Tumor Microenvironment–Derived Proteins Dominate the Plasma Proteome Response during Breast Cancer Induction and Progression

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Abstract

Tumor development relies upon essential contributions from the tumor microenvironment and host immune alterations. These contributions may inform the plasma proteome in a manner that could be exploited for cancer diagnosis and prognosis. In this study, we employed a systems biology approach to characterize the plasma proteome response in the inducible HER2/neu mouse model of breast cancer during tumor induction, progression, and regression. Mass spectrometry data derived from approximately 1.6 million spectra identified protein networks involved in wound healing, microenvironment, and metabolism that coordinately changed during tumor development. The observed alterations developed prior to cancer detection, increased progressively with tumor growth and reverted toward baseline with tumor regression. Gene expression and immunohistochemical analyses suggested that the cancer-associated plasma proteome was derived from transcriptional responses in the noncancerous host tissues as well as the developing tumor. The proteomic signature was distinct from a nonspecific response to inflammation. Overall, the developing tumor simultaneously engaged a number of innate physiologic processes, including wound repair, immune response, coagulation and complement cascades, tissue remodeling, and metabolic homeostasis that were all detectable in plasma. Our findings offer an integrated view of tumor development relevant to plasma-based strategies to detect and diagnose cancer.

Introduction

The tumor microenvironment and host factors play a major role in the establishment, progression, and metastatic dissemination of cancer (1). Stromal fibroblasts, myofibroblasts, endothelial cells, and immune cells provide growth factors, proteases, and angiogenic potential to support tumor growth (2, 3). Dendritic cells and recruited T and B cells, regulatory T cells, macrophages, and other myeloid-derived suppressor cells, contribute both positively and negatively to multiple stages of tumor progression (4–7). Tumors can also recruit progenitor cells from the bone marrow to promote angiogenesis and to establish premetastatic niches in distal organs such as the lung (8–10). To date, these tumor–host interactions have been primarily studied at the cellular and tissue levels, and it remains unclear up to what extent they contribute to the changes in the plasma proteome, particularly at early stages of tumor development. Proteomic advances currently allow in-depth quantitative profiling of biological fluids, such as plasma spanning of more than 6 logs of protein abundance (11, 12). Application of proteomics to mouse models provides a number of distinct advantages to interrogate the systems biology of cancer (13, 14).

We utilized a conditional transgenic neu-induced mouse model of mammary cancer to determine the extent and source of changes in the plasma proteome at predetermined stages of tumor development (15). The neu is an activated rat homologue of the human epidermal growth factor receptor 2 (ErbB2) gene. ErbB2 is a receptor tyrosine kinase amplified and overexpressed in more than 25% of human breast cancers, and signaling from this oncogene is a central driver in breast cancer development. When induced with doxycycline, bitransgenic MMTV-rtTA/TetO-NeuNT mice synchronously develop invasive mammary carcinomas that recapitulate the morphologic, pathologic, and molecular features of ErbB2-positive human breast cancer (16). Doxycycline withdrawal results in...
transgene de-induction and tumor regression, mimicking responses of tumors to targeted therapy. We applied in-depth quantitative proteomic profiling to this model and identified a set of proteins that changed in relative abundance in plasma during tumor induction, progression, and regression. Database mining, cross-referencing proteomic data with transcript profiles, and confirmatory cell-type localization studies revealed that tumor cells, the microenvironment, and systemic responses contribute to the dynamic changes in the plasma proteome with tumor development and progression.

Materials and Methods

Tissue collection
Bitransgenic FVB MMTV-rtTA/TetO-NeuNT "case" and monotransgenic FVB TetO-NeuNT "control" mice (15) were paired at weaning and maintained in the same cage. Doxycycline (2 mg/mL) was added to drinking water starting at 8 weeks of age. Mice were palpated every other day to detect mammary tumor growth. Each pair of case and control mice was euthanized on the same day by CO2 inhalation and plasma was collected as described (13).

