Folate receptor alpha (FRA) expression in breast cancer: identification of a new molecular subtype and association with triple negative disease

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Abstract
Given that several targeted therapies directed towards folate receptor alpha (FRA) are in late stage clinical development, the sensitive and robust detection of FRA in tissues is of paramount importance relative to patient selection, prognosis and prediction. In the present study we undertook an immunohistochemical evaluation of expression of FRA in breast cancer samples using formalin-fixed, paraffin-embedded (FFPE) tissues, primarily invasive ductal carcinomas, using a newly described monoclonal antibody, 26B3. Samples assessed included both tissue microarrays (TMA) and whole tissue sections from archival tissue blocks. Normal breast shows a highly restricted expression of FRA to luminal membrane staining of secretory ductal cells, consistent with FRA secretion into milk. In early stage (stages I-III) invasive ductal carcinomas, FRA staining was observed in approximately 30% of all samples, independent of molecular subtype (estrogen receptor (ER)/progesterone receptor (PR)/human epidermal growth factor receptor type 2 (Her2)). However, FRA expression was shown to associate with ER/PR negative tumors relative to ER/PR positive tumors (p = 0.012) and perhaps more importantly, with triple negative breast cancers (TNBC; p < 0.0001). FRA immunoreactivity was also shown to be retained in stage IV metastatic breast cancer samples from diverse anatomic sites including lymph node and bone. In metastatic breast cancer, FRA was shown to be expressed in 86% of TNBC patients. Taken together, these data suggest that FRA expressing breast cancer represents a novel molecular subtype and, further, may represent a new therapeutic target for this devastating disease.

Keywords: Folate receptor alpha, FRA, Breast cancer, Triple negative breast cancer, Immunohistochemistry, IHC

Introduction
According to Global Cancer Facts & Figures 2nd Edition, in 2008 the estimated worldwide new cases for breast cancer were 1,383,500 with a projected 458,400 deaths and a mortality rate of approximately 33%. In the U.S., 229,060 new cases of breast cancer and 39,920 deaths from this disease are expected in 2012 (Siegel & Naishadham 2012). Treatment for breast cancer is currently tailored according to cellular protein expression. Estrogen receptor/progesterone receptor (ER/PR) expressing breast cancers are treated with endocrine therapy. The treatment armamentarium of human epidermal growth factor receptor 2 (Her2) overexpressing breast cancers includes an anti-Her2 agent. The triple negative breast cancers (TNBC) that do not express ER, PR or Her2 are treated with traditional cytotoxic chemotherapy alone. New therapeutic approaches for this poor prognosis breast cancer subtype are sorely needed.

Folate, or vitamin B9, is an essential cofactor in the synthesis of purines and pyrimidines and other cellular methylation reactions including DNA, proteins and lipids (Elnakat & Ratnam 2004). The folate receptors (folate receptor alpha, FRA; beta, FRB, gamma, FRG; and delta, FRD) constitute a family of proteins that, at least in part, mediate accumulation of folate into cells, regulate folate homeostasis and may have effects on cellular proliferation (Elnakat & Ratnam 2004; Kelemen 2006). FRA, a glycosylphosphatidylinositol (GPI)-anchored cell surface glycoprotein, has a very limited tissue distribution. In normal tissue, FRA is mainly expressed on the apical surface of a

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subset of polarized epithelial cells including parotid, kidney, lung, thyroid and breast (Weitman et al. 1992a; Weitman et al. 1992b; O'Shanessy et al. 2011; O'Shanessy et al. 2012). Previous studies have also reported FRA to be expressed on carcinomas of the ovary and endometrium, non-small cell lung adenocarcinoma, clear cell renal carcinoma, colorectal carcinoma, and breast carcinoma (Weitman et al. 1992a; Weitman et al. 1992b; O'Shanessy et al. 2011; O'Shanessy et al. 2012; Franklin et al. 1994; Ross et al. 1994; Wu et al. 1999; Bueno et al. 2001; Parker et al. 2005; Shia et al. 2008).

The limited tissue distribution of FRA and its specific expression on certain malignancies makes FRA an attractive target for directed therapies. Indeed, the potential to exploit the differential expression of FRA for targeted cancer therapy has long been appreciated. Two primary approaches have been explored, one involving targeted drug delivery via folate-conjugated therapeutic compounds that binds both FRA and FRB (Low & Kularatne 2009; Dosio et al. 2010), while another approach involves direct targeting and tumor cell death via humanized anti-FRA monoclonal antibodies (Ebel et al. 2007; Konner et al. 2010; Spannuth et al. 2010). Both approaches have advanced to late-stage clinical development in ovarian cancer.

