Thrombospondin modulates melanoma–platelet interactions and melanoma tumour cell growth in vivo

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Summary

In this study we have investigated the role of thrombospondin (TSP) as a possible ligand playing a key role in the human M₃Da melanoma cell interaction with platelets and in tumour growth. TSP is secreted (80 ± 6 ng TSP 10⁻⁶ cells) and bound to the surface of M₃Da. cells via receptors different from CD36, as shown by biosynthetic labelling and immunofluorescence studies. The levels of TSP binding to M₃Da. cells evaluated by binding studies, using an anti-TSP monoclonal antibody (MAB LY8), show 367 000 ± 58 000 (mean ± s.d.) TSP binding sites per cell with a dissociation constant (K_d) of 67 nm. TSP binding to M₃Da. cells shows 400 000 ± 50 000 TSP binding sites per cell with a K_d of 10 nm. The capacity of anti-TSP MAB (LY8) to inhibit M₃Da. platelet interactions was followed on an aggregometer and evaluated by electron microscopy studies. The biological role of TSP binding to M₃Da. cells was investigated by implanting subcutaneously the M₃Da. cell line in nude mice and following the size and site of in vivo tumour growth. Reducing the availability of TSP by using an anti-TSP MAB (LY8) resulted in a significant decrease in platelet aggregates interacting with M₃Da. melanoma cells. Using an enzyme-linked immunosorbent assay, purified α₃β₁ was shown to bind TSP. Moreover, LY8-coated M₃Da. cells showed a reduced capacity to form tumours in vivo. M₃Da. cells were observed to attach and spread on human platelet TSP-coated plastic wells. This attachment by M₃Da. cells was inhibited in a similar way by LY8 and an anti-α₃β₁ MAb (LYP8). The results obtained in this study show that TSP secreted and bound to the surface of a human melanoma cell line (M₃Da.) acts as a link between aggregated platelets and the M₃Da. cell surface.

Moreover, these results show that TSP can modulate tumour growth in vivo. Reagents such as MABs directed against TSP and peptides derived from TSP could not only be used as a new therapeutic approach in the control of tumour metastasis of melanoma, but may also contribute to elucidation of the role of TSP in cancer biology.

Keywords: thrombospondin; melanoma; monoclonal antibodies; platelet aggregation; vitronectin receptor

Thrombospondin (TSP) is a high molecular weight (450 kDa) glycoprotein (GP) that is released from the α-granules of platelets during activation (Lawler, 1986). In the presence of a physiological concentration of calcium, TSP binds to the surface of stimulated platelets and plays an active role in promoting platelet aggregation (Leung, 1984; Boukerche and McGregor, 1988; McGregor and Boukerche, 1993). TSP is also secreted by a variety of cells, such as pneumocytes, endothelial cells, macrophages, fibroblasts, smooth muscle cells, chondrocytes and mesangial cells, as well as by a number of tumour cell lines, including melanomas and carcinomas (Varani et al., 1986, 1989). In addition to its role in blood coagulation, TSP has been reported to promote adhesion and motility of several of these cell types (Walz, 1992). Released TSP will bind to the surface of activated platelets or cells via a number of glycoprotein receptors, such as the integrins GP Ib–IIa (α₂β₁) and α₅β₃ (Karczewski et al., 1989; Tuszyński and Kowalska, 1991), the vitronectin receptor α₅β₃ (Lawler et al., 1988), GPIV (also called CD36) (Asch et al., 1987; McGregor et al., 1989), the integrin-like receptor (105 80 kDa) (Yabkowitz and Dixit, 1991) and the heparan sulphate proteoglycans (Roberts, 1988). Recently, it was shown that tumour formation and the metastatic spread of lung tumours in mice was increased by TSP and was significantly reduced by use of TSP peptide CSVTCG or a TSP cDNA antisense expression vector (Tuszyński et al., 1987, 1992; Castle et al., 1991). These studies suggest that TSP plays a major role in cell adhesion and cell–cell interactions in the metastatic process.

However, so far very little is known about the precise biological role of TSP in tumour–platelet interactions and tumour cell growth. Moreover, the identity of the TSP receptor(s) mediating tumour cell–platelet interactions and tumour growth remains to be elucidated. We have previously reported that a monoclonal antibody (LYP8), generated against human blood platelet glycoprotein Ib–IIa (α₂β₁), immunoprecipitated two proteins from a tumorigenic human melanoma cell line (M₃Da.) immunologically related to the vitronectin receptor (α₅β₃) (Boukerche et al., 1989a,b). When bound to the melanoma cell surface, LYP8 dramatically inhibited melanoma–platelet interactions and the growth of melanoma cells in nude mice. However, at that stage of our work we had not yet investigated possible ligands involved linking M₃Da. receptors, such as α₂β₁, and platelets or the basement membrane extracellular matrix (ECM) components. Results presented in this study show that TSP secreted and bound to the surface of a human melanoma cell line (M₃Da.) acts as a link between aggregated platelets and the M₃Da. cell surface. Moreover, these results show that TSP can modulate tumour growth in vivo.

