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Extracellular Isoforms of CD6 Generated by Alternative Splicing Regulate Targeting of CD6 to the Immunological Synapse

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The great majority of mammalian genes yield multiple transcripts arising from differential mRNA processing, but in very few instances have alternative forms been assigned distinct functional properties. We have cloned and characterized a new isoform of the accessory molecule CD6 that lacks the CD166 binding domain and is expressed in rat and human primary cells. The novel isoform, CD6Δd3, results from exon 5 skipping and consequently lacks the third scavenger receptor cysteine-rich (SRCR) domain of CD6. Differential expression of the SRCR domain 3 resulted in a remarkable functional difference: whereas full-length CD6 targeted to the immunological synapse, CD6Δd3 was unable to localize at the T cell:APC interface during Ag presentation. Analysis of expression of CD6 variants showed that, while being more frequent in coexpression with full-length CD6, the CD6Δd3 isoform constituted the sole species in a small percentage of T cells. In the rat thymus, CD6Δd3 is less represented in double-positive thymocytes but is detectable in nearly 50% of single-positive CD4 or CD8 thymocytes, suggesting that CD6 switching between full-length and Δd3 isoforms may be involved in thymic selection. Strikingly, CD6Δd3 is markedly up-regulated upon activation of T lymphocytes, partially substituting full-length CD6, as evaluated by RT-PCR analysis at the single-cell level, by immunoblotting, and by flow cytometry using Abs recognizing SRCR domains 1 and 3 of human CD6. This elegant mechanism controlling the expression of the CD166 binding domain may help regulate signaling delivered by CD6, through different types of extracellular engagement. The Journal of Immunology, 2007, 178: 4351–4361.

The TCR recognition of peptide-MHC complexes expressed on APCs involves the formation of a tight cell-to-cell contact area, the immunological synapse (IS), where individual proteins are selectively partitioned (1, 2). The model of IS, mediating early and late T cell-APC interactions required aggregation of the small adhesion receptors CD2 and CD58, which anchor the region of the contact and allow the TCRs to scan MHC-peptide complexes (3). With time, stably engaged TCR complexes coalesce into a central area (cSMAC; central supramolecular activation clusters), surrounded by a ring (pSMAC; peripheral supramolecular activation clusters) enriched in the integrin LFA-1 (2, 4). In the mature synapse, cSMAC are also enriched in the costimulatory molecule CD28 and the CD4 or CD8 coreceptors (3, 5). It has been suggested that the formation of the synapse is favored by receptor-ligand interactions of small auxiliary molecules such as CD2-CD58 and CD28-CD80 that stabilize the cell contact by the formation of low-affinity interactions, in a size dependent-manner. According to this model, the small adhesion pairs cluster at a tight membrane region, thus segregating larger glycoproteins such as CD43 or CD45 (1, 6).

CD6 is a 100- to 130-kDa surface glycoprotein expressed primarily on medullary thymocytes and mature T lymphocytes (7). CD6 has been regarded as a costimulatory molecule as Ab-mediated CD6 cross-linking can potentiate proliferative T cell responses (8–10). Moreover, a subpopulation of CD6-negative T cells displayed lower alloreactivity in MLRs compared with normal CD6+ T cell populations (11). CD6 could additionally play a role in thymocyte maturation, because CD6 Abs have been shown to partially block the adhesion of thymocytes to thymic epithelial cells (12). This observation also provided the first evidence that CD6 had a cell surface ligand. A requirement for ligand engagement for costimulatory effects of CD6 was shown by inhibition of Ag-specific and CD3 mAb-induced T cell responses by soluble CD6 (13, 14). Recently, CD6 has been reported to accumulate at the IS, mediating early and late T cell-APC interactions required for IS maturation and cell proliferation (13, 15).

Upon CD3 engagement CD6 becomes phosphorylated on tyrosine residues, suggesting that interactions with Src homology (SH)2 domain-containing intracellular effectors may occur (16).
An interaction with the SH2 domain of SLP-76 has been shown to be critical for the costimulatory effects of CD6 (17). The cytoplastic domain of CD6 is unusually long and also contains two well-conserved proline-rich sequences, which are potential binding sites for SH3 domain-containing proteins well suited for signal transduction (16, 18). Indeed, the rat homolog of CD6 has been shown to associate with protein tyrosine kinases of different families, namely Src-family kinases Lck and Fyn, Zap70 of the Syk family, and the Tec-family kinase Itk (19). CD6 has additionally been shown to associate at the surface of T cells with CD3 (13) and with the structurally related receptor CD5 (19, 20). However, there are still no data proposing a functional role for these interactions.