In order to further support the development of such therapeutic strategies, it is important to identify patients who may benefit from FRA-targeted therapy, particularly in cancers where the frequency and degree of expression is not ubiquitous, such as in breast cancer (Franklin et al. 1994; Shia et al. 2008; Stein et al. 1991). Additionally, reports that FRA expression levels may be associated with disease stage or survival in ovarian cancer or non-small cell lung cancer, suggest that FRA may be a useful prognostic marker (O'Shanessy et al. 2012; Toffoli et al. 1997; Toffoli et al. 1998; Iwakiri et al. 2008). Specific and sensitive methods to detect FRA expression in biological samples such as tissue are essential if it is to be pursued as a potential anti-cancer therapy.

Here we describe and define the expression pattern of FRA in breast cancer using a recently reported monoclonal antibody, MAb 26B3 (O'Shannessy et al. 2011) at 2.5 μg/mL diluted in Antibody Diluent (Dako, Carpenteria, CA) or Universal Negative Control [mouse ready-to-use negative control antibody (Dako, for negative isotype tissue)] for 60 min at RT. After washing, slides were incubated with purified MAb 26B3.F2 (O'Shanessy et al. 2011) at 2.5 μg/mL diluted in Antibody Diluent (Dako North America, Inc., Carpenteria, CA) or Universal Negative Control [mouse ready-to-use negative control antibody (Dako, for negative isotype tissue)] for 60 min at RT. After washing, slides were incubated with MACH4 Mouse Probe Primary Antibody Enhancer for 15 min, followed by Polymer-HRP reagent for 20 min, developed with a 3,3'-diaminobenzidine tetrahydrochloride (DAB) solution (Dako) for 5 min and counterstained with hematoxylin (Dako) for 2 min, all incubations being performed at RT. Scoring for staining was performed by a single board-certified pathologist, using customary scoring for intensity and the percent of the tumor stained at each intensity.

Materials and methods

Tissues

Commercial tissue microarray (TMA)

The breast cancer TMA [breast invasive ductal carcinomas (catalog # BR1503a; 72 cases, duplicate cores)] was obtained from US Biomax, Inc. (Rockville, MD). Demographic details for this TMA can be found at: http://www.biomax.us/. These samples were predominantly from women under the age of 60 (range = 19-69 years).

Whole section FFPE slides

Individual formalin-fixed, paraffin-embedded (FFPE) slides were obtained from the archives of Genzyme Genetics. No information is available on the treatment regimens for the patients that contributed the samples analyzed in the present study.

Immunohistochemistry

IHC was performed using FFPE specimens (TMA or whole sections) and a MACH4 Universal HRP-Polymer Detection Kit (Biocare Medical, Concord, CA). FFPE specimens were sectioned at 5 um onto positively-charged glass slides and heated for approximately 60 min at 60°C. Slides were deparaffinized in three sequential baths of xylene for 3 min each, transferred to three sequential baths of 100% alcohol for 3 min each, followed by three sequential baths of 95% alcohol for 3 min each and then rinsed for 5 min in deionized (DI) water. Slides were then pretreated in Diva heat-induced epitope retrieval solution (Biocare Medical) diluted 1:10 in DI water and placed inside a pressurized decloaking chamber already filled with 500 mL of DI water. For antigen retrieval, slides were incubated for 15 min inside the decloaking chamber in which pressurized incubation reaches a maximum of 125°C at 16 PSI for 30 sec and then cooled for 15 min down to 95°C. After cooling to RT, slides were washed in three sequential baths of Tris Buffered Saline/0.1% Tween-20 wash buffer (TBST) for 3 min each and subsequently placed into Peroxidase-1 blocking solution for 5 min at RT. After washing in TBST as above, Background Sniper (Biocare Medical) serum-free universal blocking reagent was applied for 10 min at RT. Slides were then incubated with purified MAb 26B3.F2 (O'Shanessy et al. 2011) at 2.5 μg/mL diluted in Antibody Diluent (Dako, Carpenteria, CA) or Universal Negative Control [mouse ready-to-use negative control antibody (Dako, for negative isotype tissue)] for 60 min at RT. After washing, slides were incubated with MACH4 Mouse Probe Primary Antibody Enhancer for 15 min, followed by Polymer-HRP reagent for 20 min, developed with a 3,3'-diaminobenzidine tetrahydrochloride (DAB) solution (Dako) for 5 min and counterstained with hematoxylin (Dako) for 2 min, all incubations being performed at RT. Scoring for staining was performed by a single board-certified pathologist, using customary scoring for intensity and the percent of the tumor stained at each intensity.
Scoring method
In this study, FRA IHC membrane and intracellular staining intensity was scored as 0, no staining; 1+, weak; 2+, moderate and 3+, strong. The percent of cells staining at each intensity in the sample was also determined. Sections were analyzed under 4x, 10x, 20x and 40x objectives. 3+ strong membrane staining was readily visualized under 4x and confirmed at 10x objective. 2+ moderate membrane staining was visible at 10x and confirmed at 20x, whereas 1+ weak staining required 20x or 40x objectives (Figure 1 a-d). In the presence of 3+ staining, the membrane was thick and occurred at apical and lateral cell borders. In tangential sections, a complete circumferential pattern was evident (Figure 1a). 2+ membrane staining was weaker in intensity and thinner than 3+ membrane staining, usually localized on the apical luminal borders and occasionally on lateral cell borders. 1+ weak membrane staining was generally limited to the luminal borders. The accompanying intracellular staining was variable, depending on the type of tumors.

