



KEY ROLE OF EGFR, ErbB2 AND PR PROTEINS IN GENE AND PROTEIN INTERACTIONS WITH VIMENTIN, FOXC1/2, FAK, AND THE BRCA1 IN PREVENTION OF TRIPLE NEGATIVE BREAST CANCER

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ABSTRACT Studies on protein–protein interactions and RNA–protein interactions between transcription factors and receptors that act as tumor suppressor proteins play key roles in designing new anticancer drugs for treatment. The BRCA1 RING domain exhibits protein–protein interactions with Vimentin, FOXC2, Focal adhesion kinase and EGFR, whereas the EGFR nucleic acid contig shows gene–protein interactions with Vimentin, FOXC2, Focal adhesion kinase and BRCA1. ErbB2 and the progesterone receptor play key roles in protein–protein interactions with the BRCA1 RING domain and ErbB2 in TNBC, and the PR nucleic acid contig shows gene–protein interactions with vimentin, FOXC2, focal adhesion kinase and the BRCA1 RING domain, clearly indicating that the transaction of proteins occurs mostly although noncovalent interactions.

KEYWORDS : BRCA1 RING domain, Vimentin, FOXC2, Focal adhesion kinase and BRCA1, Protein–protein interactions, Gene–protein interactions

INTRODUCTION:

BRCA1 (breast cancer gene 1) and BRCA2 (breast cancer gene 2) repair damaged DNA, and usually, offspring inherit the protein from their parents, with each gene being from their paternal or maternal partner. Mutated BRCA1 and BRCA2 are responsible for more than 60% of breast or ovarian cancers in inherited individuals. One of the important factors to be considered is genetic predisposition to the disease, and studies on mutations that genetically predispose BRCA1 and BRCA2 to mutations can prevent cancer-related deaths¹.

The nongenetic factors that are likely to contribute to breast cancer include early menopause, alcohol and tobacco consumption, exposure to radiation, obesity, decreased physical activity, urbanization, a sedentary lifestyle, a high-fat diet, frequent spontaneous miscarriages, lack of breastfeeding, hormone replacement therapy, aging, geographical location, socioeconomic conditions, reproductive events, exogenous hormones, breast density, and family history of breast cancer or other cancers^(2,3,4-14).

Human epidermal growth factor receptor 2 (HER2) is overexpressed in approximately 20-30% of cancers and can result in more aggressive cancers with a high recurrence rate and increased mortality. Trastuzumab is a HER2 receptor blocker used to treat most HER2-positive cancers, and emerging patterns of trastuzumab resistance have been recorded earlier; however, the development of therapeutic agents, such as monoclonal antibodies and other targeted therapies, can overcome trastuzumab resistance and help in the treatment of most HER2-positive breast cancers^{15,16}.

The progesterone receptor modulates estrogen receptor expression, and PR expression depends strongly on estrogen and can help improve the prognosis in most TNBC patients with a positive PR17. The estrogen receptor and progesterone receptor are mostly expressed in approximately 15-30% of luminal epithelial cells, and studies have shown that the PR also contains portions of ER α . Studies by Rachel Schiff revealed that ER+/PR- tumors share gene profiles with both the ER+/PR+ and ER-/PR- luminal subtypes, and patients with these genotypes have a poor prognosis concluded by Perou¹⁸.

METHODOLOGY:

Protein–protein interactions were studied using the PDB ids of RCSB through the H dock server, and RNA–protein interactions were studied by developing nucleic acid contigs from the protein sequence and through the H Dock server by selecting ssRNAs. A model with a low positive free energy is selected from the top 10 predicted models, and

receptor–ligand interface data with negative free energy are captured.

1. The amino acid contig of EGFR:

SGSGEAPNQALLRLKETEFKKIKVLGSGAFGTVYKGLWIPE
GEKVKIPVAIKELREATSPKANKEILDEAYVMASVDNPHVCR
LLGICLTSTVQLITQLMPFGCLLDYVREHKDNIGSQYLLNWC
VFVVCVQIAKGMNYLEDRRHRDLAARNVLKTPQHVKITDFGL
AKLLGAEKEYHAEGGKVPIKWMALESILHRIYTHQSDVWS
YGVTVWELMTFGSKPYDGIPASEISSILEKGERLPQPICTIDV
YMIMVMACWMIDWGDERMHLPSPDTSNFYRALMDEEDMD
DVVDADDEYLIPQQG

