Dendritic cell maturation and antigen presentation in the absence of invariant chain

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ABSTRACT In immature dendritic cells (DCs), major histocompatibility complex class II molecules accumulate in peptide-loading compartments and, during DC maturation, are exported to the cell surface in response to inflammatory stimuli. Moreover, it has recently been proposed that DCs have specific mechanisms of antigen uptake and delivery into major histocompatibility complex class II-loading compartments. B cells bearing a genetically disrupted invariant chain gene (li 0–/) show alterations in the transport and function of class II molecules. We herein report that DCs derived from li 0–/– H2k but not li 0–/– H2b mice undergo normal maturation in response to tumor necrosis factor α and show a high degree of class II surface expression. Class II molecules are accumulated in cathepsin D- and H2-M-positive compartments in immature li 0–/– DC and, during DC maturation, are exported to the cell membrane as compact dimers. li 0–/– DCs present putative li-dependent hen egg lysozyme-derived epitopes to T cells. These data support the existence of li-independent molecular requirements for class II transport and peptide loading in DCs.

Presentation of exogenous antigens to CD4+ T lymphocytes requires antigen internalization and processing in endocytic compartments (1–4). Newly synthesized α and β chains of class II heterodimers in the endoplasmic reticulum (ER), with invariant chain (li) as a third partner. This complex is transported to specialized compartment(s) along the endocytic route where the loading with antigenic peptides derived from exogenous proteins occurs (5–12). This event is facilitated by the presence of the molecule H-2M, in mice (13–17) and by HLA-DM in humans (18–20), involved in the dissociation of li complexes are retained into the ER; (ii) few complexes reach the cell surface do not have the “compact” conformation indicative of tight peptide binding (6, 27). Furthermore, li is required for the efficient assembly of class II dimers in allelic variants only: li 0–/– B cells from mice with an H-2b background mostly accumulate free α and β chains in the ER, although li 0–/– class II dimers from H-2k and H-2d background are efficiently assembled (28).

The general health status and growth rate of li 0–/– mice and wild-type (wt) litter mates are similar (24). Although primary IgM responses against nominal antigens were impaired in li 0–/– mice, recall vaccinations induced the production of antigen-specific IgG at levels comparable with those of wt mice (ref. 24 and unpublished results). The production of IgG upon in vivo challenge with antigen requires efficient antigen presentation to class II-restricted CD4+ T cells. Data from several laboratories point out to an essential role of dendritic cells (DCs) in priming and maintenance of T cell responses (29–32). DCs originate from bone marrow CD34+ progenitors (33–35), which include DC colony-forming units that yield homogeneous DC colonies under particular in vitro conditions (36). DC precursors enter the blood and reach peripheral organs where they develop to immature DCs. Immature DCs are capable to capture soluble antigens via macrophagocytosis (37, 38) and particulate antigens through phagocytosis (39, 40), to process them to form epitopes that will be subsequently expressed at the cell surface in association with major histocompatibility complex (MHC) class II molecules (41–44). To fully perform their antigen-presentation functions, DCs residing in nonlymphoid tissues need to be activated by stimuli that promote their maturation, including the expression of costimulatory molecules and the rearrangement of the actin-based cytoskeleton that allow their migration to T cell areas of lymphoid organs (45–50), where the priming of naive T lymphocytes occurs (31, 50). Once DCs have interacted with T cells, they complete the differentiation process (50, 51), which is believed to terminate by DC apoptosis (49).

We have recently shown that class II molecules in immature murine DCs are mainly located in internal vesicles and colocalize with the H-2M molecule (49), which have been convincingly implicated in facilitating the loading of antigenic peptides onto class II molecules (14–17) and whose expression has been traced throughout the endocytic pathway, along both conventional and unconventional class II containing compartments (17). Maturation of DCs is characterized by profound changes in antigen-presenting capacity (49, 50, 52), possibly due to modifications of li and class II distribution and trans-
port (53–55). This phenomenon possibly reflect the requirement for migratory DCs to retain antigens from the peripheral tissues to the lymphoid organs, where they actually perform the antigen-presentation function (31, 50). In this study we have investigated whether the requirements of Ii to regulate the antigenic peptide loading and presentation by MHC class II molecules apply to DCs from H-2k mice, either freshly purified from the spleen or during their in vitro maturation.

