Title
CD36 repression activates a multicellular stromal program shared by high mammographic density and tumor tissues

Permalink
https://escholarship.org/uc/item/3qb8b3pp

Journal
Cancer Discovery, 2(9)

ISSN
2159-8274

Authors
DeFilippis, RA
Chang, H
Dumont, N
et al.

Publication Date
2012-09-01

DOI
10.1158/2159-8290.CD-12-0107

Peer reviewed
CD36 Repression Activates a Multicellular Stromal Program Shared by High Mammographic Density and Tumor Tissues
ABSTRACT
Although high mammographic density is considered one of the strongest risk factors for invasive breast cancer, the genes involved in modulating this clinical feature are unknown. Tissues of high mammographic density share key histologic features with stromal components within malignant lesions of tumor tissues, specifically low adipocyte and high extracellular matrix (ECM) content. We show that CD36, a transmembrane receptor that coordinates multiple protumorigenic phenotypes, including adipocyte differentiation, angiogenesis, cell–ECM interactions, and immune signaling, is greatly repressed in multiple cell types of disease-free stroma associated with high mammographic density and tumor stroma. Using both in vitro and in vivo assays, we show that CD36 repression is necessary and sufficient to recapitulate the above-mentioned phenotypes observed in high mammographic density and tumor tissues. Consistent with a functional role for this coordinated program in tumorigenesis, we observe that clinical outcomes are strongly associated with CD36 expression.

SIGNIFICANCE: CD36 simultaneously controls adipocyte content and matrix accumulation and is coordinately repressed in multiple cell types within tumor and high mammographic density stroma, suggesting that activation of this stromal program is an early event in tumorigenesis. Levels of CD36 and extent of mammographic density are both modifiable factors that provide potential for intervention.

INTRODUCTION
Many tissues are comprised of an epithelium associated with a stroma that is composed of extracellular matrix (ECM) proteins, adipocytes, fibroblasts, endothelial cells, neuronal cells, and immune cells. Fibroblasts, a major stromal cell type, produce many components of the ECM, as well as proteases that remodel it (1). In addition, fibroblast signaling is a critical determinant of epithelial and stromal cell fate during development and differentiation (2), tissue homeostasis, and wound healing (1).

Stromal changes associated with malignant lesions are heterogeneous and range from tumor suppressing to tumor promoting. Tumors, subtyped by marker expression in epithelial cells, can be further categorized by their stromal signature, which dictates good or poor outcome (3). Fibroblasts within the stroma of malignant lesions, called carcinoma-associated fibroblasts (CAF), differ from their counterparts in disease-free tissues (4). CAFs can stimulate tumor progression of initiated nontumorigenic epithelial cells, both in vitro and in vivo, whereas normal fibroblasts cannot (5). CAFs increase proliferation and decrease apoptosis in adjacent epithelial cells and promote angiogenesis by recruiting endothelial progenitor cells into tumors (5, 6). All other stromal components, including immune cells and endothelial cells, also participate in malignant progression (4).

The stroma within and immediately adjacent to a malignant lesion may exhibit a range of histologic alterations, called desmoplasmia. The alterations range from a predominantly cellular stroma, containing fibroblasts, vascular cells, and immune cells with little ECM, to a dense tissue with a minimum of cells and a maximum content of matrix (4, 7). In addition, tumor stroma exhibits a reduction in the size and number of adipocytes (8). The most odious stromal changes include extreme ECM deposition and remodeling, along with aberrant vasculature and fibroblast and immune cell infiltration. Classically, participation of the stroma has been viewed as a reactive process, in which signals from malignant epithelial cells recruit and stimulate these stromal components.

Interestingly, some desmoplastic features are seen in cancer-free tissues of women at high risk for breast cancer, namely in the context of wound healing, radiation response, pregnancy-associated involution, and high mammographic density (4, 8–10). Mammographic density is of particular interest, as almost one third of breast cancers are thought to be attributable to phenotypes associated with high breast density, and mammographic density is a strong risk factor with high prevalence (11, 12). Mammographic density is determined by the relative amounts of radiolucent material (fat) and radiodense material (epithelial cells, fibroblasts, and connective tissue) within the breast on a mammogram, either of which may occupy anywhere between 0% and 100% of the gland. Radiodense areas exhibit several histologic characteristics of...
stoma associated with malignant epithelial cells, specifically low adipocyte content and high ECM and stromal cell content (9, 10). Epidemiologic studies suggest that women with high mammographic density have a 4- to 6-fold increase in risk for invasive breast cancer compared with women with low mammographic density (11, 13–15).

Studies comparing pairs of monozygotic and dizygotic twins attributed 53% to 67% of the variation in mammographic density to heritable factors, with the remaining portion modulated by environmental or physiologic factors (16–21). For example, postmenopausal hormone therapy increases mammographic density in some women, whereas tamoxifen treatment decreases mammographic density in both pre- and postmenopausal women and subsequently modulates breast cancer risk (19, 21). This modulation by exogenous factors provides an exciting opportunity for intervention and prevention.