[RNA contig of EGFR generated with seq id 7EAI (RCSB PDB):
GGUUCUGGUGAAGCUCCAAUGAAGCUCUUCUUAAGAAU
UCUUAACAACAAUAAUUUUAAAAAAUUAAGUUCUUG
GUAGUGGUGCUUUUGGUACUGUUUAUAAAGGUCUUUGG
AUUCCUCAAGGUCAAAAAGUUAUUUCCUGUUGCUAA
AACAACUUCGUCAAGCUACUAGUCCUAAAGCUAAUAAAA
CAAAUUAAGAUAAGCUUAUGUUUAUGGCUAGUGUUGA
UAAUCCUCAUGUUUGUCUGCUUCUUGGUUUUUGUCUUA
CUUAUGUACUGUUGAACUUUUACUGAACUUAUGCCUU
UUGGUUGUCUUCUUGAUUAUGUUCGUCAACAUAAUUAU
AAUAUUGGUUCUGAAUAUCUUUUAAACUGGUGUCGAGA
UCGCCAAGGGCAUGAACUACCUACCUACAGGAUCGACGG
CUAGUGUUCUUCUAAAA

2. Amino acid contig of HER2/erbB2:

MKFLVNVALVFM-YYISYIYADYKDDDDKHHHHHHHHHHHL
EVLFQGPYPYDVPDYATQVCTGTDMKRLRPASPETHLDMLR
HLYQGCQVQGNLEYLPTNASLSFLQDIQEVQGYVLIHNNQ
VRQVPLQRLRIVRGTLQFEDNYALAVLDNGDPLNNTTPVTG
ASPGGLRELQRLSLTEILKGGVLIQRNPQLCYQDTILWKDIFH
KNNQLALTLIDTNRSRACHPCSPMCKGSRCWGESSEDCQSLT
RTVCAGGCARCPALPTDCCHEQCAAGCT

{RNA Contig sequence of HER2/erbB2 generated with SNP id:9QBF:RCSBPDB: Due to the large size of the contig, I limited the writing of the RNA contig to first 276 aminoacids (AUGAAUUU CUUGUUAUUGUCGUCUCUGUAUUCUUGUUGUUAUUAAGUUAACUUAUGCCGAUUACAAGGACGACGACGAUA AACACCACCAUCAUACACCACCAUCAUACACCUACAAG UCCUCUUUGAAGGUCCUUAACCAUACGUUCCAGACUUAUG CCACGAAGUGUGUACUGGUACAGACAGAAACUUGCGGCC UACCUGCAAGUCCACAAACGCACUUAUGUUGCGACAUCU UUACGAAGGAUGUGAGGUAGUAGACGGAAACCUACAGC

UAAACAUUCUGCCAACAAACGCCAGCUUCCUCGAAGACA
UAGAACAGGUUGAAGGAUACGUGCUGAUAGCUCACAAC
GAAGUUCGUGAAGUCCUCUAGCGACUGCGUAUGCGUA
UUGUACGGCACAGAACUCUUUCAGGACAACUAUUGUGC

3. Amino acid contig of the PR:

GQDIQLIPPLINLLMSIEPDYAGHDHHDNTKPDTSLLTSLN
QLGERQLSVVWWSKSLPGFRNLHIDDQITLIQYSWMSLMVF
GLGWRYSYKHVSGQMLYFAPDLILNEQRMKESSFSYSLCTMW
QIPQEFVKLQVSQEFLCLLLLLNTIPLEGLRSQTQFEEMRSS
YIRELIKAIGLRQKGVVSSSRFYQLTKLLDNLDLVKQLHLHY
CLNTFIQSRALSVEFPEMMSEVIAAQLPKILAGMVKPLLFHK
K.