By contrast, the molecular basis of the interaction between CD6 and the extracellular physiological ligand, CD166, has been comprehensively investigated in human and murine models (21). CD166 is a widely expressed glycoprotein containing five Ig superfamily domains, of which the N-terminal domain has been shown to bind to the third scavenger receptor cysteine-rich (SRCR)-type domain of CD6, the membrane-proximal domain (22, 23). The residues of CD6 involved in the contact with CD166 are located in the E-F loop, a region that, in the particular case of the third domain of CD6, is most divergent from all known SRCR domains (24). The resulting interaction between CD6 and CD166 is therefore unusual, because most other T cell surface proteins mediating cell-cell interactions bind through their N-terminal domains. However, affinity and kinetic measurements revealed that, in solution, binding occurs with a mildly strong compared with most other leukocyte adhesion pairs, and that removal of each of these exons would result in a protein sequence lacking the SRCR domain 3. This CD6 isoform, CD6Δd3, is present in all T lineage cells studied and is up-regulated upon T cell activation, paralleling a decline in the expression of full-length CD6 (Cd6Δd3 binding to CD166 highlights the role of domain 3 induced upon T cell activation reveals a rare mode of positional control of cell surface receptors dependent on alternative mRNA splicing.

Materials and Methods

Cells and cell lines

Rat thymocytes, splenocytes, and cervical lymph node cells were from 9- to 12-wk-old Lewis male rats (Charles River Laboratories). Human peripheral blood lymphocytes were isolated by centrifugation on a Ficoll gradient followed by negative depletion on magnetic beads (T cells, negative isolation kit; Dynal Biotech), or using the RosetteSep human T cell enrichment mixture (StemCell Technologies), where indicated. Cell lines COS7 (31) and Raji (32) were maintained in RPMI 1640 supplemented with 10% FCS, 1 mM sodium pyruvate, 2 mM l-glutamine, penicillin G (50 U/ml), and streptomycin (50 μg/ml).

Abs and reagents

Rat mAbs used were CD6-OX52 (33) (a gift from Dr. Horák, University of Oxford, U.K.), TCR-734 (34), and CD28-J313 (35) (gifts from T. Hünig, University of Würzburg, Germany). Conjugated mAbs used for cell sorting were as follows: biotinylated CD4-W3/25 (obtained from Serotec); CD6-OX52 FITC-labeled, CD3-1F4, and CD45SRC-OX22, conjugated with PE, biotinylated CD8β-341 and CD134-OX40, and CD4-OX35 conjugated with allophycocyanin (obtained from BD Biosciences). Biotinylated Abs were detected with streptavidin complexes conjugated with PE, PE-Cy7, or allophycocyanin. Human mAbs were as follows: OKT6 (17), specific for human CD6 domain 3, FITC-labeled UMC6D (9), anti-domain 1 of CD6 (obtained from Ancell); MEM-98 (36) (anti-CD6 domain 1; a gift from V. Hofejš, Academy of Sciences, Prague, Czech Republic), CD166-3A6 (37) (BD Pharmingen), and CD71 (DakoCytomation). Goat anti-mouse peroxidase-conjugated Abs were purchased from Molecular Probes Europe and rabbit anti-mouse FITC-labeled from DakoCytomation. Biotinylated proteins were detected in Western blotting with ExtrAvidin peroxidase (Sigma-Aldrich).

cDNA cloning and plasmids

Rat CD6 cDNAs were cloned as described previously (19). Full-length rat CD6 (CD6FL) cDNA (19) was extracted from pcR2.1-TOPO vector (Invitrogen Life Technologies) using SpeI and ApaI and blunt-end cloned into the expression vector pEF-BOS (38). CD6Δd3/pEF-BOS was obtained by digestion with SpeI and BamHI of a sequence flanking the alternatively spliced exon 5 from the rCD6Δd3/pCR2.1-TOPO template. The vector encoding rat CD6FL-YFP was produced as follows: rat CD6 cDNA was amplified from the original clone rCD6/pCR2.1-TOPO (19) by PCR using as the reverse primer 5′-GCAGGATTCGAAAGCTTGCTGCCTAATGTCGAG-3′. The PCR product was then cloned into the pEFYFP-N1 vector (Clontech Laboratories) in frame with the N-terminal sequence of the yellow fluorescent protein (YFP), in the HindIII restriction sites included in the primers. The rCD6Δd3/pEYFP-N1 construct was obtained by digestion as described above and cloned in frame with pEYFP-N1. A cytoplasmic deletion mutant of rat CD6 fused to YFP was produced by PCR from rCD6/pCR2.1-TOPO using as reverse primer 5′-CTCTAATGCTTTGCTCGTATGGTGC-3′ and the reverse primer in the exon 7 (TM-coding sequence) of human Cd6 5′-GGGATCCCTCTAGATGTTGCTCTTCTCGGATCCTGTATTG-3′ and reverse primer in the exon 7 (TM-coding sequence) of human Cd6 5′-GGGATCCCTCTAGATGTTGCTCTTCTCGGATCCTGTATTG-3′. The PCR fragments were cloned into pcR2.1-TOPO vector. The sequences were confirmed by sequencing.

