Human FDC express PrPc in vivo and in vitro

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Prion diseases are fatal neurodegenerative disorders caused by accumulation of abnormal prion protein (protease-resistant prion, PrPres). PrPres accumulation is also detected in lymphoid organs after peripheral infection. Several studies suggest that follicular dendritic cells (FDC) could be the site of PrPres retention and amplification.

Here we show that human follicular dendritic cells can express normal cellular prion protein (PrPc) both in situ and in vitro. When tonsillar cryosections were treated with anti-PrP antibody, the label was found on some very delicate cell extensions inside the lymphoid follicles, especially in the germinal centres. These extensions react with DRC1 antibody, used frequently to label FDC. Other structures labelled with anti-PrP antibody were the keratinocytes.

To confirm the ability of FDC to synthesise PrPc, we isolated FDC by a non-enzymatic procedure and cultured them. By cytochemistry and flow cytometry it was clearly shown that FDC do produce PrPc.

Keywords: Follicular Dendritic Cells (FDC), germinal centre, human tonsil, immunolabelling, prion protein (PrPc)

INTRODUCTION

Prion diseases are characterised by accumulation of infectious particles of protease-resistant prion (PrPres) (Prusiner et al., 1982). This pathological isoform results from modification of normal cellular protein called PrPc. This protein is widely distributed in the brain and other organs notably in the lymphoreticular system.

The symptoms of prion diseases always occur after accumulation of PrPres in the nervous system, leading to observable pathological lesions. After peripheral infection, however, PrPres is detected during the pre-clinical stages in the spleen and other lymphoid organs. Abundant PrPres was recently detected, before the appearance of the clinical signs of the disease, in the germinal centres of tonsils (Hill et al., 1997) and appendixes (Hilton et al., 1998) from patients affected with vCJD.

The prion diseases are now widely studied in experimental models. Inoculation of PrPres into wild-type mice induces the different symptoms, whereas injection into PrP-deficient mice leads neither death nor to infectivity. This demonstrates the major role played by PrPc (Bueler et al., 1993).

Results obtained with immunodeficient mouse models show that lymphoid organs are involved in various steps of disease development after peripheral
inoculation (Cesbron et al., 1998; Aucouturier et al., 1999). It is not clear, however, which immune system cells play a key role in pathogenicity. Spleen fractionation studies have shown that infectivity is associated with the stromal fraction and with lymphocytes (Clarke and Kimberlin, 1984; Raeber et al., 1999). The incubation period in peripherally infected mice is unaffected by whole body irradiation (Fraser and Farquhar, 1987). Thus, the prion replication depends on PrPc and notably on a radiation-resistant fraction of the cell population of lymphoid tissues. Follicular dendritic cells (FDC) belong to this radio-resistant fraction and are stromal cells (Kinet-Denoël et al., 1982). SCID mice lacking functional B and T cells and also functional FDC are resistant to peripheral infection by scrapie agent but develop the disease after a bone marrow reconstitution (Fraser et al., 1996). Reconstitution restores the presence of mature T and B cells and induces differentiation of functional follicular dendritic cells (Kapasi et al., 1993).

Immunohistochemical staining with anti-PrP antibodies reveals an extraordinary accumulation of PrPres in the lymphoid follicles, especially in the form of a delicate network (Kitamoto et al., 1991; McBride et al., 1992; Klein et al., 1998; Jeffrey et al., 1999). These results strongly suggest that the follicular dendritic cells might support scrapie agent accumulation and replication in the peripheral lymphoid organs.

Herein the aim was to determine whether human FDC can express PrPc. At this end, we have immunolabelled cryosections of human tonsils. Furthermore, since FDC are located solely in the germinal centres in intimate contact with B cells, we isolated them and analysed their capacity to react with anti-PrP antibodies just after purification and after several days of culture. We report here that human FDC do indeed synthesize PrPc.

RESULTS

FDC In Situ

On sections stained with MoAb:DRC1, a well-known marker of follicular dendritic cells, the network of cell extensions was intensely labelled (Fig. 1). After anti-PrP immunolabelling of cryosections with 3F4 antibody, the keratinocytes lining the crypts mainly at the level of basal cells were stained densely as it was previously reported by Pammer (Pammer et al., 1998), however, the germinal centres displayed a very low labelling. In the germinal centres, fine peroxidase reagent deposits were found along the intercellular spaces of the constituent cells (Fig. 2 A-B-C).

Control cryosections, reacted with normal mouse serum instead of anti-PrP antibody then with StrepAB Complex HRP and AEC didn't display any deposit in the germinal centres or along the crypts.

