Embryonic chicken fibroblast collagen binding proteins: distribution, role in substratum adhesion, and relationship to integrins

ROY C. OGLE*, A. JEANNETTE POTTS, MARSHALL YACOE and CHARLES D. LITTLE†

Department of Anatomy and Cell Biology, School of Medicine, Box 439, Medical Centre, University of Virginia, Charlottesville, VA 22903, USA

*Present address: Department of Anatomy and Cell Biology, Medical University of South Carolina, Charleston, SC 29425, USA
†Author for correspondence

Summary

Collagen binding proteins (CBP) are hydrophobic, cell surface polypeptides, isolated by collagen affinity chromatography. Antibodies to CBPs inhibit the attachment of embryonic chicken heart fibroblasts to native type I collagen fibrils in a dose-dependent manner. The CBP antibodies also induce rounding and detachment of cells adherent to a planar substratum. This process of antibody-mediated substratum detachment resulted in a clustering of CBP and cell-associated extracellular matrix at the cell surface, and the rearrangement of filamentous actin. Other functional studies showed that cells grown within a three-dimensional gel of type I collagen cannot be immunostained at the cell surface with CBP antibodies. However, treatment

Introduction

Cells produce, modify and adhere to their surrounding extracellular matrix (Hay, 1984). These cell surface ECM interactions are dynamic processes mediated by membrane-associated receptors that recognize specific features of extracellular macromolecules. One group of these adhesion molecules, the integrins, has been identified on a variety of cell types from a wide range of invertebrate and vertebrate organisms (Hynes, 1987). The integrins have conserved their structure and function during evolution (Buck and Horwitz, 1987; DeSimone and Hynes, 1988). Most members of the integrin superfamily of receptors, thus far described, bind to RGD peptide sequences in the ECM ligand (Ruoslathi and Pierschbacher, 1986).

The integrins, present on a variety of avian cell types, attach to fibronectin and laminin through an extracellular ligand-binding domain (Tamkun et al. 1986). The receptor spans the plasma membrane and interacts with microfilament bundles through a cytoplasmic domain (Chen et al. 1985; Burridge, 1986; Buck and Horwitz, 1987). Thus, these receptors appear to form a transmembrane linkage between the matrix surrounding the cell and its cytoskeletal network. Regulation of cell morphology and motile activity, processes known to require the participation of the cytoskeleton and the extracellular matrix, occur through the integrated activity of matrix receptors.

Characterization of these matrix receptors has involved the production of antibodies that perturb adhesion to a particular matrix molecule (Greve and Gottlieb, 1982; Neff et al. 1982), or more recently, affinity chromatography with a specific matrix molecule (Pytela et al. 1985; Rubin et al. 1986). The latter technique is useful for identifying receptors with high affinities for their ligand(s). The human fibronectin receptor present on osteosarcoma cells was isolated by affinity chromatography on a fragment of fibronectin containing the binding domain (Pytela et al. 1985). This receptor has, therefore, a relatively high affinity for its ligand. Another high-affinity human fibronectin receptor, VLA-5, is present in placental tissue (Hemler et al. 1987). Also present in placental tissue is a low-affinity fibronectin receptor, VLA-3. Identifying and isolating low-affinity ECM receptors is a challenge, since conventional affinity chromatography is not effective.

We previously reported that several embryonic chicken glycoproteins bind to collagen with what appears to be moderate affinity (Ogle and Little, 1989). These glycoproteins, termed collagen binding proteins (CBPs), were bound to a collagen I affinity matrix under low ionic
strength conditions, and then eluted with moderately high (0.5M) levels of NaCl. The CBPs possess the characteristics of receptors in that: (1) they are present at the cell surface as evidenced by radioiodination; (2) they are integral membrane proteins, which resist extraction by urea and salt washes; (3) they are hydrophobic; and (4) they contain N-linked carbohydrate (Ogle and Little, 1989). The CBPs appear to be distinct, individual polypeptides, on the basis of differences in apparent electrophoretic mobility on SDS-PAGE and unique antibody staining patterns on cells and tissues. Two of the CBPs are potential members of the integrin receptor family, on the basis of sensitivity to elution from a collagen column with the synthetic peptide sequence GRGDS, and electrophoretic mobilities on reduced and non-reduced SDS-polyacrylamide gels (Ogle and Little, 1989).

In this study we examine the biological properties of the CBPs through the use of polyclonal antisera raised against the CBP complex. In addition, we investigate the potential relationship of CBP 1 and CBP 2 to avian integrins by the use of antibodies that define the chicken fibronectin receptor.