MATERIALS AND METHODS

Mice. The C57BL/6 and B10.BR inbred mice bearing a genetically disrupted Ii gene were generously provided by D. Mathis and C. Benoist (Institut de Génétique et de Biologie Moléculaire et Cellulaire, Strasbourg, France) (24). The absence of Ii was tested at the gene level by Southern blot analysis (24) and at the protein level by Western blot analysis using anti-cytoplasmic Ii antibodies (53).

Antibodies. The anti-MHC class II (IAk) 10.2.16, anti-mouse FcγRII/III receptor CD32 2.4G2, anti-DEC205 NLCDC-145, anti-Mac-1, and anti-CD11c N-418 hybridomas were all from the American Type Culture Collection. The 10.2.16 mAb is capable of precipitating class II dimers regardless of the association of the α/β complexes with stabilizing peptides, with intact Ii, or with Ii cleavage products (3). The anti-cytoplasmic domain Ii (56), anti-cathepsin D, and anti-H2-M β chain cytoplasmic domain antibodies were provided by N. Barois (Centre d’Immunologie de Marseille-Luminy, Marseille, France). The anti-CD45R B220 mAb was from Pharmingen. The second-step reagents were from Jackson Immunoresearch.

Cells. Spleen cells were derived from physical dissociation of the organs (57). After collagenase digestion, an adhesion step was performed (37°C, 2 hours). B cells were retrieved from the nonadherent cells after T lymphocyte depletion (anti-Thy, J11; anti-CD4, RL172Y; anti-CD8, 3I1M) and were 85–95% positive for the CD45R marker. Adherent cells were incubated overnight at 37°C. Spleen DCs were obtained from the adherent cell population, after centrifugation over a 50% discontinuous Percoll gradient (Pharmacia) and were 75–90% positive for the DC marker N418. CD34+ precursors were grown in Iscove’s medium (Sigma) supplemented with 30% NIH 3T3 supernatant containing recombinant murine granulocyte–macrophage colony-stimulating factor (20 ng/ml; Genzyme) (49). At day 3, the granulocyte–enriched nonadherent cell populations were discarded. To induce DC terminal maturation, 15-day cultured cells were treated for 48 hours with recombinant tumor necrosis factor α (TNF-α) (100 units/ml; Genzyme) before characterization (34, 58). The IAα-restricted T cell hybridomas (3A9, 3B11) (59–61) were provided by L. Adorini (Roche, Milan, Italy). 3A9 cells are specific for the epitope of residues 46–61 of hen egg lysozyme (46–61 HEL) (59), which associate in B cells with newly synthesized IAk molecules only when they are targeted to the loading compartment by Ii. 3B11 T cells recognize the epitope of residues 34–45 of HEL (34–45 HEL) that binds to recycling class II molecules (61). The CTL2-2 cells were from the American Type Culture Collection.

Flow Cytometry. The samples were stained as described (62). Briefly, the cells were incubated with primary antibodies (1 μg/ml) and secondary reagents (fluorochrome-coupled antibodies or streptavidin) in the presence of 2.4G2 mAb and normal mouse serum to prevent FcγRII/III binding. The samples were analyzed with a FACScan apparatus (Becton Dickinson).