Materials and methods

Antibodies

Monoclonal antibody (MAB) LYP8 was produced in our laboratory and was previously shown by crossed immunoelectrophoresis and affinity chromatography to be directed against platelet thrombospondin 1 (TSP1). It recognises a determinant associated with the intact conformation of the antigen, since it failed to bind to a Western blot of SDS-electrophoresed TSP (Boukerche and McGregor, 1988).
No difference in the binding of LYP8 to TSP in an enzyme-linked immunosorbent assay (ELISA) could be observed in the presence of 2 mM Ca\(^{2+}\) or 5 mM EDTA. The LYP8 epitope lies within the 140 kDa non-heparin binding fragment. LYP10 and LYP12 anti-TSP MAb s were produced in our laboratory and are respectively directed against the 70 kDa trypsin-resistant core region and the heparin-binding domain of TSP1 (Catimel et al., 1992). An anti-GPIIIb MAb (OKM5) was a generous gift from Ortho Pharmaceutical. MAb G\(_A\) directed against a melanoma cell membrane antigen was purchased from ImmunoTech.

**Tumour cells**

The M\(_3\)Da. (= M\(_3\)Dau.) cell line was established from an achromic skin metastasis of a patient with malignant melanoma (Jacobovitch et al., 1984). The cells were cultured as monolayers in RPMI-1640 supplemented with 10% fetal bovine serum and have a doubling time of 28 h (Boukerche et al., 1989a). Cultures were routinely checked and found to be free of mycoplasma, using the Hoechst 33258 fluorescence staining procedure. Cell monolayers, used in platelet aggregation studies, were detached by EDTA and resuspended in HBSS (Hanks’ balanced salt solution) containing 0.2% calf serum albumin (BSA) as previously described (Boukerche et al., 1989a). Cell viability, assessed by trypan blue exclusion, was consistently 95% throughout the experiments. Immunofluorescence studies on these cells were performed as follows: cells were gently washed three times with Ca\(^{2+}\)/Mg\(^{2+}\)-free HBSS, detached with EDTA and resuspended in phosphate-buffered saline (PBS), pH 7.2. The cells were then incubated initially with a 1:50 dilution of the monoclonal antibody LYP8, LYP18 or OKM5 for 30 min at room temperature. Cells were washed with PBS, pH 7.2, and stained for 30 min with a 1:50 dilution of fluorescein-conjugated goat anti-(mouse IgG) F(ab’)\(_2\). Following three additional washes in PBS, pH 7.2, cells were analysed by standard fluorescence microscope techniques.

**Effect of an anti-TSP MAb (LYP8) on melanoma–platelet interaction**

Blood from healthy donors was drawn into heparin anticoagulant and centrifuged at 160 g for 20 min to obtain platelet-rich plasma (PRP). Aggregation studies were performed with platelets in PRP adjusted to 3 \(\times\) 10\(^5\) cells ml\(^{-1}\) and melanoma cells preincubated for 1 h at 37°C before addition to 0.4 ml of PRP (Boukerche et al., 1989a). Melanoma cells (4 \(\times\) 10\(^5\) cells) were preincubated with saturating concentrations of LYP8 (24 \(\mu\)g 10\(^{-6}\) cells) or LYP18 (14 \(\mu\)g 10\(^{-6}\) cells), washed three times with HBSS containing 0.2% BSA then added to PRP. Platelet aggregation by melanoma cells was quantitated by measuring the peak height of the aggregation curve as previously described (Boukerche et al., 1989a).

**Electron microscopy of platelet–melanoma interaction**

At the end of the aggregation curve, platelet–melanoma suspensions were fixed with 0.15% glutaraldehyde in 0.1 mol l\(^{-1}\) cacodylate buffer and then filtered through 0.22 \(\mu\)m filters (Millipore, France). The fixed material was processed for electron microscopy as previously described using a Siemens ELM 102 transmission electron microscope (Boukerche et al., 1989a).

**Immunoprecipitation**

Confluent melanoma cells were metabolically labelled with \(^{35}S\) for 24 h and chased for 18 h in the absence of labelled methionine. The Triton X-100-extracted melanoma chase media were immunoprecipitated with an anti-TSP MAb (LYP8) or a non-immune mouse serum IgG and analysed by sodium dodecyl sulphate–polyacrylamide gel electrophoresis (SDS–PAGE) as previously described (Boukerche et al., 1989b).