[RNA contig sequence of the progesterone receptor was built with a sequence id: 1A28: RCSBPDB: GGCGAGGACAUUGAACUAAUA
CCACCGCUAAUAAUACUGAUGUCAAUACAACCUGAC
GUCAUAAUAGCCGGACACGAUAAACACUAAACCAGAUAC
UUCUUCUUCUCCUACUCACCUCUGAACGAGCUGGGGCC
AACGUGAACUACUCAGCGUUGUAAAUGGAGCAAAAGU
CUCUUGGCUUUCGCAUUCUACAUAUAAUAGACGAAAU
AACACUAAUAGAAUAAUAGUUGGAUGUCUCUAAUGGUCU
UUGGCCUAGGCUGGCGCUCAUAAACAUGUUUCCGGC
GACAUGUAAUACUUUGCCCCGACCUCUACUAAUAAUCAA
GAACGUAUGAAGCAAUCUCCUUCUAAUUCUCUAAUGUCU
AACAAUGUGGGAAAUUCCUGAACAGUUUGUAAACUAG
AGGUAUCAGAACAACAAUCCUCUGUAUGAAGGUUAUC
UCCUGCUAAAUACUAAUCCA

RESULTS:

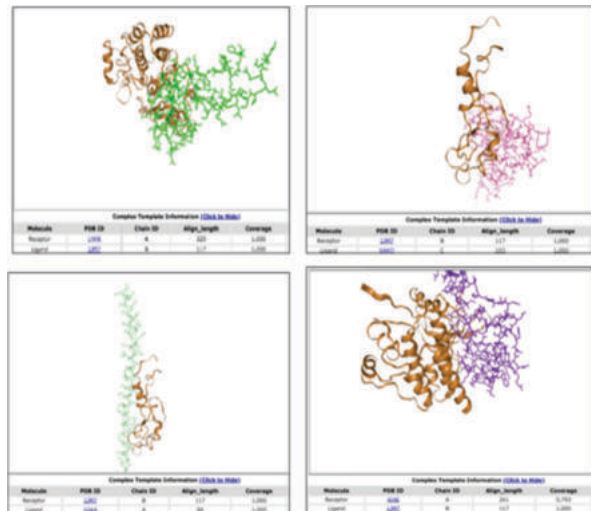


Figure 1 Protein-protein interactions between the BRCA1 RING domain and EGFR (A), FOXC2 (B), vimentin (C) and focal adhesion kinase (D).

Figure 1A shows that the BRCA1 RING domain participates in protein-protein interactions with epidermal growth factor receptor (EGFR) at the receptor-ligand interface with the amino acid pair ASP-ASN; ASN-PRO, ASN-VAL; ARG-PHE, ARG-CYS; SER-CYS; SER-PRO; PHE-PRO; PHE-ILE; and GLU-ARG, which are involved in major van der Waals interactions and ionic interactions. The amino acid pairs with low energy values were recorded to reduce the ambiguity in the data. Figure 1B shows the protein-protein interactions between BRCA1 and FOXC2 for the amino acid pairs CYS-SER, PRO-ARG, VAL-SER, VAL-PHE, CYS-GLY, THR-ASN, PRO-GLN, ASN-ASP, LEU-ASN, LEU-ARG LEU-ASP, HIS-ARG, and HIS-PRO at the receptor-ligand interface with van der Waals and ionic interactions. Figure 1C shows that the vimentin and BRCA1 ligand-receptor interfaces contain amino acid pairs such as ARG-VAL, ARG-GLU, LYS-TYR, ARG-GLU, ARG-TYR, LEU-TYR, LEU-HIS, ASP-GLU, GLU-ILE, GLU-MET, LYS-ARG, GLU-ARG, and ARG-ASN with van der Waals and ionic interactions. Figure 1D Focal adhesion kinase (FAK) shows protein-protein interactions with BRCA1 at interfaces containing amino acids such as ASP-ILE, PRO-ILE, GLN-ARG, VAL-ILE, LYS-ILE, TYR-ASN, ASP-ARG, GLN-ASP, ARG-LEU, SER-LYS, ALA-ARG, ARG-LYS, and GLN-GLN, which exhibit major van der Waals interactions and ionic interactions with few recorded hydrophobic interactions.

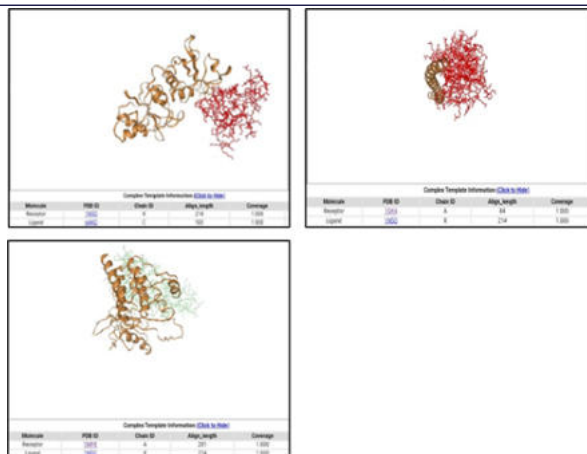


Figure 2: Protein-protein interactions of the total BRCA1 contig with FOXC2, vimentin and focal adhesion kinase.