Boyden and colleagues examined the molecular composition of low and high mammographic density tissues (9, 10) and reported higher levels of insulin-like growth factor-I and tissue inhibitor of metalloproteinase 3 in high mammographic density tissue. Recent studies found additional changes including differences in estrogen and prostaglandin metabolism and TGF-β signaling (22–24). Although these studies begin to define the molecular constituents of mammographic density that may explain the basis of high mammographic density and its association with increased breast cancer risk, they are correlative and thus cannot show that these genes or proteins mechanistically modulate phenotypes of mammographic density.

The observation of multiple desmoplastic phenotypes in tissues of high mammographic density, in the absence of a tumor, strongly suggests that these stromal phenotypes are not simply part of a reactive response to existing tumor cells but also represent a program involving multiple stromal components that may create a proactive milieu for the emergence or progression of cancer. Therefore, we hypothesized that the phenotypes observed in both high mammographic density and tumor stroma may be controlled by a common molecular program and thus their stromal components would share phenotypic, molecular, and functional characteristics.

Here we show that the level of expression of a single molecule, CD36, is necessary and sufficient to simultaneously control adipocyte content and matrix accumulation, 2 phenotypes shared by desmoplasia and high mammographic density. CD36, a widely expressed transmembrane receptor, modulates cell type- and ligand-specific phenotypes, including adipocyte differentiation, angiogenesis, apoptosis, TGF-β activation, cell-ECM interactions, and immune signaling (25). We found that CD36 expression was negligible in multiple cell types of tumor stroma compared with surrounding histologically disease-free tissue. Strikingly, high mammographic density tissues, devoid of any malignancy, also showed reduced CD36 levels in multiple cell types compared with normal mammographic density tissues. The low level of CD36 expression shared by high-risk (but cancer-free) and malignant tissues suggests that it constitutes a causal and very early event in generating the distinctive characteristics of tumor stroma.

RESULTS

Desmoplasia and Tissues with High Mammographic Density Share Histologic Characteristics

The appearance of stroma associated with malignant epithelial cells is strikingly different from the appearance of most stroma associated with disease-free ductal tissue. Increases in ECM, endothelial and immune components, and the absence of adipocytes are obvious upon inspection (Fig. 1). Strikingly, similar stromal alterations are observed histologically in disease-free tissue of women with high mammographic density as compared with women with low mammographic density, specifically low adipocyte and high ECM content (Fig. 1). We examined fibroblasts from tissue of high and low mammographic density, as well as from tumor tissue, for phenotypic similarities that may underlie their shared histologic appearance.

HDAFs and LDAFs Recapitulate Phenotypes of High and Low Mammographic Density Tissues, Respectively, In Vitro

To determine whether fibroblasts obtained from low and high mammographic density tissues and propagated in vitro could recapitulate the 2 prominent in vivo phenotypic differences described above, low-density associated fibroblasts (LDAF) and high-density associated fibroblasts (HDAF) were purified from disease-free breast biopsies from women with 25% to 50% mammographic density and women with more than 75% mammographic density, respectively (Supplementary Table S1). LDAFs and HDAFs were placed under proliferative or adipocyte differentiation conditions for 6 days.

![Figure 1](https://cancerdiscovery.aacrjournals.org/content/12/9/828/F1.large.jpg)

**Figure 1.** Desmoplasia and tissues with high mammographic density share histologic characteristics. Representative bright field images (original magnification X5) of paraffin sections stained with hematoxylin and eosin (nuclei: blue, ECM: pink, adipocytes: white). Left: desmoplastic tissue and histologically normal adjacent tissue from a patient with invasive ductal carcinoma. Right: cancer-free tissue from one woman with low mammographic density and one woman with high mammographic density. MD, mammographic density.
CD36 Modulates Phenotypes of Breast Density and Desmoplasia

before assessing their ability to accumulate fat by Oil Red O staining (Fig. 2A). Fat accumulation was quantitated on a cell-by-cell basis as previously described (Supplementary Fig. S1; ref. 26). This analysis showed that, although both LDAFs and HDAFs accumulated fat under differentiation conditions (10.0-fold and 5.6-fold, respectively, $P < 10^{-8}$), HDAFs accumulated significantly less fat (3.1-fold, $P < 10^{-8}$) than LDAFs (Fig. 2B). This decrease in the average amount of fat accumulation per cell was due to a decrease in both the percentage of cells that contain fat and the amount of fat per cell (Supplementary Fig. S2).

To determine whether LDAFs and HDAFs differed in their adipocyte differentiation capacity, rather than simply fat storage, the expression levels of CD36 and leptin (LEP; previously reported to be upregulated during adipocyte differentiation; ref. 27) were measured by quantitative PCR (QPCR) in cells grown under proliferative or adipocyte differentiation conditions for 2 weeks. Both LDAFs and HDAFs could undergo adipocyte differentiation; however, LDAFs showed higher levels of expression of CD36 and LEP compared with HDAFs (CD36: 2.7-fold, $P = 0.03$; LEP: 6.8-fold, $P = 0.05$; Fig. 2C). These data showed that differences in fat accumulation between LDAFs and HDAFs are due to differences in their adipocyte differentiation capabilities.