**Isolation of thrombospondin and vitronectin receptors**

Thrombospondin (TSP) was purified from the supernatant of ionophore-activated human platelets on a heparin–Sepharose CL-6B column in the presence of 2 mM calcium following the method of Lawler et al., (1988). Vitronectin receptor (VnR) was isolated from melanoma cells by LYP18-affinity chromatography as previously described (Boukerche and McGregor, 1988). The VnR was more than 95% pure when analysed by SDS–PAGE.

**Binding studies**

Human melanoma cells grown to confluence in serum-free RPMI were detached from culture plates by brief treatment with trypsin–EDTA and resuspended in RPMI-1640 containing 0.35% BSA. Thrombospondin (TSP) or isolated antibody was labelled with \(^{125}\)I as previously described (Boukerche and McGregor, 1988). Labelled TSP or antibody was separated from free \(^{125}\)I on a PD-10 column (Phadebas G-25 M). Increasing concentrations of \(^{125}\)I-labelled TSP (0.4–24 \(\mu\)g ml\(^{-1}\)) or \(^{125}\)I-labelled LYP8 (0.4–24 \(\mu\)g ml\(^{-1}\)) were added to 10\(^6\) cells and incubated overnight at 4°C. At the end of the incubation time, aliquots of the melanoma–antibody or melanoma–TSP mixture were layered in triplicate in 400 \(\mu\)l Eppendorf tubes containing 20% sucrose, 2% bovine serum albumin prepared in PBS, pH 7.2. The amount for \(^{125}\)I-labelled TSP or \(^{125}\)I-labelled LYP8 bound per melanoma cell was determined by counting the radioactivity of the cut-off centrifuge tips. Non-specific binding was measured by incubating cells with a 100-fold excess of unlabelled TSP or monoclonal antibody. This non-specific binding (less than 10–15%) was subtracted from the total binding. The number of binding sites per cell and the dissociation constant were obtained by double-reciprocal plots and least-squares regression analysis. The values reported are the mean of three experiments. Binding of TSP to vitronectin receptor was studied using an ELISA as described previously (Boukerche and McGregor, 1988). Briefly, purified \(\alpha_b\) (2–3 \(\mu\)g) diluted in PBS, pH 7.2, containing 2 mM calcium chloride was adsorbed on each well of the microtitre plate for overnight incubation at 4°C. Wells were then washed in PBS containing 1 mM calcium chloride supplemented with 0.05% Tween 20, and increasing concentrations of TSP or antibody (0.1–1.5 \(\mu\)g) diluted in PBS containing 2 mM calcium chloride were added to wells for overnight incubation at 4°C. Plastic wells were then washed and anti-TSP polyconal antibody was added for 1 h at 37°C to detect bound TSP. After additional washes, goat anti-rabbit antibody conjugated to horseradish peroxidase was added and bound anti-TSP polyclonal antibody was quantitated by the addition of the substrate orthophenyldiamine.

**Quantification by ELISA of secreted thrombospondin by melanoma cells**

Melanoma cells in complete medium were plated at 3.5 \(\times\) 10\(^4\) cells per well and grown for 1 day. The cells were then extensively washed with serum-free RPMI and incubated for 6 h in serum-free RPMI containing 0.2% BSA. Cells were centrifuged at 200 g for 10 min at 4°C and TSP secreted into the culture medium was assayed as previously described (Risser et al., 1988) using a TSP ELISA kit (Stago, France).

**Cell adhesion assays**

The cell adhesion assay was performed as previously described (Varani et al., 1986). Briefly, 2–3 \(\mu\)g of TSP or BSA diluted in PBS, pH 7.2, containing 1 mM calcium chloride and 0.5 mM magnesium chloride was adsorbed on each well of the microtitre plate for 1 h at 37°C. After washing, the wells were blocked with serum-free minimum essential
medium (MEM) supplemented with 0.2% BSA to minimise non-specific tumour cell adhesion. Anti-TSP MAb (LYP8) was then added to the wells and incubated for 20 min at 37°C. Melanoma cells were briefly harvested from culture by trypsinisation or by replacing the medium with PBS, pH 7.2, containing 2.5 mM EDTA, washed in serum-free MEM, and resuspended in serum-free MEM containing 0.2 mg ml\(^{-1}\) BSA. Cells (10\(^4\)) were then added to coated wells for 60–90 min at 37°C. After gently washing, wells were then fixed with glutaraldehyde and stained with 1.5% Giemsa. Attached and spread cells were counted microscopically. In other experiments, cells were preincubated for 20 min at 37°C with anti-α\(_1\)β\(_1\) MAB (LYP18) (14 μg ml\(^{-1}\)) before being added to TSP-coated wells. Cells adhering to albumin-coated wells were not significant (less than 2%) compared with cells adhering to TSP-coated wells.