C-terminal deletion mutants of human CD6 flxed to GFP was produced as follows: amplifying a full-length sequence from the clone CD6Δd3/pEYFP-N1 (18) by PCR, using as the forward primer 5′-GGGATCCCTCTAGATGTTGCTCTTCTCGGATCCTGTATTG-3′ and the reverse primer 5′-CTCTAATGCTTTGCTCGTATGGTGC-3′ and the reverse primer in the exon 7 of human CD6 including a naturally occurring EcoRI restriction site 5′-GTACTAAGATCCAGATGAGGAGTGGAGGAGGATCGGCACTAGATCTTTTTCTTCTG-3′. The PCR product was then cloned into the BamHI restriction site of the pEGFP-N1 vector (Clontech Laboratories) producing CD6FL/pEGFP-N1. The isoform CD6Δd3 was amplified from the vector pCR2.1-TOPO using as primer M13-R from the forward primer, and a reverse primer in the exon 7 of human CD6 including a naturally occurring EcoRI restriction site 5′-GTACTAAGATCCAGATGAGGAGTGGAGGAGGATCGGCACTAGATCTTTTTCTTCTG-3′. The PCR product was inserted in the CD6FL/pEGFP-N1 producing CD6Δd3/pEGFP-N1. A cytoplasmic deletion mutant of human CD6 fused to GFP was produced by PCR from CD6Δd3/pEYFP-N1 with the forward primer 5′-CTCCAGACATGGCACGTCCTG-3′ and the reverse primer 5′-GCTTTATCTCTGTTGGAAGAAA-3′, terminating just before the first tyrosine residue codon of the cytoplasmic domain. The PCR product was cloned into the TOPO cloning site of pcDNA3-1/CT-GFP-TOPO, using GFP Fusion TOPO TA Expression Kits (Invitrogen Life Technologies; version I), to produce CD6CSY5/pDNA3.1/CT-GFP-TOPO.

Rat CD6 cDNA was obtained from total RNA isolated from Lewis rat lungs and reverse transcribed using a gene-specific primer 5′-CC AGGCACGCTTAGAGGAT-3′. Amplification of the full-length molecule was performed using as forward primer 5′-TGTAGAAGACCTC
Cell transfections

Rat CD6FL and CD6Δd3 in pEF-BOS, or the empty vector, were transiently transfected in COS7 cells following the procedures described previously (39).

Transient transfections of primary T lymphocytes (5 × 10⁶ cells) were performed with 5 µg of rat CD6FL/pEYFP-N1, CD6Δd3/pEYFP-N1 or CD6Δd3/pEGFP-N1 or human CD6FL/pEGFP-N1, CD6Δd3/EGFP-N1 or CD6CY5/pcDNA3.1-CT-GFP-TOPo plasmids using the Human T cell Nucleofector kit (Amaxa), and T cells were used 24 h posttransfection. Transfection of Raji B cells was performed as follows: 10⁷ cells were centrifuged, resuspended in 900 µl of prewarmed complete medium, mixed with 20 µg of rat CD66FL/pECFP-N2 plasmid, and transferred into a GenePulse cuvette (Bio-Rad). Electroporation was performed in a Bio-Rad Gene Pulser II electroporator at 950 µF and 260 V. Cells were maintained in complete medium at 37°C and used 48 h after transfection.

Cell surface biotinylation, immunoprecipitations, and Western blotting

Cell surface biotinylation, immunoprecipitations and detection of biotinylated Ags, and immunoblotting were performed as described previously (40).

Single-cell RT-PCR analysis

RT-PCR was performed on single cells purified on the basis of their expression of selected markers after two rounds of cell sorting using a FACSAvantegde equipped with an automatic deposition unit (BD Biosciences). The characteristics and sensitivity of the RT-PCR method have been previously described in detail (41). Briefly, cells were lysed and the RNA reverse transcribed using a Cds-specific primer, 5′-GAGTCTCT ATCTTCAGGCT-3′, and the resulting cDNAs amplified in a two-step nested PCR. The 5′ primers used were as follows: 5′-ATCCACCGTAC CAAGTGA-3′ and 5′-AGACCACTACTCGGCACCA-3′ (nested), and the same reverse primer was used for the reverse transcription reaction. None of the primer combinations amplifies genomic DNA.

