Specific inhibition of GPI-anchored protein function by homing and self-association of specific GPI anchors

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The functional specificity conferred by glycophosphatidylinositol (GPI) anchors on certain membrane proteins may arise from their occupancy of specific membrane microdomains. We show that membrane proteins with noninteractive external domains attached to the same carcinoembryonic antigen (CEA) GPI anchor, but not to unrelated neural cell adhesion molecule GPI anchors, colocalize on the cell surface, confirming that the GPI anchor mediates association with specific membrane domains and providing a mechanism for specific signaling. This directed targeting was exploited by coexpressing an external domain-defective protein with a functional protein, both with the CEA GPI anchor. The result was a complete loss of signaling capabilities (through integrin–ECM interaction) and cellular effect (differentiation blockage) of the active protein, which involved an alteration of the size of the microdomains occupied by the active protein. This work clarifies how the GPI anchor can determine protein function, while offering a novel method for its modulation.

Introduction

Many cell surface proteins are attached to the membrane by a glycoprophosphatidylinositol (GPI) anchor, which consists of a conserved central structure (Low, 1989) with variable carbohydrate and lipid peripheral components (Homans et al., 1988). GPI anchors can determine protein functional specificity, just as switching a transmembrane (TM) domain for a GPI anchor can result in novel function caused by association with new signaling elements located in a shared membrane microdomain (Shenoy-Scaria et al., 1992, 1993). Membrane rafts, originally defined by their insolubility in cold, nonionic detergents such as Triton X-100 (Simons and Ikonen, 1997), are small, heterogeneous aggregations of cholesterol and sphingolipids on the cell surface (Pralle et al., 2000; Pike, 2004) that concentrate GPI-anchored proteins, but also contain other proteins. Although the existence of membrane rafts in vivo has been questioned (Munro, 2003), recent studies using a variety of methods have provided evidence for raftlike membrane microdomains (Friedrichson and Kurzhailia, 1998; Varma and Mayor, 1998; Pralle et al., 2000; Dietrich et al., 2002; Gaus et al., 2003; Sharma et al., 2004). Such microdomains may act as signaling scaffolds, determining the identity of a subset of signaling elements, as proteomic analyses have found a high concentration of such proteins in purified rafts (von Haller et al., 2001; Foster et al., 2003), with GPI-anchored proteins involved in activating this signaling (Robinson, 1997; Solomon et al., 1998). The existence of heterogeneous raft populations has been inferred from studies showing that different GPI-anchored proteins exist in separate rafts (Madore et al., 1999; Wang et al., 2002; Li et al., 2003). External rafts with different proteins may each have a defined set of associated cytoplasmic proteins, whereby aggregation of GPI-anchored proteins by external domain self-binding or by multivalent ligand binding could cluster specific rafts, resulting in downstream signaling (Harris and Siu, 2002).

Carcinoembryonic antigen (CEA), and the closely related CEACAM6, are GPI-anchored, cell surface glycoproteins that block cellular differentiation (Eidelman et al., 1993) and inhibit the apoptotic process of anoikis (Ordonez et al., 2000; Duxbury et al., 2004b), effects that appear to be caused by the activation of specific integrins (Duxbury et al., 2004a; Ordonez et al., 2006). CEA is up-regulated in many human malignancies (Hinoda et al., 1991; Iliantiz et al., 1997), implying a similar role in human cancer, whereas the TM-anchored CEACAM1 (CC1) may act as a tumor suppressor (Kunath et al., 1995; Luo et al., 1997).
Most CEA family members mediate intercellular adhesion by antiparallel self-binding (Zhou et al., 1993), which, together with parallel binding on the same cell surface (Taheri et al., 2003), may result in clustering of rafts containing CEA (Benchimol et al., 1989). Deletion of the last two thirds of the CEA N-terminal domain (∆NCEA) abrogates its adhesive ability, which leads to a loss of differentiation-blocking activity (Eidelman et al., 1993). The method of membrane anchorage determines CEA family member activity, as genetically fusing the GPI anchor of CEA to CC1's external domain creates a differentiation-blocking molecule, whereas a chimera consisting of the external domain of CEA attached to the TM domain of CC1 does not block differentiation (Screaton et al., 2000).

Deletion of the last two thirds of the CEA N-terminal domain (∆NCEA) abrogates its adhesive ability, which leads to a loss of differentiation-blocking activity (Eidelman et al., 1993). The fact that GPI-anchored neural cell adhesion molecule (NCAM) does not block differentiation, but can be converted to a differentiation-blocking molecule, suggests that the CEA GPI anchor harbors the specificity for the differentiation-blocking function and that the external domains merely function to cluster the molecules, and thus, the associated rafts (Screaton et al., 2000).