Materials and methods

Materials

For biosynthetically labeling CBP, [35S]methionine was purchased from Amersham, as was iodine-125 for radioiodination (16.7 mCi/125 mCi). Collagen binding proteins were purified from CHF plasma membranes as described (Ogle and Little, 1989).

Production and characterization of antibodies

Polyclonal antibodies to avian CBPs were raised in a rabbit using approximately 20 μg of CBP (purified by two cycles of collagen affinity chromatography) emulsified in Freund’s complete adjuvant. The immunogen was injected into the surgically exposed popliteal lymph nodes of each hind leg (Boudie et al. 1966). The rabbit was boosted with 100 μg CBP in incomplete adjuvant by subcutaneous injection at 4-week intervals and bled every 3 weeks after the second injection. Antibodies were affinity purified on CBP (Little and Chen, 1982) or on CBP transferred to nitrocellulose (described below).

Inhibition of attachment assay

A supply of embryonic chicken (11-day) heart fibroblasts in the third passage were frozen in liquid nitrogen; all experiments came from this pool of cells. Collagen substrata were prepared as described (Chen and Little, 1982) in 35 mm plastic dishes (Falcon Plastics). The CBP antiserum was dialyzed against DMEM (without serum) and filter sterilized (0.22 μm). The day prior to an attachment assay, cells were thawed and plated in 100 mm dishes with DMEM plus 10% fetal bovine serum.
The next day the subconfluent cultures were trypsinized and suspended in DMEM (with and without FBS, similar results were obtained), and the number of cells was determined with a hemocytometer. Cells were adjusted to approximately \(3.0 \times 10^5\) cells ml\(^{-1}\) and 0.5 ml samples were placed in collagen-coated dishes along with the appropriate dilution of CBP antibodies, in a 0.5 ml sample. The total volume per 35 mm dish was 1.0 ml. The number of cells in several identical samples were counted with the aid of a hemocytometer to confirm the concentration of the cell suspension. Cultures were incubated in a humidified CO\(_2\) incubator for 2 h, then the dish was flooded with 3 % paraformaldehyde and held for photography. The number of cells in the medium, after the 2 h incubation, was determined and then expressed as the percentage of the total number of cells originally placed in culture. In one set of experiments the CBP was passed over a gelatin-Sepharose column to remove rabbit fibronectin; no differences in the experimental results were noted. Control experiments with pre-immune serum were conducted exactly as the CBP anti-serum experiments.

**Immunoblots**

The antigen samples were subjected to electrophoresis in 7.5 % SDS-polyacrylamide gels, then transferred to nitrocellulose paper (Schleicher and Schull) or nylon membranes (Immobilon) with an American Bionetics graphite electrode using a 6-aminohexanoic acid buffer system. Membranes were blocked with primary antibodies for 1 h at room temperature, washed and incubated with horseradish peroxidase-conjugated goat anti-rabbit IgG for 1 h then developed with l-chloro-4-napthol as a chromogen.

**Other methods**

Indirect immunofluorescence staining, microscopy and photography were carried out as described (Little and Chen, 1982). Electrophoresis was performed as described (Laemmli, 1970) and proteins were visualized with Coomassie Blue R-250 or by autoradiography of dried gels exposed to Kodak XAR-5 film employing Dupont Cronex lightning plus intensifying screens.

**Results**

**Production and characterization of antibodies to CBP**

Affinity-purified CBP were used to immunize a rabbit. Immunoprecipitation reactions with antisera from early bleeds showed activity against bands designated CBP 1, 2 and 4 of \(M_e\) 120,000, 100,000, and 47,000, respectively (Fig. 1). After several boosts the antiserum recognized CBP 3, \(M_e\) 63,000. The migration positions of CBP 1 and 2, previously shown to be sensitive to GRGDS elution, are similar to the molecular weights of avian integrin subunits (Knudsen et al. 1985; Akiyama et al. 1986; Tamkum et al. 1986; Marcantonio and Hynes, 1988).

**Inhibition of adhesion to collagen fibrils by anti-CBP**

Chicken heart fibroblasts (CHF) were assayed for their ability to attach and spread on a substratum of native collagen I fibrils in the presence of increasing amounts of antiserum to CBP. Fig. 2A shows CHF were inhibited from attaching to collagen substrata in a dose-dependent fashion by the CBP antisera. The panel of micrographs (Fig. 2B) shows the progressive inhibition of cellular spreading on native collagen fibrils that occurs concomitantly with the decrease in adhesive activity.