Flow cytometry

Flow cytometry was performed as described previously (40).

Cellular activation

A cell suspension was obtained from rat spleen, and 2 × 10⁶ cells/ml were activated for 72 h in vitro by incubation with anti-TCR (R73) plus anti-CD28 (JJ319) mAbs used at 2–10 µg/ml, or left untreated. Cells were incubated at 37°C with 5% CO₂ and, after the indicated period, collected, labeled for OX40, sorted or lysed, and analyzed by immunoblotting.

Human primary T cells were activated by incubating 3 × 10⁶ cells/ml with PHA-p at 5 µg/ml in RPMI 1640, at 37°C with 5% CO₂, or left untreated. After 72 h, cells were collected and analyzed by flow cytometry or immunoblotting.

Conjugate formation and fluorescence analysis

Raji B cells were incubated with a mix of superantigens (staphylococcal enterotoxin A (SEA), SEB, and SEC3, 200 ng/ml each; Toxin Technologies) and plated on poly-γ-lysine-coated glass coverslips for 30 min at 37°C. T cells were added to APCs and then incubated at 37°C for 45 min. Cells were fixed with 4% paraformaldehyde in PBS for 10 min and washed several times with PBS before analysis. Where indicated, T cells were preincubated, for 30 min at 4°C, with the mAbs OX126 or UMCD6-FITC, and Raji B cells preincubated with 3A6 (CD166), all at 10 µg/ml, or left untreated. Immunofluorescence and transmission light images were acquired on an Eclipse TE300 inverted microscope (Nikon) equipped with a cooled CCD camera (CoolSNAPFx; Roper Scientific). Images were acquired and analyzed using the Metamorph software (Roper Scientific). Conjugate formation and synapse localization of CD6 or CD166 were quantified with blind scoring, counting a minimum of 50 productive conjugates in each of two or more experiments, and each experiment was observed by two to three examiners.

Results

A novel isoform of rat CD6 devoid of the CD166 binding domain arises from alternative splicing

Human and mouse CD6 possess several isoforms characterized by cytoplasmic tails of variable lengths, resulting from alternative splicing of exons coding for the intracellular domain (18, 28–30). Subsequent to our recent cloning of a cDNA of rat CD6 (19) containing all 13 exons homologous to the mouse and human sequences, we proceeded to a systematic search for novel rat CD6 isoforms. Using total RNA isolated from Lewis male rat spleens, cDNAs were obtained by reverse transcription followed by PCR using primers complementary to sequences immediately before and after the coding sequence. We obtained five cDNA species displaying distinct mobility on agarose gels (Fig. 1A). Sequencing confirmed the longest product as full-length CD6. Interestingly, sequencing of the third largest product (indicated by an arrow) revealed a novel isoform lacking not a cytoplasmic-coding exon, but missing instead the sequence corresponding to exon 5 (GenBank accession no. AY683561). Alignment of the corresponding amino acid sequence with that of wild-type rat CD6 shows that this cDNA codes for a novel isoform lacking the third SRCR domain, which contains the binding site for the ligand, CD166 (Fig. 1, B–D).

CD6Δd3 is expressed at the cell surface

To verify whether the novel isoform, CD6Δd3, could be correctly folded, transported, and expressed at the cell surface, we cloned the corresponding cDNA into the expression vector pEF-BOS and used it for transfecting COS7 cells. In parallel, COS7 cells were transfected with rCD6FL/pEF-BOS, encoding full-length rat CD6, and also with an empty vector. Forty-eight hours after transfection, cells were analyzed for membrane expression of CD6 by immunoprecipitation, using the CD6-specific mAb OX52, and streptavidin-peroxidase detection from lysates of surface biotinylated cells. As shown in Fig. 2, detection of CD6Δd3 confirmed its correct folding and expression at the cell surface. The apparent molecular mass of this isoform is 112 kDa, compared with 126 kDa of full-length CD6. No product was detected in the lane corresponding to empty vector.