Plasma proteomic analysis
The plasma proteomics workflow has been previously described (17). In brief, case and control plasma pools (250–315 µL/pool) from 5 to 11 mice were immunodepleted (3 abundant proteins, MS-3; Agilent). Samples were concentrated and reduced with dithiothreitol. Isotopic intact protein labeling of cysteine residues was done with control and reduced and alkylated with iodoacetamide and fractionated by reversed-phase chromatography (Poros R2/10). Anion-exchange chromatography (Poros HQ/10; Applied Biosystems) used an 8-step elution (0–1,000 mmol/L NaCl). Anion-exchange fractions were separated by reversed-phase chromatography (Pores R2/10). Lyophilized fractions were digested with trypsin and 96 fractions per tumor stage were analyzed by a LTQ-Orbitrap (Thermo) mass spectrometer and NanoLC-1D (Eksigent). Each tumor stage analysis yielded approximately 4,000,000 spectra, which were searched against the mouse International Protein Index (IPI) database (v.3.48) by using X!Tandem with ProteinProphet. Protein ratios were computed from peptides identified with a PeptideProphet probability greater than 0.2 were submitted to ProteinProphet. Proteins were filtered using greater than 5% error rate. Protein ratios were computed from peptides with PeptideProphet probability greater than 0.75. Proteins with multiple IPIs and single gene name were combined (log2 average of peptides) to give 1 ratio per gene name.

For proteins with multiple paired MS events of heavy and light acrylamide, a 2-tailed Student’s t test was used to calculate a P value for the mean case/control ratio of the protein. Second, the probability for the case/control ratio for each MS event was calculated from the distribution of ratios in control/control experiments in which the same sample was labeled with heavy and light acrylamide. On the basis of the control/control experiments, if the P value for each individual event was less than 0.05, the overall protein ratio was considered significant.

Breast cancer cell line profiling
The cell line methods are previously described (24). MCF7, MDA-MB-231, and SKBR3 cell lines were obtained from American Type Culture Collection (ATCC), cultured for less than 6 months, and were not reauthenticated. Cells were cultured for 7 passages in SILAC DMEM [Invitrogen; dialyzed FBS, 13C-lysine; (25)]. After 48 hours, secreted proteins were obtained from media. For total cell extract, cells were sonicated and centrifuged to remove debris. For cell surface proteins, cells were biotinylated, extracted, sonicated, and captured by us an avidin column. One milligram of each of secreted, total cell extract, and cell surface proteins was reduced and alkylated with iodoacetamide and fractionated by reversed-phase chromatography. Fractions were digested with trypsin and subjected to liquid chromatography/tandem mass spectrometry (LC/MS-MS) as described above.

Immunohistochemistry
Tissues were processed and stained with calreticulin (Calr; Abcam), Lgals1 (R&D Systems), lactotransferrin (Ltf; Abgent), S100a9 (NovusBio), Pdz and limb domain protein (Pdlim1; Abcam), smooth muscle actin (SMA; Thermo), Timp1 (Abbiotec), Vasp (Abcam), or Igfbp5 (Santa Cruz) antibodies by using a standard 3-step technique (26). Collagen staining was done using the Sigma Trichrome Kit. The University of Washington Histology Core conducted CD3 and F4/80 staining. Twelve tumors were stained. For each antibody all tumors showed similar staining patterns.

Results
Histologic evaluation of mammary tumor induction, progression, and regression
Cohorts of MMTV-rtTA/TetO-NeuNT "case" and TetO-NeuNT "control" mice were sacrificed at 4 time points during tumor development (Fig. 1A). After 5 weeks on doxycycline and prior to clinically detectable lesions, mammary duct hyperplasia, intraepithelial neoplasia, and microinvasive carcinoma were apparent (Fig. 1B). By 8 to 12 weeks, 0.5-cm tumors consisted of hyperplastic nodules of intraepithelial...
neoplasia and areas of adenocarcinoma. By 14 to 16 weeks, 1-cm breast tumors were classified as mammary adenocarcinomas with areas of local invasion. Neoplastic cells were supported by vascular stroma and fibrous connective tissue, indicating a desmoplastic reaction. Inflammatory cell infiltrates included lymphocytes, neutrophils, macrophages, plasma cells, and mast cells. Doxycycline was withdrawn from mice bearing approximately 1-cm tumors and plasma samples were taken 3 days later, when tumors had visibly decreased in size. Histologic evaluation showed regressing mammary trees with ductal ectasia, inflammation, and fibrosis. Plasma from case and control mice at each of these 4 tumor stages was pooled and subjected to differential isotopic labeling, extensive protein fractionation, and LC/MS-MS analysis of tryptic digests from individual fractions (17). The same plasma pools were analyzed using Rules Based Medicine Luminesix/Rules Based Medicine, which measured 58 proteins representing cytokines, growth factors, and other signaling proteins.