Figure 1 FRA IHC scoring criteria. a Strong 3+ membrane staining is clearly visible at 10x. The membrane staining is intense, thick and circumferential (x10). b 2+ moderate membrane staining on the luminal borders of malignant cells in a poorly differentiated ductal carcinoma (x20). c 1+ weak membrane staining is seen on the luminal border of poorly differentiated ductal carcinoma cells. It requires 40x objective to visualize the thin, incomplete membrane staining (x40). d No membrane staining is seen in this well-differentiated ductal carcinoma (x20).

Statistical analyses
All statistical analyses were performed using GraphPad Prizm 5 (GraphPad Software Inc., La Jolla, CA).

Positive staining result and TMA core rejection
A sample (TMA core or FFPE slide) was considered positive for FRA expression if the percentage of the tumor area considered by the reading pathologist (Y-S Fu) to be positive for membranous staining was greater than or equal to 5% at any intensity. A TMA core was rejected and therefore not included in the analyses if the reading pathologist determined it was either missing entirely (empty core), was composed of necrotic tissue or was deemed to represent normal tissue. Histopathologic diagnosis of cores was made by the reading pathologist.

The M-Score – a semi-quantitative staining algorithm
The M-score (O’Sheannessy et al. 2012), a metric for staining of each sample was defined as follows:

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Mi = \frac{\sum_{j=1}^{3} w_j x_{ij}}{\sum_{j=1}^{3} w_j} = \frac{\sum_{j=1}^{3} w_j x_{ij}}{6}
\]

In the equation, \(x_{ij}\) is the percentage of tumor stained at intensity \(j\) for patient \(i\) and \(w_j\) is the absolute value of the intensity. The metric has a theoretical range from zero (no positive staining) to 50 (100% 3+). As such, the M-score is a weighted score of FRA IHC tumor cell membrane staining that captures both the proportion of FRA positive cells and staining intensity.

The M-scores for each patient/sample were averaged over duplicate TMA cores, where appropriate. If a determination (core) was void of results, i.e. no tumor present or necrotic tissue, the M-score was assigned to the non-void determinations.

The expression rate for FRA expression was calculated as the proportion of tumors that were stained positive according to the definition of a positive result (≥5% tumor cell membrane staining). This procedure was also applied within specific histology subgroups. Differences for mean values were determined using Fisher’s exact test or one-way ANOVA with post hoc tests controlling for overall type I error.

Results
As previously described (O’Sheannessy et al. 2011), MAb 26B3 is a unique, high affinity antibody shown to be highly specific for FRA with no cross-reactivity to the other three members of this receptor family, namely FRB (folate receptor beta), FRG (folate receptor gamma) or FRD (folate receptor delta). MAb 26B3 has been shown to recognize FRA on FFPE sections of various normal tissues, including breast (O’Sheannessy et al.
Importantly, the staining pattern of FRA by MAb 26B3 was consistent with a membranous localization (Figure 1), although diffuse intracellular staining was also observed. In the absence of membrane staining, intracellular staining was rarely present. Intracellular staining for FRA is expected given that the receptor cycles, carrying folates with it, to the intracellular compartment while remaining membrane associated, by an endocytotic mechanism (Elnakat et al. 2009).