Based on the aforementioned model, it should be possible to inhibit the biological functions of CEA (and, by implication, that of any GPI-anchored molecule whose function is regulated by a similar mechanism) by interfering with clustering. This has been achieved for CEA by mutating regions in its N-terminal external domain responsible for self-binding or by the addition of peptides or monovalent mAbs that target these regions (Taheri et al., 2003). We test a second strategy which exploits the specificity of the CEA GPI anchor; if “shank-defective” or “shankless” CEA GPI anchors that were incapable of self-association and clustering were introduced, they could occupy the same rafts as CEA, and thus, possibly interfere with its clustering. We show that nonfunctional ∆NCEA inhabits the same membrane microdomains as NCB, as both have the same GPI anchor, and is capable of completely inhibiting NCB’s CEA-like differentiation-blocking activity.

**Results**

**Design and construction of experiments**

To test the hypothesis that the functional specificity of GPI anchors could be exploited to specifically inhibit the activity of GPI-anchored proteins, cells expressing a functional GPI-anchored protein were supertransfected with a shank-defective molecule with the same GPI anchor, and assessed for effects on function. The former functioning molecule was NCB, which has NCAM self-binding external domains linked to the CEA GPI anchor (Screaton et al., 2000); the defective molecule was ∆NCEA, which has the same GPI anchor, but external domains that are defective in self-binding (Fig. 1 A; Eidelman et al., 1993). Because ∆NCEA cannot bind to the external NCAM domain of NCB (Zhou et al., 1990), this combination allowed a study focused on the potential interaction between their GPI anchors.

∆NCEA and NCB exist in close proximity

CEA and NCAM appear to exist in separate membrane regions, potentially explaining their opposite biological effects (Screaton et al., 2000). If the GPI anchor alone determines cell surface

**Figure 1.** Surface expression of CEA and NCAM proteins on L6 myoblasts. (A) Schematic representation of proteins used in this study, with their ability to mediate intercellular adhesion and to inhibit myogenic differentiation shown (as + or −). (B) FACS profiles, after staining with mAbs D14 (anti-CEA) or 123C3 (anti-NCAM) and FITC-conjugated secondary antibody, demonstrate cell surface levels of indicated proteins with fluorescence means in parentheses. In coexpressing cells, D14 staining is shown by shaded profiles.
localization, then molecules with the same GPI anchor should exist in close proximity, whereas those with different anchors should not. Thus, NCAM and NCB would be expected to have different cell surface distributions, with NCB showing a distribution similar to that of CEA. To test this hypothesis, we examined whether ∆NCEA existed in close proximity on the cell surface to NCB, but not to NCAM, using confocal microscopy to examine the cell surface localization of these proteins. The relative surface distribution of NCAM and NCB compared with ∆NCEA was determined after indirect immunofluorescent staining. ∆NCEA showed substantial, although incomplete, colocalization with NCB, whereas ∆NCEA and NCAM showed essentially no colocalization (Fig. 2 A). Because the incubations were performed at room temperature, the antibodies used for detection may have caused partial clustering of the proteins. This, however, should not affect the heterophilic association in question, as clustering of rafts containing both proteins should not change the final amount of colocalization seen. Indeed, fixing the cells before antibody incubation to avoid clustering resulted in very similar patterns of colocalization to what is shown in Fig. 2 A (not depicted). This therefore suggests that the GPI anchor of CEA is sufficient to determine cell surface localization of a protein.

To verify these results, L6 cotransfectants were treated at 4°C (to limit protein diffusion) with the chemical cross-linker DTSSP. Nonreducing Western blots demonstrated similar cross-linking patterns for NCB and NCAM, consisting of dimers, trimers, and higher molecular weight complexes, both alone and in the presence of ∆NCEA (Fig. 2 B). To determine the cellular distribution of ∆NCEA relative to NCAM and NCB, immunoprecipitation (IP) studies of extracts from cross-linked cells expressing similar amounts of these proteins were performed. The cross-linking approach was undertaken, rather than using detergent lysis because of the potential effects of detergents on membrane raft structure. IP of extracts from untreated cells did not result in any coIP (Fig. 2 C), confirming the expected antibody specificity. However, IP with an anti-CEA mAb of extracts of DTSSP-treated cotransfectants resulted in the coIP of a considerable amount of NCB, but, importantly, not of NCAM (Fig. 2 D). Similarly, IP with an anti-NCAM mAb of extracts of cross-linked cotransfectants showed coIP of ∆NCEA only in the case of NCB, but not of NCAM (Fig. 2 D). The low proportion of coimmunoprecipitated protein can likely be explained by the lack of interaction between the external NCAM and CEA protein domains, the requirement for close (<12 Å) apposition to
be cross-linked, the presence of large levels of monomeric proteins even after cross-linking (Fig. 2 B), and the incomplete colocalization seen by confocal microscopy (Fig. 2 A). These results demonstrate that proteins with GPI anchors of the same type can exist in close proximity, providing a rationalization for specific interference with protein function.