Changes in the plasma proteome prior to clinical evidence of cancer

Plasma samples from preclinical tumor-bearing mice yielded 49 proteins with increased and 17 with decreased concentrations in case compared with control mice (>1.5-fold change, \( P < 0.05 \); Fig. 1C and Supplementary Table S1). These proteins were grouped by function, biological process, and cellular localization. Proteins involved in acute-phase response, immune cell signaling—including cytokines, chemokines, and macrophage and neutrophil signaling proteins—and cytoskeletal and extracellular matrix (ECM) functions were increased (Fig. 2). Response to wounding was the most significant biological process altered in the proteome of mice with preclinical neoplasia (Supplementary Fig. S1A). Analysis of the subset of proteins known to be extracellular identified the acute-phase response and the coagulation and complement systems as the most significantly altered pathways (Supplementary Figs. S1B and S2). Among the altered plasma proteins with known intracellular...
localization, those involved in glycolysis, fatty acid, and amino acid metabolism predominated (Supplementary Table S2).

Of the 49 plasma proteins that increased in relative abundance in mice with preclinical neoplasia, 22 were identified in proteomic profiling experiments from 3 human breast cancer cell lines, MCF7, MDA-MB-231, and SKBR3 (Supplementary Table S2). SKBR3 cells overexpress ErbB2 and have been utilized extensively to study ErbB2 signaling (27, 28). Immunohistochemical (IHC) staining of preclinical tumor sections showed cytoplasmic and/or nuclear localization in tumor cells for all 5 proteins for which antibodies were available (Fig. 3), suggesting that release from tumor cell turnover could contribute to increased plasma levels of these proteins. Calr is a calcium-binding chaperone protein reported to have prognostic significance in breast cancer based on tissue expression (29). LgalS1 is induced by ErbB2 (30), binds to H-ras, and mediates cell transformation (31). Ltf is an iron-binding protein that is secreted by breast tissue. Calgranulin B (S100A9) is a calcium-binding protein present in a wide range of cells and overexpressed in neutrophils, monocytes, and tumor cells (32). The Pdlim1 associates with actin filaments and stress fibers.

To determine the extent to which protein changes detected at this early time point may be specifically associated with tumor development, we cross-referenced these data to similar plasma proteome profiling from mouse models of subacute and chronic inflammation and angiogenesis (33). Of the 49 proteins with increased concentration in plasma from mice with early-stage breast cancer, 12 were increased in the plasma of 1 or more of the confounding mouse models (Supplementary Table S2), indicating these 12 proteins are not breast cancer specific. Levels of S100a9, Pdlim, Ltf, Calr, and LgalS1 that were detected in primary tumors by IHC (Fig. 2) were not increased in the confounder models, thereby increasing their potential cancer specificity (see Supplementary Table S2 for additional examples). Interestingly, a greater percentage (~50%) of proteins with decreased concentration in preclinical plasmas were decreased in confounding condition models, indicating that reduced proteins are less likely to be cancer specific. Thus, even at a relatively early stage of tumor development, changes were observed in the plasma proteome reflecting contributions from tumor cells, the microenvironment, and the host response. Proteins involved in functions relevant to

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**Figure 2. Wound-healing response components: acute-phase, immune cell, and ECM proteins.** Plasma proteins that increased in tumor-bearing mice fell into major functional categories. A, acute-phase response proteins, mainly liver-derived. B, proteins from tumor-infiltrating immune cells (macrophages and neutrophils). C, proteins involved in cell movement and tumor microenvironment, for example, cytoskeletal, ECM, and tumor stroma proteins. Red indicates case/control ratios greater than 1.5 (P < 0.05), black indicates no change between case and control plasma, and white indicates that the proteins were not measured. For the breast cancer cell lines, red indicates that the protein was identified by proteomic profiling and white indicates that the protein was not identified. Unless otherwise indicated, levels were otherwise indicated, levels were measured by LC-MS. *, measured by Lumines/Rules Based Medicine. **, measured by both LC-MS and Lumines/Rules Based Medicine.
wound healing, inflammation, metabolism, and ECM remodeling predominated.