**Tumorigenicity assays**

Melanoma cells were harvested by trypsinisation and washed three times with serum-containing medium, then resuspended in PBS, pH 7.2. Two hundred microlitres of M\(_{3}D\)a cells (1 × 10\(^5\)) were grafted subcutaneously (s.c.), as previously described (Boukerche et al., 1989b), on the ventral surface of nude mice, after preincubation for 20 min with purified LYP8 (24 μg ml\(^{-1}\)), LYP18 (7 μg ml\(^{-1}\)), a non-immune mouse serum IgG, or G\(_{\alpha}\) (directed against a melanoma cell-surface antigen other than the vitreoretinal receptor α\(_1\)β\(_1\) or thrombospondin). Tumor sizes at different time points were expressed as the mean of the sum of two perpendicular diameters. In these experiments, five mice were used for each time point.

**Assays for effect of LYP8 on tumour cell proliferation and nucleic acid synthesis**

**Tumour cell proliferation** Aliquots (6.5 × 10\(^4\)) of melanoma cell suspension preincubated for 10 min at room temperature with a saturating concentration of MAB LYP8 (24 μg ml\(^{-1}\)) or a non-immune mouse serum IgG were seeded to each of the 96 wells of the flat-bottomed tissue culture plate. At selected time, cells were washed with Dulbecco’s PBS, trypsinised and counted. Data are the mean of six well counts.

**Nucleic acid synthesis** Aliquots (15 × 10\(^3\)) of melanoma cell suspension preincubated with saturating concentration of MAB LYP8 (24 μg ml\(^{-1}\)) or a non-immune mouse serum IgG were added to each of the 24 wells of the flat-bottomed tissue culture plate and cultured for 1–2 days. The cells in each well were pulsed for 12 h with [\(^3\)H]thymidine (0.0185 MBq per well, 37 GBq mmol\(^{-1}\), Amersham). Cells were then washed with Dulbecco’s PBS, trypsinised and radioactivity incorporated in the cells counted. Data are the mean of 12 well counts.

**Results**

**Cell-surface expression of thrombospondin (TSP) by M\(_{3}D\)a**

An ELISA, performed on the conditioned media of cultured cells, showed that M\(_{3}D\)a secreted 80 ± 6 ng TSP 10\(^4\) cells (mean ± s.d., n = 3). LYP8, a monoclonal antibody (MAB) directed against platelet TSP and a potent inhibitor of platelet aggregation induced by thrombin and collagen (Boukerche and McGregor, 1988), immunoprecipitated from chase media of metabolically labelled M\(_{3}D\)a melanoma cells a protein having the same apparent molecular weight (mol. wt) as TSP with the characteristic changes in mobility on reduction (Figure 1a). This MAB was observed by immunofluorescence staining to bind to the surface of both M\(_{3}D\)a melanoma cells (Figure 1b). This staining appeared uniformly distributed over the entire surface of M\(_{3}D\)a cells, with high fluorescence intensity in some areas of the membrane, suggesting the presence of clusters of TSP. Alternatively, this clustering may be an artefact due to multivalent interaction of the first antibody with cell-surface TSP. Similarly, as previously reported, M\(_{3}D\)a cells stained with an anti-α\(_1\)β\(_1\) MAB LYP18 (Figure 1b) (Boukerche et al., 1989b). In contrast, an anti-CD36, polyclonal to MAB OKM5, did not bind to M\(_{3}D\)a cells (Figure 1b). To determine the number of TSP molecules expressed on the surface of melanoma cells, binding studies using \(^{125}\)I-labelled LYP8 were performed. LYP8 binding to M\(_{3}D\)a melanoma cells was specific, concentration-dependent and saturable with 367 000 ± 58 000 (mean ± s.d., n = 3) LYP8 binding sites per cell and a dissociation constant (K\(_d\)) of 67 nM (Figure 2). Binding of labelled LYP8 to M\(_{3}D\)a cells was reduced by 90% in the presence of a 100-fold excess of cold LYP8. Inhibition of TSP binding to M\(_{3}D\)a cells was concentration dependent over the range of 0.4 to 24 μg ml\(^{-1}\) with saturation 400 000 ± 50 000 TSP binding sites per cell (mean ± s.d., n = 3) and a dissociation constant (K\(_d\)) of 10 nM. In the presence of an excess of cold TSP, binding of \(^{125}\)I-labelled TSP was reduced to less than 85%. LYP8 (10 μg ml\(^{-1}\)) did not affect the binding of labelled TSP to M\(_{3}D\)a cells or to platelets stimulated with collagen (0.4 U ml\(^{-1}\)) (results not shown). In human blood platelets and in certain tumour cell lines, CD36 was shown to act as one of the TSP receptors (Asch et al., 1987; McGregor et al., 1989; Silverstein et al., 1992). These results clearly demonstrate that M\(_{3}D\)a synthesise and secrete into the culture medium a protein immunologically related to TSP that binds to the cell surface via receptors different from CD36.