Figure 2A shows the results of protein interaction studies between the BRCA1 total amino acid contig and FOXC2, which revealed interactions between amino acids at the binding surface with low energy values recorded at HIS-ASN, HIS-LEU, THR-ASN, GLU-ASN, ASP-GLN, ASP-LYS, TYR-GLN, LEU-SER, and PRO-TYR, revealing ionic and van der Waals interactions. Figure 2B shows that the BRCA1 and vimentin pairs had low energy values for amino acid pairs such as LEU-LYS, THR-ASN, LEU-GLU, MET-MET, GLU-ARG, GLU-THR, PHE-GLY, GLU-GLN, TYR-LYS, ILE-GLN, ILE-VAL, GLN-VAL, and GLN-LYS through noncovalent interactions such as hydrophobic interactions, van der Waals forces and ionic interactions. As shown in Figure 2C, BRCA1 interacts with focal adhesion kinases at the amino acid positions ARG-HIS, ILE-HIS, GLN-LYS, GLN-HIS, LEU-HIS, THR-ARG, THR-HIS, GLY-ARG, GLU-ARG, VAL-SER, ARG-LEU, SER-THR, SER-ASN, and SER-LEU at the binding interface with vascular forces more strongly than with ionic interactions.

Figure 3A shows that BRCA1 interacts with EGFR at the 21st position Lys with the c1 position of adenine (588), the C1 position of uracil at the 654th position with ARG (24), the 27th position of ARG with the C4 position adenine, the 33rd position Arg with the N3 position cytosine (504), the 37th position Cys with cytosine at N2 (536) and the C3 position cytosine (537). At position 72, Ile with OP2 was adenine (547), and at position 83, Cys was adenine at position C6 (628).

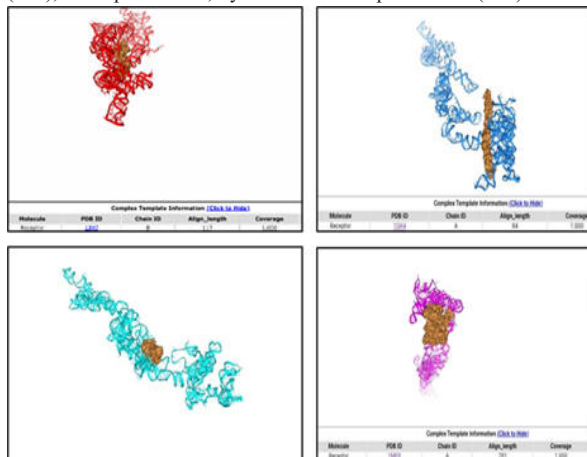


Figure 3 Protein-gene interactions between the BRCA1 ring domain and the EGFR contig (A), between vimentin and the EGFR contig (B), between FOXC2 and the EGFR contig (C) and between FAK and EGFR (D).

Figure 3B shows that EGFR interacts with vimentin at 31 Tyr with 28 cytosine residues, 35 Ile with 13 guanine residues and 14 cytosine residues, and the 38th position of LEU with 782,783 adenine and uracil residues. 39, Gln with 15 and 783 positions of uracil; 45, MET with 713 positions of adenine; 72, Thr with 887 positions of uracil; 75, Lys; and 76, Leu with 888 adenine and 867 guanine. The 76 position Leu with the 887 position uracil in the gene.

As shown in Figure 3C, EGFR interacts with FOXC2 at the 45th position of Gln with the 666th position uracil, at the 52nd position Arg with 160 and 161th positions cytosine and uracil, at the 53rd position His with adenine (136) and guanine (667), at the Ser (56) and Leu (57) with guanine (125) and at the Ser (56) with uracil (162), at the Leu (57) with uracil (135) and guanine (164), at the Glu (59) with uracil (134) and guanine (165), at the Lys (63) with uracil (162) and cytosine (163), at the Asp (82) with adenine (183), at the Arg (97) with uracil (171) and guanine (184) of the nucleic acid sequence.