FRA expression on the breast cancer TMA
The distribution of histologies present on the breast cancer TMA are shown in Table 1, the majority (83%) of the cases represented being identified as invasive ductal carcinoma (IDC). The TMA included two normal breast samples, and as previously described (O’Shannessy et al. 2011; O’Shannessy et al. 2012), both were positive for FRA expression as determined by MAb 26B3. Membrane staining of normal breast is restricted to the luminal borders of secretory cells while myoepithelial cells in the outer layer of the duct are negative (Figure 2a). The staining of normal breast is not unexpected in that FRA is secreted into breast milk and believed to be a source of bound folates for the developing embryo (Elnakat & Ratnam 2004).

Of the 71 evaluable cases on the TMA, 21 (30%) were shown to be FRA(+) using the criteria of ≥5% of tumor cells exhibiting membrane staining. Two of three fibroadenoma cases (67%), 0/2 cystosarcoma cases (0%) and 1/6 ductal carcinoma in situ (DCIS) cases (17%) were FRA(+) (Figure 2b). The single invasive lobular carcinoma (ILC) was FRA(−). The small number of samples represented by these histologies precludes any definitive statement with respect to FRA expression rates and further work is warranted. However, of the 59 IDC samples 18 (31%) were FRA(+) (Table 2). In this sample set, no significant differences were noted for FRA expression in IDC relative to stage, nodal status or grade (Table 2), although it should be noted that the number of FRA(+) samples in this analysis is relatively small.

Of the 18 FRA(+) IDC cases, the majority (89%) were Her2(−) suggesting that FRA expression is negatively correlated with Her2 expression in breast cancer (Table 2). Further, of the 18 FRA(+) IDC cases there was a statistically significant difference between ER/PR(+) and ER/PR(−) cases (p = 0.012; Table 2). In addition, of the 4 FRA(+) samples that were also ER/PR(+), all were Her2(−). A statistically significant difference was seen for FRA(+) samples relative to expression of either ER, PR or Her2 versus ER/PR/Her2(−), or triple negative disease (p < 0.0001). Sixty-seven percent of the 18 TNBC specimens express FRA (Figure 3). Taken together, these data support the claim that FRA expression is enriched in the TNBC subtype and may represent a novel molecular subtype of breast cancer.

| Histology | FRA positive | FRA negative | Total |
|-----------|--------------|--------------|-------|
| Normal    | 2 (100%)     | 0 (0%)       | 2     |
| Fibroadenoma | 2 (67%)     | 1 (33%)      | 3     |
| Cystosarcoma | 0 (0%)      | 2 (100%)     | 2     |
| DCIS – Ductal carcinoma in situ | 1 (17%) | 5 (83%) | 6 |
| ILC – Invasive lobular carcinoma | 0 (0%) | 1 (100%) | 1 |
| IDC – Invasive ductal carcinoma | 18 (31%) | 41 (69%) | 59 |
| Total samples | 21 (30%) | 50 (70%) | 71 |

FRA folate receptor alpha, TMA tissue microarray.
The TMA described above was composed primarily of early stage breast cancers: stage I, 6/60 (10%); stage II, 44/60 (73%); stage III, 10/60 (17%); no metastatic breast cancer cases were represented on the TMA (Table 3). Therefore, to confirm and extend the results obtained on the TMA, we identified 61 archival FFPE tissue blocks from stage IV (T4) Her2(−) breast cancers with known ER/PR expression ranging from 0-100% positive. Importantly, all 61 of these samples were from metastases, not the primary tumor.

FRA expression was found in 22/61 (36%) of these patients, demonstrating that the percent of FRA(+) specimens/tumors determined in early stage disease is retained in late stage metastatic disease, at least in a Her2(−) population (TMA expression rate = 35%; stage IV metastatic disease = 36%). Of the 22 FRA(+) stage IV metastatic cases, only three (14%) showed any expression level for ER/PR which tended to be in the low range (up to 30%). As such, 19/22 (86%) FRA(+) patients were of the TNBC molecular subtype. As with the data obtained in early stage disease on the TMA, triple negative samples are overrepresented in the FRA(+) population. However, both data sets support the assertion that FRA staining associated more strongly with triple negative disease. In metastatic disease, there was a significant difference between FRA(+) patients that were also ER/PR(+) or ER/PR(−) (p = 0.0054), as seen on the TMA, but also a significant difference in FRA expression by grade of disease (grade 1 or grade 2 versus grade 3, p = 0.037).