\( \Delta \text{NCEA restores differentiation to NCB-expressing cells} \)

Having demonstrated specific colocalization of \( \Delta \text{NCEA} \) and NCB on the cell surface, the effect of this defective protein on NCB’s ability to block differentiation was examined. NCB levels in the \( \Delta \text{NCEA} \) coexpressing L6 transfectants were actually higher than those in NCB-only transfectants, thus, validating comparisons between NCB alone and in the presence of \( \Delta \text{NCEA} \) (Fig. 1 B; FACS means of 124 vs. 63, respectively). NCB completely blocked differentiation, whereas coexpression of \( \Delta \text{NCEA} \) with NCB resulted in an almost complete restoration of differentiation, with a fusion index of 78% of that seen for \( \Delta \text{NCEA} \) alone (Fig. 3 A and Table I; \( P < 0.0001 \)). As a control, coexpressing NCAM had no effect on the differentiation block imposed by CEA (Fig. 3 A), despite the differentiation-enhancing effects of NCAM (Dickson et al., 1990). To confirm this result, up-regulation of myosin, a biochemical differentiation marker, was examined. \( \Delta \text{NCEA} \) induced myosin production in two independent populations of NCB-expressing cells, as shown by Western blot, whereas NCB alone showed no myosin expression (Fig. 3 B), confirming the previous results. Because of the length of the differentiation assay (10 d total), it was possible that a loss of NCB expression caused the differentiation restoration in these cotransfectants. However, no decrease in NCB levels was seen in differentiated \( \Delta \text{NCEA} \) + NCB cultures, as Western blots showed higher expression levels in both cell populations than the NCB-alone transfectants for
which differentiation was blocked (Fig. 3). Thus, ∆NCEA expression interfered markedly with the differentiation-blocking function of NCB, presumably via their common feature, the GPI anchor.

**Effects of CEA-like proteins on binding to ECM**

∆NCEA releases NCB’s block of differentiation, suggesting that it is interfering with downstream signaling by NCB.

CEA signaling has been found to involve activation of the integrin α5β1 in rat myoblast and human colonic cell lines (Ordonez et al., 2006) and the integrin αvβ3 in neuronal cells (unpublished data). We assessed NCB signaling by incubating single-cell suspensions, prepared from exponential cultures, with plates coated with the ECM components fibronectin (Fn), vitronectin (Vn), and collagen I. Either CEA or NCB expression increased binding to both Fn and Vn, relative to L6 parental cells (Fig. 4 A; P < 0.004). Cells expressing ∆NCEA + NCB (and ∆NCEA alone) showed no such increase, demonstrating a complete loss, in the presence of ∆NCEA, of the NCB-mediated increase in ECM binding (P < 0.0001). As a control, no difference in binding to collagen I was seen between any of these cell lines (Fig. 4 A). In addition, LR-73 (LR) transfectants were tested for binding to Fn, and these cells showed a similar loss of NCB-mediated effects upon coexpression of ∆NCEA (Fig. 4 B; P < 0.01). The total cell levels of α5 and β1 integrins in the L6 transfectants were assessed by Western blot (Fig. 4 C), and cell surface levels were assessed by FACS (α5 only; not depicted) and showed only minor differences between transfectants, confirming that the ability of cells to adhere to Fn, rather than changes in integrin surface expression level, was the source of the observed difference (Ordonez et al., 2006; unpublished data).

![Image of Figure 4](image_url)

Table I. Differentiation of L6 transfectants

| Cell line            | Fusiona |
|----------------------|---------|
| L6 parental         | 71      |
| L6 (CEA)            | 0       |
| L6 (∆NCEA)          | 64      |
| L6 (NCB)            | 0       |
| L6 (∆NCEA + NCB)    | 50      |
| L6 (CEA + NCAM)     | 0       |

aFusion index was measured as the number of cells containing three or more nuclei divided by total nuclei in the field and expressed as a percentage. Values are the mean of three different experiments, with three fields scored from each experiment.
Thus, the ability of \( \Delta \text{NCEA} \) to interfere with the NCB-mediated differentiation blockade is correlated with interference of enhanced integrin–ECM interaction promoted by NCB.