**Inhibition of platelet–melanoma interaction by anti-thrombospondin (TSP) MAB LYP8**

Since M\(_{3}D\)a cells synthesise TSP and bind LYP8, the role of TSP in tumour–platelet interaction was investigated. M\(_{3}D\)a cells irreversibly aggregated human platelets in heparinised PRP (Figure 3). Platelets washed by the technique of Mustard et al. (1972) were not aggregated by M\(_{3}D\)a melanoma cells (results not shown). Addition to platelets of LYP8 (24 μg ml\(^{-1}\) cells) inhibited platelet aggregation induced by M\(_{3}D\)a (Figure 3). If M\(_{3}D\)a cells are preincubated with a saturating concentration of LYP8 (24 μg ml\(^{-1}\) cells), washed three times with HBSS–BSA, then added to platelets, aggregation is blocked (Figure 3). Anti-TSP MABS, LYP10 or LYP12, used at the same concentration or a non-immune mouse serum IgG, had no effect on platelet aggregation induced by melanoma cells. As previously shown, an anti-α\(_1\)β\(_1\) MAB (LYP18) added to M\(_{3}D\)a cells significantly inhibited platelet aggregation induced by M\(_{3}D\)a cells (Figure 3) (Boukerche et al., 1989c).

**Electron microscopy of melanoma–platelet interactions**

Electron microscopy studies showed that melanoma cells closely interacted with platelets (Figure 4a). Melanoma cells at the site of their interaction with platelets formed extrusions or processes which penetrated into the platelet aggregate (Figure 4b) (Boukerche et al., 1989c). Addition to a platelet–tumour cell mixture of anti-TSP MAB LYP8 resulted in a significant decrease in the size of platelet aggregates (Figure 4c). Moreover, platelet–melanoma cell interactions could not be observed, confirming the aggregometry results (Figure 4c). Moreover, in the presence of LYP8, the tumour cell surface did not show cytoplasmic extrusion or processes (Figure 4c).

**Effect of LYP8 on melanoma tumour growth in nude mice**

The biological role of TSP binding to M\(_{3}D\)a cells was investigated by implanting subcutaneously the M\(_{3}D\)a cell line in nude mice and following the size and time of in vivo tumour growth over a period of 6 weeks. In the presence of LYP8 (24 μg ml\(^{-1}\) cells), M\(_{3}D\)a cells are inhibited from growing into full-sized tumours as observed in control animals (Figure 5a). Similar inhibition was observed with an anti-α\(_1\)β\(_1\),
MAb LYP18 (Figure 5a). The inhibition of tumour cell growth by LYP8 or LYP18 extends over a period of 40 days. Over that period M2 Da. cells gave rise to small tumours growing at a lower rate than in the controls. Other anti-TSP MAbs (LYP10, LYP12, directed respectively against the 70 kDa trypsin-resistant core region and the heparin-binding domain of TSP) had no effect on tumour growth. Combination of the two MAbs (LYP8 + LYP18), used at saturating concentrations, gave similar results as that obtained with anti-α,β3 MAb LYP18. To rule out the possibility that LYP8 might have a direct cytotoxic effect, melanoma cells were preincubated with excess LYP8 (24 μg ml⁻¹) and monitored for cell viability and cell growth. No loss of melanoma cells viability was shown by trypan blue exclusion. Furthermore, control and LYP8-treated cells showed a similar degree of [³H]thymidine uptake and cell growth in vitro (Figure 5b and c). The lack of effect caused by the antimalanoma MAb G3A3, suggests that the observed inhibition by LYP8 or LYP18 in vivo was not due to complement-dependent cytotoxicity or opsonisation. Moreover, in the presence of fresh rabbit or nude mice sera containing complement, LYP8 did not support lysis of melanoma cells (results not shown). As previously reported, natural killer cells are not involved in M2 Da. growth in nude mice (Jacubovich et al., 1984).