Full-length CD6, but not CD6Δd3, targets to the IS upon T cell-APC conjugate formation

An accumulation of human CD6 at the IS has been recently reported in Jurkat-Raji cell conjugates (20). We thus tested the dependence on CD166-binding interactions for CD6 recruitment to the synapse, by using both CD6FL and CD6Δd3 isoforms. T cells were transfected with constructs coding for either CD6FL-YFP or CD6Δd3-YFP fusion proteins. In isolated T cells, the distribution of both isoforms is homogenous at the plasma membrane (Fig. 3A), a result which additionally confirms that CD6Δd3 can be efficiently expressed at the plasma membrane. Incubation of T cells with Raji B cells primed with superantigen, but not expressing rat CD166, did not change the pattern of CD6FL or CD6Δd3 distribution at the cell surface (Fig. 3B). However, when Ag-primed Raji expressed rat CD166 (shown in green) and T cells expressed CD6FL, both molecules were able to concentrate at the Raji-T cell contact zone (Fig. 3C, left panels, and E). Interestingly, the CD6Δd3 isoform did not efficiently relocalize to the synapse, and, moreover, did not induce colocalization of CD166 as well (Fig. 3C, right panels, and E). This result indispensably proves that the effective CD166-dependent CD6 recruitment to the synapse relies on the presence of the third SRCR domain of CD6. Conversely, rat CD166 targets to the synapse when its T cell ligand rat CD6, expressing the SRCR domain 3, is expressed at the T cell surface.

To verify whether intracellular interactions with other signaling intermediates or with the cytoskeleton could also be involved in the
translocation of CD6 to the IS, T cells were induced to express CD6CY5, a truncated form of the protein retaining only 5 aas of the cytoplasmic domain, fused to YFP. These cells were incubated with Ag-primed Raji cells, expressing rat CD166 (Fig. 3). The truncated form of CD6 was found to localize in the IS as efficiently as the full-length form (Fig. 3, D and E), excluding an exclusive role for cytoplasmic domain interactions on CD6 targeting to the contact zone.

Cd6 expression in different lymphoid tissues and individual cells

Given a potential distinct role of the two isoforms, we conducted a detailed analysis on the profiling of expression of Cd6 in different lymphoid organs, specifically addressing the frequency of Cd6Δd3. We performed PCR from mRNA from thymus, spleen, and lymph nodes amplifying only the sequences corresponding to the extracellular and transmembrane domains, thus excluding cytoplasmic domain isoforms. Two major isoforms were obtained in all three organs, and at similar proportions (Fig. 4A). The heaviest and most abundant product matched the entire sequence, whereas the smallest product, 300 nt shorter, matched the isoform lacking exon 5. This experiment was reproducible and PCR products from several trials were extensively sequenced and confirmed the

FIGURE 1. A novel CD6 isoform lacks the CD166 binding domain. A, Total RNA from Lewis rat spleens was used for RT-PCR. The cDNA products were analyzed on gels, isolated, and sequenced. One mRNA species, indicated by an arrow, corresponded to an isoform lacking the coding sequence of the SRCR domain 3. B, Alignment of CD6FL and Cd6Δd3 peptide sequences from the N terminus up to the transmembrane domain. The boundaries of the SRCR domains are indicated by arrows, according to Hohenester et al. (24). Residues involved in binding to CD166 are gray shaded (54) and are missing in the sequence Cd6Δd3 (dashed lines). Asterisks denote identical residues. C, Organization of the rat Cd6 locus. Introns are shown as thin lines connecting exons, represented by □. The □ indicate the exons coding for the three SRCR domains. Alternative splicing across exon 5 is indicated by solid lines. D, Scheme of the structure obtained for the two CD6 protein isoforms.
expression of only these two isoforms but not of others, for example excluding exons 3 or 4.

The pattern of expression of the CD6 gene in bulk cellular preparations suggested that both isoforms could be found in all tissues analyzed. However, it did not clarify whether each isoform was the sole species in a given cell, or whether both isoforms could be coexpressed in the same cell. We sorted different cell populations and addressed Cd6 expression at the single-cell level. The target populations for this study were developing thymocytes at different maturation stages as well as diverse subsets of T lymphocytes (Fig. 4B). Cells were sorted based on the protein expression of specific markers, and expression of each isoform was assayed by single-cell RT-PCR. All sorted populations were gated on CD6\textsuperscript{high}-expressing cells, and thus Cd6 amplification directly attested for plating efficiency, with no need for parallel amplification of an internal control.

PCR amplification was performed using primers flanking the spliced exon, rendering products of 452 bp for full-length Cd6 and 148 bp for Cd6\textsuperscript{H9004d3}. Fig. 4C shows an analysis of PCR products obtained from CD3\textsuperscript{+}CD4\textsuperscript{+}CD8\textsuperscript{−} individual thymocytes. All possible CD6 phenotypes were present in this population, with many cells expressing only full-length CD6, a few expressing exclusively the Cd6\textsuperscript{H9004d3} isoform, whereas others coexpressed both. In mature T cells, the frequency of Cd6\textsuperscript{H9004d3} expression was remarkably consistent, regardless of the cell type studied. Nearly 40% of total CD4 or CD8 T cells expressed Cd6\textsuperscript{H9004d3} at detectable levels (Fig. 4D, left panel).
addition, CD4⁺CD45RChigh and CD4⁺CD45RClow splenocytes, which have been associated with a naive and memory phenotype, respectively, showed no significant variation in the percentage of cells expressing Cd6d3 (Fig. 4D, left panel), although the ratio between the number of cells expressing only Cd6Δd3 and cells where coexpression was observed was the highest overall in memory cells and the lowest in naive cells.