According to Figure 3D, EGFR interacts with focal adhesion kinase at sites such as Tyr (165) with adenine (760), Lys (176) with guanine (748), Ile (189) with uracil (749), and Arg (192) with guanine (748), and positions with a free energy change less than 2.0 are recorded and mentioned here.

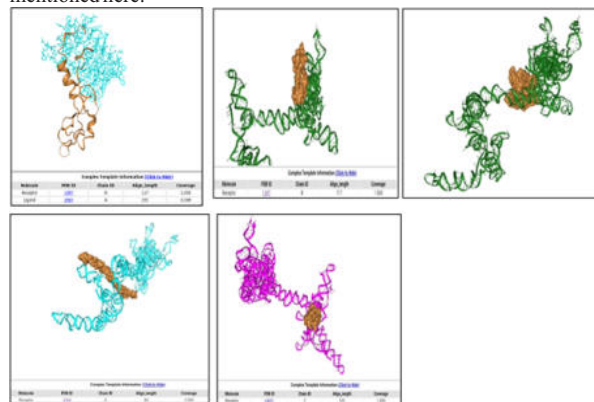


Figure 4 Protein-protein interactions between BRCA1 and HER3 (seq id:3P0Y) (A) and gene-protein interactions between erb2 and BRCA1 with seq.id 2A91 (B), between erbB2 and FOXC2 with seq.id 2A91 (C), between erbB2 and vimentin with seq.id 2A91 (D) and between erbB2 and FAK with seq.id 2A91 (E).

Figure 4A shows that HER3 proteins interact with the BRCA1 ring domain at amino acid pairs such as Cys (41) - Asn (417), Asp (97) - Gln (425), Asp (97) - His (449), Glu (100) - His (416) & Asn (417), Pro (103) - Leu (415) & Asn (417), Arg (104) - Val (332), and Phe (108) - Leu (415) at the receptor-ligand interface through ionic interactions followed by van der Waals and dipole-dipole interactions.

As shown in Figure 4B, the ErbB2 gene interacts with the BRCA1 protein at various sites, such as Phe (45) with uracil (347), Cys (46) and Pro (59) with guanine (346), Thr (63) with guanine (367), Trp (66) with cytosine (750), Asn (88) with guanine (751), His (91) with adenine (344), and Asn (93) with uracil (343).

As shown in Figure 4D, the ErbB2 gene interacts with vimentin at sites such as Arg (15 and 18) at cytosine (162), Glu (27) at uracil (12), Tyr (31) at uracil (11), Tyr (31) at adenine (255) and guanine (256), and Gln (39) at guanine (260).

Figure 4E shows that ErbB2 and FOXC2 have gene-protein interactions at Tyr (165) with adenine (760), Lys (176) and Arg (192) with guanine (748), and Ile (189) with uracil (749). As shown in Figure 4C, Focal adhesion kinase interacts with ErbB2 at the nucleotide positions Phe (191) and uracil (554), Asn (224) and Asp (225) and adenine (449), Arg (229) and adenine (447), Glu (231) and uracil (555), Glu (234) and adenine (321) and uracil (446), and Met (238) and Tyr (247) with adenine (323), Tyr (256) and uracil (555).

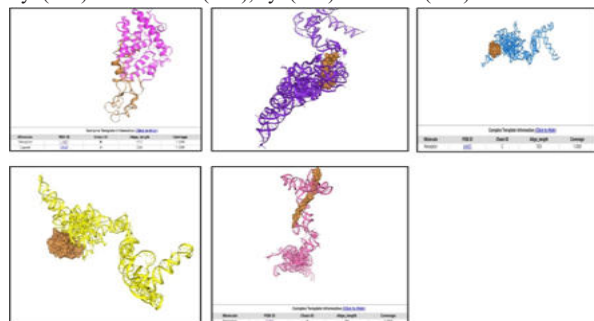


Figure 5: Protein-protein interactions between BRCA1 and

progesterone receptor sequence 1A28 (A), and gene-protein interactions between PR and BRCA1 (B), between PR and FOXC2 (C), between PR and FAK (D) and between PR and vimentin (E).

Figure 5A shows that BRCA1 interacts with the progesterone receptor at positions 89 (Leu) with Tyr (23), Phe (45) with Asn (108) and Arg (111), Cys (46) with Ile (106), Pro (59) with Pro (103), Val (60) with Asn (28) Thr (63) with Tyr (23) Leu (89) with Asn (100), His (91) with Val (94), and Ser (96) with Met (98) predominantly through van der Waals interactions and dipole-dipole interactions.

