Decay-accelerating factor (DAF, CD55) in normal colorectal mucosa, adenomas and carcinomas

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Summary
Decay-accelerating-factor (DAF, CD55), a phosphatidyl-inositol anchored glycoprotein, is a member of the cell membrane bound complement regulatory proteins that inhibit autologous complement cascade activation. DAF was found expressed on cells that are in close contact with serum complement proteins, but also on cells outside the vascular space and tumour cells. Using CD55(BRIC110) and CD55(143-30) we show here that DAF(CD55) is only sporadically expressed on the luminal surface of normal colonic epithelium. However, 5/20 adenomas expressed DAF(CD55) on the cell surface of all tumour cells, 5/20 adenomas were completely negative, 10/20 adenomas expressed DAF(CD55) in various amounts. DAF(CD55) was expressed in various intensities on almost all tumour cells of the colon carcinoma cell line HT29. In 5/8 colorectal carcinomas DAF(CD55) was localised on the apical cell surface of all tumour cells, 31/88 were completely negative, 52/88 expressed DAF(CD55) in parts of their neoplastic populations. There was no correlation between the tumour grading, staging and location and the mode of DAF(CD55) expression, but DAF(CD55) was found more often in mucinous carcinomas (P = 0.007).

Although the mode of DAF(CD55) expression is not correlated with tumour prognostic parameters, the upregulation of DAF(CD55) in a subset of adenomas and carcinomas needs further investigation concerning protection of tumour cells against complement cytotoxicity.

Material and methods

Tissues and cells
Tissue samples from patients who underwent tumour resection of the colon or rectum reached our laboratory within 1 h after removal. Tissue samples were collected from the cancerous lesion, from the unaffected mucosa and from adenomas found in the removed tissue. They were quick frozen in liquid nitrogen and stored at –70°C until sectioning. Serial sections of 4 to 6 µm thickness were cut, extensively air dried, fixed in acetone for 10 min at room temperature, then stained immediately or stored at –20°C for a short time. The collection comprised 20 tissue samples of unaffected mucosa, 20 adenomas, and 88 carcinomas. The tumours, whose primary site and metastatic spread at the time of operation were well documented, have been typed, graded, and staged according to the International Union Against Cancer (UICC) classification (Dukes & Bussey, 1958; Hermanek & Sobin, 1987; Jass & Sobin, 1987).

Ten carcinomas (11.4%) were grade I, 66 (75.6%) were grade II, and 12 (13.6%) were grade III; 23 (26.1%) were mucinous and 65 (73.9%) were nonmucinous adenocarcinomas. According to the Dukes’ staging there were 34 stage A patients (38.6%), 27 stage B patients (30.7%), 26 stage C patients (29.5%), and one patient with liver metastasis. Twenty-four (27.3%) carcinomas were located in the right hemilolon, 64 (72.7%) in the left hemilolon. The colon carcinoma cell line HT29 (ATCC, Rockville, MA) was raised in RPMI 1640 medium (Gibco, Paisley, Scotland, UK) containing 10% foetal calf serum, sodium pyruvate and L-glutamine. Cells were detached with 0.25% ethylen-diamine-tetraacetate (EDTA), centrifuged at 1000 r.p.m. for 5 min, and washed in RPMI 1640. Cytoskins were made, air-dried, fixed in acetone for 10 min, and stained immediately or stored at –20°C.

Immunohistochemistry

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UK, mAb 143-30 by R. Vilella, Barcelona, Spain, on the occasion of the Non-Lineage/Natural Killer Section Meeting of the IVth International Workshop and Conference on Leucocyte Differentiation Antigens held in Vienna, February 1989. To underline that we are dealing in our study with those DAF epitopes recognised by the CD55 monoclonal antibodies BRIC110 and 143-30, we use the term 'DAF' (CD55) whenever presenting and discussing own data while using the term 'DAF' when we cite other authors using unclustered mAbs. Monoclonal antibody binding was detected with a polyclonal biotinylated sheep antibody to mouse immunoglobulins (Amersham, High Wycomb, UK) and a streptavidin-biotinylated peroxidase complex (Amersham). 3-Amino-9-ethylcarbazole (AEC) and N'-dimethylformamid (DMF) were obtained from Sigma Chemical Co. (St Louis, MO). The mAbs were diluted 1:1000 in phosphate-buffered saline (PBS), biotinylated sheep antibody to mouse immunoglobulin was diluted 1:50 in PBS, and the streptavidin peroxidase complex was diluted 1:100. Incubation times were 1 h at room temperature for the primary antibody and 30 min for the second and third step reagents. Using AEC as the chromogen (0.4 mg ml⁻¹ in 0.01% H₂O₂ for 30 min), the peroxidase reaction caused an intense red precipitate. The sections were rinsed in tap water, counterstained in Harris’ hematoxylin and mounted with glycerol gelatin. Isotype-matched controls with irrelevant mAb were carried out on a limited number of normal mucosae and colon carcinomas and revealed no isotype-associated side reaction in or on epithelial cells. Each frozen section series contained a negative control without the primary reagent; staining was observed solely in granulocytes whose endogenous peroxidase was not blocked for the benefit of optimal antigenicity, and to a much lesser extent in some epithelial areas due to endogeneous biotin.

