1	Title: Aberrant epigenetic and transcriptional events associated with breast cancer risk

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24 ABSTRACT

Background: Genome-wide association studies have identified several breast cancer susceptibility loci. However, biomarkers for risk assessment are still missing. Here, we investigated cancer-related molecular changes detected in tissues from women at high risk for breast cancer prior to disease manifestation.

Disease-free breast tissue cores donated by healthy women (N=146, median age=39 years) were processed for both methylome (MethylCap) and transcriptome (Illumina's HiSeq4000) sequencing. Analysis of tissue microarray and primary breast epithelial cells was used to confirm gene expression dysregulation.

Results: Transcriptomic analysis identified 69 differentially expressed genes between women at either high and 31 32 those at average risk of breast cancer (Tyrer-Cuzick model) at FDR<0.05 and fold change 2. The majority of the identified genes were involved in DNA damage checkpoint, cell cycle, and cell adhesion. Two genes, 33 FAM83A and NEK2, were overexpressed in tissue sections (FDR<0.01) and primary epithelial cells (p<0.05) 34 35 from high-risk breasts. Moreover, 1698 DNA methylation aberrations were identified in high-risk breast tissues (FDR<0.05), partially overlapped with cancer-related signatures and correlated with transcriptional changes 36 $(p < 0.05, r \le 0.5)$. Finally, among the participants, 35 women donated breast biopsies at two time points, and age-37 related molecular alterations enhanced in high-risk subjects were identified. 38

39 Conclusions: Normal breast tissue from women at high risk of breast cancer bears molecular aberrations that 40 may contribute to breast cancer susceptibility. This study is the first molecular characterization of the true 41 normal breast tissues and provides an opportunity to investigate molecular markers of breast cancer risk, which 42 may lead to new preventive approaches.

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14 Keywords: Cancer risk, transcriptome, DNA methylation, normal breast

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H7 BACKGROUND

Genetic and epigenetic alterations in breast cancer (BC) have been widely investigated. However, when, during the carinogenesis process, these events first emergeremains unknown. The identification of molecular aberrations associated with BC development can provide a conceptual framework for a deeper understanding of this complex disease.

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Genome-wide association studies (GWAS) have detected more than 170 genomic loci harboring common 53 variants associated with BC risk including modifier alleles with high (e.g., BRCA1, BRCA2, TP53, PTEN) to 54 55 moderate penetrance (e.g., BRIP1, CHEK2, ATM, and PALB2) (1-4). Nevertheless, many variants are located in noncoding or intergenic regions and their functional role in cancer transformation remains largely unknown. 56 Recently, transcriptome-wide association studies were used to integrate GWAS and gene expression datasets 57 58 and identified 154 genes whose predicted expression associated with the risk for BC (5-9). However, these studies drew data from the Genotype-Tissue Expression (GTEx) project, where the use of autopsy-derived 59 normal breast tissues may make the breast-specific transcriptome profilings questionable. The relative lack of 50 51 molecular profiling of normal breast tissue from subjects who are disease-free makes the field challenging.

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Many studies searching for cancer biomarkers have identified gene expression signatures, epigenetic signatures, loss of heterozygosity and allelic imbalance resulting from the development of malignancy (10). Among the molecular processes linked with cancer, DNA methylation has a keyrole in early cancer development through a process known as epigenetic reprogramming (11), potentially leading to silencing and loss of expression of tumor suppressor genes (12), and genomic instability (13).

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Here, we performed an integrated analysis of DNA methylation and transcriptome profiling of cancer-free breast tissues donated by healthy women at either average or high risk for BC. In addition to early epigenetic events, we identified two molecules overexpressed in high-risk breasts independently from DNA methylation changes and, therefore, potential markers of BC susceptibility. Moreover, using a subcohort of repeated breast

tissues donation by the same donors, we confirmed that the molecular changes identified in high-risk subjects are age-independent. These findings will lead to a deeper understanding of BC susceptibility and also provide the scientific community with the molecular profiling of a true normal breast tissue.

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77 **RESULTS**

78 Study cohort used to investigate molecular aberrations in association with breast cancer (BC) risk

To identify transcriptomic and epigenetic differences linked with BC risk, we analyzed cancer-free breast tissue cores donated by 146 healthy women (median age: 39 years) including 112 Caucasian, 24 African American and 10 Asian subjects (**Additional File 1: Table S1**). Out of 146 participants, 117 were pre- and 29 postmenopausal women. Tyrer-Cuzick model was employed to estimate the lifetime risk of developing BC and allocated the subjects to either high- (score≥20%, N=68) or average-risk group (score<20%, N=78) (**Fig. 1A**, **Table 1** and **Additional File 1: Table S1**).