Freshly Isolated FDC

FDC prepared by enzymatic digestion or mechanical dissociation showed engulfment of germinal centre
lymphoid cells by their cytoplasmic extensions. FDC always surrounded 3 to 20 lymphoid cells. These FDC-clusters could easily be distinguished from lymphocyte aggregates, vascular or epithelial fragments and tingible body macrophages. A strong positive reaction with DRC1 antibody was found on the cytoplasmic projections of these FDC, which were frequently bi- or multinucleated (Fig. 3).

PrPc expression was detected only on FDC-clusters isolated without enzymatic treatment. Weak staining with MoAbs:3F4, 3B5 and 12F10 was observed on the cytoplasmic projections of the FDC (Fig. 4 A-B). The percentage and intensity of positive FDC-clusters depended to some degree on the monoclonal antibody used; 12F10 and 3F4 were recommended.
Cultured FDC

During several hours of primary culture, the FDC extended thin cytoplasmic processes and adhered to the plastic surface. The engulfed lymphocytes eventually died by apoptosis. These FDC showed no proliferative activity and possessed one to ten nuclei.

After 4 days in culture, most of FDC reacted with 12F10 and 8G8 monoclonal antibodies in immunolabelling experiments (Fig. 5 A-B). Expression of PrPc was detected over the entire cell surface. Staining appeared more intense than when freshly prepared FDC-clusters were used.

PrPc expression was also analysed by cytometry on FDC cultured for 6 days. Since FDC in vitro adhere to the substrate, it was necessary to use trypsin to
detach them. The intensity of PrPC expression on freshly trypsinised FDC was low for both 3B5 and 8G8 monoclonal antibodies. To confirm that the FDC synthesise PrPC, we cultivated the trypsinised FDC in cell suspension conditions for an additional 15 h. This procedure allowed a re-expression of PrPC by FDC (Fig. 6 A-B). PrPC was detected with both antibodies and labelling was intense as with Jurkat cells (Fig. 7).

**DISCUSSION**

*In vivo* findings of this and other laboratories (Kitamoto et al., 1991; McBride et al., 1992; Klein et al., 1998) indicate the presence of PrPC in low quantity along the cytoplasmic extensions of follicular dendritic cells. Since FDC make intimate contact with lymphoid cells and since lymphoid cells express PrPC (Cashman et al., 1999; Mabbott et al., 1997; Dodelet and Cashman, 1998), these findings do not constitute conclusive evidence that FDC synthesise PrPC.

In the present study, we thus chose to work on FDC isolated from human tonsils. We isolated the FDC in two different ways: by enzyme treatment and mechanical disruption. The former method has the disadvantage of removing PrPC from the surface of the FDC, as shown here and with Jurkat cells (data not shown). The latter method yielded FDC-clusters in which the FDC did appear to express PrPC on their surface. The problem with these results is again the presence of lymphocytes entrapped in the dendritic processes of the FDC, leaving open the possibility that the PrPC might actually be synthesized by the lymphocytes, then transferred to the FDC.

The question was thus: can FDC produce PrPC by themselves? To answer this question, we cultured these isolated FDC-clusters so as to eliminate all engulfed lymphocytes. In the culture system developed previously in this laboratory, FDC adhere to the culture substrate. The FDC stretch out and take a fibroblastic-like morphology. The lymphocytes eventually disappear by apoptosis so that only flat, firmly attached FDC remain in the culture system. On these attached FDC, we again detected PrPC by immunohistochemistry. Secondly, to analyse these cultured FDC by cytometry, we detached them with the aid of trypsin. Since this treatment removes the surface PrPC, as seen by cytometry and as reported by Caughey...
We further cultured the detached FDC in cell suspension to give them time to synthesise PrPc and express it at their surface. The cultured FDC were PrPc-positive in absence of any lymphoid cells or stimulant. Thus, FDC appear to express PrPc independently of any stimulation or interaction with other cells. It is noteworthy that cultured lymphoid cells can also synthesise PrPc. Antoine et al. (submitted) working on cultured and freshly isolated tonsillar cells observed by cytometry higher surface expression of PrPc on the former than on the latter. Thus, in vitro expression of PrPc is independent of cell activation and may be related to stress or to loss of the in vivo microenvironment. In this work, furthermore, we observed different reactions according to the antibody used. These differences might reflect epitope accessibility related to protein folding or its glycosylation.

By long-term culturing of FDC, we have thus shown that these cells can express PrPc. In the future, this model will enable us to study PrPres replication and infectivity in FDC in vitro.

**MATERIALS AND METHODS**

**Isolation and Purification of FDC-clusters**

Palatine tonsils were surgically removed from children aged 3 to 10. FDC-clusters were isolated by two different procedures, enzymatically for culturing and non-enzymatically for immunostaining on cytopspins.

