumed to be homozygous for wild-type ADA. ADA activity of the control and R142Q T cell extracts were 1,750 and 592 nmol/h per mg protein, respectively; they expressed approximately equal levels of CD26 by flow cytometry (not shown). After FPLC fractionation of lysates of these cells, ~1.6-fold more control (wild-type) than R142Q ADA activity eluted as monomer, whereas 45-fold more control than R142Q ADA activity coeluted with DPPIV activity, resulting in ratios of ADA units bound per unit CD26 of 1.8 for the control and 0.03 for the R142Q T cells (Fig. 7 B).

**Discussion**

Murine ADA has a parallel α/β architecture with an eight-stranded central β barrel and eight peripheral α helices; the active site containing an essential Zn²⁺ ion lies at the COOH-terminal end of the β barrel (44; Fig. 3). Human ADA is 83% identical in sequence and presumably has the same overall structure. Using a panel of recombinant human–mouse hybrids that retain catalytic activity, we have identified the COOH end of the peripheral α2 helix as the primary determinant of the difference in the ability of mouse and human ADA to bind CD26/DPPIV. Replacing the entire 18-residue segment in human ADA with murine residues 126–143 modestly reduced the expression of ADA activity in E. coli, but abolished stable binding to rabbit and human CD26. Both effects could largely be achieved by replacing Arg142 of human ADA with Gln, the residue found at this position in murine ADA. At the resolution of our screening assays, human to mouse substitutions at the other four nonidentical positions within this segment did not reduce CD26 binding significantly, and in various combinations with R142Q they had no greater effect than the R142Q substitution alone. Making the converse Q142R substitution in mouse ADA, or introducing the entire human 126–143 segment, conferred some ability to bind rabbit CD26. No other regions of human ADA appear to be involved in CD26 binding, including the...
11 COOH-terminal residues, which are absent in murine ADA.

Residues 294 (Leu) and 340–343 (Leu-Val-Ala-Arg) of human CD26 have been identified as essential for binding ADA (22, 23). It was postulated that stable ADA–CD26 binding is due to contacts between these CD26 residues and a nonconserved hydrophobic surface of human ADA, possibly involving residues Leu346, Ala350, and Gly352, which are respectively Arg, Glu, and Gln in mouse ADA (23). This cannot be correct because the human–mouse ADA hybrid h1–247/m248–352 binds CD26 (Fig. 2). Moreover, residues 141–143 are Glu-Arg-Asp in human and bovine ADA, each of which binds CD26, and Glu-Gln-Ala in mouse ADA, which does not, suggesting that charged residues of ADA are critical for binding CD26. This type of contact between proteins, involving mainly hydrophobic residues of one partner and charged residues of the other, is not unique, and has recently been observed in the complex of hen egg white lysozyme with the antilysozyme mAb HyHEL-63 (49). Stable ADA–CD26 binding may also arise from interactions of the critical hydrophobic CD26 and charged ADA residues, not with each other, but with other partners that will be identified when the 3D structures of human ADA and a CD26–ADA complex have been determined.

**Significance of the R142Q Mutation.** Most patients with ADA deficiency have SCID, but 15–20% have a milder immune deficit with a later clinical onset (37, 45, 50). Screening of populations and studies of the relatives of patients with SCID has identified some healthy individuals, most of African ancestry, with low or absent ADA activity in red cells, but 5–70% of normal activity in nucleated cells; this “partial ADA deficiency” has been shown to result from several different missense mutations (37, 39, 51). Red cells of subjects with partial ADA deficiency show no (or minimal) accumulation of dAdo nucleotides (dAXP), whereas a massive increase occurs in patients with SCID, and a more modest elevation in those with milder immune deficiency (3, 41, 45, 52). This correlation, and evidence that dATP is toxic to lymphoid cells and can induce apoptosis by p53- and caspase-dependent mechanism(s) (53–59), has implicated dAdo-induced dATP pool expansion as the principal cause of lymphopenia in ADA deficiency.

With this background, the finding of <20% of normal red cell ADA activity in the healthy Somali father of a child with SCID led to our discovery that he had transmitted an ADA allele with a codon 3 nonsense mutation to his affected daughter, and that his second ADA allele carried the R142Q mutation (38). We postulated that the R142Q mutation, which is distant from the active site, reduced cellular ADA activity by impairing enzyme folding or stability, but that residual activity due to the R142Q allele was sufficient to catabolize dAdo, preventing dATP accumulation and immune deficiency. Our present studies with the 1C5 anti-ADA mAb show that the R142Q protein has a specific activity per molecule close to that of wild-type human ADA. In our original report of the R142Q allele, we also identified SCID patients who were homozygous for a codon 142 nonsense mutation (38). Both the R142Q and R142X mutations arose from a CGA (Arg) codon. We