Figure 5C shows that the PR interacts with the FOXC2 protein via gene-protein interactions at position 3 (Lys) with guanine (405 & 406), with Pro (5) with uracil (357), with Tyr (8) with guanine (405), with Phe (39) with guanine (406), with Asn (43) with uracil (407), with Gly (46) and Trp (47) with guanine (406), with Asn (49), with Arg (52) with cytosine (307), with His (53) with cytosine (305 & 306), with Leu (57) with cytosine (305), with Glu (88) with uracil (396), and with Arg (97) with uracil (395).

As shown in Figure 5B, the progesterone receptor interacts with BrCA1 at Arg (24) with uracil (443) and Asn (30) with cytosine (444), Ile (31) with uracil (707), Arg (33) with uracil (453) and adenine (706), Gln (81) with cytosine (455) and uracil (456), Ser (84) with adenine (705), Arg (87) with uracil (446) and adenine (705), Asn (88) with uracil (704), Leu (95) with adenine (701) and Asp (97) with adenine (689).

As shown in Figure 5D, the progesterone receptor interacts with focal adhesion kinase, which results in gene-protein interactions at various positions, such as Pro (39) with cytosine (16), Asp (57) with adenine (323), Arg (136) with adenine (337), Asp (153) with guanine (6), and Glu (266) with guanine (253).

Figure 5E shows that nucleic acid interactions with the progesterone receptor occur with the protein vimentin at Glu (2) with uracil (695), Glu (27) with uracil (490), Ala (29), Asp (33) and Gln (32) with cytosine (664), Asn (30) with uracil (490 and 491), Tyr (30) with uracil (490), Thr (34) with uracil (491), Arg (37) with uracil (491 and 492), Leu (53) and Gln (57) with adenine (551), Tyr (56) with uracil (505) and cytosine (506).

DISCUSSION:

The roles of progesterone hormone and progesterone receptor in cancer treatment have been debated over the past several years, and by the early 20th century, it was determined that the presence of estrogen receptor alone cannot be a solution and that the PR is one of the targets of estrogen and the first baseline hormonal therapy for breast cancer¹⁸. The human epidermal growth factor receptor 2 (HER2) receptor is a membrane protein tyrosine kinase that affects proliferation and growth when activated. The HER2 oncogene is located at chromosome number 17q12, and HER2 gene amplification is one of the major mechanisms responsible for HER2 overexpression, which is associated with most current HER2-positive cancers¹⁹.

BRCA1 is a tumor suppressor gene that plays a key role in repairing DNA damage, regulating the cell cycle, maintaining genome stability and regulating several other physiological responses²⁰. Women with BRCA1 mutations can develop not only most aggressive breast cancers, such as triple-negative breast cancer but also a high risk of exposure to other cancers, such as ovarian cancer²⁰.

Declarations:

Ethics approval: NA

Consent to participate: NA

Consent for Publication: NA

Availability of data and material:

Figure 1

FIGURE 1a: BRCA1-EGFR

ACCESSION NUMBER: 1JM7- BRCA1

ACCESSION NUMBER: 1XKK- EGFR

URL: RCSB PDB - 1JM7: Solution structure of the BRCA1/BARD1 RING-domain heterodimer

URL: RCSB PDB - 1XKK: EGFR kinase domain complexed with a quinazoline inhibitor- GW572016

FIGURE 1B: BRCA1-FOXC2:

ACCESSION NUMBER: BRCA1- 1JM7

ACCESSION NUMBER: FOXC2 - 6AKO
 URL: RCSB PDB - 1JM7: Solution structure of the BRCA1/BARD1 RING-domain heterodimer
 URL: RCSB PDB - 6AKO: Crystal Structure of FOXC2 DBD Bound to DBE2 DNA

FIGURE 1C: BRCA1 VIMENTIN:
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: VIMENTIN-1GK4
 URL: RCSB PDB - 1JM7: Solution structure of the BRCA1/BARD1 RING-domain heterodimer
 URL: RCSB PDB - 1GK4: HUMAN VIMENTIN COIL 2B FRAGMENT(CYS2)

FIGURE 1D: BRCA1-FAK:
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: FAK-4I4E
 URL: RCSB PDB - 1JM7: Solution structure of the BRCA1/BARD1 RING-domain heterodimer
 URL: RCSB PDB - 4I4E: Structure of Focal Adhesion Kinase catalytic domain in complex with hinge binding pyrazolobenzothiazine compound.