Adhesion assays

In order to study the mechanism allowing an anti-TSP MAb (LYP8) to inhibit tumour growth, we looked at the ability of TSP to support M2 Da. adhesion, and the effect of LYP8 on such an interaction. M2 Da. cells attached to TSP-coated wells with 20% of cells spreading (Figure 6a). LYP8 had no effect on the attachment of M2 Da. cells to TSP but decreased significantly the number of cells spreading on TSP (Figure 6a and b). Extending the incubation time of tumour cell adhesion on TSP with LYP8 from 60 to 90 min gave similar results (results not shown). TSP secretion does not promote shedding of antibody since [¹²⁵I]-labelled LYP8 bound to TSP-

Figure 1 (a) Immunoprecipitation by LYP8 monoclonal antibody (MAb) of thrombospondin (TSP) from chase medium of metabolically labelled (³⁵S)methionine) M2 Da. (= M2 Dau.) human melanoma cells. Immunoprecipitates were applied to 5–15% exponential gradient SDS–polyacrylamide gels and electrophoresed under non-reducing (NR) or reducing (R) conditions. Lanes A. chase medium; lanes B and E, immunoprecipitates with LYP8 MAb; lanes C and D, immunoprecipitates with non-immune mouse serum IgG. (b) Immunofluorescence staining of M2 Da. human melanoma cells by an anti-TSP MAb LYP8. M2 Da. cells were incubated with LYP8 or an anti-α,β3 MAb (LYP18) or an anti-CD36 MAb (OKM5). Rabbit anti-mouse IgG F(ab')₂ antibody conjugated to fluorescein isothiocyanate (FITC) was then added to cells.
Biological role of thrombospondin in melanoma cell interactions
H Boukerche et al

Figure 2 Binding of 125I-labelled anti-thrombospondin (TSP) monoclonal antibody (MAb) (LYP8) to M1Da human melanoma cells. Human melanoma cells grown to confluence in serum-free RPMI were detached by trypsin–EDTA or EDTA and resuspended in RPMI-1640 containing 0.35% BSA. Increasing concentrations of 125I-labelled LYP8 were added and incubated overnight at 4°C. The insert shows a double-reciprocal plot of the same data with $K_d = 66.6$ nM and the maximum number of binding sites of $4 \times 10^5$ per cell. Non-specific binding of labelled LYP8 was obtained by using a 100-fold excess of unlabelled LYP8.

Figure 3 Effect of an anti-thrombospondin (TSP) monoclonal antibody (MAb) (LYP8) on aggregation of platelets in PRP induced by M1Da. ( = M1Dau.) human melanoma cells. Typical aggregation curves of platelet in heparinised PRP induced by the addition of M1Da melanoma cells (4 x 10^5). Controls were either HBSS–BSA buffer or a non-immune mouse serum IgG. LYP8 (24 μg 10^-5) or LYP18 (14 μg 10^-5) were added to platelets before the addition of M1Da melanoma cells. Similar results were obtained with M1Da melanoma cells pretreated with LYP8 (24 μg 10^-5), washed three times with HBSS–BSA, then added to platelets.

Figure 4 Ultrastructural analysis of M1Da human melanoma–platelet interactions. At the end of the aggregation curve, platelet–melanoma suspensions were fixed with glutaraldehyde and prepared for electron microscopy as described in Materials and methods. (a) Platelets aggregates (p) in direct contact with M1Da melanoma cells (m) ( x 3400). (b) At higher magnification, melanoma (m) showed extrusions (arrows) at the site of their interaction with platelets (p) ( x 12 500). (c) Platelets (p) not interacting with M1Da melanoma cells (m) when platelets were preincubated with LYP8 ( x 3400). Bar = 2 μm.
coated wells in the presence of M₁₂₅ Da, cells was not released into the supernatant (less than 1% of the total TSP bound LYP8 was recovered in the supernatant). Similarly, an anti-α₁β₃ MAb, LYP18, had no effect on attachment but inhibited spreading of M₁₂₅ Da, cells on TSP (Figure 6b). To examine further the specificity of TSP binding to α₁β₃, an ELISA was performed. Preliminary experiments performed with Triton X-100 melanoma cell lysate using LYP18 and LYP8 in a double-antibody sandwich ELISA showed that TSP binds to α₁β₃ (results not shown). Further experiments were then performed with purified α₁β₃. Purified α₁β₃ was added to wells of the microtiter plate followed by the addition of increasing concentrations of TSP or albumin. Binding of TSP to α₁β₃ bound to LYP18 was significant compared with albumin (Figure 7). These results show that TSP in M₁₂₅ Da, interacts with α₁β₃ receptors and exclude the possibility that the observed binding may be due to minor contaminants present in the sample.