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**Figure 6.** Flow cytometry analysis of CD26 and ADA on the surface of AlNe cells. (A) CD26 expression. Shown are reactivity of AlNe cells with PE-conjugated anti-CD26 mAb Ta1 (shaded histogram) and control mAb IgG1-PE (open histogram). (B and C) ADA. Anti-ADA mAb 1C5 binding to AlNe cells that had been washed after incubation with lysate of untransformed SO3834 (B) or with lysate of SO3834 expressing wild-type human ADA (400 nmol/min per ml of medium) (C). (D) Binding of wild-type (wt) human ADA (circles), R142Q (triangles), and E217K (squares) ADA mutants. AlNe cells were incubated with 0.3, 1.7, 8.3, 42, 209, and 400 nmol/min per ml of recombinant wild-type and R142Q ADAs. The amounts of E217K-expressing SO3834 lysate protein used (0.024–30 μg) were made equal to wild-type ADA lysate protein. After washing, cell surface–associated ADA was determined by reactivity of mAb 1C5 and flow cytometry. For clarity, the horizontal axis shows only units of ADA activity. Data shown are from one of two experiments.
ADA activity; triangles, Tris-HCl, pH 8.4, 150 mM NaCl (see Materials and Methods). Circles, that the Superose 12 column was equilibrated and eluted with 50 mM cells were lysed, fractionated on Superose 12, and assayed as in A, except from AlNe T cells. 10^7 AlNe T cells in 1 ml medium were incubated 2 h CD26. (A) Binding of ADA from erythrocyte lysates (RBC) to CD26

speculated that during murine evolution there might have been some advantage in replacing this Arg codon, which contains a mutagenic CpG dinucleotide, with a CAA Gln codon (38). Our finding that Arg 142 is the “keystone” of the bridge between ADA and CD26 permits an additional speculation: that at the protein level, a Gln142 residue might have conferred some evolutionary advantage on mice by reducing interaction between ADA and CD26.

Our studies of CD26 binding by the R142Q and E217K mutants, derived from ADA-deficient individuals with distinctive phenotypes, are relevant to theories implicating CD26 in the pathogenesis of the immune defect in ADA deficiency. These have postulated either: (a) that ADA binding per se (unrelated to catalytic activity) is crucial to an essential function of CD26 in lymphocyte development or activation (27, 60); or (b) that CD26-bound “ecto-ADA”, by controlling the level of extracellular Ado, regulates signaling through cell surface Ado receptors (assumed to be critical for lymphocyte development or function [15, 24, 27, 47]). The first hypothesis ignores evidence, discussed above, of the correlation between mutant ADA catalytic activity and the metabolic and clinical phenotypes of ADA-deficient individuals (3, 41, 61). The ADA active site mutation E217K was identified in a patient with early onset SCID, whose other ADA allele is not expressed (40). We have shown that the catalytically inactive E217K protein is stable and binds to CD26 on the surface of T cells in a manner similar to wild-type human ADA, suggesting that ADA–CD26 binding is not sufficient to preserve immune function.

The second hypothesis, that CD26-bound “ecto-ADA” regulates levels of extracellular Ado, has been supported by in vitro studies showing lesser effects of exogenous Ado (0.1−10 mM) on proliferation and IL−2 production by CD26-transfected compared with untransfected Jurkat cells (47). Flow cytometry with anti-ADA antiserum indicated that only the former possessed cell surface–associated ADA (although CD26 on these cells was not saturated with ADA). Whole cell lysates of the transfected and untransfected cells had the same total ADA, as estimated by Western blotting. The physiologic relevance and interpretation of these observations may be questioned. The Ado concentrations used were at least 100-fold higher than occurs in normal plasma and at least 10-fold higher than has been found in plasma of patients with ADA deficiency (3). The differences in proliferation and IL−2 production observed were attributed to different rates of Ado deamination (despite identical total ADA levels), but this was not measured. This result, if true, would be surprising because ADA substrates (at physiologic levels) can equilibrate efficiently across the cell membrane via a ubiquitous, nonconcentrative nucleoside transporter (62). As a result of this transport, transfused ADA" erythrocytes (63) and polyethylene glycol–modified ADA acting in plasma (64) can correct metabolic abnormalities in cells of ADA-deficient individuals.

Ado receptor–mediated signaling is functional in mice despite the lack of ADA–CD26 binding. In addition, a strain of Fischer 344 rats is homozygous for a CD26/DPPIV missense mutation that abolishes both peptidase catalytic activity and cell surface expression of the protein (65, 66). Studies of these rats have indicated a role for DPPIV in renal and intestinal hydrolysis and absorption of prolyl dipeptides (67, 68). Neither CD26/DPPIV-deficient rats nor normal mice manifest the purine metabolic abnormalities and profound immunodeficiency (or nonlymphoid pathology) associated with ADA deficiency in humans and mice.

Advocates of ADA–CD26 hypotheses may consider rodents to be poor models for ADA deficiency in humans (24). In this regard our observations have special relevance. There is good evidence that ADA bound to CD26 on human T cells is not derived from within these cells, but is acquired from the extracellular space (possibly after release from senescent cells [47]). In the range of ADA activity found in normal human plasma (48), we observed ~10−
fold less binding of recombinant R142Q than wild-type human ADA to CD26+ ADA− AlNe T cells. The defect in CD26 binding was even greater with ADA from erythrocytes and T cells obtained from the R142Q-expressing subject (Fig. 7). It is reasonable to expect that his lymphocytes and other tissues also have a markedly reduced level of ADA–ligated CD26 in vivo. The existence of a healthy adult with defective ADA–CD26 binding suggests that interaction of these proteins is not essential for the development or maintenance of immune function in humans, as is certainly the case in mice and perhaps some other species.

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