35 Characterization of the transcriptome alterations in high-risk breast

We performed a transcriptome analysis of the fresh frozen disease-free breast tissue donated by all the 36 participants. Differential expression analysis was performed using DESeq2. From a total of 22,344 genes, the 37 differential expression analysis between high- and average-risk breasts revealed 1,874 transcripts to be 38 significant at 5% false discovery rate (FDR). Of these, 1,798 transcripts also passed the cutoff of t-test p-value \leq 39 0.05 (Additional File 1: Table S2). Sixty-nine genes, including 51 upregulated and 18 downregulated genes, Э0 were identified with a fold change ≥ 2 (**Table 1**). Because both groups consisted of non-malignant breast tissue, Э1 a limited number of differentially expressed genes was expected (14). Canonical pathway analysis revealed Э2 ЭЗ enrichment in pathways related to kinetochore signaling (p=1.3E-05), DNA damage checkpoint (p=0.0005), Э4 granulocytes adhesion (p=0.002), and the IL17 pathway (p=0.004) (Figure 1B, Additional File 1: Table S3). Our data further confirm the previously described impact of dysregulated DNA damage in breast carcinogenesis Э5 Э6 (15). Molecular network analysis showed an enrichment in functional categories involved in cell cycle, DNA

replication and repair (Figure 1C, Additional File 1: Table S3). One of the major molecular networks
regulating cell cycle is centered around AKT and the transcription factor FOXM1 (16).

Except for DCX, the transcriptional changes detected between high- and average-risk breasts listed in Table 2

30 are independent of both racial background and menopausal status of the tissue donors (Additional File 1: Table

S4 and **Additional File 2: Fig. S1**).

Among the 69 differentially expressed genes, FAM83A and NEK2, showed the highest gene)2 amplification rate (number of cases with gene amplification>200) and genetic alteration frequency (>10%), as)3 well as overexpression in BC (p>0.001) and Oncoscore >50, as detected through METABRIC database,)4 cBioportal, UALCAN and Oncoscore analyses FAM83A and NEK2 (Table 1 and Additional File 2: Fig. S2))5 (17-19). The expression of FAM83A and NEK2 in the breasts of high- and average-risk women is shown in)6)7 Fig. 1D. We detected a 4.5-fold increase in FAM83A and 2.2-fold increase in NEK2 expression in primary epithelial cells isolated from the breast of high-risk women when compared with cells isolated from breast)8 tissue of average-risk women (Fig. 1E). Overexpression of both targets was detected also in a dataset of)9 LO hTERT-immortalized epithelial cells as compared with the isogenic primary cells (20) (Fig. 1F). Moreover, immunostaining of a breast tissue microarray showed a 1.4 fold increase in FAM83A protein levels in the breast Ι1 tissues from women at high risk of BC as compared with the breast tissues from subjects at average risk L2 (p<0.0001, Fig. 1G, Additional File 2: Fig. S3A and Additional File 1: Table S5). FAM83A overexpression L3 in normal breast tissues was associated with parity (p < 0.001), tobacco use (p = 0.01) and family history of BC L4 (p=0.02) (Additional File 1: Table S6). On the contrary, NEK2 staining showed no difference in protein levels ۱5 between the two groups (Fig. 1G). No difference in Ki67, estrogen receptor alpha (ER α), FOXA1 and GATA3 ۱6 L7 staining between high- and average-risk breasts was observed (Additional File 2: Fig. S3B and Additional File 1: Table S5). This data shows that FAM83A expression changes are specific to breasts of women at high risk of ٢8 ٤9 developing breast cancer.