Confocal Microscopy. Intracellular immunofluorescence was performed on fixed and permeabilized cells as described (49). The confocal laser scanning microscopy was carried out using a Leica TCS 4D instrument (Leica): focal series of four horizontal planes of section were simultaneously monitored for fluorescein isothiocyanate and Texas Red. Unless otherwise specified, a plane of focus 0.8 mm above DC contact to the coverslip was selected. The micrographs were visualized and printed as reported (49). To identify fluorochrome colocalization unambiguously, correlation maps were made by using a statistical method based on the calculation of the local joint moment of standardized images (63). These correlation maps discriminate coincident fluorescence distributions from the superimposition of noncorrelated fluorescence profiles on a local basis. Each pair of red–green images is standardized for contrast and local variation in fluorescence. This correction and the joint moment of red–green images were calculated by considering a 19 × 19 pixel Gaussian window with a half-height width of 5 pixels corresponding to a resolution of 0.5 mm (63).

RESULTS AND DISCUSSION

In the H-2k haplotype, the absence of Ii does not compromise class II IAα/β assembly (28); however, the transport of class II dimers to the peptide-loading compartment is greatly affected in Ii−/− B cells (28). We therefore used H-2k II−/− mice to analyze class II distribution and peptide loading in immature and mature DCs. The DCs and B cells were purified from whole spleen of wt and II−/− mice, and their class II expression was monitored. Freshly purified N418+ II−/− splenic DCs expressed high amounts of membrane class II molecules; on the contrary, as expected, the B220+ II−/− B cells had an impaired membrane expression of IAk molecules (Fig. 1a).

The analysis of spleen sections in H-2k II−/− mice confirmed the presence of highly class II positive DCs in T cell areas (data not shown). Moreover, II−/− DCs were able to stimulate IAα-restricted 3A9 T hybridoma cells upon incuba-
tion with native HEL protein, whereas B cells from the same mice were not (Fig. 1b).

Splenic DCs mainly consist of mature DCs (29). To investigate the role of Ii during DC maturation, we derived immature DC from wt and Ii−/− antigen-presenting cells were purified from the spleen and immediately analyzed. (a) Flow cytometric analysis of splenic CD45R+ (B220 mAb) B cells and CD11c+ (N418 mAb) DCs labeled with the 10.2.16 mAb to reveal IAk surface expression (solid histograms). Control stainings (open histograms) were performed with the second-step reagent only. (b) Antigen presentation by B cells and DCs of HEL using the 46–61 epitope-specific 3A9 T cell hybridoma. The assay was performed at the indicated ratios of T cells and antigen-presenting cells, preincubated (○) or not (●) with HEL at 1 mg/ml.

FIG. 1. Splenic DCs overcome the need for Ii for class II surface expression and antigen presentation. wt and Ii−/− antigen-presenting cells were purified from the spleen and immediately analyzed. (a) Flow cytometric analysis of splenic CD45R+ (B220 mAb) B cells and CD11c+ (N418 mAb) DCs labeled with the 10.2.16 mAb to reveal IAk surface expression (solid histograms). Control stainings (open histograms) were performed with the second-step reagent only. (b) Antigen presentation by B cells and DCs of HEL using the 46–61 epitope-specific 3A9 T cell hybridoma. The assay was performed at the indicated ratios of T cells and antigen-presenting cells, preincubated (○) or not (●) with HEL at 1 mg/ml.

FIG. 2. DC maturation induces high surface levels of class II molecules that are partially compact in the absence of Ii. (a) Immature DCs derived from wt and Ii−/− BMDCs were cultured for 14 days in vitro and characterized by their expression of Mac1, CD11c, and DEC-205 markers (anti-Mac1, N418, and NLDC-145 mAbs) (solid histograms). (b) Immature BMDCs (14 days) were analyzed for class II expression immediately and after 24 and 48 hours of incubation with TNF-α (100 units/ml), which induces class II surface expression (solid histograms). Control stainings (open histograms) were performed with the second-step reagent only. (c) After surface iodination, class II dimers were immunoprecipitated from immature and 48-hour TNF-α-treated BMDCs with the 10.2.16 mAb. Iodinated compact forms (CF) migrating as 55-kDa heterodimers were revealed in nonboiled samples, whereas α and β chains were only detected after boiling.
This feature apparently depends on the H2 haplotype of the strain investigated, because class II expression was reduced in Ii−/− BMDCs derived from H2b mice (data not shown). A small percentage of H2b BMDCs exhibited high levels of class II molecules (Fig. 2b) and may represent fully mature DCs. When BMDCs from H2b mice were cultured for 48 hours in the presence of TNF-α at 100 units/ml, [i.e., a cytokine that induces DCs exhaustive maturation (33, 37, 47, 49)], the level of class II membrane expression was enhanced, regardless of the expression of Ii (Fig. 2b).