**Plasma proteome changes associated with tumor progression**

To identify plasma protein changes associated with tumor progression, we profiled the plasma proteome of mice bearing 0.5- and 1-cm breast tumors. The overall numbers of proteins that showed altered plasma levels in case versus control mice progressively increased with tumor growth (Fig. 1C). In mice bearing 0.5-cm tumors, 38 proteins showed a significant increase and 105 a decrease in abundance, whereas in mice bearing 1.0-cm tumors, 70 proteins were increased and 114 were decreased (>1.5, \( P < 0.05 \); Supplementary Table S2). As in the preclinical samples, proteins that increased in mid- and late-stage tumor-bearing mice included those identified in proteomic profiling of human breast cancer cells, as well as proteins involved in the acute-phase response, complement, and coagulation cascades, chemokines, macrophage and neutrophil signaling, and metabolic proteins (Fig. 2 and Supplementary Fig. S2). A number of cytoskeletal, ECM, and stroma-associated proteins including Loxl1, Tnc, matrix metalloproteinase (Mmp) 9, Timp1, Dsg2, Fbln2, Col15a1, Pxdn, Prg4, and Vwf increased in their levels. Interestingly, only 1 of 30 of these tissue-remodeling proteins was identified as increased in plasma from the confounder mouse models (Fig. 2), suggesting a substantial contribution of the stroma to plasma protein changes associated with tumor development.

It is noteworthy that a greater proportion of plasma proteins were reduced in abundance, rather than increased in plasma from mice with mid- and advance-stage tumors (Fig. 1C). Decreased proteins that were unique to tumor-bearing mice included metabolic proteins and 10 components of the proteasome (Psma1, Psma2, Psma4, Psma7, Psmb1, Psmb10, Psmb6, Psmc5, Psmc4, and Psmc1; Supplementary Table S2). Approximately a quarter of the proteins with reduced levels in mice with clinically apparent breast cancer were decreased in plasma from mice with inflammation or angiogenesis (Fig. 4). These data indicate that decreased protein abundance associated with cancer can be attributed to metabolic changes, reduced proteasomal components, as well as host responses that are not cancer specific.

We next examined the extent and potential contribution of altered gene expression in both primary tumor and nontumor host tissues to changes in the plasma proteome. Gene expression profiling was conducted on blood buffy coat, thymus, spleen, liver, and breast tumors from mice with 1-cm tumors, using an independent cohort of case and control mice (34). Of the 31 proteins that were increased in plasma from tumor-bearing mice and identified in proteomic profiling of breast cancer cells, 25 (81%) showed a corresponding increase in...
expression in HER2/neu breast tumors as compared with normal breast tissue (Supplementary Fig. S3). Twelve of these proteins (Flna, Vasp, Cnn2, Coro1a, Capl1, Pxdn, Dsg2, Tpm4, Pfn1, Cotl1, Zyx, and Ctsd) are involved in cytoskeletal function or ECM remodeling, processes that are necessary for tumor cell invasion. None of these proteins increased in concentration in plasma from the confounder models. The concordance between protein levels in plasma, protein expression in breast cancer cells, and mRNA expression in HER2/neu-driven tumors for these proteins indicates a contribution of tumor cells to their increased levels in plasma.