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21 Genome-wide DNA methylation analysis reveals 1698 aberrant DNA methylation sites in normal breast

22 tissue of high-risk women

With the goal of identifying alterations in regulatory regions leading to BC susceptibility, we performed 23 a methylome analysis using the MethylCap-seq approach. Differential analysis of the methylated regions 24 detected in the breasts from average-risk women and those from women at high risk of cancer revealed a wide 25 chromosomal distribution of the epigenetic aberrations (Fig. 2A). DNA methylation changes with a $\Delta Z \ge 1$ 26 (hypermethylated) or ≤ -1 (hypomethylated) were selected. We identified 1698 regions methylated that 27 differentiate the breast tissue of high-risk women from that of women at average risk (FDR $\square \le \square 5\%$), mapping 28 29 to 1115 unique genes (Additional File 1: Table S7). The twenty most hypermethyated and hypomethylated regions are shown in Fig. 2B and Table 2. Interindividual variability in DNA methylation can be observed 30 within each experimental group. 98.9% of the DNA methylation aberrations consisted of hypermethylated loci 31 $(p=\Box 9\Box \times \Box 10^{\Box 8};$ Fig. 2C). More than 90% of hypermethylated loci localized in regulatory regions including 32 the promoter, untranslated region, and introns, while only 41% of hypomethylated loci localized in these 33 regions (Fig. 2C). However, hypomethylated regions were localized predominantly in the gene body (59%), a 34 35 phenomenon that has been linked with the activation in cancers of aberrant intragenic promoters that are 36 normally silenced (21, 22).

Pathway analysis revealed the involvement of cell adhesion (aka synaptogenesis, p=1.2E-06), ErbB 37 (p=3.7E-04) and protein kinase A (p=4.8E-04) signaling pathways (Fig. 2D, Additional File 1: Table S8). 38 Notably, one of the molecular networks showed ESR2 as the central molecule (Fig. 2E). Although ESR2 39 expression decreased in high-risk breasts (fold change=0.82), the intronic ESR2 hypermethylation showed no 10 11 inverse correlation with ESR2 expression (r=-0.03, p=0.4; Additional File 2: Fig. S4A-C). One of the 12 hypomethylated genes, MUC1 (ΔZ =1.4, FDR=1.6E-17) is reported to be aberrantly overexpressed in over 90% 13 of breast tumors (23, 24) (Additional File 2: Fig. S4D). However, no significant difference in MUC1 14 expression was observed between high- and average-risk breasts (Additional File 2: Fig. S4E). In the analyzed

cohort, DNA methylation was not highly affected by either racial background or menopausal status of the tissue
donors (FDR>0.05; Additional File 1: Table S9). Finally, we found overlap between DNA methylation
aberrations in high-risk breasts and breast cancer-related DNA methylation signatures such as those identified
by Saghafinia *et al* (4%, 25/666, (25)), Chen *et al* (6%, 10/174, (26)), de Almeida *et al* (9%, 25/285, (27)), and
Xu et al (9%, 37/414, (28)) (Additional File 1: Table S10).

50 DNA methylation and gene expression changes in high-risk breast show a weak correlation

To identify potential epigenetically regulated genes linked with BC risk, we performed a Pearson's 51 correlation test on paired DNA methylation and gene expression data (Fig. 3). Among the 69 genes in Table 2), 52 the expression of eight genes was associated with aberrant intronic DNA methylation, including six genes 53 54 showing a direct correlation (APELA, DIO2, FEZF2, LPAR3, UNC5D and PRSS51) and two genes (PROK2, 55 and SULT1C2) with a negative correlation (Fig. 3A). Furthermore, among the DNA methylation aberrations in **Table 2,** only the intronic hypermethylation of PHACTR1 ($\Delta Z = 1.88$, FDR= 1.0E-31) was negatively 56 57 correlated with PHACTR1 expression (fold change=0.77, FDR=0.006, r=-0.21) (Fig. 3B). However, the 58 correlations identified were weak (r: -0.2,-0.5) suggesting that other regulator events (chromatin modifications, 59 gene amplification, nucleotide variants), rather than DNA methylation aberrations, may be the determinants of the transcriptomic changes observed in the high-risk breasts as compared with average-risk breasts. 50