**NCB raft association is unaltered by the presence of \( \Delta \text{NCEA} \)**

Signaling by GPI-anchored proteins requires intact membrane rafts (Stulnig et al., 1997). One possible mechanism for the effects of \( \Delta \text{NCEA} \) on NCB functional properties could be by expulsion of NCB from rafts. When this possibility was examined, however, NCB remained primarily insoluble in cold Triton X-100 after coexpression of \( \Delta \text{NCEA} \) in either L6 or LR cells (Fig. 5, A and B, respectively). Complete cellular lysis was demonstrated by the fact that the integrin \( \alpha_5 \) chain, an integral membrane protein, was localized in the soluble fractions. As confirmation, isopycnic sucrose gradient ultracentrifugation, where raft-associated proteins migrate to the lower density regions of the gradient, was performed on cold Brj-98 lysates of L6 transfectants. Again, no obvious difference was noted between NCB alone and NCB coexpressed with \( \Delta \text{NCEA} \) (Fig. 5, C and D), as almost all of the NCB was present in the low-density fractions in both cases. Under these conditions, the \( \alpha_5 \) integrin chain showed partial raft association for both transfectants, demonstrating that an alteration of \( \alpha_5 \) localization was not responsible for the lack of NCB function. The distribution of \( \Delta \text{NCEA} \) was also found to be essentially the same as that of NCB, as expected for two GPI-anchored proteins. As controls, the \( \alpha_2 \) integrin chain was found solely in higher density fractions, whereas the raft lipid GM1 was entirely in the low-density fractions. Thus, NCB retained membrane raft association in the presence of \( \Delta \text{NCEA} \), so that this could not explain the loss of NCB function.

**Effect of expression of \( \Delta \text{NCEA} \) on NCB-mediated intercellular adhesion**

The proteins of the CEA family mediate intercellular adhesion, as does NCAM (Soroka et al., 2003), by external domain self-binding. Such self-binding is required for the differentiation-inhibitory activity of CEA, presumably to affect raft clustering (Eidelman et al., 1993; Taheri et al., 2003). If \( \Delta \text{NCEA} \) interfered with NCB clustering, one might predict a reduction in the ability of NCB to mediate intercellular adhesion. NCAM was used as a control, as it inhabits different rafts from CEA, presumably to affect raft clustering. Triton X-100 solubility assays of L6 (A) and LR (B) cells at 4°C showed no alteration in membrane raft association of NCB ± \( \Delta \text{NCEA} \) transfectants with the majority of the protein in the insoluble (P), i.e., raft-associated, fraction. As a control for efficiency of lysis, the integrin \( \alpha_5 \) chain showed essentially no insolubility, with all protein found in the soluble (S) fraction for both L6 and LR parental and transfectant cell lines. Isopycnic sucrose density gradient ultracentrifugation showed similar membrane raft association, as seen by flotation at lower density fractions of NCB in both L6 (NCB; C) and L6 (\( \Delta \text{NCEA} \) + NCB; D) transfectants. Integrin \( \alpha_5 \) had partial raft association, whereas integrin \( \alpha_2 \) was not found in lower density fractions. The raft marker GM1 was found exclusively in lower density fractions, whereas \( \Delta \text{NCEA} \) showed similar distribution to NCB.

**Figure 5.** NCB membrane raft association is unaltered in the presence of \( \Delta \text{NCEA} \).

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**\( \Delta \text{NCEA} \) alters the size of NCB-containing rafts**

One mechanism whereby \( \Delta \text{NCEA} \) could interfere with NCB clustering is by altering the structure of the rafts it is associated with. Therefore, the size of the rafts that NCB occupied was approximated by lysing the cells under conditions identical to those used for isopycnic separation on sucrose density anchored proteins is believed to involve the formation of large, zipperlike structures through the aggregation of multiple proteins and rafts, creating stabilized platforms (Harris and Siu, 2002). The ability of \( \Delta \text{NCEA} \) to interfere with NCB-mediated adhesion is thus consistent with models invoking interference with NCB clustering.
gradients and separating the lysate by velocity sedimentation through a uniform 12.5% sucrose solution. Under these conditions, NCB was almost entirely raft associated (Fig. 5); therefore, this technique should provide a measure of the size of the rafts inhabited by these proteins. The fractions, which were collected from the top (fraction #1), were assessed by Western blot for protein localization, using equal volumes of each fraction. NCB was found to be shifted to fractions farther from the top when \( \Delta \)NCEA was coexpressed, indicating that it was present in larger complexes under these conditions (Fig. 7 A). The distribution of NCAM, on the other hand, was found to be similar whether \( \Delta \)NCEA was present or not (Fig. 7 B), demonstrating that the size of the NCAM complexes was not altered in the presence of \( \Delta \)NCEA. The distribution of \( \Delta \)NCEA was very similar to that of NCB in shifting to larger complexes when coexpressed with NCB, while remaining in smaller complexes when coexpressed with NCAM (unpublished data). A significant (P < 0.05) difference in NCB distribution (Fig. 7 C) relative to NCAM distribution (Fig. 7 D) upon coexpression with \( \Delta \)NCEA was demonstrated by densitometric analysis of three independent experiments. This suggests that the presence of \( \Delta \)NCEA specifically alters the rafts containing NCB. To confirm that this was a raft-specific effect, cells were pretreated with methyl-\( \beta \)-cyclodextrin (M\( \beta \)CD) to sequester cholesterol and disrupt raft structure. Initially, sucrose gradient ultracentrifugation was performed on lysates of these treated cells, to confirm the disruption of the rafts. The distribution of NCB demonstrated that this treatment partially disrupted the rafts, as a portion of the NCB was now present in higher density fractions (Fig. 7 E; compare to Fig. 5). When these samples were tested for the size of the complexes that NCB was localized to, it was found that NCB, both alone and coexpressed with \( \Delta \)NCEA, remained in the first few fractions after velocity sedimentation (Fig. 7 F). Thus, treatment with M\( \beta \)CD