Having shown that cultured LDAFs and HDAFs recapitulated their respective in vivo adipocyte phenotypes, we examined whether they could also recapitulate their in vivo matrix accumulation phenotypes in a short-term assay. LDAF and HDAF cultures were probed after 5 days of propagation for accumulation of matrix proteins known to be increased in desmoplasia: collagen 1A1 (COL1A1), fibronectin (FN1), osteopontin (OPN), and tenasin C (TNC; ref. 1; Supplementary Fig. S3). Cell-by-cell quantitative immunofluorescence analysis (Supplementary Fig. S1) revealed heterogeneity within the cultures with an overall small but statistically significant greater amount of COL1A1, FN1, and OPN (each 1.1-fold, $P < 10^{-8}$) in HDAFs compared with LDAFs (Fig. 2D). However, HDAFs accumulated a statistically significant lesser amount of TNC (1.1-fold, $P < 10^{-8}$) than LDAFs, suggesting that there is not a global increase in all matrix protein accumulation (Fig. 2D). One could speculate that this modest increase in matrix accumulation over a prolonged period of time (years), in contrast to our 5-day assay, could account for the dramatic differences observed in vivo.

CAFs Recapitulate Phenotypes of Desmoplasia In Vitro

CAFs have been shown to promote ECM deposition in vitro and in vivo, but little is known about their ability to modulate adipocyte differentiation (1). To this end, human mammary fibroblasts purified from invasive cancer tissues (CAFs) and from reduction mammaplasty tissues (RMF; Supplementary Table S2) were assayed as above for fat accumulation (Fig. 2E, left panel). Cell-by-cell quantitation of Oil Red O staining (Supplementary Fig. S1) showed that, although both RMFs and CAFs accumulated fat under differentiation conditions (27-fold and 8-fold, respectively, $P < 10^{-8}$), CAFs were deficient in fat accumulation compared with RMFs (5.8-fold, $P < 10^{-8}$; Fig. 2E, right panel).

CD36 Expression Is Decreased in Fibroblasts from High Mammographic Density and Tumor Tissues

To gain insight into the molecular basis for these phenotypes, gene expression profiling was used to identify genes differentially expressed between LDAFs and HDAFs (GEO GSE38506). CD36 showed the largest difference, its expression being downregulated 3.7- to 4.3-fold in HDAFs relative to LDAFs (Supplementary Table S3). To validate these microarray results, CD36 transcript and protein levels were measured in LDAFs and HDAFs by QPCR and Western blot analysis. Both CD36 gene expression (8.6-fold, $P = 0.002$) and protein levels were decreased in HDAFs relative to LDAFs (Fig. 3A and B).

Given the striking downregulation of CD36 in fibroblasts of high mammographic density, we examined CD36 expression in fibroblasts isolated from malignant tissue and in microarrays of tumor tissue. CD36 expression, measured by QPCR, was decreased 4.3-fold ($P = 0.001$) in CAFs relative to RMFs (Fig. 3C). Similarly, analysis of 6 independent public gene expression datasets (Supplementary Table S4) consistently showed a statistically significant decrease in CD36 expression in invasive ductal carcinoma (IDC) tissues compared with normal tissues (3.6- to 12.3-fold). In light of the striking repression of CD36 in fibroblasts from high mammographic density and tumor tissues, we examined the role of CD36 in modulating the 2 prominent in vivo phenotypes shared by these tissues: decreased adipocyte content and increased matrix accumulation (4, 8-10).

CD36 Expression Is Necessary and Sufficient to Modulate Adipocyte Differentiation and Matrix Accumulation In Vitro