51 Age-related molecular changes in cancer-free breast tissues in relation with cancer risk

Age is the strongest demographic risk factor for most human malignancies, including breast cancer. Age-related transcriptome and DNA methylation aberrations were investigated on breast tissues cores donated by 35 women at two separate time points (**Additional File 1: Table S11**). Differential expression analysis (FDR<0.05) between the two donation time points revealed the dysregulation of 317 genes involved in LXR/RXR activation (p=7E-04), immune response (p=2E-03) and senescence (p=7E-03) (**Additional File 1: Tables S12 and S13**). Forty-eight age-related transcriptomic changes with a fold change (fc) \geq 2 and FDR <0.05 included two upregulated genes, CETP (fc:2.4;FDR=0.04) and HP (fc=2.3, FDR=0.03); and downregulation of

five genes, SLC5A1 (fc=0.4; FDR=0.03), SLCO1A2 (fc=0.4; FDR=0.03), GRIA4 (fc=0.4; FDR=0.01), IL22RA2 (fc=0.4; FDR= 0.01), and CHRM1 (fc=0.4; FDR=0.03) (**Additional File 2: Fig. S5**). Furthermore, age-dependent dysregulation of the following five genes was enhanced in breast tissues from high-risk women: NEURL1, USP50, GRIA4, SPDEF and DNM3 (**Fig. 4A**). Notably, the expression of GRIA4 (r=-0.43, p=0.04) and DNM3 (r=-0.47, p=0.03) showed a negative correlation with their DNA methylation pattern, thus suggesting a potential epigenetic regulation for these two molecules (**Fig. 4B**).

Age-dependent DNA methylation aberrations affected 301 loci corresponding to 280 unique transcripts 75 76 (Additional File 1: Table S14). As previously reported (29), age-related DNA methylation alterations were predominantly hypermethylation events (85.4%) and affected the intronic regions (Fig. 4C). DNA methylation 77 measurements were previously used to develop epigenetic biomarkers of aging, otherwise known as "DNA 78 79 methylation age" or the "epigenetic clock" (30, 31). We observed a limited overlap between the 301 DNA methylation aberrations and the epigenetic clocks described by Horvath et al (1.4%, (31)), while 73 genes 30 associated with the bin in our dataset overlapped with age-associated DNA methylation alterations reported by 31 32 Johnson et al (24.2%,(29)) (Fig. 4D). Finally, we identified age-related DNA methylation aberrations enhanced in high-risk breasts, localized on four genes: PTPRM, SPOCK1, KCNH1, and CFAP43. (p<0.001, Fig. 4E and 33 Additional File 1: Table S14). In contrast, both transcriptomic and DNA methylation aberrations affecting 34 35 FAM83A and NEK2 resulted age-independent.

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37 **DISCUSSION**

This study aimed to define the distinct features of cancer-free breast tissues from women at high risk for BC and, thus, identify molecular markers that could potentially screen for women susceptible to cancer. We conducted transcriptome and methylome analyses using breast tissue cores donated by healthy women. The participants were divided into two cohorts based on their risk of developing breast cancer, according to the Tyrer-Cuzick lifetime risk assessment score: high-risk (\geq 20%) and average-risk (<20%) (32). Among the genes upregulated in high-risk breast, we identified two promising markers of BC susceptibility, FAM83A and NEK2.

Furthermore, when investigating DNA methylation aberrations in high-risk breasts, we detected 4-10% overlap with cancer-related signatures.

Our transciptomic analysis of high- and average-risk breasts revealed significant changes in 69 genes Э6 Э7 (FDR<0.05). Pathway analysis suggested the activation of cell cycle and cell adhesion in the high-risk breasts. 98 Furthermore, one of the molecular networks including the differentially expressed genes showed the involvement of FOXM1 signaling. FOXM1 itself is upregulated 1.6 fold in high-risk breasts (p=0.001). The)9 transcription factor FOXM1 regulates the transcription of cell-cycle genes essential for exit from the G1/S)0 phase into the G2/M phase such as cyclin A2, JNK1, ATF2 and CDC25A phosphatase as well as genes critical)1 for chromosome segregation and cytokinesis (33). FOXM1 is overexpressed and plays critical role in)2 tumorigenesis, metastasis, and drug resistance in a broad range of human cancer types, such as lung, gastric, and)3)4 breast cancers (16). Compounds targeting FOXM1 expression or activity are under investigation (16). Our results suggest that the transcriptional dysregulation detected in high-risk breasts may be driven by FOXM1.)5