**Fig. 1.** Antiserum to CBP was used to precipitate reactive bands from a detergent lysate of cells metabolically labeled with \(^{35}\)S-methionine. A prominent band at approximately 120K is observed along with bands at 100K and 45K (lane a). A control immunoprecipitation reaction with non-immune serum (lane b). The arrowheads indicate the position of molecular weight markers (from the top: 205, 116, 97, 66, 45K) separated on this 7.5 % gel.

**Fig. 2. A.** Inhibition of cellular attachment to collagen I fibrils. CHF were incubated with increasing amounts of CBP antiserum (○●○●), which had been dialyzed against DMEM, or dialyzed pre-immune serum (▲). Attachment was inhibited in a dose-dependent fashion by antibodies against CPBs. B. Phase-contrast images of cell after 2 h of culture in the dilution of CBP antiserum indicated. Native collagen fibrils are visible. Cells show less spreading in response to increasing amounts of CBP antibodies. Control (pre-immune) cultures resembled the upper micrograph.

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Fig. 3. Progressive release of embryonic chicken heart fibroblasts (CHF) from substratum by antibody treatment. Confluent CHF were incubated in the presence of 5% antiserum to total CBP and photographed at 15-min intervals (as shown). Cell–substratum rather than cell–cell adhesion was perturbed most by this treatment. Monospecific antibodies to CBPs 1 and 2 gave identical results (see Fig. 4B). The process could be accelerated by a higher concentration of antiserum.

Release of adherent cells by antibody treatment
Confluent cultures of CHF were treated with medium containing 5% heat-inactivated CBP antiserum or pre-immune serum. Over the course of 1 h cells were observed to become detached from the substratum, to round up and to be released into the medium (Fig. 3). Most cells were released from the plastic or glass substratum prior to detaching from neighbouring cells, suggesting that cell–substrate, rather than cell–cell adhesions were perturbed. Purified monospecific antibodies to CBP 1 and CBP 2 had similar effects on substratum adhesion (see Fig. 4B). Pre-immune antiserum had no discernable effect (data not shown).

Cell cultures undergoing this process of retraction and release were examined for the distribution of CBP, collagen and microfilament bundles. Fig. 4A shows cells examined for the co-distribution of CBP and f-actin. The embryonic fibroblasts were fixed at 15 min and 60 min after the addition of CBP antiserum. The immunostaining is intensified at the cell periphery where retraction is occurring. These same areas show a focal concentration of f-actin. Similar cultures were incubated with affinity-purified antibodies to CBP 1 and CBP 2 (Fig. 4B). These specific antibodies caused the retraction and release of most cells, although some cells appeared to resist the antibody effect. In experiments not shown, immunoreactive clumps of collagen I and fibronectin were also drawn into these retraction processes at the cell periphery after the addition of anti-CBP. Eventually, the retraction and rounding process resulted in most of the cells being released completely from the substratum. If these 'released' cells were briefly treated with trypsin (0.1% for 15 s), they readily became re-attached to a new substratum (data not shown).

Localization of collagen binding proteins on CHF grown in three-dimensional collagen gels
CHF grown in native three-dimensional type I collagen gels were examined by immunofluorescence. Repeated attempts to immunolabel cells, grown within three-dimensional collagen gels with CBP antibodies, ended in failure (Fig. 5B). Positive immunological control experiments with anti-tubulin showed that IgG could diffuse through the gel and not only reach the cell surface, but also enter the cytoplasmic compartment and label microtubules (data not shown; see Tomasek and Hay, 1984). Failure to label the cell surface with CBP antibodies, therefore, was not due to limited movement of IgG through the collagen gel. Treatment of the cultures with highly purified bacterial collagenase (free of non-collagenous proteolytic activity) hydrolyzed the collagen and rendered CBPs accessible to antibody (Fig. 5D). Furthermore, immunoreactive sites unmasked by the collagenase were distributed evenly over the entire surface of the CHF; including the fine filopodial projections characteristic of fibroblastic cells cultured in three-dimensional collagen gels (Fig. 5C,D).

Distribution of the collagen binding proteins on CHF
Monospecific affinity antibodies were isolated on individual bands of CBP 1–4 bound to nitrocellulose (see Materials and methods). Immunolocalized studies on embryonic chicken heart fibroblasts revealed that cells labeled independently with antibodies to either CBP 1 or CBP 2 displayed indistinguishable fluorescence staining patterns. Moreover, no differences in the CBP 1 or CBP 2 staining patterns were observed after the cells were made permeable with detergent. Fig. 6A and B shows cells stained with anti-CBP 1, before and after detergent treatment. Cells labeled with antibodies to CBP 3 had sparse clumps of fluorescence; permeabilization resulted in bright perinuclear staining (not shown). Immunostaining for CBP 4 showed an unremarkable diffuse staining (not shown).