Little is known about the basis of decreased adipocyte content in high mammographic density and desmoplasic tissues. However, induction of CD36 expression is necessary for proper adipocyte differentiation in preadipocytes (28) and serves as a marker for terminal differentiation (27). To determine whether CD36 expression could modulate adipocyte content, we genetically modulated CD36 levels in RMFs selected for their intermediate baseline expression level of CD36. To determine whether CD36 expression was necessary for fat accumulation, CD36 expression was repressed in RMFs by transduction of lentiviral particles expressing short hairpin RNA to CD36 (shCD36; Supplementary Fig. S4). Cell-by-cell quantitation of Oil Red O staining (Supplementary Fig. S1) showed that although both control (shLuc) and shCD36 cells accumulated fat under differentiation conditions (8.9-fold and 3.1-fold, respectively, $P < 10^{-8}$), control cells accumulated significantly more fat than shCD36 cells (6.1-fold, $P < 10^{-8}$; Fig. 4A). These data showed that CD36 is necessary for fat accumulation. To determine whether CD36 expression was sufficient for fat accumulation, CD36 expression was increased in RMFs by transduction of lentiviral particles expressing CD36 (CD36 OE; Supplementary Fig. S4). Cell-by-cell quantitation of Oil Red O staining (Supplementary Fig. S1) showed that, although both control (vector) and CD36 OE cells accumulated fat under differentiation conditions (2.2-fold and 4.7-fold, respectively, $P < 10^{-8}$), CD36 OE...
Figure 2. Cultured fibroblasts recapitulate multiple phenotypes associated with their respective tissues of origin. A, three LDAFs and 3 HDAFs were placed under proliferative (ctl) or adipocyte differentiation (+PJ2) conditions for 6 days and assessed for fat accumulation by Oil Red O staining. Representative bright field images (original magnification ×10) of the fibroblasts under differentiation conditions. B, average and SEM of Oil Red O staining per cell. C, four LDAFs and 4 HDAFs were placed under proliferative (ctl) or adipocyte differentiation conditions (+PJ2) for 2 weeks and assessed for adipocyte differentiation by measuring the expression of genes upregulated in adipogenesis. Average and SEM of CD36 and LEP expression by QPCR (n = 4 for each of 4 LDAFs and 4 HDAFs). GUSB, beta-D-glucuronidase. D, representative fluorescent images (original magnification ×10) highlighting differential ECM protein deposition in 6 LDAFs and 6 HDAFs grown for 5 days and assessed for accumulation of matrix proteins COL1A1, FN1, OPN, and TNC. E, left: representative bright field images (original magnification ×10) of 1 RMF and 2 CAFs under proliferative (ctl) or adipocyte differentiation (+PJ2) conditions for 7 days and assessed for adipocyte formation by Oil Red O staining. Right: average and SEM of Oil Red O staining per cell.
CD36 Modulates Phenotypes of Breast Density and Desmoplasia

**Figure 3.** CD36 expression is decreased in HDAFs and CAFs relative to LDAFs and RMFs, respectively. 

- **A**, average and SEM of CD36 expression measured by QPCR (n = 3 for each of the 7 LDAFs and 7 HDAFs).
- **B**, Western blot analyses for CD36 (top), where the arrows indicate CD36 glycosylated state and actin (bottom). Positive control for CD36 expression: RMF under proliferative (ctl) or adipocyte differentiation (+PJ2) conditions for 7 days.
- **C**, average and SEM of CD36 expression measured by QPCR (n = 3 for each of the 8 RMFs and 8 CAFs).

Cells accumulated significantly more fat than control cells (3.3-fold, P < 10^{-8}; Fig. 4B). These data showed that CD36 is sufficient for fat accumulation. Therefore, the decrease in CD36 expression in HDAFs and CAFs could account for their reduced ability to accumulate fat compared with LDAFs and RMFs, respectively.

To determine whether CD36 expression could also modulate matrix accumulation, the fibroblasts with genetically modulated levels of CD36 described above were probed for the accumulation of matrix proteins known to be increased in HDAFs: COL1A1, FN1, and OPN (Fig. 4C). Cell-by-cell quantitative immunofluorescence analysis

**Figure 4.** CD36 expression is necessary and sufficient to modulate adipocyte differentiation and matrix accumulation in vitro. 

- **A and B**, left: Representative bright field images (original magnification x10) of shLuc, shCD36, vector, or CD36 OE cells, under proliferative (ctl) or adipocyte differentiation (+PJ2) conditions for 7 days and assessed for adipocyte formation by Oil Red O staining. 
- **A and B**, right: Average and SEM of Oil Red O staining per cell.
- **C**, representative fluorescent images (original magnification x10) of shLuc, shCD36, vector, and CD36 OE cells grown for 5 days and assessed for accumulation of matrix proteins COL1A1, FN1, and OPN.
- **D**, average and SEM of staining per cell expressed as signal x10^4.

Published OnlineFirst July 9, 2012; DOI: 10.1158/2159-8290.CD-12-0107
accumulation and increase in matrix accumulation observed in tumor tissues can result in the coordinated decrease in fat CD36 expression seen in high mammographic density and tumors can result in the coordinated decrease in fat accumulation and increase in matrix accumulation observed in vivo.

Cd36 Knockout Mice Phenocopy Human High Mammographic Density and Desmoplastic Tissues

To determine whether the modulation of fat and matrix accumulation by CD36, which we observed in vitro, also occurs in vivo, we compared characteristics of fat and matrix accumulation in mammary glands of wild type (WT) and Cd36 knockout (Cd36 KO) mice (29). Paraffin sections of the #4 mammary glands of WT or Cd36 KO mice were stained with Masson Trichrome, imaged (Fig. 5A) and quantitated on a cell-by-cell (for fat analysis) or image-by-image (for matrix analysis) basis (Supplementary Fig. S1). Cd36 KO mice exhibited a statistically significant decrease (1.2-fold, \( P < 10^{-8} \)) in fat accumulation (fat cell area; Fig. 5B) and a statistically significant increase (2.1-fold, \( P < 10^{-8} \)) in matrix accumulation (Fig. 5C) relative to WT mice. Paraffin sections were stained for COL1A1 and FN1 (Fig. 5D and E, left panels), 2 matrix proteins modulated by Cd36 in vitro (Fig. 4C and D). Image-by-image quantitative analysis (Supplementary Fig. S1) revealed that Cd36 KO mice accumulated a statistically significant greater amount of COL1A1 (1.6-fold, \( P < 10^{-8} \), and FN1 (2.8-fold, \( P = 5.5 \times 10^{-7} \)) in matrix accumulation compared with WT mice (Fig. 5D and E, right panels). Therefore Cd36 KO mice exhibited 2 prominent phenotypes of high mammographic density and desmoplastic tissues in women: decreased fat accumulation and increased matrix accumulation, showing causality between Cd36 expression and modulation of both phenotypes in vitro and in vivo.