Access of class II dimers to the peptide-loading compartment leads to peptide association and a stabilization of the class II complex that is visualized as a 50-kDa dimer in SDS/polyacrylamide gels under reducing conditions (64). Immature BMDCs from wt and Ii−/− mice were iodinated and membrane class II molecules were immunoprecipitated with the anti-IAk specific 10.2.16 mAb, which recognizes both and membrane class II molecules were immunoprecipitated with the anti-IAk specific 10.2.16 mAb, which recognizes both the “compact” class II dimers associated with antigenic peptides and empty or unstable dimers (3). SDS/PAGE of the immunoprecipitates demonstrated that class II dimers from immature BMDCs can reach the plasma membrane in a compact conformation, even in the absence of Ii (Fig. 2c). The maturation of BMDCs in response to TNF-α amplifies the proportion of compact dimers in wt and Ii−/− DCs. Of interest, in the absence of Ii, H2b MHC class II complexes that reach the plasma membrane in a SDS-resistant conformation display a wide range of molecular weights between 50 and 120 kDa. This result is reminiscent of the binding between class II α/β dimers and long polypeptides that takes place in H2-M or cathepsin D and contains vesicles and, therefore, identified the vesicles in which class II and H2-M or cathepsin D were colocalized. Colocalized vesicles were detectable to similar extents in both wt and Ii−/− DCs, thus confirming that Ii is not required for class II accumulation in the putative peptide-loading compartments of immature DCs. Class II molecules were exported to the cell surface in both wt and Ii−/− DCs after treatment with TNF-α (Fig. 3b), suggesting that class II export during DC terminal maturation does not require Ii expression.

Ii−/− DCs from H2b mice are capable of efficiently presenting epitopes derived from the processing of exogenous antigens to class II-restricted T cells. In particular, Ii−/− immature DCs (but not Ii−/− B cells) efficiently present epitopes from the internalized HEL protein to 3A9 T cells (3, 59, 60) (Fig. 4a). On the contrary, antigen presentation of the 34–45 HEL epitope to 3B11 T cells (3, 60, 61) was not affected in Ii−/− DCs and B cells (Fig. 4a). In immature DCs, exogenous antigens are retained for a long time before efficient MHC class II loading and presentation (66). A pulse of at least 1 hour is required to induce the degradation of the internalized antigen and no presentation of exogenous antigens is detectable before 8 hours of processing (ref. 66, Fig. 4b, and data not shown). In Ii−/− immature BMDCs, presentation is even more delayed: it is not detectable before 4 hours of pulse at high DC/T cell ratios (1:1) and before 6 hours of pulse at lower DC/T cell ratios (1:4–1:10) (Fig. 4b). This delay in the kinetics of antigen presentation could be associated with a difference in the efficiency of H2b MHC class II peptide loading. It is likely that the long polypeptides associated with the MHC class II in H2b Ii−/− DCs are differentially resistant to H2-M-catalyzed peptide exchange. Indeed, in the absence of Ii, HLA-DM expression influences the array of peptides presented by HLA-DR molecules (70). The fact that no difference was observed after long incubation times is most
Fig. 4. Immature li−/− BMDCs can present a larger array of HEL epitopes than B cells. (a) wt (■) or li−/− (□) B cells, immature BMDCs, and TNF-α-elicited mature BMDCs were incubated with serial dilutions of HEL protein or HEL 46–61 synthetic peptides and tested for their ability to stimulate the 46–61 epitope-specific 3A9 T cell hybridoma, which requires li in B cells, or the li-independent 34–45 epitope-specific 3B11 T cell hybridoma. (b) wt (■) and li−/− (□) immature BMDCs were pulse-labeled with HEL protein at 1 mg/ml for 30 min and 1, 2, 4, and 6 hours (x axis) before extensive washing and then tested for their ability to present the 46–61 epitope to increasing number of 3A9 T cells for 24 hours (ratios tested, 1:1, 1:4, and 1:10). The y axis is the emission at 490 nm. (c) wt (solid bars) and li−/− DC (open bars) were preincubated with 20 μM BFA for 15 min before and during 6 hours of pulse labeling with various amounts of the HEL protein (10 μg/ml and 100 μg/ml). After extensive washing, cells were tested for their ability to present the 46–61 HEL epitope to the 3A9 T cells and the 34–45 HEL epitope to the 3B11 T cells. The results are expressed as percentage of inhibition of the BFA treatment calculated as follow: 100 − [(OD490 emission in the presence of BFA × 100)/OD490 emission in the absence of BFA].