Figure 6. \( \Delta \)NCEA interferes with NCB-mediated intercellular adhesion. (A) LR surface expression (fluorescence means are shown in parenthesis) of NCAM-like proteins (unshaded profiles) and \( \Delta \)NCEA (shaded profiles) after staining with mAbs 123C3 and D14, respectively. (B) Adhesion assays of LR (NCB) and LR (\( \Delta \)NCEA + NCB; right) and LR (NCAM) and LR (\( \Delta \)NCEA + NCAM; left). Note the 20% decrease (*, P < 0.001) in NCB-mediated adhesion in the presence of \( \Delta \)NCEA, which is not seen in NCAM versus NCAM + \( \Delta \)NCEA transfectants. Values represent the mean ± the SEM for four independent experiments. (C) Aggregate assays of cells from B, showing that the size of multicellular aggregates created by NCB was significantly reduced in the presence of \( \Delta \)NCEA. The number of cells per multicellular aggregate was determined, after 1 h of incubation, in four separate fields per experiment. Data represent the mean ± SEM of three independent experiments (*, P < 0.0001).
abrogated the difference seen for NCB complex size after ∆NCEA coexpression, confirming that the difference seen was a raft-mediated effect. Although cellular lysis with detergents at low temperatures can affect raft structure (Shogomori and Brown, 2003), the fact that the ∆NCEA-dependent sedimentation difference is seen for NCB, but not for NCAM transfec-
tants, suggests that it represents a valid increase in raft size. This would indicate a dilution of the NCB concentration in membrane rafts, as an increase in the size of a raft containing the same number of NCB molecules would cause a relative concentration decrease. This would thus reduce the incidence of cis-interactions between the proteins, which are necessary for clustering, explaining the decrease in intercellular adhe-
sion (Fig. 6 B), and the loss of biological function (Figs. 3 A and 4 A).

Antibody cross-linking restores function of NCB inhibited by ΔNCEA

As the mechanism of inhibition of NCB function by ΔNCEA appears to involve interference with clustering, NCB func-
tion should be restored by artificial clustering with antibodies. Antibody cross-linking of cell surface proteins induces signaling events, including restoring the defective differentiation-blocking function of ΔNCEA (Taheri et al., 2003) through integrin activation manifested by increased cellular binding of Fn (unpublished data). ΔNCEA and NCAM coexpressed with ΔNCEA both appear to be nonfunctional because of defects in protein clustering, so clustering of NCB with antibodies should have a similar effect to what has previously been seen for ΔNCEA. To test if NCB retained the potential to modulate ECM binding, in spite of the deactivating effects of coexpressed ΔNCEA, cells in monolayer culture were treated with mAbs directed against the NCAM external domains of NCB, along with secondary antibodies to enhance clustering, and binding of soluble Fn was measured. Several mAbs were used, including J22, which binds to internal CEA domains and, as such, remains ca-
pable of clustering ΔNCEA; D13, which is a control mAb that has an epitope in the region deleted from ΔNCEA; and 123C3, which binds to the NCAM external domains of both NCAM 125 and NCB. As expected, cross-linking ΔNCEA with J22, but not with D13, resulted in a significant increase in bound Fn (Fig. 8). Similarly, cross-linking of NCB, alone and in the presence of ΔNCEA, increased bound Fn (Fig. 8). Cross-linking NCAM, which does not normally modulate integrins, with the NCAM-specific antibody did not lead to an increase in bound Fn levels, demonstrating the specificity of this effect. The lack
of difference in Fn binding between NCB and parental cells, unlike that seen in Fig. 4, in monolayer culture is likely caused by the intact ECM surrounding the cells in this assay. This would provide the ligands for integrins that have previously been activated, so that these integrins would not bind to the Fn added to the culture medium. Thus, NCB, in the presence of ∆NCEA, remained capable of altering Fn interaction after antibody cross-linking, which is consistent with the hypothesis that a defect in NCB clustering is created upon introduction of ∆NCEA.

Discussion

The GPI anchor of CEA contains specific information determining protein function (Screaton et al., 2000) when attached to an adhesive extracellular protein domain. We investigated the hypothesis that interference with the GPI anchor could cause specific inhibition of function using a defective CEA GPI anchor–bearing molecule, ∆NCEA, coexpressed with NCB, a chimera of the NCAM external domain attached to the CEA GPI anchor. Coexpression of ∆NCEA with NCB resulted in a complete loss of the latter’s biological function.