CD36 Expression Is Decreased in Multiple Cell Compartments of High Mammographic Density Tissues Compared with Low Mammographic Density Tissues

CD36 is expressed on the surface of many cells that reside within tissue stroma, but not on epithelial cells (25). Given that microarray analysis of whole tumor tissues showed a dramatic reduction in Cd36 compared with disease-free tissues (Supplementary Table S4), we determined whether cellular components other than fibroblasts also exhibited a repression of Cd36. Tissue sections from core or excisional biopsies, obtained from cancer-free women with either low mammographic density or with high mammographic density (Supplementary Table S5) were stained for Cd36 (Supplementary Fig. S5) and imaged (Fig. 6A, left panels). As expected, adipocytes, endothelial cells, macrophages, and fibroblasts each stained positive for Cd36, whereas epithelial cells did not. However, a striking decrease in the intensity of Cd36 staining in the high mammographic density tissues was observed in every cell type noted above, indicating that Cd36 repression reflects a widespread multicellular program. Cd36-positive staining was measured on a cell-by-cell basis as previously described (ref. 26; Supplementary Fig. S1). The average amount of Cd36 staining per cell was lower in high mammographic density tissues compared with low mammographic density tissues (4.5-fold, \( P < 10^{-8} \); Fig. 6A, right panel) due to a decrease in both the percentage of Cd36-positive cells and the intensity of Cd36 staining per cell (Supplementary Fig. S6).

CD36 Expression Is Decreased in Multiple Cell Types within the Stroma of Malignant Lesions Compared with Histologically Normal Adjacent Tissues

Having observed that Cd36 expression is decreased in multiple cell types within tissues of high mammographic density, we investigated whether the same would be true in tumor tissues obtained from women with estrogen receptor (ER)/progesterone receptor (PR)-positive, HER2-positive, or ER/PR/HER2-negative (triple negative) IDC (Supplementary Table S6). Tissue sections from this cohort were stained for Cd36 and imaged in both the tumor field and the histologically normal adjacent areas (Fig. 6B). In all tumor subtypes, Cd36 expression was strikingly absent in the tumor field compared with the normal adjacent tissue (Fig. 6B). Cell-by-cell quantitation of Cd36 staining (Supplementary Fig. S1) revealed that Cd36 expression was reduced 14.7-fold (\( P < 10^{-8} \)) in tumor tissue compared with normal adjacent tissue (Supplementary Fig. S7). This was due to a decrease in both the percentage of Cd36-positive cells and the intensity of Cd36 staining per cell (Supplementary Fig. S6). A similar dramatic decrease in Cd36 expression was observed in each tumor subtype (Fig. 6C). Notably, it has been reported that women with high mammographic density have a higher propensity to develop ER-negative tumors compared with women with low mammographic density (30). Strikingly, normal adjacent tissue from triple negative (ER-negative) tumors exhibited significantly lower levels of Cd36 expression compared with ER-positive or HER2-positive tumors (Fig. 6C). Taken together, these data suggested that low expression levels of Cd36, as seen in tissues with high mammographic density, might predispose a subset of women to develop triple negative tumors, a tumor subtype associated with poor disease outcome.