As distinct from mature T cells, levels of Cd6Δd3 expression fluctuated noticeably between different thymocyte populations. Cd6Δd3 was least abundant in double-positive (DP) thymocytes, being expressed in just 30% of the cells (Fig. 4D, right panel). By contrast, nearly 50% of single-positive (SP) CD4 or CD8 thymocytes expressed Cd6d3, and no differences were registered between these two subpopulations.

**Cell activation increases rat Cd6Δd3 expression**

We next analyzed the pattern of expression of rat Cd6 isoforms upon cell activation. Activated T cells were generated by incubating rat splenocytes with anti-TCR plus anti-CD28 mAb over 3 days. The percentage of CD4⁺ T cells with an activated phenotype, as determined by OX40 expression (42), increased from 20 to 91% (Fig. 5A). CD4⁺OX40⁺ cells were sorted as individual cells and compared with the nonactivated phenotype. RT-PCR analysis of individual cells revealed a clear modification in the pattern of expression of Cd6 isoforms following activation, with an increase in the number of cells coexpressing both isoforms (Fig. 5B). The relative levels of CD4 T cells expressing the Cd6Δd3 mRNA increased significantly from 42 to 86% upon stimulation (Fig. 5C).

Analysis of Cd6 expression at the protein level was also performed by immunoblotting. Two isoforms of Cd6 with molecular masses of ~130 and 122 kDa (indicated by dashed arrows) were clearly detected in resting splenocytes. These two isoforms should be products of alternative splicing, but still retain domain 3, given that the size is very close to that displayed by full-length rat Cd6 in COS7 cells (126 kDa), whereas the Cd6Δd3 mutant expressed in COS7 cells had a molecular mass of 112 kDa (see Fig. 2). Splenocytes were stimulated with TCR and CD28 mAb, and after 72 h cells were collected, lysed with detergent, and CD6
was immunoprecipitated and detected by immunoblotting. Interestingly, after 72 h of activation two smaller isoforms of 106 and 99 kDa were expressed (Fig. 5D, solid arrows). It is plausible that these new isoforms correspond to protein products missing domain 3.

**T cell activation controls expression of the human SRCR domain 3 of CD6**

We obtained from human PBMC, as well as from the Jurkat cell line E6.1 (data not shown), cDNAs coding for the full-length molecule and also several PCR products with sizes ranging from 2,000 down to 1,600 bp (Fig. 6A, middle lane). We then tested for the existence of the CD6Δd3 isoform in human cells, and following the same strategy as for the rat CD6 gene, performed RT-PCR amplifying just the extracellular domain-coding sequences. A prominent band of 1,200 bp and also a 300 nt shorter product were clearly detected (Fig. 6A, right lane, compare with Fig. 4A).

Next, we gel-purified the cDNA fragments, subcloned them, and sequenced multiple clones. We could confirm that the message coding for the extracellular domain isoform CD6Δd3 (GenBank accession no. DQ786329), lacking exon 5, was present in both PBMC and E6.1 Jurkat cells. An additional mRNA species (GenBank accession no. DQ786330) was detected in the cells analyzed, lacking not only exon 5 but also exon 6, which codes for a linker between the membrane proximal SRCR domain and the transmembrane stretch (Fig. 6B).

The previous results suggested that, as in the rat, multiple CD6 isoforms coexist in human T cells. We compared the expression of...
the putative CD6 extracellular isoforms between resting and PHA-activated T cells. For detection of CD6, we used mAbs specific for CD6 domain 3, OX126, or CD6 domain 1, UMCD6 (22). In purified resting T lymphocytes, both SRCR domains 1 and 3 of CD6 are expressed at similar levels, a result consistent with resting T cells expressing mostly CD6FL (Fig. 6C, left panel). In contrast, 3 days following stimulation with PHA, the expression of domain 3 was significantly down-modulated, compared with the relatively unchanged labeling of domain 1 (Fig. 6C, right panel).