Determination of specific protein function by GPI anchors

TM- and GPI-anchored NCAM isoforms exhibit different effects on myoblast fusion (Peck and Walsh, 1993). Genetically attaching the CEA GPI anchor to the external domains of CC1 and NCAM, two adhesive cell surface molecules that do not inhibit cellular differentiation, results in chimeras that block differentiation, demonstrating functional anchor specificity (Screaton et al., 2000). Anchors may determine associated signaling proteins, as GPI-anchored CD59 can signal through kinases after having been exogenously administered to cells (van den Berg et al., 1995). Attaching this anchor to GFP also recreates the same signaling events upon antibody cross-linking (Hiscox et al., 2002). Membrane rafts concentrate a wide variety of different proteins on both sides of the plasma membrane; thus, GPI anchor-mediated targeting of a protein to rafts containing specific signaling molecules, in conjunction with clustering through external domain interactions, would explain how the CEA anchor can determine specific protein function.

Existence of distinct membrane raft domains

To have functionally distinct membrane rafts, these domains should contain different molecules; indeed, the segregation of gangliosides GM1 and GD3 into different domains has been observed (Gomez-Mouton et al., 2001, 2004; Vyas et al., 2001). This may help determine signaling specificity, as a close and specific association occurs between some gangliosides and certain signaling molecules (Kasahara et al., 1997; Iwabuchi et al., 1998). Similarly, GPI anchors are heterogeneous, as differences in both hydrophobic and hydrophilic GPI anchor regions have been documented (Ferguson et al., 1988; Armesto et al., 1996). These raft and anchor variations may be sufficient to create GPI anchor–specific membrane domains with different protein repertoires. In support of this idea, human folate receptor and placental AP do not exist in close proximity (Wang et al., 2002), whereas Thy-1 and prion protein associate with different domains (Madore et al., 1999) that show major differences in lipid composition (Brugger et al., 2004). CEA and NCAM do not colocalize on the cell surface (Screaton et al., 2000), which was confirmed by IP and colocalization results presented in this study (Fig. 2). Replacing the GPI anchor signal sequence of NCAM for that of CEA, however, produces a chimera (NCB) partially located in close proximity to ∆NCEA (Fig. 2). This incomplete colocalization is likely a result of the small size of membrane rafts before activation, as it has been suggested that under resting conditions GPI-anchored proteins are primarily found as monomers, with a proportion found in small clusters (Sharma et al., 2004). NCB affects cell–Fn interaction similarly to CEA (Fig. 4 A), but unlike NCAM, suggesting that this altered localization affects signaling. Different Ras isoforms inhabit separate cytoplasmic leaflet compartments (Prior et al., 2003), showing that differences in protein distribution on the cell surface may be mirrored on the cell interior. We suggest that the C-terminal primary amino acid sequence can direct the addition of a specific type of GPI anchor, which determines both membrane localization and function through associated signaling elements.
Role of membrane raft clustering in signaling

Intense study of membrane raft dynamics has been undertaken in T cells, where these domains have a key role in signaling upon activation of the T cell receptor. Costimulation of T cells by CD28 is caused by raft redistribution, which alters internal tyrosine phosphorylation patterns (Viola et al., 1999), whereas antibody cross-linking of raft lipids replicates the membrane re-arrangements that occur upon receptor activation (Janes et al., 1999). CEA mediates intercellular adhesion by antiparallel binding (Zhou et al., 1993) and differentiation inhibition by both antiparallel and parallel binding (Taheri et al., 2003), both of which may cause clustering of the rafts inhabited by CEA. The discovery of CEA mutants that retain intercellular adhesive ability but do not block differentiation indicates the key role, in conjunction with antiparallel adhesion, of parallel interactions in CEA's biological function (Taheri et al., 2003). NCAM-mediated adhesion involves intermolecular cis- and trans-interactions (Rao et al., 1992; Soroka et al., 2003), so at a threshold cell surface density, CEA and NCAM could create large cell surface raft aggregates leading to signaling (Harris and Siu, 2002). CEACAM7, a weakly adhesive member of the CEA family, poorly activates integrin α5β1; however, attaching its anchor to the stronger adhesive external domain of CC1 results in increased integrin activation (unpublished data). Previous work has demonstrated that treatment of CEA-expressing cells with fragment antigen-binding antibody fragments (Fabs) to reduce protein clustering is sufficient to release the CEA block of differentiation (Taheri et al., 2003). Similarly, reducing initial cis-clustering, through the addition of molecules with defective nonadhesive extracellular domains attached to the same GPI anchor, could also interfere with effective protein and raft clustering, causing substantial effects on downstream signaling. ∆NCEA coexpression resulted in a reproducible decrease in intercellular adhesive strength of NCB-expressing cells (Fig. 6 B) and an observed loss of cell–Fn interaction (Fig. 4 A). ∆NCEA expression increased the size of the membrane rafts containing NCB (Fig. 7 A), which would dilute the NCB molecules within the raft compartment. Thus, a larger raft with the same number of NCB molecules would lead to less efficient cis-clustering, decreasing the resulting adhesive strength. Consistent with this model, mimicking the clustering seen upon adhesion by antibody-mediated cross-linking of the NCB molecules restored its effects on integrin–Fn interaction (Fig. 8). Therefore, it would appear that treatment with Fabs or nonfunctional GPI anchors leads to similar effects on clustering and protein activity.