To identify the cellular basis of the reduced Cd36 staining, we examined the cells that typically express Cd36 in histologically normal tissues (adipocytes, endothelial cells, and macrophages). Although scarce, adipocytes within tumor tissue not only showed decreased expression of Cd36 but were also smaller in size compared with adipocytes adjacent to the tumor (Supplementary Fig. S7). Staining of serial sections for endothelial and macrophage markers CD31 and CD68, respectively, revealed that...
Figure 5. CD36 knockout mice phenocopy human high mammographic density and desmoplastic tissues. Representative bright field images (original magnification x10) of #4 mammary gland paraffin sections from either WT (n = 7) or CD36 KO (n = 5) mice stained with Masson Trichrome (A), or for COL1A1 (D, left) and FN1 (E, left). Average and SEM of fat cell area (B) or percent area of either matrix staining (C), COL1A1 staining (D, right) or FN1 staining (E, right).
**Figure 6.** CD36 expression is coordinately regulated in multiple cellular compartments. **A,** left: representative bright field images (original magnification ×20) of paraffin sections from 13 low mammographic density and 14 high mammographic density tissues stained for CD36. Right: average and SEM of CD36 signal per cell. **B,** representative bright field images (original magnification ×5) of paraffin sections with tumor and normal adjacent tissue from 8 ER-positive tumors, 6 HER2-positive tumors, and 6 triple negative tumors stained for CD36. **C,** average and SEM of CD36 signal per cell for each tumor subtype. **D,** left: representative bright field images (original magnification ×20) of paraffin sections from 21 low mammographic density and 14 high mammographic density ER-positive IDC patients stained for CD36. Right: average and SEM of CD36 signal per cell. CA, tumor; HD, high mammographic density; LD, low mammographic density; NA, normal adjacent, TN, triple negative.
endothelial cells and macrophages were still present within the field of tumor tissue, albeit with repressed CD36 expression (Supplementary Fig. S7). In contrast, in the normal adjacent tissue, endothelial cells and macrophages positive for CD31 and CD68, respectively, also stained positive for CD36 (Supplementary Fig. S7). Together with the results in Fig. 3C, these data showed that CD36 expression is actively repressed in fibroblasts, adipocytes, endothelial cells, and macrophages within tumor tissue compared with normal adjacent tissue. This documents that CD36 repression in tumor tissue, also seen in high mammographic density tissue, reflects a widespread, coordinated, and multicellular program.

The repression of CD36 in multiple cell types in high mammographic density tissue compared with low mammographic density tissue, in the absence of a tumor, suggests that this repression can be an early event that precedes overt tumor formation. If this were the case, one might predict that CD36 levels would be repressed in histologically normal tissue directly adjacent to the tumor compared with histologically normal tissue distal to the tumor. To test this hypothesis, multiple tissue sections containing both tumor and histologically normal adjacent tissue or histologically normal distal tissue from 3 mastectomies stained for CD36 were obtained from 3 women with invasive cancer (Supplementary Table S7 and Fig. S8). Tissue sections were stained for CD36, imaged in the tumor, normal adjacent, and normal distal fields (Fig. 7A–C, left panels) and quantified for CD36 expression on a cell-by-cell basis (Supplementary Fig. S1). In accordance with the results reported above, CD36 expression was reduced in the tumor compared with normal adjacent and normal distal tissues in all 3 women (4.7-fold to 113-fold and 6.8-fold to 159-fold, respectively, \( P < 10^{-8} \); Fig. 7A–C, right panels). Consistent with the hypothesis above, we observed in 2 of 3 cases that the normal adjacent tissue exhibited CD36 repression compared with its matched normal distal tissue in all 3 women (4.7-fold to 113-fold and 6.8-fold to 159-fold, respectively, \( P < 10^{-8} \); Fig. 7A and B). In further support of this interpretation, it has been reported that invasive cancers often develop in regions previously measured as radiodense for many years before tumor detection (31). Interestingly, one sample did not show this difference between normal adjacent and normal distal tissues (Fig. 7C).
A more detailed analysis of this heterogeneity is currently under investigation.

**Clinical Outcome Is Associated with CD36 Expression**

Given the dramatic universal repression of CD36 observed within the stroma of all tumor subtypes examined (Fig. 6B and C), we determined whether the initial CD36 levels in nondiseased breast tissue (i.e., mammographic density) would stratify tumors for their ultimate CD36 repression and subsequent clinical phenotypes. Tissue sections containing tumor tissue obtained from women with low mammographic density and women with high mammographic density diagnosed with ER-positive IDC (Supplementary Table S8) were stained for CD36 and imaged (Fig. 6D, left panels). As expected, CD36 staining was very low in all tumor tissues. However, using cell-by-cell quantitation (Supplementary Fig. S1), the average amount of CD36 staining per cell was lower in the tumor tissues of women with high mammographic density (1.7-fold, \( P < 10^{-6} \); Fig. 6D, right panel). This was due to a decrease in both the percentage of CD36-positive cells and the intensity of CD36 staining per cell (Supplementary Fig. S6).

Women with high mammographic density who develop invasive cancer have an increased risk for aggressive disease, in that they develop larger tumors of more advanced stage and higher grade compared with women with low mammographic density (30, 32, 33). Because CD36 expression is reduced in tissues of high mammographic density, we examined whether CD36 expression was correlated with tumor grade and size. We interrogated CD36 expression in 2 public gene expression datasets (GSE6532 and GSE9195) for which molecular and associated clinical data were available for tumor samples from a total of 398 treatment-naive IDC patients (34, 35). We analyzed tumor grade, tumor size, and age as a function of CD36 expression using linear regression analyses (Supplementary Table S9). For the purpose of this analysis, we transformed CD36 levels so that they approximated a standard normal distribution (mean = 0, SD = 1). These analyses revealed a statistically significant inverse relationship between CD36 expression and tumor grade and, independently, tumor size. CD36 expression was reduced 0.38 SD (\( P = 0.007 \)) in poorly differentiated tumors (grade 3) compared with well-differentiated tumors (grade 1), after adjusting for the association between CD36 expression and tumor size. In addition, CD36 expression was reduced 0.15 SD (\( P = 0.003 \)) for every 1 cm increase in tumor size, after adjusting for the association between CD36 expression and tumor grade. There was no significant association between CD36 expression and age.

