Expression of fibronectin, fibronectin isoforms and integrin receptors in melanocytic lesions

PG Natali¹, MR Nicotra², F Di Filippo¹ and A Bigotti¹

¹Regina Elena Cancer Institute and ²Institute of Biomedical Technologies CNR, 00161 Rome, Italy.

Summary In vitro studies have demonstrated that fibronectin (FN) can deliver a mitogenic signal to quiescent human melanoma cells and that the α, β-integrin receptor mediates this stimulus. In view of this finding we have analysed the in vivo expression of FN, and of ED-A and ED-B FN isoforms, in benign and malignant lesions of melanocytic origin. In the same specimens the expression of fibronectin integrin receptors was evaluated. The results demonstrate that, while detection of FN does not correlate with transformation and tumour progression, the expression of the two isoforms is associated with transformation and that only the ED-A variant is found in metastases. Integrin phenotyping disclosed that α, β expression is associated with tumour progression, α, β is a marker of transformation. α, β is rarely expressed and α, β is expressed by about 50% and 30% of the primary and metastatic lesions respectively. Taken together, the results of this study demonstrate that transformation and tumour progression of the melanocyte lineage are associated with modulation of expression of FN isoforms and FN integrin receptors. Furthermore, the expression of α, β-integrin in a considerable percentage of primary and metastatic lesions indicates that FN may deliver a proliferative stimulus to melanoma cells in vivo.

Keywords: fibronectin; integrin; melanoma

Interactions between epithelia and connective tissues play a major functional role during embryogenesis, tissue repair and tumour growth (Bissel et al., 1982; Bronner-Fraser, 1986; Korhonen et al., 1990; Albelda, 1993). More recently these interactions have been shown to be mediated mainly by the integrin superfamily of molecules (Albelda et al., 1990; Hemler, 1990; Hynes, 1992). These heterodimeric transmembrane glycoproteins can modulate gene expression and differentiation by interacting with cytoskeletal components (Horvitz et al., 1986; Menko et al., 1987; Werb et al., 1989; Adams et al., 1990; Albelda et al., 1990), thus acting as signal transducing molecules. Although a number of in vitro studies have extensively documented that malignant transformation results in qualitative as well as quantitative changes in integrin expression (Juliano et al., 1990) a more complete appreciation of the biological relevance of integrins in tumour growth and metastasis requires in vivo analysis of autochthonous tumours. Malignant cutaneous melanoma is a human malignancy which, because of its development in well-defined stages (Clark et al., 1986) and its relative ease of propagation in culture (Herlyn et al., 1985), offers the possibility of a critical comparison of in vitro and in situ integrin expression. Recent in vitro studies have demonstrated that fibronectin (FN) possesses mitogenic activity in melanoma cells and that this stimulus is mediated by the α, β heterodimer (Mortarini et al., 1993). In the present study we have analysed the expression of FN and two FN isoforms in benign and malignant lesions of the melanocyte lineage. This analysis has been compared with the expression of the major FN receptors of the integrin family.

Materials and methods

Tissue specimens, immunohistochemistry

Surgical biopsies of naevocytic naevi (nine cases) and primary and metastatic melanoma (15 and 26 cases respectively), were obtained from the Surgical Pathology Division of the Regina Elena Cancer Institute. Tissue samples were immediately snap frozen in liquid nitrogen. From each specimen 4 µm cryostat sections were obtained and were fixed in absolute acetone for 10 min. Sections were either immediately used in immunohistochemical assays or kept frozen at -20°C with no loss of immune reactivity. Fixed sections stained with 1% toluidine blue were used to evaluate the histological features of the lesions. Histological diagnosis was done according to Clark (1972). Tumour thickness was evaluated according to Breslow (1975).

Indirect immunoperoxidase stain employed primary MABs at concentrations ranging from 10 to 30 µg ml⁻¹ and a commercially available avidin–biotin staining kit (Vector Laboratories, Burlingame, CA, USA). Negative controls consisted of tissue sections from which incubation with primary antibody was omitted. Selected tissue substrates were used as positive controls. Reaction product was detected using 3-amino-9-ethylcarbazole as a chromogenic substrate and Mayer’s haematoxylin as nuclear counterstain.

Monoclonal antibodies

The murine MABs IST4, IST9 (Borsi et al., 1987), BC-1 (Carmemolla et al., 1989) recognising total tissue fibronectin (FN-T), ED-A (FN-A) and ED-B (FN-B) FN isoforms respectively were kindly provided by Dr L Zardi (Laboratory of Cell Biology, Istituto Nazionale Tumori, Genoa, Italy). MAB M-kid2 specific for the α, β heterodimer was produced as described previously (Bartolazzi et al., 1991). MABs to α, β (M142) and α, β, (M609) integrins were purchased from Chemicon Laboratories, (Temecula, CA, USA). MABs to α, β (10P49d) and to α, β (10F49e) chains were obtained from Immunotech (Marseille, France).