**Evaluation**

A semiquantitative evaluation system was used to determine the antigen expression in normal, adenoma, and carcinoma tissue and cells. Antigen expression was scored ‘+’ whenever specific staining was detectable, and ‘-’ when no antigen was detectable. To evaluate the amount of stained cells, a semiquantitative scoring system was established: ‘+’ indicates that stained cells clearly outnumbered the unstained cells; ‘+’ indicates that positive and negative cells were found in equal proportions; ‘-’ means that unstained cells outnumbered the stained cells. According to this system, the antigen expression was correlated with tumour grade, type, stage and location of the tumour along the large bowel. The Fisher’s exact test was applied for statistical analysis.

**Flow cytometry**

For flow cytometry 1 × 10⁶ cells of HT29 were used per probe. Cells were suspended in FACS-medium containing RPMI, FCS, NaH₄ and HEPESS buffer. HEA 125, a monoclonal antibody which detects an epithelium-specific glycoprotein, Egp34, in normal and neoplastic transformed epithelial cells (Momburg et al., 1987) was used as positive control. CD53 (H177), a monoclonal antibody which detects the leucohistiocytic population and a probe without the primary antibody were used as negative controls. Cells were incubated with the CD55 antibody BRIC 110 diluted 1:200, and after three washing steps with the polyclonal FITC-coupled goat-anti-mouse antibody (Dianova Hamburg, Germany) diluted 1:50. The incubation time for each antibody was 1 h. After three washing steps propidium jodide (10 μg ml⁻¹ diluted in FACS-medium) was used for gating out the living cells. Flow cytometry was performed on a FACS-cant® (Becton Dickinson) with the LYSYS II software program.

**Results**

The expression of DAF(CD55) in normal colonic mucosa, in adenomas and carcinomas was immunohistochemically studied with mAb BRIC110. Previous studies found DAF expressed in the epithelium of the lower gastrointestinal tract (Medof et al., 1987), we found DAF(CD55) only sporadically localised in the epithelium of 3/20 normal colonic mucosae with CD55 BRIC110. In these three cases DAF(CD55) was expressed in small foci on the luminal surface of the epithelium in the upper parts of the crypts. A control study was performed with the mAb 143-30, another CD55 antibody which was also characterised on the IVth Workshop on Leucocyte Typing (Hadam, 1989). MAb 143-30 like BRIC110, revealed that DAF(CD55) was only sporadically expressed in the normal colonic epithelium (Figure 1a,b). In the normal gut wall mAb BRIC110 detected DAF(CD55) in fibrillar structures and fibroblasts, particularly in the submucosa, in nerve fibres, in reticular cells of lymph follicles and weakly in some endothelial cells; smooth muscle cells of the gutwall and vessel wall, ganglion cells, B cells of lymph follicles, T cells and plasma cells were negative. MAb 143-30 showed similar results; in addition, endothelial cells were somewhat stronger positive and B cells of lymph follicles were strongly positive. It is not excluded that, in analogy to other DAF mAbs (Kinoshita et al., 1985), BRIC110 and 143-30 although both reacting with the DAF molecule (Hadam, 1989) recognise different epitopes.

In colorectal adenomas mAb BRIC110 showed DAF(CD55) in the whole epithelium of 5/20 adenomas, 5/20 adenomas were completely negative. 10/20 adenomas focally showed...
DAF(CD55) expression, three of which had more positive cells than negative ones. Two further cases contained positive and negative tumour cells in about equal proportions and five others had more negative tumour cells than positive ones (Table I). DAF(CD55) was essentially localised on the luminal cell surface (Figure 2a); occasionally, apico-lateral and baso-lateral cell surfaces were positive, too (Figure 2b,c). MAb 143-30 showed a similar DAF(CD55) expression in adenomatous epithelium.

In colorectal carcinomas DAF(CD55) was expressed on the cell surface of all tumour cells in 5/88 carcinomas; 31/88 were completely negative (Table I). 52/88 showed DAF(CD55) expression only focally; 15 had more positive cells than negative ones, nine showed positive and negative tumour cells in about equal proportion, and 28 were predominantly negative. Representative cases stained with MAb 143-30 showed a staining pattern similar to that of BRIC110. However, there was also some reactivity within the cytoplasm. The stroma surrounding the tumour nodules strongly expressed DAF(CD55). The location of DAF(CD55) expression in colorectal carcinoma was the same as described for adenomas; most tumours expressed DAF(CD55) on the apical cell surface (Figure 3c). In the stroma there was strong expression of DAF(CD55) in peritumorous fibrillar structures and fibroblasts (Figure 3b). There was no statistical correlation between presence vs absence of DAF(CD55) and the tumour grading or staging. However, DAF(CD55) was more frequently expressed in mucinous carcinomas (P = 0.007, Fisher's exact test). Finally, there was no statistical correlation between the mode of DAF(CD55) expression and the tumour location (right versus left hemicolon).