To determine the contributions of the tumor microenvironment to increased levels of proteins in plasma, we used IHC to examine the cellular and stromal landscape of HER2/neu mammary tumors. In addition to tumor cells, HER2/neu tumors contained macrophage, neutrophil, and lymphocyte infiltrates, fibroblasts, extensive collagen deposition, and prominent blood vessels (Fig. 5). Plasma proteins that could be derived from tumor cell interactions with these recruited cell types and the local environment are also displayed according to function (immunity, ECM, cytoskeletal, and angiogenesis). Note the examples of Timp1 and Vasp that were elevated in plasma and expressed in tumor cells.

For 14 proteins that were increased in plasma of tumor-bearing mice and that are known to be involved in myeloid cell function, we compared the corresponding mRNA expression levels in thymus, spleen, and blood buffy coat fractions that are rich in white blood cells. Of these 14 proteins, 13 had increased levels of mRNA expression levels in 1 or more of these tissues from tumor-bearing versus control mice (Supplementary Fig. S4). Lcn2, S100a9, S100a8, neutrophilic granule protein (Ngp), myeloperoxidase (Mpo), and peptidoglycan recognition protein (Pglyrp1) were the most significantly increased in both plasma and immune cell compartments, suggesting the increased levels of these proteins in plasma reflect an immune cell response to the developing tumor. S100a9 and Lcn2 are examples of proteins identified as increased in both immune cells and tumors. A set of 27 genes showed significantly increased mRNA expression in the thymus in case versus control mice (P < 0.05; ref. 34). Proteins corresponding to 7 of the 27 genes were increased in the plasma, suggesting a contribution of distal immunologic tissues to their increased plasma levels.

**Figure 5.** HER2/neu mammary tumor microenvironment schematic. In addition to the tumor cells, Her2/neu tumors contain cancer-associated macrophages (F4/80), CAFs (α-SMA), lymphocytes (CD3), blood vessels (BV; not shown), and extensive collagen deposition (trichrome). Also shown are IHC results of HER2/neu tumors for Timp1 and Vasp, which were elevated in plasma from tumor-bearing mice. Additional proteins elevated in plasma from tumor-bearing mice are listed according to functional categories (boxes).
Reversion of the plasma proteome toward baseline after oncogene de-induction and tumor regression

To identify plasma protein changes associated with early-stage tumor regression, we quantitated protein changes in plasma from mice 3 days after doxycycline withdrawal and oncogene de-induction (Fig. 1). At this time point, only 24 proteins had increased levels and 28 decreased levels in case versus control mice (Supplementary Table S2). Many proteins increased in tumor-bearing mice were undetected or returned to baseline levels after oncogene de-induction. Of the 12 that remained elevated, 8 were immune cell or acute-phase proteins (Hp, C8g, Ccl8, EG214403, Mpo, Cd40l, Orm3, and Hpx). Changes in acute-phase proteins and complement have also been observed in plasma from epirubicin/docetaxel-treated breast cancer (35).

Comparative analysis of the plasma proteome at different tumor stages

Comparison of the plasma proteome from mice with 0.5- and 1.0-cm tumors revealed a significant positive correlation (Pearson correlation coefficient \( r = 0.67 \)) in relative protein abundance, indicating many of the same proteins are either reduced or increased at both tumor stages (Fig. 4). Some proteins were quantified at all 4 tumor stages. Papln, Ltf, Igfbp5, S100a8, Lrg1, Pcsk9, and Ccl8 are examples of proteins that showed progressive increases in relative abundance during tumor progression and diminished to near control levels after tumor regression (Fig. 6). Of these, Ltf and Igfbp5 were expressed in tumor cells as assessed by IHC. A set of proteins progressively decreased in their levels during tumor development but rebounded after tumor regression (Fig. 6). These proteins are mainly involved in acute-phase response, and many were also decreased in the confounder inflammation models. Pathway analysis was used to assess the biological processes behind these dynamic changes in the plasma proteome that accompanied tumor development. Proteins involved in wound healing, the complement pathway, and metabolism were altered at multiple tumor stages (Supplementary Fig. S2).