The semi-quantitative M-score (O'Shannessy et al. 2012) was also used to analyze the pattern of FRA staining (intensity and percent of tumor cells) of metastatic breast cancer samples. While a significant difference could be demonstrated between the ER/PR(+) versus the ER/PR(−) populations (Figure 4a; p = 0.0029), no such difference was evident for grade of disease either across the entire population (Figure 4b) or within the FRA(+) population (Figure 4c). These data are consistent not only with FRA expression overall, but also with the data described for the early stage TMA samples. Taken together, the present data demonstrate a strong association between FRA expression and TNBC and further support the notion that FRA(+) breast cancer may represent a new molecular subtype of this disease.

Analysis of individual slides from Her2(−) metastatic breast cancer patients

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The samples represented in the FFPE blocks from stage IV metastatic disease were obtained from a

| Variable                  | FRA positive | FRA negative | Total | P valuea |
|---------------------------|--------------|--------------|-------|----------|
| Marker                    | N (%)        | N (%)        |       |          |
| ER/PR(+)                  | 4 (14%)      | 24 (86%)     | 28    |          |
| ER/PR(−)                  | 14 (45%)     | 17 (55%)     | 31    | 0.012    |
| Her2(+)                   | 2 (15%)      | 11 (85%)     | 13    |          |
| Her2(−)                   | 16 (35%)     | 30 (65%)     | 46    | 0.307    |
| ER/PR/Her2(−)             | 12 (67%)     | 6 (33%)      | 18    | <0.0001  |
| TNM Classification        |              |              |       |          |
| T1                        | 3 (43%)      | 4 (57%)      | 7     |          |
| T2                        | 10 (26%)     | 29 (74%)     | 39    |          |
| T3                        | 5 (63%)      | 3 (37%)      | 8     |          |
| T4                        | 0 (0%)       | 5 (100%)     | 5     |          |
| Nodal Status              |              |              |       |          |
| NO                        | 18 (35%)     | 33 (65%)     | 51    |          |
| N1/N2b                    | 0 (0%)       | 8 (100%)     | 8     | 0.092    |
| Tumor Grade               |              |              |       |          |
| Grade 1                   | 1 (14%)      | 6 (86%)      | 7     |          |
| Grade 2                   | 12 (36%)     | 21 (64%)     | 33    | 0.393    |
| Grade 3                   | 5 (26%)      | 14 (74%)     | 19    | 0.6465c  |

IDC invasive ductal carcinoma, TMA tissue microarray, FRA folate receptor alpha, ER estrogen receptor, PR progesterone receptor, Her2 human epidermal growth factor receptor type 2, TNM tumor node metastasis.

a P values calculated via 2X2 contingency table analysis using Fisher’s exact test.

b 4/8 (50%) of N1/N2 samples were Her2(+).

c Grade 1 versus grade 3.

http://www.springerplus.com/content/1/1/22
number of metastatic sites including lymph node, bone, skin and liver, as well as fluid and fine needle aspirate (FNA) samples obtained primarily from pleura and paracentesis. Several of these ‘fluid biopsies’ were stained positive for FRA (Figure 5) suggesting the general applicability of the described IHC methodology to multiple samples types. However, given the small sample numbers in the present study, additional work on the suitability of FNAs as a FRA IHC diagnostic sample source are warranted. It should be noted, however, that FNAs have also been demonstrated to be positive for FRA expression by IHC in non-small cell lung adenocarcinoma (O’Shannessy et al. 2012).

Discussion

Currently, breast cancer is grouped into three subtypes that dictate therapy: ER and/or PR expressing, Her2 expressing, or TNBC which implies the absence of these three receptors. Hormone responsive breast tumors represent approximately two-thirds of all breast cancers (Bauer et al. 2007). Treatment at some point during the course of therapy will necessarily include hormonal agents. Endocrine therapy has been shown to both prolong life and decrease treatment related morbidity. Her2 overexpressing breast cancer represents approximately 15-20% of breast cancers (Bauer et al. 2007). Although treatment with trastuzumab has vastly improved survival in patients with Her2 positive tumors, the prognosis for this group remains worse than for their hormone receptor expressing counterparts. The balance of breast cancers lack all three of these receptors and are referred to as triple negative.

It is unclear from the current investigation if the FRA (+) subset represents a particular TNBC histotype. It is conceivable that the FRA expressing cancers may fall into the basal-like category that are also known to sometimes be BRCA(+), characterized high grade infiltrating ductal carcinomas that can show necrosis (Metzger-Filho et al. 2012), or by contrast the luminal androgen receptor subtype that often present with bone and lymph node involvement but are more indolent by nature (Gelmon et al. 2012).