The colon carcinoma cell line HT29 expressed DAF(CD55) in almost all tumour cells (Figure 4a). DAF(CD55) was in varying intensities detectable on the cell surface. The cell line had subpopulations of strongly positive tumour cells with positive cytoplasm and of faintly positive tumour cells. The fluorescence histogram revealed that almost all tumour cells expressed DAF(CD55). The peak of DAF(CD55) was broader than that of the positive control Egp34 (HEA125), a broadly expressed but epithelium specific glycoprotein (Momburg et al., 1987), because of the heterogeneous antigen density of DAF(CD55) on the tumour cell population (Figure 4b).

Discussion

Although DAF was formerly found only on cells with close contact to serum proteins, the extensive study of Medof et al. (1987) showed that DAF is broadly expressed in cells, tissues and fluids outside the vascular space. We investigated the expression of DAF(CD55) in 20 normal colonic tissues by an immunohistochemical technique using the murine mAb BRIC110 (Spring et al., 1987; Hadam, 1989). BRIC110 recognises a glycoprotein which was formally found on erythrocytes, leucocytes, platelets and several haematopoietic cell lines. This glycoprotein carries Cromer-related blood group antigens on normal erythrocytes and is absent or altered on erythrocytes of patients with the Inab phenotype (Spring et al., 1987). We observed strong DAF(CD55) reactivity in fibrillar structures, especially in the submucosa, while DAF(CD55) was only sporadically detected in the epithelium of 3/20 colon specimens. In these three cases DAF(CD55) was expressed in small foci of epithelial cells carrying the molecule on the luminal cell surface. In a recent study (Medof et al., 1987) reporting on DAF expression in the

| Score | Adenomas | Carcinomas |
|-------|-----------|------------|
|       | n = 20    | n = 88     |
| +     | 5          | 5          | 5.7        |
| + > - | 3          | 15.0       | 15         | 17.1       |
| +     | 2          | 10.0       | 9          | 10.2       |
| => +  | 5          | 25.0       | 28         | 31.8       |
| -     | 5          | 35.0       | 31         | 35.2       |

Notes: ‘A’ > ‘B’, cells with staining modality A clearly outnumbered those with modality B; ‘A’, ‘B’, + and − cells were found in equal proportions.

Figure 2 (a) Expression of DAF(CD55) in a well-differentiated adenoma. DAF(CD55) lines as a thin band only the apical cell surface of the epithelium. The scarce tumour stroma shows DAF(CD55) positive fibrillar-reticular structures (×62.5). (b) Occasionally DAF(CD55) was found on the apico-lateral and baso-lateral cell surfaces in addition to the thin-banded apical cell surface in colon adenomas (×125). (c) The adenoma of B is localised in the neighbourhood of a moderately differentiated carcinoma. In contrast to the overall expression of DAF(CD55) on the cell surface of the adenoma, DAF(CD55) was expressed in small foci on the cell surface of the malignant transformed epithelium. Staining was done with mAb BRIC110 (×125).
epithelium of the lower GI-tract, pooled anti-DAF mAbs were used. As various epitopes on the DAF molecule have been defined by different mAbs (Kinoshita et al., 1985), it is not excluded that the CD5BRIC110 fails to recognise an epitope of DAF which is expressed in normal colonic epithelium and is detected by pooled mAbs. However, a control staining with CD55(143-30), another mAb directed against the DAF molecule, showed a reactivity corresponding to that of BRIC110; the colonic epithelium was likewise negative. Despite the lack of DAF(CD55) in normal colonic epithelium, DAF(CD55) was focally expressed in most adenomas and carcinomas at the luminal cell surface. We show here that DAF(CD55) tends to be overexpressed in neoplastic colon epithelia, especially in mucinous carcinoma. In vitro DAF(CD55) expression appears to be a common feature of carcinoma cell lines. Apart from HT29, detection of DAF(CD55) was recently found on CaCo2 and SK-CO15 human intestinal cell lines (Lisanti et al., 1989). DAF was also found expressed in several other carcinomas, HeLa cells (Medof et al., 1987) and breast carcinomas (Cheung et al., 1988). The possible functions of DAF on human tumour cells have so far been investigated on a very few carcinomas, namely on renal carcinoma cells and melanomas (Cheung et al., 1988; Terachi et al., 1991). Membrane deposition of autologous antibodies against these tumour cells was found to sensitise them for complement-mediated cytotoxicity. In these experiments some tumour cells underwent complement-mediated lysis while others did not. The expression of DAF on the cell surface was found to be responsible for complement resistance, and anti-DAF mAb rendered complement-sensitive cell lines into sensitive ones (Cheung et al., 1988). Thus, DAF served as a tumour cell protecting protein on renal cell carcinomas and melanomas. Recent studies suggested the application of anti-DAF Fc-antibodies in conjunction with autologous tumour cell antibodies as one way of tumour effective killing (Cheung et al., 1988; Terachi et al., 1991), although the wide distribution of DAF in various normal cells and tissues is likely to be a severe restriction factor for a systemic application of anti-DAF. Nevertheless, the immunohistochemical determination of DAF expression in individual tumours can help to select carcinomas which are susceptible for agents that modify DAF expression or function (Cheung et al., 1988).

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