In sum, stromal characteristics before tumor formation, in this case reflected by extent of mammographic density and CD36 expression levels, are associated with clinical outcome.

**DISCUSSION**

Using a combination of *in vitro* cell assays and *in vivo* screening of human and murine mammary tissue samples, we show that the level of expression of a single molecule, CD36, is necessary and sufficient to simultaneously control adipocyte content and matrix accumulation, 2 phenotypes that histologically define mammographic density. This is the first report of a gene shown to mechanistically modulate selected phenotypes of mammographic density. An important and novel finding of our study is that repression of CD36 is observed in both stroma associated with a malignant lesion and in breast tissue with high mammographic density in the absence of malignancy. Equally important is our demonstration that repression of CD36 is not limited to a specific cell type but rather is observed in all stromal components, as documented here for adipocytes, endothelial cells, macrophages, and fibroblasts, thus highlighting that expression of CD36 is part of a complex, coordinated, multicellular program. Taken together, these data suggest that expression of this stromal program is not only a reactive response to existing tumor cells but also represents a physiologic state that increases cancer risk. Our data suggest a functional role for this coordinated program because the more repressed CD36 expression is, the more aggressive the tumor.

CD36 is uniquely poised to coordinate orchestrating distinct protumorigenic phenotypes involving multiple cell types. In endothelial cells, in which CD36 blocks VEGF-induced proliferation, migration, and sprouting as well as induces apoptosis in response to ligand binding, its reduced expression could promote a proangiogenic phenotype (36, 37). In activated macrophages, in which CD36 promotes the formation of foam cells and atherosclerotic plaques, triggers the release of inflammatory cytokines and reactive oxygen species (ROS), and inhibits migration (25), its reduced expression could induce multiple, potentially protumorigenic, phenotypes. In support of this, macrophages from CD36 KO mice have reduced levels of ROS, iNOS, and inflammatory cytokines (including IFN-\( \gamma \) and TNF-\( \alpha \)) and increased arginase activity (38), indicative of a shift from an M1 proinflammatory/antitumorigenic state to an M2 anti-inflammatory/protumorigenic state (39). In dendritic cells, in which CD36 normally mediates the uptake of apoptotic cells and the cross-presentation of tumor-specific antigens to cytotoxic T cells (25), its reduced expression could allow the tumor to evade the immune system (40). The decreased adipocyte differentiation [with reduced adiponectin (Apn)] and increased matrix accumulation [with increased collagen (Col1a1)] that we observed with reduced CD36 expression in *in vitro* and in *vivo* have been shown to cause an increase in tumor burden, angiogenesis, and metastasis in a murine mammary tumor model (MMTV-PyMT mice crossed with Apn KO or Col1a1m\textsuperscript{ink} mice, respectively; refs. 41, 42). Finally, increased matrix accumulation could stiffen the microenvironment, thereby potentially altering mechanosensory networks involved in differentiation and malignancy (43, 44). In light of the central role of CD36 in modulating many functions in multiple cell types, one would predict that the reduction of CD36 expression could have widespread consequences.

The pro-oncogenic tissue state created by repression of CD36 goes far beyond merely supporting the malignant program of a transformed epithelial cell; it creates an interactive...
milieu that actively participates in tumorigenesis. Hence, our study provides novel insights into a molecular program that controls several stromal phenotypes underlying tumorigenesis. The "single hit, multiple target" nature of CD36, similar to miRNAs, makes it a very attractive therapeutic target. Moreover, in light of its role in mammographic density, strategies to modulate CD36 have the potential to prevent cancer progression in women who are at high risk. In support of this, we show that increased expression of CD36 can restore stromal phenotypes associated with low-risk tissues (Fig. 4). Promisingly, expression of CD36 can be increased in vitro by treatment with aspirin, dexamethasone, statins, or adalimumab and in vivo by treatment with tamoxifen (45–49). Finally, mammographic density can also be modulated by a variety of exogenous agents such as hormones, tamoxifen, and diet. As such, mammographic density represents a highly promising target for intervention as well as a biomarker that can assess response to prevention interventions (19, 21).

METHODS

Isolation and Propagation of Human Mammary Fibroblasts

Fibroblasts were isolated from reduction mammoplasty, low and high mammographic density, or tumor tissues and assessed for purity as described (5).

Microarray and Quantitative PCR Analysis

Total RNA, isolated from fibroblasts subjected or not to adipocyte differentiation, was used to generate cDNA using TaqMan Reverse Transcription Reagents (Applied Biosystems). The cDNA was used for QPCR (TaqMan) using the standard curve method with primer probe sets for CD36 and leptin. Expression of beta-glucuronidase (GUSB) was used to normalize input cDNA. For microarray experiments, mRNA, amplified and biotinylated from total RNA obtained from LDAFs and HDAFs using the MessageAmp II Kit (Ambion), was fragmented and hybridized to HU133 plus 2 chips (Affymetrix).