Results

Immunohistochemical evaluation of the three forms of FN in normal skin samples from different body sites demonstrated that, while no B isoform is expressed at detectable level, total and A FN display an overlapping distribution. The two molecules are distributed with an interrupted pattern along the dermal–epidermal junction and the basement membrane of sebaceous and sweat glands. Scattered reaction product is observable in the papillary and reticular dermis. The results of the immunohistochemical evaluation of the expression of FN, FN isoforms and integrin receptors in benign and malignant lesions of melanocyte origin are summarized in Table I.
Table 1 Patterns of expression of total FN (FN-T), FN isoforms (FN-A, FN-B) and integrin receptors in benign and malignant lesions of the melanocyte lineage

| Naevi | FN | a | A | B | a | a | a |
|-------|----|---|---|---|---|---|---|
|       |    | a | + | - | ± | - | ± |
| 1     |    | 2 | - | NT | ± | - | ± |
| 3     |    | 4 | b | - | - | ± | - | ± |
| 5     |    | 6 | b | - | - | - | + | ± |
| 7     |    | 8 | b | - | - | - | - | ± |
| 9     |    | b | - | - | - | - | ± | ± |

Primary melanomas (Clark's level: growth phase)

|       |    | a | b | NT | ± | - | - | ± |
| 1     |    | a | b | NT | ± | - | - | ± |
| 3     |    | a | b | NT | ± | - | - | ± |
| 4     |    | a | b | NT | ± | - | - | ± |
| 5     |    | b | b | NT | ± | - | - | ± |
| 6     |    | b | b | NT | ± | - | - | ± |
| 7     |    | b | b | NT | ± | - | - | ± |
| 8     |    | b | b | NT | ± | - | - | ± |
| 9     |    | b | b | NT | ± | - | - | ± |

Metastatic melanomas

|       |    | a | b | - | ± | - | - | ± |
| 1     |    | a | b | - | ± | - | - | ± |
| 2     |    | a | b | - | ± | - | - | ± |
| 3     |    | a | b | - | ± | - | - | ± |
| 4     |    | a | b | - | ± | - | - | ± |
| 5     |    | a | b | - | ± | - | - | ± |
| 6     |    | a | b | - | ± | - | - | ± |
| 7     |    | a | b | - | ± | - | - | ± |
| 8     |    | a | b | - | ± | - | - | ± |
| 9     |    | a | b | - | ± | - | - | ± |

Patterns of expression of TN and TN isoforms: a, homogeneous and or heterogeneous cellular stain; b, stain of minor interstitial septa with occasional stain of single cells and or small cell nests; -, no stain or stain limited to the major interstitial septa; NT, not tested; RGP, radial growth phase; VGP, vertical growth phase. Patterns of expression of integrin receptors: -, no stain; +, weak homogeneous stain; ±, homogenous stain; +, strong homogeneous stain; v, stain of variable intensity; is, stain of isolated areas accounting for 20% of the lesion. In naevic lesions the stain was mainly restricted to A type cells.

Total FN was detectable in all types of lesions with two major distribution patterns. The glycoprotein was distributed either homogeneously or with a stippled pattern around small single-cell nests or was mainly present in minor interstitial septa which surround cell nests of variable size. While the latter pattern was predominantly observed in benign naevocytic naevi and in primary tumours (Figure 1a), the former was frequently seen in metastatic melanoma (Figure 1b and c). Staining of the same lesions with MAb to FN-A and FN-B isoforms demonstrated that the two molecules are detectable only in malignant lesions. While both isoforms could be detected in a small portion of primary tumours the metastases almost invariably displayed expression of the FN-A variant only. The FN promiscuous receptors exhibit the following distribution patterns. Expression of the $\alpha_5\beta_3$ heterodimer appeared to increase with the degree of invasiveness of primary tumours (Figure 1d–f).