There are no currently available targeted therapies for TNBC and their prognosis remains decidedly worse than those with hormone receptor expression. Recently, PARP inhibitors were assessed in this TNBC patient population with the hope of improving survival and outcome. However, a phase III randomized clinical trial

Table 3 Distribution of FRA expression relative to hormone receptor status and tumor grade in a Her2(−) metastatic breast cancer cohort

| Variable           | FRA positive | FRA negative | Total | P value<sup>b</sup> |
|--------------------|--------------|--------------|-------|--------------------|
| Marker             | N (%)        | N (%)        |       |                    |
| ER/PR(+)           | 3 (14%)      | 20 (86%)     | 23    | 0.0054             |
| ER/PR/Her2(−)      | 19 (50%)     | 19 (50%)     | 38    |                    |
| Tumor Grade        |              |              |       |                    |
| Grade 1            | 3 (30%)      | 7 (70%)      | 10    | 1.0                |
| Grade 2            | 11 (28%)     | 28 (72%)     | 39    |                    |
| Grade 3            | 8 (67%)      | 4 (33%)      | 12    | 0.037              |
| Total Samples      | 22 (36%)     | 39 (64%)     | 61    |                    |

FRA folate receptor alpha, Her2 human epidermal growth factor receptor type 2, ER estrogen receptor, PR progesterone receptor.

<sup>a</sup> All samples were whole tissue FFPE (formalin-fixed, paraffin-embedded) slides.

<sup>b</sup> P values calculated via 2X2 contingency table analysis using Fisher’s exact test.
failed to show benefit (O’Shannessy et al. 2011). Treatment for this molecular subtype of breast cancer remains untargeted cytotoxics.

FRA is an interesting and relevant biomarker in cancer therapy. This GPI-anchored protein at least in part serves to bind and transport folate, primarily 5-methyltetrahydrofolate, the predominant plasma folate, into cells. However, its expression is not necessary for cytosolic folate accumulation and purine and pyrimidine synthesis as there are multiple portals of folate entry, primarily the reduced folate carrier (RFC), ubiquitous on all mammalian cells (Elnakat & Ratnam 2004). Hence, blocking or disrupting FRA will not a priori deprive a cell of necessary folate and kill it. However, FRA has been reported to impart a growth advantage to cells expressing the receptor, especially in low folate environments (Luhrs et al. 1992). Furthermore, FRA’s high expression on epithelial malignancies, compared to normal tissues, provides the rationale for its use as a potential targeted cancer therapy. FRA targeted therapies are currently in clinical investigation in both ovarian and lung cancers.
and it is interesting to speculate on the potential value of such an approach in FRA expressing breast cancer. It is worth repeating that FRA is expressed in normal breast, restricted to the luminal borders of secretory cells, consistent with the secretion of FRA into breast milk. No other breast cell type has been shown to express FRA. Such an expression pattern in normal breast tissue was seen in the present study although it should be noted that not all samples, especially TMA cores, contain normal tissue. The expression of FRA in carcinomas may reflect the cellular origin of the carcinoma and may, at least in part, explain the approximately 30% incidence in IDC demonstrated here.

Hormone receptor positive tumors appear to have a lower incidence of FRA expression. FRA expression has been shown to be regulated by steroid hormones, particularly estrogens (Rochman et al. 1985; Kelley et al. 2003). Specifically, 17β-estradiol has been demonstrated to down-regulate FRA expression by direct action of the estrogen receptor on the P4 promoter of FRA suggesting a negative correlation between the expression of ER and FRA. The data presented herein is in line with these findings in that there was a clear association between FRA(+) and ER(+) samples, i.e. ER(−) samples were significantly more likely to be FRA(+).

A recent report (Hartmann et al. 2007) on a cohort of 63 invasive breast cancers, roughly equally distributed between good and poor outcome, demonstrated that strong FRA staining was highly associated with poor outcome. While the authors did not report on Her2 status of these patients, it is interesting to speculate that the poor outcome group may have included a significant TNBC population. Studies are underway to assess the prognostic significance of FRA expression in breast cancer and any relationship to molecular subtype and histotype.

Our data, although limited by a small sample size, demonstrate that approximately 30% of breast cancers express FRA and suggest that as many as 70–80% of stage IV metastatic TNBC tumors express this receptor. FRA expressing breast cancer may, therefore, represent an important and clinically significant subset of breast cancer and in particular triple negative disease. FRA targeted therapies, alone or in combination with cytotoxics, may represent a novel approach to treatment for this disease with a high unmet medical need.

Further work is clearly needed to confirm and extend the present findings and to provide a clinical correlation with FRA expression. Additional correlations with disease progression and response to treatment would be interesting and potentially valuable in disease monitoring and response assessments.