Adipocyte Differentiation Assay

Fibroblasts, plated at near confluence and grown for 48 hours, were treated with 10 μmol/L 15-deoxy-Δ12,14-Prostaglandin J2 (PJ2; Cayman Chemical) to induce adipocyte differentiation. Lipid accumulation was assayed by Oil Red O staining. Adipocyte differentiation was assessed by monitoring CD36 and leptin expression after 2 weeks of treatment.

Matrix Accumulation

Fibroblasts, plated at near confluence into 8-well glass chamber slides, were grown for 5 days. Matrix accumulation was assessed by monitoring expression of COL1A1 (Abcam), FN1 (BD Biosciences), OPN (Abcam), and TNC (R&D) with primary antibodies all diluted 1:100.

Human and Mouse Tissue Preparation and Immunohistochemistry

Mouse sections (5 μm) were stained with Masson Trichrome using standard protocols. Human sections (4 μm) and mouse sections were subjected to antigen retrieval: microwaving in EDTA buffer (pH = 8.0) for 1 minute for CD36, incubation in Citrate buffer (pH = 6.0) at 80°C for 1 hour for CD31 and CD68, or microwaving in citrate buffer (pH = 6.0) for 10 minutes for COL1A1 and FN1. Sections were incubated for 1 hour with antibodies to CD36 (Sigma-Aldrich), COL1A1 (Abcam), or FN1 (Epitomics) diluted 1:100, 1:20, 1:200, 1:500, and 1:250, respectively. Human sections were incubated for 20 minutes with Primary Antibody Enhancer (Thermo Scientific). All sections were incubated for 30 minutes with HRP Polymer (Thermo Scientific) and 5 minutes with diaminobenzidine, and then counterstained in Mayer hematoxylin to visualize nuclei.

Analysis of Published Microarray Data

Microarray datasets GSE6532 and GSE9195 were analyzed using the robust multiarray average method (50). Expression values for each of the 5 CD36 probe sets were centered by the median, then divided by the SD. Linear regression analyses were used to test the associations between the expression of each CD36 probe set and tumor grade, tumor size, and age. Data from the probe set that were most strongly associated with tumor grade and tumor size are presented in Supplementary Table 59.

Detailed experimental procedures are included in Supplementary Materials and Methods.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Authors’ Contributions

Conception and design: R.A. DeFilippis, N. Dumont, Y.-Y. Chen, H.K. Berman, K. Kerlikowske, T.D. Tlsty


Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): R.A. DeFilippis, J.T. Rabban, Y.-Y. Chen, M.L. Gauthier, J. Zhao, J.J. Marx, J.A. Tjoe, M. Febrasio, K. Kerlikowske, T.D. Tlsty


Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): R.A. DeFilippis, G.V. Fontenay, J. Zhao, J.A. Tjoe, E. Ziv, K. Kerlikowske

Study supervision: R.A. DeFilippis, Y.-Y. Chen, T.D. Tlsty

Acknowledgments

The authors thank members of the Tlsty laboratory, particularly Drs. Colleen Fordyce and Philippe Gascard, for thoughtful discussions, the Nikon Imaging Center (UCSF) for imaging support, and Dr. Alfred Au (UCSF), the breast surgeons (Aurora Health Care), and Carol Halliday, RN (Aurora Health Care), for providing tissue.

Grant Support

This work was supported by NIH/NCI PO1 CA107584 to T.D. Tlsty, K. Kerlikowske, E. Ziv, and B. Parvin (under LBNL contract No. DE-AC02-05CH11231), California Breast Cancer Research Program grant 14OB-0165 to T.D. Tlsty, and NIH/NCI U54 CA143803 to T.D. Tlsty and B. Parvin (UC Riverside).

Received March 12, 2012; revised June 28, 2012; accepted July 2, 2012; published OnlineFirst July 9, 2012.
CD36 Modulates Phenotypes of Breast Density and Desmoplasia


CD36 Repression Activates a Multicellular Stromal Program Shared by High Mammographic Density and Tumor Tissues

Rosa Anna DeFilippis, Hang Chang, Nancy Dumont, et al.


Updated version
Access the most recent version of this article at:
doi:10.1158/2159-8290.CD-12-0107

Supplementary Material
Access the most recent supplemental material at:
http://cancerdiscovery.aacrjournals.org/content/suppl/2012/07/09/2159-8290.CD-12-0107.DC1

Cited articles
This article cites 50 articles, 14 of which you can access for free at:
http://cancerdiscovery.aacrjournals.org/content/2/9/826.full#ref-list-1

Citing articles
This article has been cited by 11 HighWire-hosted articles. Access the articles at:
http://cancerdiscovery.aacrjournals.org/content/2/9/826.full#related-urls

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, use this link http://cancerdiscovery.aacrjournals.org/content/2/9/826.
Click on "Request Permissions" which will take you to the Copyright Clearance Center's (CCC) Rightslink site.