Modulation of raft lipid content

The function of membrane proteins can be altered by modulating membrane lipid content. Depletion of cellular cholesterol inhibits signaling from membrane raft domains (Incardona and Eaton, 2000), whereas administration of exogenous gangliosides displaces GPI-anchored proteins from membrane rafts (Simons et al., 1999). Treatment of T cells with polyunsaturated fatty acids interferes with tyrosine kinase activation and calcium release upon stimulation (Stuhl et al., 1998), which appears similar to this study, where signaling by NCB is inhibited via its GPI anchor. The key difference lies in the GPI-anchor specificity because exogenous administration of lipids likely causes a global alteration in raft structure, whereas GPI anchors contain information that targets specific subdomains. Thus, ∆NCEA is not found in close proximity to NCAM, but changing the anchor of NCAM for that of CEA alters the localization of this protein, such that it now colocalizes with ∆NCEA (Fig. 2). This colocalization explains how ∆NCEA, which shows no effect on differentiation, can restore differentiation to NCB-expressing cells, whereas NCAM, which accelerates differentiation, does not restore fusion to CEA-expressing cells (Fig. 3 A). It is intriguing to note that ∆NCEA expression has such a significant effect on NCB despite the fact that the proteins are found on the cell surface at similar levels. This suggests that once a threshold number of proteins have been inhibited, the biological function is completely lost, as any remaining functional proteins may be incapable of clustering sufficiently to lead to effective signaling activation.

We therefore suggest that the GPI anchor plays a key role in protein function by directing localization to a specific subset of membrane rafts, which determines the associated signaling molecules. Exploiting this biological specificity by competition with functionally specific GPI anchors attached to nonfunctional external protein domains results in a complete loss of biological activity. This therefore confirms the key biological role of the GPI anchor, and suggests a novel method for the manipulation GPI-anchored proteins.

Materials and methods

Constructs and antibodies

∆NCEA is a CEA deletion mutant that has the last 75 amino acids of the N domain deleted, such that it is no longer biologically active (Zhou et al., 1993). The NCAM splice variant used in this study, p125, is a human GPI-anchored NCAM isoform containing the muscle-specific domain (Barton et al., 1988). NCB is a chimera of the NCAM p125 external domain genetically fused to the CEA GPI anchor signal sequence (Screaton et al., 2000). The mAbs J22 and D14 bind to internal CEA domains (Zhou et al., 1993), whereas the epitope of D13 is in the portion of the CEA N domain that is deleted in ∆NCEA, and rabbit polyclonal anti-CEA binds to all CEA external protein domains. The mAb 123C3 (Santa Cruz Biotechnology, Inc.) recognizes human NCAM, whereas antibodies H-293, H-104, and M-106 (Santa Cruz Biotechnology, Inc.) recognize the α2, α5, and β1 integrins, respectively. The mAb 47A (De Giovanni et al., 1993) binds to myosin. C20 is a goat polyclonal anti-Fn antibody (Santa Cruz Biotechnology, Inc.).

Cell culture and differentiation assay

Cells were grown attached to tissue culture plastic surfaces (Nunc), as previously described (Screaton et al., 2000). In brief, CHO-derived LR-73 fibroblasts were grown in αMEM with 10% FBS. Rat L6 myoblasts were grown in DME containing 10% FBS (GM), and were subcultured before reaching confluency to avoid selecting for nonfusing variants. Cell concentrations were determined using a particle counter (Beckman Coulter). For myoblast differentiation, 10^6 L6 cells/cm^2 were seeded in 60-mm dishes. After 3 d, the media was switched to DME with 2% horse serum (DM), 4–7 d later, cultures were assessed for differentiation by hematoxylin (Sigma-Aldrich) staining and microscopic examination (Screaton et al., 1997), or by lysing and assessing myosin levels by Western blotting.

Transfections

100-mm dishes were seeded with 2 or 4 × 10^5 cells/plat for LR or L6, respectively. 24 h later, cells were cotransfected by calcium phosphate coprecipitation with 5 µg of cDNA, 0.5 µg of pSV2neo, and 10 µg of carrier DNA isolated from LR-73 cells. Double transfections were performed in the same manner, either by cotransfecting both cDNAs at once (for LR cells)
with pSV2(neo) or with 0.5 μg of pBabe(puro) to supertransfect L6 transfectants; transfectants were isolated by selection with 400–600 μg/ml neomycin (G418; Invitrogen) or 1 μg/ml puromycin (Sigma-Aldrich). After 10–14 d, resistant clones were pooled and sorted for high expression by FACS using mAbs J22 or 123C3. After several pools, two independent transfections of L6 cells were performed to ensure no clonal variation occurred, with identical assay results. Note that the L6 (NCB) cells were pooled colonies resistant to both G418 and puromycin, and that although data from L6 and LR-73 parental cells is shown, no difference between these cells and pooled G418-resistant clones transfected with the pSV2(neo) alone has been noted (Ordonez et al., 2006).