Discussion

This study indicates that one of the adhesive ligands playing a key role in human melanoma (M₁₂₅ Da,) cell interaction with platelets is thrombospondin (TSP). Moreover, TSP also appears to play a crucial role in the control of M₁₂₅ Da, tumour growth. The site on TSP to which monoclonal antibody (MAb) LYP8 is directed appears, in contrast to other anti-TSP MAb's (LYP10, LYP12) binding to different epitopes, to play an important role in melanoma—platelet interaction and tumour growth. Several lines of evidence back the above statements:

(1) TSP binds with a high affinity to the surface of M₁₂₅ Da, via receptors that differ from CD36, not expressed by M₁₂₅ Da.
cells is directly involved in mediating tumour cell–platelet interactions. A similar observation was reported with LYP8 and other MAbS directed against GPIIb and IIIa, which together inhibited platelet aggregation without affecting fibrinogen binding (Newman et al., 1987; Boukerche and McGregor, 1988). These results suggest that additional post-TSP binding events such as conformational changes and/or clustering of cell-surface molecules may be required to support tumour–platelet interaction (Peerschke and Zucker, 1981).

Such a role of TSP in mediating cell–cell interactions was also shown with thrombin-activated platelets binding to a melanotic cell line (USMC-4, McGregor et al., 1991; Silverstein et al., 1992). In contrast to these findings, platelets in our system are activated by ADP and released by M3Da. cells, under conditions in which presumably no secretion from platelets takes place (Boukerche et al., 1989a). Recent results from our laboratory indicate that ADP is present in appreciable amounts (180 ± 10 pmol 10⁶ cells) in the cell supernatant of M3Da. cells as shown by high-performance liquid chromatography (HPLC) (Boukerche et al., 1994). Previous observations have shown that platelets activated by ADP in the presence of physiological levels of Ca²⁺, as present in heparinised PRP, aggregate but do not release their α-granule content (Mustard et al., 1972). Under these conditions, TSP released by M3Da. cells will not bind to platelets that have not undergone release (Leung, 1984; Boukerche and McGregor, 1988). In view of these results, it appears that ADP-activated platelets need to interact with cell-surface receptors presumably via aβ₃, to undergo complete degradation as observed in electron–micrographs obtained in this study and in a previous work (Boukerche et al., 1989a). Platelet–melanoma interactions may therefore be initiated by: (1) the release of ADP by M3Da. cells and (2) mechanical stimulation induced by the M3Da. cell surface. Melanoma cells, by interacting with the platelet surface, may promote agonist and ADP release from platelets, leading to degradation and formation of larger platelet–tumour aggregates. TSP coating of degraded platelet is known to help in cementing platelet aggregates (Leung, 1984, Boukerche and McGregor, 1988). The ability of tumour cells to induce platelet activation in vitro has been used as indirect evidence to show the role of platelets in the dissemination of tumour cells (Gasic et al., 1973). TSP linking the tumour–platelet aggregates may provide the vehicle for transport and dissemination of tumour cells. Experiments are under way to determine the metastatic capacity of M3Da. binding TSP.

Tumour growth is the result of a complex interaction between tumour cells and the basement membrane extracellular matrix components (Dvorak et al., 1991). TSP appears to play a crucial role in the control of tumour growth. The finding that anti-TSP MAb LYP8 did not completely inhibit tumour growth suggests that other glycoprotein receptors and adhesive proteins contribute to the full expression of the tumorigenic phenotype of the cells. Adhesive proteins such as TSP binding to melanoma cells may in vivo modulate cellular proliferation as previously shown for normal and transformed cells (Majack et al., 1988; Abbadia et al., 1993). Alternatively, MAb LYP8 binding to TSP may block critical interactions between melanoma cells and stromal matrices that are vital for successful angiogenesis (Tolsma et al., 1993; Dameron et al., 1994).

TSP is a multidomain glycoprotein that binds to a number of adhesive receptors (Asch et al., 1987; Lawler et al., 1988; Roberts, 1988; Karzewska et al., 1989; McGregor et al., 1989; Yabkowitz and Dixit, 1991). One of its receptors, CD36, which binds to the 68 kDa TSP fragment via the type I repeat (CSVTCG) in the absence of Ca²⁺, is not expressed by platelet cells (Asch et al., 1992; Caras et al., 1992). In the current study, M3Da. melanoma cells attached and spread on TSP in a similar way as reported by Roberts et al. (1987) for melanoma G361. LYP8 and LYP18 MAbS inhibited M3Da. spreading on TSP. Purified aβ₃ was shown to bind TSP. These results and previous work (Lawler

(2) Reducing the availability or functional level of TSP by using an anti-TSP MAb (LYP8) resulted in a significant decrease of platelet aggregates interacting with melanoma.