Comparison of CBP 1 with integrin
Previous work (Ogle and Little, 1989) suggested biochemical similarities between integrins and CBPs 1 and 2 (e.g. their sensitivity to elution from a collagen-Sepharose column with a synthetic GRGDS peptide). Here we prepared whole lysates from surface radioiodinated cells for analysis by immunoprecipitation in order to compare anti-CBP activity with 140K, a polyclonal antiserum.
Fig. 4. A. Cells treated with antibodies to CBP as described for Fig. 3 were fixed at 15 min and 60 min following addition of antibodies to cultures. The distribution of actin microfilament bundles (ACT) and CBPs were examined by fluorescence microscopy, and compared with untreated cells (UNT). As CBP antibodies cause cells to retract from the substratum, f-actin in microfilament bundles is drawn into clusters. These clusters of actin-rich material co-distributed in CBPs at sites that remained unattached to the substratum. Bar, 50 μm. B. Cells treated with affinity-purified antibodies to CBP 1 and CBP 2 (approximately 200 μg ml⁻¹) were incubated for 30 min, then fixed for DIC microscopy. Most cells retracted or were released; however, some cells resisted the antibodies, which may be due to the heterogeneous nature of early-passage CHF.

raised against the β subunit of the avian fibronectin receptor. Fig. 7A shows that polypeptides immuno-reactive with anti-CBP (lanes 1, 2) or anti-140K (lanes 3, 4) display remarkably similar electrophoretic mobilities when analyzed by SDS–PAGE; both without disulfide reduction (lanes 1, 3), and with disulfide reduction (lanes 2, 4).

The possibility that CBPs 1 and 2 are immunologically related to the integrin family of ECM receptors was tested in two ways (Fig. 7B). First, affinity-purified collagen binding proteins were separated on SDS–polyacrylamide gels, transferred to nylon membranes, and then probed with either CBP antibodies (lane 1) or 140K antibodies (lane 2). Second, immunoaffinity-purified avian integrins were prepared using a CSAT antibody column. This material was also separated by SDS–PAGE, electroblotted, and probed with the CBP antibodies or 140K antibodies (lanes 3 and 4). Collagen binding proteins

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show an immunoreactive polypeptide band with an apparent molecular weight of 100,000 when probed with either anti-CBP (lane 1) or anti-140K (lane 2). Similarly, CSAT-purified avian integrins show immunoreactive bands at approximately 100,000 when reacted with anti-CBP (lane 3) or anti-140K (lane 4). Antibodies to CBP detected higher molecular weight complexes when reacted with collagen binding proteins (lane 1), and 140K antibodies detected high molecular weight complexes within the CSAT-purified integrin fraction (lane 4). This higher molecular weight material is possibly β dimers or integrin complexes similar to those reported by Buck and co-workers (1986).

Discussion

Role of collagen binding proteins in substratum adhesion

Antibodies to CBP inhibited the attachment and spreading of cells on collagen fibrils and caused the detachment of cells from untreated tissue-culture surfaces. Monospecific antibodies to CBP 1 and CBP 2 also caused retraction of cells from untreated planar substrata. In view of the apparent crossreactivity of the CBP antibodies with CSAT-purified avian integrins, and vice versa, it is likely that we prepared a 'generic' anti-integrin β subunit antibody. Therefore, when anti-CBP is applied to embryonic chicken heart fibroblasts (CHF) grown on planar surfaces, all integrin-mediated extracellular matrix (ECM) adhesions are probably perturbed. In contrast, anti-CBP antibodies also inhibit cell attachment to a homogeneous scaffold of native collagen fibrils. This more specific inhibition activity suggests the presence of collagen integrins. Furthermore, it should be noted that the original immunogen was isolated on native collagen affinity columns (shown to be free of fibronectin).

The cell shape changes that occur concomitantly with antibody treatment show a perturbation of the actin-containing microfilament bundles. These observations also suggests the possibility that some of the collagen binding proteins may function similarly to integrins, by mediating concerted interactions between the cytoskeleton and the extracellular matrix (Chen et al. 1985; Horwitz et al. 1986; Burridge, 1986).