FACS analysis

Cells were collected with PBS-citrate containing 4 mM EDTA (PBSCe) for NCAM-expressing transfectants (because of the sensitivity of the NCAM external domain to trypsinization; Screaton et al., 2000) or 0.063% trypsin in PBS-citrate for CEA transfectants. 2.5 × 10^5 cells were resuspended in ice-cold PBS with 2% FBS (PBSF). Cells were incubated for 30 min with mAb at a dilution of 1:50–100, washed with PBSF, and incubated with FITC-conjugated goat anti-mouse antibody (Jackson ImmunoResearch Laboratories) diluted 1:100. After an additional 30 min, cells were pelleted, resuspended in PBSF, and analyzed using a FACScan instrument (Becton Dickinson).

Binding to purified ECM components

Cells were seeded at 10^5 cells/cm^2 on day 0. On day 2, subconfluent cultures were collected with PBSCe, resuspended in GM, and incubated at 37°C for 30 min. Cells were washed with serum-free media and resuspended in serum-free media at 4 × 10^5 cells/ml. 100 μl/ml of this suspension was added to wells from a 96-well plate coated with Fn, Vn, or collagen I (CHEMICON International, Inc.) and incubated for 1 h at 37°C. Wells were washed with PBS containing Mg^2+, and adherent cells were stained with crystal violet. Wells were washed again with PBS, and the bound stain was solubilized with 0.05 M NaOH-P04, pH 4.5, plus 25% ethanol. Staining was quantified with a microplate reader (Bio-Tek Instruments) at 570 nm. Statistical significance was determined using a t test (http://www.physics.csbsju.edu/stats/t-test bulk_form.html).

Immunoblotting

Proteins in cellular lysates were resolved by SDS-PAGE and transferred electrophoretically to a 0.45-μm PVDF membrane (Millipore). Immunoblotting was performed as previously described (Screaton et al., 2000), with antibody binding detected using ECL Plus reagent (GE Healthcare).

Triton X-100 solubility and isopycnic sucrose gradient ultracentrifugation

Triton X-100 solubility was determined as previously described (Screaton et al., 2000). In brief, subconfluent cell cultures were rinsed with PBS, collected with PBSCe, and rendered single-cell suspensions by passing through a 27-gauge needle. 10^5 cells/ml were resuspended in ice-cold lysis buffer containing 1% Triton X-100 and the protease inhibitors aprotonin (Roche), leupeptin (Roche), and PMSF (Sigma-Aldrich). Lysates were syringed with a 27-gauge needle to obtain single-cell suspensions, and allowed to aggregate at 37°C with stirring at 100 rpm using a small magnetic stirring bar. Aliquots were removed at the indicated times, and the cells were counted with a hemocytometer to determine the percentage of single cells. For the aggregate assay, cells were prepared as for adhesion assays, but the number of cells present in each of ~50 multicellular aggregates was scored after 1 h in suspension.

Soluble Fn-binding assay

On day 0, 10^5 cells/well were seeded in 96-well plate. 2 d later, cells were washed with PBSF and incubated for 30 min at 37°C in DME with 50 μl of 10 μg/ml human Fn (BD Biosciences), along with, where indicated, 5 μg/ml primary mAb and 30 μg/ml donkey anti–mouse secondary antibody (Jackson ImmunoResearch Laboratories) to further cross-link CEAs to NCAM constructs. Cells were then washed three times with PBSF, and fixed with 4% paraformaldehyde. Bound Fn was determined by incubation with anti-Fn antibody, C20, at a dilution of 1:100 in 3% BSA (Sigma-Aldrich) in PBSF for 90 min at RT, having blocked nonspecific binding by incubation for 1 h at RT with PBSB. Cells were washed with PBSB, and incubated with HRP-conjugated rabbit anti–goat secondary antibody (Jackson) at a dilution of 1:2,500 in PBSB for 1 h. After incubation with a hydrogen peroxide solution containing ABTS (Sigma-Aldrich), bound Fn was determined with a microplate reader at 405 nm, with a reference wavelength of 490 nm.

We thank Jacynthe Labbé for technical assistance with confocal microscopy. This work was supported by the Cancer Research Society and by a research contract to C.P. Stanners from NovoThera Biotechnologies, Inc. T.B. Nicholson was the recipient of a Natural Sciences and Engineering Research Council of Canada scholarship.

Submitted: 1 May 2006
Accepted: 16 October 2006

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