Confirmation of LC/MS-MS detected changes in protein abundance by independent antibody-based methods

Four proteins that span a wide range of concentrations in normal plasma and that exhibited altered plasma levels in tumor-bearing mice on the basis of LC/MS-MS analysis were subjected to validation by ELISAs applied to samples from independent cohorts of mice. Protein concentrations of Igfbp5, Lcn2, Timp1, and Ccl8 were found to progressively increase during tumor development and return to normal after regression, concordant with MS data (Fig. 7). We also quantified levels of 37 proteins using bead-based assays applied to plasma samples from all 4 tumor stages. Thirteen of these showed increased levels in case versus control mice for at least 1 tumor stage (Supplementary Table S3). These included Cd40 ligand, monocyte and macrophage signaling proteins, and stromal proteins Mmp9, Timp1, and Vwf. Seven of these proteins were quantified by LC/MS-MS, and again there was substantial agreement between the 2 measurement methods.
Discussion

The cancer-associated plasma proteome is derived from the developing tumor, the tumor microenvironment, and host tissues

In this study, we present a dynamic analysis of the plasma proteome that accompanies HER2/neu-induced mammary tumor establishment, malignant progression, and regression. Our findings show that the plasma proteome increasingly deviates from normal as tumors develop and progress. This presented as an escalation in the total number of altered proteins as well as a progressive increase or decrease in the relative abundance of a number of specific proteins with advancing tumor stage. Most of these tumor-associated protein changes rapidly reverted upon oncogene downregulation and incipient tumor regression. Thus, the plasma proteome responds very early, that is, shortly after tumor induction, changes both qualitatively and quantitatively during tumor progression, and rapidly reverts to normal following oncogene downregulation and tumor regression.

We used several methods to ascertain the specificity and tissue(s) of origin of the plasma protein signature that occurred in the context of breast tumor development. First, comparisons with mouse models of inflammation and angiogenesis were used to identify those proteins that were not cancer specific. Approximately 25% of the elevated and 50% of the reduced proteins overlapped between tumor and confounder plasma proteomes. This indicates that the proteins that decrease in abundance in tumor-bearing mice are more likely attributable to a host response that is not specific to cancer. In the future, analysis of other confounder models such as wound-healing or tissue-specific inflammatory conditions will be useful to further refine the cancer-specific signature. Second, we compared plasma proteins identified in tumor-bearing mice to proteomic and gene expression analyses of human breast cancer cells and HER2/neu-induced breast tumors. Among all the proteins that were increased in plasma from tumor-bearing mice, more than half were identified in the proteomic profiling of human breast cancer cell lines and approximately 80% of this concordant set showed a corresponding increase in gene expression in HER2/neu breast tumors. Third, a subset of those proteins found to be altered in the plasma of tumor-bearing mice was expressed in tumor cells as detected by IHC. These findings point to the tumor itself...
as the likely source of a subset of proteins found to be altered in plasma. In addition to the secretion of proteins by tumor cells, tumor cell lysis may also lead to detectable levels of cancer-specific proteins in the blood. Finally, microarray gene expression analysis of known inflammatory proteins identified a correlation between protein levels in plasma and gene expression in distal immunologic tissues such as thymus and spleen. Collectively, these findings show that changes in the plasma proteome that arise during cancer development derive from multiple sources: nonspecific systemic reactions, for example, acute-phase proteins such as those produced by the liver; other host responses that are more specific to breast cancer, but not directly tumor derived, for example, proteins involved in immune cell recruitment and signaling, and metabolic pathways; and proteins that originate directly from the tumor and the local microenvironment (Fig. 5). Some proteins are expressed in multiple tissues (e.g., S100a9, Lcn2, metabolic, and other housekeeping proteins) and unambiguous ascertained of tissue of origin is more difficult.