Seventy-three per cent of the metastatic lesions were $\alpha_5\beta_3$ positive. Expression of the $\alpha_5\beta_3$ receptor, while associated with transformation (i.e. differential expression between benign and malignant lesions), did not display a significant relationship with degree of the tumour invasiveness in the series of primary tumours studied (Figure 1g–i). Eighty-five per cent of the metastases were $\alpha_5\beta_3$ positive. The analysis of the distribution of the FN receptors which utilise the $\alpha_5$ chain, i.e. $\beta_1$ and $\beta_3$, demonstrated that these heterodimers characterise the phenotype of the melanocyte lineage with no apparent modulation in the various lesions studied. The evaluation of the FN highly specific integrin receptors disclosed that the one which utilises the $\alpha_5$ heavy chain is rarely expressed at detectable levels in the melanocytic lineage. On the contrary, the $\alpha_5$-containing receptor which is expressed with no apparent relationship with transformation and tumour progression can be detected in about 50% and 30% of the primary and metastatic tumours respectively. The results reported in Table II clearly demonstrate that the above described phenotype is maintained in multiple metastases analysed in three patients.
Discussion

A combination of *in vitro* and *in situ* studies has disclosed that melanoma cells express a panoply of integrin receptors (Kramer et al., 1991) but that distinct integrin repertoires characterise transformation and progression of this neoplasm (Hart et al., 1991). This multistep process seems likely to be mediated by multiple adhesive interactions with the extracellular matrix as well as by growth-modulating stimuli. The latter can be delivered from different ligands either directly (Panayotou et al., 1989) or indirectly, owing to the complexing of extracellular matrix (ECM) macromolecules with

![Figure 1](image-url)

**Figure 1** Immunohistochemical analysis of the expression of FN, FN isoforms and FN receptors in melanocyte lesions as detected by indirect immunoperoxidase stain on 4 μm cryostat sections. In a primary tumour FN is distributed in the minor interstitial septa which surround tumour cell nests of variable size (arrows) (a). FN-A (b) and FN-B (c) are highly expressed in a case of metastatic melanoma. While α, β, is barely detectable in an intradermal naevus (d, arrow; the dark stain is produced by melanin granules) the heterodimer is strongly expressed in an invasive primary (2.4 mm thick) (e) and a metastatic (f) tumour. α, β, is not detectable in an intradermal naevus (g, asterisk) but is expressed at high level in a primary (h) and a metastatic (i) melanoma. ep = epidermis (original magnification × 100).
growth factors (Nathan et al., 1991; Yaoi et al., 1991). Fibronectin has recently been shown to convey a mitogenic signal in vitro to human melanoma cells through binding to the α, β3 highly specific receptor (Mortarini et al., 1993). In view of this finding it was of interest to evaluate in melanoma cells in situ the integrin phenotype that might mediate interactions in vivo with this abundant ECM component. To this end we performed an extensive immunohistochemical analysis of benign and malignant melanocyte lesions aimed at establishing the in situ expression of FN and two of its isoforms and the integrin receptors.

We show here that total FN is widely expressed in melanocytic lesions with no apparent relationship with transformation, tumour progression and metastatic phenotype. The evaluation of the FN-A and FN-B isoforms on the other hand has confirmed that their expression is enhanced by malignant transformation (Carnemolla et al., 1989) and has demonstrated that only the FN-A variant is expressed in metastatic lesions. Although the biological basis of this differential expression of the two isoforms is presently unclear, the demonstration that melanoma cells can produce transforming growth factor beta (TGF-β) (Rodeck et al., 1994), which has the ability to increase expression of the isoform of the FN-A variant (Balza et al., 1988), may account for this finding. It is interesting in this context that increase in expression of the ED-A mRNA has been found to be closely associated with invasiveness of hepatocarcinomas (Oyama et al., 1989). The study of the third known FN isoform, the IIICS variant (Borsi et al., 1991), could not be performed because of the present lack of specific antibody.

The analysis of the expression of the integrin receptors for FN has revealed a complex repertoire. The α5, β1 heterodimer, which has been shown to modulate adhesion of melanoma cells to FN in vitro (Mould et al., 1991) and to control melanoma metastasis in an ‘in vivo’ model (Qian et al., 1994), is rarely expressed by human melanoma cells in vivo. Expression of α5 does not appear to be significantly associated with transformation and tumour progression. On the other hand, α2 characterises the phenotype of about 50% and 30% of the primary and metastatic lesions respectively, thus confirming the in vitro finding (Mortarini et al., 1993) that a subset of tumours may be susceptible to the mitogenic activity of FN. This finding may have some practical implications since the mitogenic activity (Mortarini et al., 1993) of FN as well as its ability to modulate the metastatic behaviour of melanoma cells (Humphries et al., 1991) has been shown to be antagonised by the hydrophilic amino acid peptides Arg-Gly-Asp located in the cell-binding domain of FN. The detection of α5 in naevi cells which display no proliferative activity raises the question of whether the ability of this heterodimer to transduce a mitogenic signal differs according to the stage of differentiation, perhaps through changes in receptor molecule density, requirement for a co-stimulatory signal or presence of variant forms of the heterodimer (Hynes, 1992).