(3) Tumour formation in vivo was also affected by the presence of bound LYP8.

(4) LYP8 and an anti aβ₃ (LYP18) inhibited in a similar way spreading of M3Da. cells to coated TSP.

(5) Purified aβ₃ binds TSP in an ELISA.

An important point to be added to the above observations is that NAK- platelets deficient in CD36 have been shown to aggregate normally in the presence of M3Da. melanoma cells (H Boukerche, B Kehrel and JL McGregor, unpublished observations) (Kehrel et al., 1993). The absence of CD36 expression on M3Da. suggests that TSP binds to this melanoma cell line via another receptor. An obvious candidate to bind TSP could be aβ₃, which is known to bind to the RGDA sequence of TSP on endothelial and melanoma cells (Lawler et al., 1988; Tuszyński et al., 1989). This receptor (aβ₃) has been shown to be expressed by M3Da. and plays, as previously indicated, a crucial role in melanoma–platelet interaction and in vivo melanoma tumour growth (Boukerche et al., 1989a,b; Marshall et al., 1991; Felding-Habermann et al., 1992). Results in this study and previous work (Lawler et al., 1988; Tuszyński et al., 1989) suggest that TSP binding to aβ₃, expressed by M3Da., is directly implicated in melanoma–platelet interaction and tumour growth formation in vivo. Our data do not exclude a possible role of CD36 present in the microenvironment of a tumour in vivo. Results in this study extend observations made by Tuszyński et al. (1987, 1992), who showed that whole TSP and a peptide (CSVTCG) derived from this adhesive ligand affect tumour metastasis of mouse melanoma. Moreover, CSVTCG and its analogue have been shown to be potent inhibitors of platelet aggregation (Byyck and McGregor, 1992). The recent discovery of four homologous forms of TSP (TSP-1, TSP-2, TSP-3 and TSP-4) encoded by distinct genes indicates that further characterisation of TSP expressed by melanomas is required (Bornstein, 1992).

An anti-TSP MAb (LYP8) significantly affects platelet aggregates interacting with melanoma cells, as shown in the electron micrograph presented in this study. However, LYP8 did not reduce TSP binding to M3Da. cells, nor did it affect TSP binding to washed platelets stimulated with thrombin. These results indicate that LYP8 may interfere by steric hindrance with a mechanism involved in linking tumour cells and platelets together and suggest that TSP coating M3Da.

![Figure 7 ELISA detection of thrombospondin (TSP)–aβ₃ complex formation. Purified aβ₃ (1–2 μg) was coated on the wells as described in Materials and methods. After incubation and washing, increasing concentrations of TSP or albumin (0.1–1.5 μg) were added to the wells and bound TSP was probed with anti-TSP polyclonal antibody followed by horseradish peroxidase conjugated to goat anti-rabbit antibody. Binding of anti-TSP polyclonal antibody was quantified by the addition of the substrate orthophenylendiamine. Data are the mean of three experiments.](image-url)
et al., 1988; Tuszynski et al., 1989) suggest that TSP interacts with αβ₃. Inhibition by LYP8 of TSP-mediated M₃Da spreading and melanoma–platelet interaction could be the result, as previously suggested in this study, of steric or conformational changes of TSP induced by LYP8 (Dixit et al., 1986). Our data do not exclude the possibility that, in addition to αβ₃, TSP may bind to other receptors (i.e., heparan sulphate proteoglycans) in view of the lack of effect of LYP8 in inhibiting cell attachment of M₃Da to TSP-coated wells (Roberts, 1988; Asch et al., 1991).

Our results are consistent with recent studies of Castle et al. (1991) showing the importance of TSP by transfected human squamous carcinoma cells with the TSP cDNA antisense expression vector and decreasing the tumorigenic phenotype of these cells. Moreover, the CSVTCG TSP peptide and its analogue were shown to block cell adhesion, platelet aggregation and tumour cell metastasis (Tuszynski et al., 1992; Byck and McGregor, 1992). Tumour metastasis is a complex sequence of events in which malignant cells enter the bloodstream and interact with various host cells, including vascular endothelial cells, before plasticating and forming secondary tumours (Nicolson, 1988). Reagents such as MAbs against TSP and peptides derived from TSP could not only be used as a new therapeutic approach in the control of tumour metastasis of malignant melanoma, but may also contribute to the elucidation on the role of TSP in cancer biology.

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