Figure 2

Figure 2A: BRCA1-FOXC2:
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: FOXC2 - 6AKO
 URL: rcsb.org/entry/1JM7/display>1JM7_1|Chain A|BREAST CANCER TYPE 1 SUSCEPTIBILITY PROTEIN|*Homo sapiens* (9606)
 URL: RCSB PDB - 6AKO: Crystal Structure of FOXC2 DBD Bound to DBE2 DNA
 FIGURE 2B: BRCA1 VIMENTIN:
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: VIMENTIN-1GK4
 URL: rcsb.org/entry/1JM7/display>1JM7_1|Chain A|BREAST CANCER TYPE 1 SUSCEPTIBILITY PROTEIN|*Homo sapiens* (9606)
 URL: RCSB PDB - 1GK4: HUMAN VIMENTIN COIL 2B FRAGMENT(CYS2)
 FIGURE 2C: BRCA1-FAK axis
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: FAK-4I4E
 URL: rcsb.org/entry/1JM7/display>1JM7_1|Chain A|BREAST CANCER TYPE 1 SUSCEPTIBILITY PROTEIN|*Homo sapiens* (9606)
 URL: RCSB PDB - 4I4E: Structure of Focal Adhesion Kinase catalytic domain in complex with hinge binding pyrazolobenzothiazine compound.

Figure 3

Figure 3 A: BRCA1-EGFR:
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: EGFR-7AEI
 URL: rcsb.org/entry/1JM7/display>1JM7_2|Chain B|BRCA1- ASSOCIATED RING DOMAIN PROTEIN 1|*Homo sapiens* (9606)
 URL: rcsb.org/entry/7AEI/display>7AEI_1|Chain A|Epidermal growth factor receptor|*Homo sapiens* (9606)
 Figure 3B: EGFR-VIMENTIN:
 ACCESSION NUMBER: EGFR-7AEI
 ACCESSION NUMBER: VIMENTIN-1GK4
 URL: rcsb.org/entry/7AEI/display>7AEI_1|Chain A|Epidermal growth factor receptor|*Homo sapiens* (9606)
 URL: RCSB PDB - 1GK4: HUMAN VIMENTIN COIL 2B FRAGMENT(CYS2)
 Figure 3C: EGFR-FOXC2
 ACCESSION NUMBER: EGFR-7AEI
 ACCESSION NUMBER: FOXC2 - 6AKO
 URL: rcsb.org/entry/7AEI/display>7AEI_1|Chain A|Epidermal growth factor receptor|*Homo sapiens* (9606)
 URL: RCSB PDB - 6AKO: Crystal Structure of FOXC2 DBD Bound to DBE2 DNA
 Figure 3D EGFR-FAK
 ACCESSION NUMBER: EGFR-7AEI
 ACCESSION NUMBER: FAK-4I4E
 URL: rcsb.org/entry/7AEI/display>7AEI_1|Chain A|Epidermal growth factor receptor|*Homo sapiens* (9606)
 URL: RCSB PDB - 4I4E: Structure of Focal Adhesion Kinase catalytic

domain in complex with hinge binding pyrazolobenzothiazine compound.