The recent observation of Fath et al. (1989) on the vitronectin integrin and fibronectin integrin shows that cells can redistribute their repertoire of receptors in response to a different ECM substratum. When cells were grown on planar substrata their integrins were localized to focal adhesions (Chen et al. 1985; Horwitz, 1986).
integrins are probed with anti-140K (lane 4), higher at approximately 100K. Collagen binding proteins show by immunoaffinity (CSAT) chromatography were separated Arrowheads indicate the position of molecular weight markers

Fig. 7. A. Lysates prepared from cell-surface radioiodinated surface, including the finest protrusions observable by fibrillar collagen gels, chicken embryo fibroblasts showed adhesions on planar substrata. that mature fibrillar collagen is not present in CHF focal

Reliability of CBP to integrins
During the biochemical characterization of the CBP complex, it became evident that CBP 1 and CBP 2 shared properties with integrin subunits. These include: susceptibility to RGD peptide elution and molecular weight shift upon reduction when analyzed by SDS–PAGE (Ogle and Little, 1989).

Identification of the first avian integrin (see Buck and Horwitz, 1987) led to the determination of the protein structure of this receptor family (Tamkum et al. 1986). Human analogues to the avian integrin complex, have under these more physiological conditions of growth, in a hydrated collagen lattice, embryonic fibroblasts do not form focal adhesions (Heath and Hedlund, 1984). Previous immunostaining studies, using frozen sections of chicken embryos, showed that CBP was concentrated on the surface of cells adjacent to extracellular collagen I (Ogle et al. 1986). Attempts to extend the colocalization studies to whole mounts of cells within collagen gels in vitro were unsuccessful. Chicken embryo fibroblasts embedded in collagen fibrils did not stain with anti-CBP. The best explanation for this observation is that the antibodies were unable to bind to their antigenic sites due to steric hindrance by receptor-bound collagen fibrils.

After removal of collagen by collagenase the CHF surface showed strong immunoreactivity. In view of the fact that the cells shown in Fig. 5C and D were previously surrounded by a collagen lattice, it is noteworthy that the ‘unmasked’ CBPs are distributed in a granular pattern. The filopodia and protrusions of the ‘liberated’ cells show distinct punctate staining, with very dense immunolabelling in the perinuclear region. This latter distribution agrees with the findings of Heath and Hedlund (1984) on their studies of CHF locomotion in fibrillar collagen gels.

The total lack of detectable surface immunolabeling of cells within three-dimensional collagen gels, even at very high antibody concentration, leads to some interesting considerations. For instance, this is not the experimental result one would anticipate in view of the generic anti-integrin nature of the CBP antibody. This lack of surface staining suggests that when CHF are grown in a three-dimensional ECM, the various integrins may be complexed in heterofunctional adhesion particles. We have no direct proof of this; however, if collagen receptors were distributed independently of other ECM receptors, one would not expect the presence of the collagen fibrils to hinder antibody access to all cell surface antigens. On the other hand, if most or all ECM receptors (integrins) were clustered in multifunctional attachment domains, then the saturating presence of a single ECM ligand (e.g. collagen fibrils) might then hinder immunolabeling of all the receptors in that particle.

As a positive control for these experiments, we showed that IgG could diffuse through the collagen gel and decorate intracellular tubulin, thus eliminating the possibility that antibodies did not have access to the CHF embedded within the hydrated collagen lattices.
provided information on the function of these receptors and their relationships to each other (Hemler et al. 1987).

Our biochemical and functional analyses of avian collagen binding proteins suggests that these cell surface molecules are involved in mediating cell adhesion to collagen. We have compared affinity-purified CBP with the integrins through immunological cross-reactivity studies. Whole CBP antisera identified the β subunit of collagen binding proteins suggests that these cell surface molecules are involved in mediating cell adhesion to collagen. We have compared affinity-purified CBP with the integrins through immunological cross-reactivity studies. Whole CBP antisera identified the β subunit of the avian integrin complex, identified a CBP at 100,000, the molecular weight of the avian β subunit. Taken together, the biochemical and cell biological studies suggest that CBP 1 and CBP 2 are related to the avian integrin complex.

At the present time we can only speculate on the ligand specificity of CBPs 1 and 2. One possibility is that either polypeptide may be a subunit of a promiscuous integrin that recognizes multiple ECM components including collagen. Alternatively, or perhaps in addition, the molecules may represent a distinct collagen integrin that shares a β subunit with other integrins. Either of these possibilities would result in immunological cross-reactivity.

The platelet collagen receptor in mammals is an example of a well-characterized collagen integrin (Hemler et al. 1988; Kunicki et al. 1988; Takada et al. 1988). Our work suggests that, like humans, birds may have a collagen integrin. The exact nature of the αβ subunits responsible for mediating cell adhesion to collagen in avian embryos is under investigation.

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