**Plasma proteome dynamics reflect breast cancer biology**

After only 5 weeks of HER2/neu oncogene activity, extensive neoplastic changes were noted throughout the mammary ductal tree. Although the lesions were clinically undetectable, significant changes in the plasma proteome were already apparent, and these reflected acute-phase responses, macrophage and neutrophil trafficking, ECM remodeling, metabolism, and potentially tumor-derived proteins. A significant fraction of these early plasma proteome alterations were not seen in models of inflammation and so are more likely to be associated with early-stage cancer. A number of the candidate tumor-derived proteins, such as Igfbp5, Ltf, Timp1, Lgals1, Mmp9, Vwf, Calr, and Pdlim1, that were increased in preclinical plasma were also seen in tumor sections and/or breast cancer cell lines based on IHC or gene expression changes. A subset of these was confirmed to be elevated in plasma from other cohorts of mice and by independent antibody-based methods. Furthermore, the plasma levels of some of these proteins increased with tumor size and rapidly fell after oncogene de-induction, indicating they effectively track tumor growth. This establishes both the sensitivity of our method to detect tumor-associated proteins very early in the neoplastic process and also the ability to monitor protein levels over the course of tumor development and during tumor regression.

Notable histopathologic changes that occurred during tumor progression included extensive remodeling of the tumor microenvironment, in particular, recruitment of immune cells, including T and B cells, macrophages, and neutrophils. This presentation resembles a wound bed and implies that the tumors actively attract these cells from the circulation. In support of this conclusion, plasma levels of multiple macrophage signaling proteins, including monocyte chemoattractant proteins (Mcps), macrophage inflammatory proteins (Mip), and other chemokines, calgranulins, and lipocalin 2, were increased. Many of these proteins were elevated even at the earliest tumor stage and progressively increased in abundance with tumor growth. Once recruited to the tumor site, tumor-associated macrophages (TAM) secrete Mmp2, Mmp9, TGFβ, cathepsins, and other factors that degrade and remodel matrix (36) as well as promote angiogenesis, tumor cell invasion, and metastasis (37). Our analysis showing elevated plasma levels of Mmp9, Ctsb, and numerous stromal components associated with TAMs extends the reach of this network to the organismal level.

Cancer-associated fibroblasts (CAF) are another significant component of the HER2/neu mammary tumor microenvironment (Fig. 5). These "activated" fibroblasts express smooth muscle actin (38) and support tumor growth, deposit ECM, and promote an inflammatory response by secreting factors such as collagens, tenascin C, Mmps, and chemokines (2, 3). Tumor cells, in turn, must reorganize cytoskeletal structures, break down the basement membrane, and degrade ECM components to invade, intravasate, and metastasize. The levels of many proteins related to these processes, including ECM, stromal, and cytoskeletal proteins, were increased in plasma, particularly from mice with late-stage tumors. Together this shows that, in addition to nonspecific host response proteins and tumor-derived proteins, alterations in signaling and effector pathways regulating tumor–TAM and tumor–CAF interactions are prominent features of the cancer-associated plasma proteome. These features bear similarities to a wound response (39), which has recently been described to parallel tumor initiation (40), and indicate the extent to which the tumor co-opts these host responses. This establishes the plasma proteome as a sensitive biomonitor of the ongoing interaction between the developing tumor and the host.

Altered plasma levels of enzymes involved in glucose, lipid, and amino acid metabolism were observed at all 3 tumor stages. Liver transcriptional profiling identified altered expression of similar metabolic enzymes in case versus control mice (34), indicating the presence of a tumor triggers systemic metabolic dysregulation, and this initiates very early during tumor establishment. It will be interesting to determine the mechanistic similarities between this response and cachexia, which is an altered systemic metabolic response to the presence of late-stage cancer (41).

In summary, this study highlights the value of integrating in-depth proteomic, transcriptomic, and tissue-wide approaches as a means to illuminate the systems biology of cancer. The identification of plasma protein networks and transcriptional responses in distal normal tissues of tumor-bearing mice indicates the host senses and responds to neoplastic growth to a greater degree than previously appreciated. The developing tumor simultaneously engages a number of innate physiologic processes including wound repair, the immune system, the blood coagulation and complement cascades, and metabolic homeostasis. This extensive conversation initiates very early, involves many cell types and tissues, accurately tracks tumor development
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over time, and mirrors the underlying biology. Further mining of the cancer-associated plasma proteome and other host tissue responses will likely identify additional regulators, mediators, and biomarkers of the evolving tumor, its microenvironment, and the systemic response to cancer. Extension of this approach to other models of cancer as well as human studies will establish its utility for early detection, diagnosis, and prognosis.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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