Two genes, FAM83A and NEK2, both upregulated in high-risk breast, showed a high Oncoscore (75.5)6)7 and 61.4, respectively), and have been reported amplified in breast cancer. FAM83A is the smallest member of the eight-member FAM83 family of proteins that share a conserved amino-terminal Domain of Unknown)8 Function (DUF1669 domain) (34). It was identified based on its transforming potential (35-37). FAM83A)9 upregulation has been detected in multiple human tumor types, including breast, lung, pancreatic and cervical LO cancer (37-44). Lee et al (45, 46) revealed the ability of FAM83A to confer resistance to epidermal growth Ι1 factor receptor/ tyrosine kinase inhibitors (EGFR-TKIs) through interactions with c-RAF and PI3K p85 in L2 breast cancer. The authors also showed that BC patients with high FAM83A expression had a worse prognosis. L3 L4 FAM83A depletion suppressed proliferation and invasiveness in vitro as well as tumor growth in vivo (36). Based on the aforementioned studies, FAM83A is considered a candidate oncogene and our findings suggest ۱5 that FAM83A may be one of the first molecules dysregulated in cancer transformation. Our team is currently ۱6 investigating the role of FAM83A in breast carcinogenesis. Moreover, our DNA methylation data, in agreement Γ ٢8 with previous literature, suggest that FAM83A overexpression is mainly driven by genomic amplification

rather than epigenetic regulation (47, 48). Additional studies such as deep whole genome sequencing of DNA
from breast tissues of high-risk women are required to support this hypothesis.

The NIMA-related kinase 2 (NEK2) protein belongs to a family of serine/threonine kinases, which have 21 22 a role in mitotic progression by initiating the separation of centrosomes (49). NEK2 overexpression was <u>23</u> previously reported in BC as result of gene amplification (47, 50). NEK2 depletion blocks cell cycle progression and tumor cell growth, making it an ideal therapeutic target (51). Notably, FOXM1 is reported to 24 both bind NEK2 promoter and interact with NEK2 directly (52, 53). Our study further suggests a role of NEK2 25 26 dysregulation in breast carcinogenesis. However, we did not observe changes in NEK2 protein levels in breast tissues of high-risk women suggesting a disconnect between mRNA and protein levels, which is not 27 uncommon. Further investigation of the role of NEK2 in breast carcinogenesis is needed. 28

<u>29</u> We observed DNA methylation changes in high-risk breasts, consisting mostly of hypermethylation (98.8%) in the intronic regions (88%). Previous studies reported aberrant hypermethylation in normal breast 30 tissue adjacent to the tumor (54). Hypermethylation in specific gene promoters is indeed 31 linked to 32 carcinogenesis through transcriptional silencing of tumor suppressor genes or regulatory regions within the genome leading to dysregulation of cell growth, cancer initiation and progression (55-57). We identified a 4-33 10% overlap between methylome aberrations in high-risk breasts and previously reported cancer-related 34 35 signatures (25-28). The limited overlap may be linked to the different technical approaches (Methyl-capture vs Infinium HumanMethylation450 array) but may also suggest that cancer-related epigenetic marks are newly 36 acquired during cancer initiation rather than being imprinted into the genome. Although the expression 37 epigenetic modifiers such as DNMTs remain unaffected, we detected the upregulation in high-risk breasts of 38 39 HASPIN (fc=1.7; FDR<0.005), a serine/threonine kinase involved into the phosphorylation of the histone H3 during mitosis (58), suggesting that other genetic and epigenetic mechanisms rather than DNA methylation may 10 drive the transcriptomic aberrations in high-risk breasts. 11

Age is the strongest demographic risk factor for most human malignancies, including BC (59). The limited size of our cohort (N=35) prevented us from subdividing the subjects by age at tissue donation.

14 Nevertheless, we identified age-related transcriptomic aberrations enhanced in high-risk breasts including GRIA4 and DNM3, which resulted potentially epigenetically regulated. In terms of DNA methylation 15 16 aberrations, we found a limited overlap between the age-related DNA methylation changes here identified and 17 epigenetic clock from Hovarth et al. (31) (Additional File 1: Table S14). However, a 24.2% overlap of our 18 dataset with age-related DNA methylation aberrations described by Johnson el at (29) was detected. The limited overlap is probably due to the different platform used for DNA methylation detection (Infinium Human 19 Methylation 450 array vs Methyl-Cap-seq) and the type of analysis (epithelium-specific deconvolution vs whole 50 51 tissue)(29, 31). Notably, we identified specific age-related DNA methylation changes, located on PTPRM, KCNH1, SPOCK1, CFAP43 gene, enhanced in the high- versus average-risk breasts. 52