Abbreviations
FRA: Folate receptor alpha, folate receptor α; FRβ: Folate receptor beta, folate receptor β; FRG: Folate receptor gamma, folate receptor γ; FRD: Folate receptor delta, folate receptor δ; IHC: Immunohistochemistry; MAb: Monoclonal antibody; FFPE: Formalin-fixed, paraffin-embedded; FNA: Fine needle aspirate; TMA: Tissue microarray; IDC: Invasive ductal carcinoma; DCIS: Ductal carcinoma in situ; ER: Estrogen receptor; PR: Progesterone receptor; Her2: Human epidermal growth factor receptor type 2; TNBC: Triple negative breast cancer; TNM: Tumor node metastasis.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
DO drafted the manuscript, developed and contributed the antibody, and performed the statistical analysis. ES conceived the project, and acquired and validated the samples. JM contributed the clinical relevance of the data. RS had the immunohistochemical staining done in his laboratory, and compiled the raw data. YF read and interpreted the immunohistochemical staining results. All authors read and approved the final manuscript.

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References
Bauer KR, Brown M, Cress RD, Parise CA, Caggiano V (2007) Descriptive analysis of estrogen receptor (ER)-negative, progesterone receptor (PR)-negative, and HER2-negative invasive breast cancer, the so-called triple-negative phenotype: A population-based study from the California Cancer Registry. Cancer 109:1721–1728
Bueno R, Appasani K, Mercer H, Lester S, Sugarbaker D (2001) The alpha folate receptor is highly activated in malignant pleural mesothelioma. J Thorac Cardiovasc Surg 121(2):225–233
Dosio F, Millis P, Cattel L (2010) EC-145, a folate-targeted Vinca alkaloid conjugate for the potential treatment of folate receptor-expressing cancers. Curr Open Investig Drugs 11(12):1424–1433
Ebel W, Routhier EL, Foley B, Jacobs S, McDonough JM, Patel RK, Turchin HA, Chao Q, Kline JR, Old LJ, Phillips MD, Nicolaides NC, Sass PM, Grasso L (2007) Preclinical evaluation of MORAb-003, a humanized monoclonal antibody antagonizing folate receptor-alpha. Cancer Immun 7:6
Elnakat H, Ratnam M (2004) Distribution, functionality and gene regulation of folate receptor isoforms: implications in targeted therapy. Adv Drug Deliv Rev 56(8):1067–1084
Elnakat H, Goniit M, Salazar MD, Zhang J, Basur V, Gunning W, Kamen B, Ratnam M (2009) Regulation of folate receptor internalization by protein kinase C alpha. Biochemistry 48(34):8249–8260
Franklin WA, Waintrub M, Edwards D, Christensen K, Prendegrant P, Woods J, Bunn PA, Kolhouse JF (1994) New anti-lung-cancer antibody cluster 12 reacts with human folate receptor present on adenocarcinoma. Int J Cancer Suppl 889–95
Gelmon K, Dent R, Mackey JR, Laing K, McLeod D, Verma S (2012) Targeting triple negative breast cancer: optimising therapeutic outcomes. Ann Oncology, Epub ahead of print
Hartmann LC, Keeney GL, Lingle WL, Christianson TJ, Varghese B, Hillman D, Obreg AL, Low PS (2007) Folate receptor overexpression is associated with poor outcome in breast cancer. Int J Cancer 121(5):938–942
Iwakiri S, Sonobe M, Nagai S, Hirata T, Wada H, Miyahara R (2008) Expression status of folate receptor alpha is significantly correlated with prognosis in non-small-cell lung cancers. Ann Surg Oncol 15(3):889–899
Kelermen LE (2006) The role of folate receptor alpha in cancer development, progression and treatment: cause, consequence or innocent bystander? Int J Cancer 119(2):243–250
Kelley KM, Rowan BG, Ratnam M (2003) Modulation of the folate receptor alpha gene by the estrogen receptor: mechanism and implications in tumor targeting. Cancer Res 63(11):2820–2828
Konner JA, Bell-McGuinn KM, Sabbatini P, Herskoy ML, Tew WP, Pandit-Taskar N, Vander Els N, Phillips MD, Schweizer C, Weil SC, Larson SM, Old LJ (2010) Farletuzumab, a humanized monoclonal antibody against folate receptor alpha, in epithelial ovarian cancer: a phase I study. Clin Cancer Res 16(21):5288–5295. doi:10.1158/1078-0432.CCR-10-0709