FIGURE: 4

FIGURE: 4A: BRCA1-ErB2
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: ErbB2- 2A91
 URL: RCSB PDB - 1JM7: Solution structure of the BRCA1/BARD1 RING-domain heterodimer
 URL: RCSB PDB - 2A91: Crystal structure of ErbB2 domains 1-3
 FIGURE 4B: BRCA1: HER2/ErbB2
 ACCESSION NUMBER: BRCA1- 1JM7
 ACCESSION NUMBER: HER2/ErbB2-9QBF
 URL: rcsb.org/entry/1JM7/display>1JM7_2|Chain B|BRCA1- ASSOCIATED RING DOMAIN PROTEIN 1|*Homo sapiens* (9606)
 URL: rcsb.org/entry/9QBF/display>9QBF_1|Chain A|Receptor tyrosine-protein kinase erbB-2, Green fluorescent protein|*Homo sapiens* (9606)
 FIGURE 4C: HER2/ErbB2: FOXC2
 ACCESSION NUMBER: HER2/ErbB2-9QBF
 ACCESSION NUMBER: FOXC2 - 6AKO
 URL: rcsb.org/entry/9QBF/display>9QBF_1|Chain A|Receptor tyrosine-protein kinase erbB-2, Green fluorescent protein|*Homo sapiens* (9606)
 URL: RCSB PDB - 6AKO: Crystal Structure of FOXC2 DBD Bound to DBE2 DNA
 FIGURE 4D: HER2/ErbB2: VIMENTIN
 ACCESSION NUMBER: HER2/ErbB2-9QBF
 ACCESSION NUMBER: VIMENTIN-1GK4
 URL: rcsb.org/entry/9QBF/display>9QBF_1|Chain A|Receptor tyrosine-protein kinase erbB-2, Green fluorescent protein|*Homo sapiens* (9606)
 URL: RCSB PDB - 1GK4: HUMAN VIMENTIN COIL 2B FRAGMENT(CYS2)
 FIGURE: 4E: HER2/ErbB2: FAK
 ACCESSION NUMBER: HER2/ErbB2-9QBF
 ACCESSION NUMBER: FAK-4I4E
 URL: rcsb.org/entry/9QBF/display>9QBF_1|Chain A|Receptor tyrosine-protein kinase erbB-2, Green fluorescent protein|*Homo sapiens* (9606)
 URL: RCSB PDB - 4I4E: Structure of Focal Adhesion Kinase catalytic domain in complex with hinge binding pyrazolobenzothiazine compound.

FIGURE 5

FIGURE 5A BRCA1:PR
 ACCESSION NUMBER: 1JM7- BRCA1
 ACCESSION NUMBER: 1A28 - PR
 URL: RCSB PDB - 1JM7: Solution structure of the BRCA1/BARD1 RING-domain heterodimer
 URL: RCSB PDB - 1A28: HORMONE-BOUND HUMAN PROGESTERONE RECEPTOR LIGAND-BINDING DOMAIN
 FIGURE 5B: BRCA1-PR
 ACCESSION NUMBER: 1JM7- BRCA1
 ACCESSION NUMBER: 1A28 - PR
 URL: rcsb.org/entry/1JM7/display>1JM7_2|Chain B|BRCA1- ASSOCIATED RING DOMAIN PROTEIN 1|*Homo sapiens* (9606)
 URL: rcsb.org/entry/1A28/display>1A28_1|Chains A, B|PROGESTERONE RECEPTOR|*Homo sapiens* (9606).
 FIGURE 5C: PR-FOXC2
 ACCESSION NUMBER: 1A28 - PR
 ACCESSION NUMBER: FOXC2 - 6AKO
 URL: rcsb.org/entry/1A28/display>1A28_1|Chains A, B|PROGESTERONE RECEPTOR|*Homo sapiens* (9606).
 URL: RCSB PDB - 6AKO: Crystal Structure of FOXC2 DBD Bound to DBE2 DNA
 FIGURE 5D: PR-FAK
 ACCESSION NUMBER: 1A28 - PR
 ACCESSION NUMBER: FAK-4I4E
 URL: rcsb.org/entry/1A28/display>1A28_1|Chains A, B|PROGESTERONE RECEPTOR|*Homo sapiens* (9606).
 URL: RCSB PDB - 4I4E: Structure of Focal Adhesion Kinase catalytic domain in complex with hinge binding pyrazolobenzothiazine compound.
 FIGURE 5E: PR-VIMENTIN
 ACCESSION NUMBER: 1A28 - PR
 ACCESSION NUMBER: VIMENTIN-1GK4
 URL: rcsb.org/entry/1A28/display>1A28_1|Chains A,

B|PROGESTERONE RECEPTOR |*Homo sapiens* (9606).
URL: RCSB PDB - 1GK4: HUMAN VIMENTIN COIL 2B
FRAGMENT (CYS2)

The first Author/Corresponding Author Dr. Eswari Beeram will be contacted to obtain the raw data.

Competing interests:

The authors declare that they have no competing interests.

Funding: NA

Limitations statement:

The amino acid sequence is encoded at only 100-300 amino acids due to the large size of the contig in the case of eErb2/HER2. When predicting SOX10 expression, the methodology used for the study was limited because the data generated represented by some protein IDs ultimately limited the study to only 8 proteins instead of 10 proteins, and I am unable to study all the proteins and genes involved in TNBC.

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