In this regard the lack of expression of distinct FN isoforms suggests that variants of the ligand are unlikely to influence the receptor’s functional activity in naevic cells.

Although our tissue phenotyping could not distinguish α5, β1 from α2, β1 heterodimer, these two highly specific FN receptors do not appear to undergo modulation in the melanocyte lineage. The results of the evaluation of highly promiscuous receptors on FN confirm the increasing expression of α5, β1 with tumour progression (Natali et al., 1993) and the transformation-associated modulation of the α5, β1 integrin (Albelda et al., 1990).

In conclusion, the results of the present study have demonstrated that FN expression remains relatively stable but alternate FN isoforms are differentially regulated during melanoma progression. This contrasts with other ECM glycoproteins which are modulated during melanoma progression (Natali et al., 1985, 1990). Although this in vivo analysis cannot establish to what extent FN is derived by melanocyte cells, this lineage is known to have the ability to synthesise and release this macromolecule (McClenic et al., 1989). Furthermore, the expression of this combination of highly promiscuous and specific integrin receptors suggests that the interactions with this ubiquitous ECM glycoprotein may, at least in part, account for the well-known property of this malignancy to metastasise to many anatomical sites (Rosai, 1981).

Finally, in view of the ability of TGF-β to bind to FN (Fava and McClure, 1987) this ECM glycoprotein appears capable of modulating tumour growth by multiple mechanisms e.g. cell adhesion by interacting with various integrin receptors, direct mitogenic activity through selected integrin molecules, or growth control stimuli (Rodeck et al., 1994) through bound growth factors. Additional work using well-defined experimental models may eventually confirm this hypothesis.

Acknowledgements

The authors kindly acknowledge the technical help of Miss Cristina Valentin and the secretarial assistance of Miss Maria Vincenza Sarcone. This study has been supported by PF-CNR ACRO, by AIRC and the Italian Ministry of Public Health.

References

ADAMS J C AND WATT F M (1990). Changes in keratinocyte adhesion during terminal differentiation: reduction in fibronectin binding precedes α5, β1 integrin loss from the cell surface. Cell, 63, 425–435.

AlBELDA SM (1993). Biology of disease. Role of integrins and other cell adhesion molecules in tumor progression and metastasis. Lab. Invest., 68, 4–17.
HUMPHRIES HYNES RO. HEMLER ME. CLARK CANNELOM A. HART BRONNER-FRASER ALBELDA BARTOLAZZI.

murine peptide functions in KOPROWSKI ELDER receptor 83-103.

Cancer RNA of analysis PANDEH lular Generation 42-44. isoforms 572-575.

sion L. mRNA generation of fibronectin isoform 192,372-379. BALZA A.

mRNA matrixdirectgene Biol., 595-600. LEPRINI A.

precursors in tumors. Expression of melanoma. Cancer, 54, 68–72.

NATALI C AND SPORN M. (1991). Cytokines in context. J. Cell Biol., 113, 981–986.

OYAMA F. HIROHASHI S. SHIMOSATO Y. TITANI K AND SEKI-GUCHI K. (1989). Deregulation of alternative splicing of fibronectin pre-mRNA in malignant human liver tumors. J. Biol. Chem., 264, 10331–10334.

PANAYOTOU G. END P. AUMAILLEY M. TIMPL R AND ENGEL J. (1989). Domains of laminin with growth activity. Cell, 56, 93–101.

RODECK U. BOSSLER A. GRAEVEN U. FOX FE. NOWELL PC. KNABB C AND KARI C. (1994). Transforming growth factor β production and responsiveness in normal human melanocytes and melanoma cells. Cancer Res., 54, 475–581.

QIAN F. VAUX DL AND WEISSMAN IL. (1994). Expression of the integrin α5β1 on melanoma cells can inhibit the invasive stage of metastasis formation. Cell, 77, 335–347.

ROSAL J. (1981). Ackerman’s Surgical Pathology. C.V. Mosby: St. Louis, MO.

WERB Z. TREMBLE P. BEHRENDTSSEN O. CROWLEY E. DAMSKY CH. (1989). Signal transduction through the fibronectin receptor induces collagenase and stromelysin gene expression. J. Cell Biol., 109, 877–889.

YAOI Y. Hashimoto K. Takahara K AND KATO I. (1991). Insulin binds to type V collagen with retention of mitogenic activity. Exp. Cell Res., 194, 180–185.