We have also analyzed the changes on CD6 isoform expression upon cell activation by immunoblotting. Nonactivated human T cells expressed a major CD6 species of 130 kDa. However, 3 days postactivation with PHA, an additional CD6 isoform with a molecular mass of 97 kDa was clearly detected by immunoblotting, whereas the expression of the larger CD6 species was decreased to a level equivalent to that of the smaller isoform (Fig. 6D). Together, the cytometry and Western blotting analysis suggest that, following activation of human T lymphocytes, the expression of

FIGURE 6. Increased expression of CD6Δd3 in human cells, upon activation. A, Total RNA from human PBMC was used for RT-PCR and amplified from the ATG up to the transmembrane region (TM) or the STOP codon. The cDNA products were analyzed on gels, isolated, and sequenced. The middle lane shows the amplification of the full-length cDNA, clearly visible at 2,071 bp, and additional shorter products between 2,000 and 1,600 bp. As can be seen on the right lane, products of amplification of the extracellular region were obtained, including the full-length isoform of 1,200 bp and a smaller band of 900 bp. B, Schematic representation of the extracellular isoforms of human CD6 protein deduced from sequencing of the RT-PCR products obtained from human PBMC and E6.1 Jurkat cells. The full-length transcript was the most abundant in both PBMC and Jurkat E6.1 cells. The CD6Δd3 isoform excluding only exon 5 and also a similar isoform excluding additionally exon 6 were also recurrent, as evaluated by the frequency in the number of clones sequenced. C, Human primary T cells isolated from peripheral blood, using the RosetteSep human T cell enrichment mixture, were analyzed by flow cytometry for the expression of CD6 extracellular isoforms by using specific Abs recognizing either domain 1, UMCD6, or domain 3, OX126. The expression is compared with the fluorescence of the control secondary Ab FITC conjugated (left panel). After 3 days of stimulation with PHA, a significant decrease in the level of expression of the isoform lacking domain 3 could be detected, as can be seen by using the same Abs (right panel). PHA-activated cells were gated based on the expression of the activation marker CD71 (data not shown). D, The expression of CD6 in resting cells, and in cells 3 days postactivation with PHA, was analyzed by Western blotting with anti-CD6 MEM-98 mAb. A decrease in the level of the higher molecular mass isoform and the appearance of a lower molecular mass species can be seen after activation.
full-length CD6 is partially down-regulated with the concomitant appearance of the CD6Δd3 isoform.

Human CD6 targeting to the IS: analysis of binding to CD166

We proceeded to confirm that human CD6 isoforms behaved similarly to those analyzed in rat, regarding translocation to the IS upon Ag recognition. Human T lymphocytes were induced to express full-length human CD6 (CD6FL), the CD6Δd3 isofrom, or the CD6CY5 mutant, all fused to GFP. Following incubation of T cells with superantigen-loaded Raji cells, which expressed endogenous CD166 at high levels on the surface (data not shown), conjugate formation and CD6 translocation to the synapse was evaluated in all conditions. Both CD6FL and CD6CY5 localized very efficiently to the IS in ~80% of total conjugates, whereas CD6Δd3 was typically dispersed throughout the cell surface (Fig. 7, A and B).

To test whether nonengineered endogenous CD6 expressed in T lymphocytes was still capable of targeting to the synapse, and that this effect was due to the binding to endogenous Raji-expressed CD166, we evaluated, through immunofluorescence, the localization of CD6 in T:Raji interfaces using blocking, as well as non-blocking Abs. Characteristically, CD6 confined to the synapse in 75% of unblocked T cell:Raji conjugates, a figure that did not significantly change when T cells had been previously incubated with UMCD6, an Ab recognizing domain 1 of CD6, and not able to block CD6-CD166 interactions (Fig. 7, C and D). However, when T cells had been incubated with the blocking, anti-CD6 domain 3, mAb OX126, CD6 localization at the synapse dropped.
dramatically to residual levels (Fig. 7, C and D). Similarly, previous incubation of Raji with the CD166 mAb, described to obstruct the interaction with CD6 (37), resulted in the noticeable reduction of the translocation of CD6 to the synapse (Fig. 7, C and D), in agreement with a previous study using blocking CD166 Abs in T cell-dendritic cell conjugates (15). Taken together, the results demonstrate that the direct interaction between CD166 and the SRCR domain 3 of CD6 is the driving force for the translocation of CD6 to the IS.