likely due to the higher efficiency of antigen presentation by DCs compared to B cells (Fig. 4a).

Fig. 4 shows that li dependence of different epitopes for H2k MHC class II loading and presentation can vary in different cell types (61). It has been previously reported that the 34–45 HEL epitope, recognized by the 3B11 T cell hybridoma, and the 116–129 HEL epitope can be efficiently presented in the absence of li by Rat-2 fibroblasts transfected with MHC class II molecules (61), but these epitopes are not presented by li−/− B cells (21). The efficient presentation of the 34–45 HEL epitope by li−/− DCs is not affected by BFA, i.e., by an agent that compromises newly synthesized class II complexes egress from the ER, supporting the hypothesis that it can be due to MHC class II complexes recycling from the plasma membrane (ref. 61 and Fig. 4c). In contrast, the 46–61 HEL epitope presentation to the 3A9 T cells is severely inhibited by BFA treatment after pulse labeling li−/− immature BMDC with HEL protein (Fig. 4c). This was not the case when the cells are pulse-labeled with the 46–61 HEL synthetic peptide (data not shown). This result implicates that newly synthesized MHC class II are involved in the presentation of 46–61 HEL epitope to T cells (71) and is in agreement with the existence of molecular constraints other than li expression in the presentation of li-dependent HEL epitopes by immature DCs.

Fully mature DC inefficiently presented the HEL protein (Fig. 4a). This is associated with the reduced ability of wt and li−/− mature DCs to internalize soluble antigens (data not shown) and with the redistribution of most of the MHC class II molecules at the cell surface (Fig. 3b). The HEL 46–61 peptide, which does not require internalization, was presented with high efficiency by both DC populations (Fig. 4a), and this event was not affected by BFA treatment (data not shown).

DCs are the most potent antigen-presenting cells for CD4+ T cells (31): to induce similar rates of T cell activation, B cells required a 100-fold higher concentration of HEL antigen (Fig. 4a). The molecular basis of this phenomenon can reside in (i) the ability of immature DCs to efficiently present soluble antigens internalized by several mechanisms to CD4+ T cells, including macropinocytosis and receptor-mediated endocytosis (37, 38), whereas B cells efficiently present only antigens internalized through their antigen receptor (72); and (ii) the higher levels of MHC class II and costimulatory molecules in DCs (ref. 49 and Figs. 1 and 3).

This report shows that in H2k li−/− DCs, class II dimers apparently overcome the requirement of li for loading with antigenic peptides but that B cells require an intact li polypeptide for efficient presentation. This implicates that at least in the H2k haplotype DCs, newly synthesized MHC class II complexes are capable of presenting antigens internalized by B cells, which requires li, and also that H2-M-positive intracellular compartment(s) in the absence of li targeting. The strict haplotype dependence of the phenomenon implicates that a more complex regulation must exist in different murine strains. This pathway is constitutively operating in li−/− H2k DCs and may underlie the conserved IgG response to nominal antigens reported in B10.BR li−/− mice.

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