Low PS, Kularatne SA (2009) Folate-targeted therapeutic and imaging agents for cancer. Curr Opin Chem Biol 13(3):256–262. doi:10.1016/j.copp.2009.03.022

Luhrs C, Raskin CA, Dubin R, Wu B, Sadatavani E, Rothenberg SP (1992) Transfection of a glycosylated phosphatidylinositol-anchored folate-binding protein complementary DNA provides cells with the ability to survive in low folate medium. J Clin Invest 90(3):840–847

Metzger-Filho O, Tutt A, de Azambuja E, Saini KS, Viale G, Loi S, Bradbury I, Bliss IM, Azim HA Jr, Ellis P, Di Leo A, Baselga J, Sotiriou C, Piccart-Gebhart M. (2012) Dissecting the heterogeneity of triple negative breast cancer. J Clin Oncol 30(15):1879–1887

O'Shannessy DJ, Somers EB, Albone E, Cheng X, Park YC, Tomkowicz BE, Hamuro Y, Kolf TO, Forsyth TM, Smaa E, Fu Y-S, Nicolaides NC (2011) Characterization of the human folate receptor alpha via novel antibody-based probes. Oncotarget 2(1):227–1243

O'Shannessy DJ, Yu G, Smale R, Fu Y-S, Singhal S, Thiell RP, Somers EB, Vachani A (2012) Folate Receptor Alpha Expression in Lung Cancer: Diagnostic and Prognostic Significance. Oncotarget 4(4):414–425

O'Shaughnessy J, Schwartzberg LS, Danso MA et al (2011) A randomized phase III study of iniparib (BSI-201) in combination with gemcitabine/carboplatin (G/C) in metastatic triple-negative breast cancer. J Clin Oncol 29(Supplement 15): Abstract 1007

Parker N, Turk MJ, Westrick E, Lewis JD, Low PS, Leamon CP (2005) Folate receptor expression in carcinomas and normal tissues determined by a quantitative radioligand binding assay. Anal Biochem 338(2):284–293

Rochman H, Selhub J, Karrison T (1985) Folate binding protein and the estrogen receptor. J Nutr 115(11):1671–1715

Ross JF, Goldenberg DM, Mattes MJ (1991) Normal tissue reactivity of four anti-folate receptor monoclonal antibodies of clinical interest. Int J Cancer 47(2):163–169. doi:10.1002/ijc.2910470202

Shaia J, Klimstra DS, Nitzkorski JR, Low PS, Conen M, Landmann R, Weiser MR, Franklin WA, Prendergast FG, Murphy L, Tang LH, Temple L, Guillem JG, Wong WO, Paty PB (2008) Immunohistochemical expression of folate receptor alpha in colorectal carcinoma: patterns and biological significance. Hum Pathol 39(4):498–505

Siegel R, Naishadham D (2012) Jemal A (2012) Cancer statistics, CA Cancer J Clin 62:10–29

Spannuth WA, Sood AK, Coleman RL (2010) Farletuzumab in epithelial ovarian carcinoma. Expert Opin Biol Ther 10(3):431–437. doi:10.1517/14712591003355069

Stein R, Goldenberg DM, Mattes MJ (1991) Normal tissue reactivity of four anti-tumor monoclonal antibodies of clinical interest. Int J Cancer 47(2):163–169. doi:10.1002/ijc.2910470202

Toffoli G, Cennigoli C, Russo A, Gallo A, Bagnoli M, Bolocchi M (1997) Overexpression of folate binding protein in ovarian cancers. Int J Cancer 74(2):193–198

Toffoli G, Russo A, Gallo A, Cennigoli C, Miotti S, Soro R, Tumolo S, Bolocchi M (1998) Expression of folate binding protein as a prognostic factor for response to platinum-containing chemotherapy and survival in human ovarian cancer. Int J Cancer 79(2):121–126

Weitman SD, Lark RH, Coney LR, Fort DW, Frasca V, Zurawski VR, Kamen BA (1992a) Distribution of the folate receptor GP38 in normal and malignant cell lines and tissues. Cancer Res 52(23):3396–3401

Weitman SD, Weinberg AG, Coney LR, Zurawski VR, Jennings DS, Kamen BA (1992b) Cellular localization of the folate receptor: potential role in drug toxicity and folate homeostasis. Cancer Res 52(23):6708–6711

Wu M, Gunning W, Ratnam M (1999) Expression of folate receptor type alpha in relation to cell type, malignancy, and differentiation in ovary, uterus, and cervix. Cancer Epidemiol Biomarkers Prev 8(9):775–782

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