Discussion

The great majority of mammalian genes yield multiple mRNA transcripts arising from diverse promoter selection, alternative splicing, and/or differential polyadenylation (43). Despite the frequency and potential impact of this phenomenon, particularly relevant in the immune system, the cases in which distinct functional properties have been assigned to different isoforms of the same molecule are scarce. In T lymphocytes, just a few examples of naturally occurring alternatively spliced-encoded isoforms of transmembrane proteins have been reported, and only in three cases with clear distinct functional consequences: CTLA4 and CD95 can become transmembrane molecules instead of being secreted (44, 45), as a consequence of the inclusion of membrane-spanning domains following cell stimulation; and CD45, upon T cell differentiation, is produced with a shorter N-terminal extracellular domain, which increases the dimerization capacity of the molecule (46). Alongside this general trend, multiple isoforms resulting from alternative splicing of cytoplasmic domain-coding exons have been described for CD6, but with no distinct functions assigned (18, 28–30). In this study, we describe a rare example of a functional consequence resulting from alternative splicing in molecules of the immune system: the localization of CD6 with respect to the IS depends on the regulated expression of the CD166 binding domain by means of alternative mRNA splicing.

Targeting of CD6 to the synapse can be driven by simple lateral diffusion and does not require cytoskeletal reorganization, because deletion of the cytoplasmic tail did not affect the ability of CD6 to localize to the synapse. Therefore, the positioning of CD6 within the synapse must be largely determined by molecular interactions established with the contacting cell. Both human and rat CD6 molecules targeted very efficiently to the IS when expressing the CD166 binding domain, provided that the APCs were equally expressing CD166. However, when using T cells expressing CD6Δd3, synaptic localization of that isoform could still be attained in a small percentage of conjugates. A putative role for domain 2 of CD6, now substituting the positioning of domain 3 in CD6Δd3, or even for the membrane-juxtaposed stalk region, could be considered, given the fact that previous studies had suggested a minor participation of these domains in the binding to CD166 (22, 23). Nevertheless, it is the presence of domain 3 that largely determines CD6 synaptic localization, and thus regulation of its expression is the key event establishing the positioning of CD6 upon conjugate formation and T cell activation.

The mode of action of CD6 appears to be significantly different from that of other accessory molecules that influence signaling at or near the IS. CD2 binding to its ligand CD58 (CD48 in rodents) contributes to enhanced signaling, because it associates with intracellular-positive mediators (47). CD28 and CTLA-4 have devised a scheme whereby sequential expression of each molecule, combined with binding to one of the alternative ligands, CD80 and CD86 (themselves also sequentially expressed during T cell-APC interactions), drives responses from costimulation to inhibition (48). Differences in the affinity and avidity of binding determine the selective recruitment of CD28 or CTLA-4 to the IS upon recognition of CD86 or CD80 (49).

Meanwhile CD6, which was reported to deliver costimulatory signals as strongly as CD28 (15), mimics CD28 distribution with respect to synaptic localization, in that it is recruited to the IS at the onset of activation, and, according to the present study, is excluded from the cellular interface at later stages. However, it may display this same pattern of localization through a completely different mechanism: expressing the SRCR domain 3, CD6 is directed to the synapse where it can control signaling; choosing not to express the SRCR domain 3, it is no longer restricted to the cellular interface and its regulatory function may be diluted.

The reason why CD6 does not simply switch off its expression is not immediately evident. However, the existence of an alternative ligand for CD6 raises new perspectives for its overall function (50). If the new ligand can interact with a different part of CD6, switching from full-length to Δd3 may allow CD6 to shift from the synapse established with the APC to either an interaction with a second adjacent cell or again to the initial interface, provided that the second ligand is present. Differential molecular interactions could then modulate the behavior of CD6.

The apparent steadiness in expression of the alternative CD6 isoforms in different mature subpopulations may suggest that CD6Δd3 can define small subsets of thymocytes and mature T cells; however, the switch to the expression of the shorter isoform observed upon stimulation of T cells is undoubtedly a consequence of a product of activation. In the resting state, the majority of the cells express mainly the full-length CD6 form; upon productive T cell stimulation, there is a partial substitution of the full-length isoform by a substantial number of CD6 molecules per cell that no longer express the CD166 binding domain and thus are not restricted to the IS. This type of regulation of splicing is not unique, because it has long been known to control the expression of CD45 isoforms (51, 52). The novel CD6 isoform reported in this study seems to have a more obvious function because it disables the interaction of the protein with its ligand. Productive TCR recognition of Ag may signal for the shorter CD6 isoform to be produced, and this response may reduce the activity of CD6 at the synapse, possibly inducing a different behavior.

The expression of the CD6FL isoform seems to be highly favored in the DP stage in the thymus. Previous observations suggest that CD6 can be involved in positive selection, where it correlates with the expression of CD69 in DP thymocytes (53). Moreover, an inverse correlation between thymocyte CD6 expression and the rate of apoptosis has been demonstrated. Our results may be indicative of a mechanism of regulation where the expression of CD6FL capable of interacting with the ligand CD166, expressed by thymic epithelial cells, is favored at the DP stage and bypassed later on during thymic selection.

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Disclosures

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References