Enzymatic digestion method was described elsewhere (Tsunoda et al., 1990); the tonsils were cut in slices (1 mm thick) and digested for 20 min at 37°C with an enzyme cocktail containing 0.1% collagenase (Type A, Boehringer-Mannheim), 0.05 % dispase II (Grade II, Boehringer-Mannheim), 0.06 % Dnase I (Grade II, Boehringer-Mannheim) in PBS with 0.4 % Bovine Serum Albumin (BSA). The freed-cells fraction was stored in cold PBS with 0.4 % BSA. The remaining tissues were digested again with fresh enzyme solution.

For non-enzymatic dissociation, we chose gross dissection and mechanical disruption of lymphoid follicles. After the enucleating the follicles under ana-
tomical microscope, we squeezed them between two glass slides to obtain a cell suspension composed mainly of germinal-centre cells. The cell preparation was filtered on gauze to eliminate collagen bundles and blood vessels. The whole procedure was carried out in PBS + 0.4 % BSA at 4°C.

After both cell dissociation procedures, the preparations were enriched in FDC-clusters by repetitive 1G sedimentation through Fetal Calf Serum (FCS) (Wekerle et al., 1980). FDC-clusters were finally enriched at approximately 96% (v/v ratio).

**FDC Culture**

The final FDC-cluster rich fraction was incubated in a plastic culture dish for 60 min with RPMI 1640 with 10 % FCS at 37°C, 5 % CO₂ to remove contaminating macrophages. Next, non-adhering cells were transferred to new wells for a 6-h incubation in fresh medium: RPMI 1640, 2 mM L-glutamine, 50 μM 2-mercaptoethanol, 100 U/ml penicillin, 100 μg/ml streptomycin, 5 % FCS. During this incubation, FDC adhered weakly to the substrate. Non-adhering population was carefully discarded. The culture medium was replaced once daily.

For immunohistochemistry, some FDC-clusters were cultured on polylysine-coated glass coverslips in 6-well culture plates.

**Immunohistochemistry**

Cryosections (8–10 μm) of tonsils and cytopsins of FDC-clusters (non-enzymatically isolated) were fixed in acetone for 10 min at 4°C. On the other hand, FDC cultured on polylysine-coated glass coverslips for 4 days were fixed in paraformaldehyde 2 % for 10 min at 4°C.

Cryosections and cytopsimens were treated with a non-specific rabbit serum for 15 min, then allowed to react for 1 h at room temperature with MoAbs: DRC1 (Dako, Denmark), 12F10, 3B5, 8G8 or 3F4 antibody at optimal dilution (1/100, 1/100, 1/100, 1/100 and 1/200 respectively). The preparations were incubated with biotinylated rabbit anti-mouse immunoglobulin (αlg, 1/200) for 1 h, followed by StrepAB Complex HRP (Zymed). Peroxidase activity was revealed with 9-ethyl-3-aminocarbazol (AEC) and H₂O₂ as substrates (Zymed). For the cryosections, we used an amplification system (Envison kit, Dako, Denmark) in place of the secondary antibody. All antibody dilutions and wash steps were performed in phosphate buffered saline, pH 7.2 (supplemented with 0.5 % Tween for MoAb 3F4 dilution only).

Negative controls consisted of preparations in which the specific antibody was replaced with non-specific mouse serum and once from which the primary antibody was omitted.

**Flow-cytometric Analysis**

After 6 days of culture, the FDC were checked for the ability to effect B lymphocytes emperiploesis. This is a reliable test for identifying FDC in vitro (Tsunoda et al., 1992). Freshly prepared tonsillar B lymphocytes were added to the monolayered FDC in vitro. Cultured FDC can entrap lymphocytes beneath their cytoplasm extensions (pseudoemperiploesis). Other FDC that had been cultured for 6 days were harvested by treatment with 0.25 % trypsin and 0.04 % ethylenediamine tetra-acetic acid in PBS and washed. Some detached FDC were stored in FCS on ice until immunostaining. Others were cultured for 15 h at 37°C as cell suspensions in RPMI containing 10 % FCS and HEPES. This was done with the Techne® system (Duxford Cambridge, UK) avoiding attachment to the substrate. Thereafter, the cells were collected and washed. Both types of FDC preparation were incubated with biotinylated monoclonal antibody (8G8 or 3B5), followed by Streptavidin-Per-Cys (Dako, Denmark). Jurkat cells (human T-cell lymphoma-cell line) constituted the positive control for PrPc. In the negative control, we used non-specific murine IgG-Per-Cys (Dako, Denmark).

The fluorescence intensity was analyzed by means of the Cell-Quest system with a FACSscan flow cytometer (Becton-Dickinson, USA).
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