This study harbors some limitations: the relatively small sample size prevented us from investigating in 53 54 details cancer-related varables such as racial background. The selection of normal breast tissue cores with high content in epithelial compartment limited the number of available samples (Additional File 2: Fig. S6). 55 Outcome data for the women at high risk for BC is not available at this time; however, this cohort is under an 56 57 ongoing annual medical follow up. Because of the faster processing time and smaller cost, we performed whole tissue analysis instead of the more epithelium-specific laser microdissection or single-cell analysis. This limits 58 the compartment specificity of the data but generates a more comprehensive view of the molecular features of 59 the entire breast tissue core. Further deconvolution analysis may overcome this limitation (60, 61). 50

51 CONCLUSIONS

The present study reveals transcriptomic and epigenetic aberrations linked with BC risk. Transcriptional targets potentially promoting cancer susceptibility were identified. The described investigation provides an avenue for deciphering the functional relevance of genes involved in BC development and a rich resource for further investigation.

56 **METHODS**

57 Study cohorts

58 Breast specimens were obtained from the Susan G. Komen Tissue Bank at the IU Simon Comprehensive Cancer Center (KTB) and donated voluntarily upon informed consent by healthy women. Subjects were 59 70 recruited under a protocol approved by the Indiana University Institutional Review Board (IRB protocols 71 number 1011003097 and 1607623663). Subject demographics and breast cancer (BC) risk factors were 72 collected using a questionnaire administered by the KTB and summarized in Additional File 1: Table S1, S5 and S11. Breast tissue cores are collected by using a needle biopsy as previously described (14). The study 73 cohort consisted of two groups: 1) For the transcriptome and methylome analyses, 146 women (median age: 39 74 75 years) were selected based on the lack of clinical and histological breast abnormalities and high content in breast epithelial compartment (cellularity>40%). Germline mutation status of the subjects was obtained from 76 KTB. Data retrieved from the LifeOmic's Precision Health 77 were Cloud platform 78 (https://lifeomic.com/products/precision-health-cloud/). Nine established breast cancer-predisposition genes (BRCA1, BRCA2, PALB2, ATM, CHECK2, BARD1, RAD51C, RAD51D, CDH1) were evaluated for variants 79 30 identified as "pathogenic" "likely pathogenic" in the ClinVar database or 31 (https://preview.ncbi.nlm.nih.gov/clinvar/) (Additional File 1: Table S1) (2, 3).

Thirty-five of these 146 women, including 10 at high risk and 25 at average risk for BC, donated their breast tissue at two time points at intervals from 1-10 years (mean: 3.2) between the tissue donations (**Fig. 1A** and **Additional File 1: Table S11**). 2) In a second analysis, paraffin-embedded breast tissue blocks related to 395 healthy women were obtained from the KTB and used to generate tissue microarrays. The cohort included 287 Caucasian, 66 African American, 49 Asian, with age ranging from 18 to 61 (Additional File 1: Table S5).

Breast cancer risk assessment

Lifetime risk of developing BC was estimated by using the Tyrer-Cuzick risk score (IBISv8) (32) and a threshold of 20% to separate high- (\geq 20%) from average-risk (<20%) individuals. The Tyrer-Cuzick model was selected over the other risk estimation tools for its accuracy and inclusion of young subjects (62).

Homeon Tissue processing and nucleic acid extraction

Э2 To limit stromal contamination, only breast tissue samples abundant in epithelial compartment (cellularity >40%) were selected and processed. Total DNA and RNA were isolated from fresh frozen breast ЭЗ Э4 tissue biopsies (80-1500mg) using AllPrep DNA/RNA/miRNA kit (Qiagen). Tissues were homogenized by Э5 using 2ml prefilled tubes containing 3mm zirconium beads (Benchmark Scientific, cat.# D1032-30), 350µl Э6 Lysis Buffer and 2-Mercaptoethanol, and BeadBug 6 homogenizer (Benchmark Scientific) in a cold room at the following conditions: 4000 rpm for 45 seconds was repeated twice with 90 seconds rest time. The concentration)7 and quality of total RNA and DNA samples were first assessed using Agilent 2100 Bioanalyzer. A RIN (RNA 98)9 Integrity Number) and DIN (DNA integrity number) of six or higher is required to pass the quality control.

Whole transcriptome analysis

cDNA library was prepared using the TruSeq Stranded Total RNA Kit (Illumina) and sequenced using)1)2 Illumina HiSeq4000. Data included 146 paired-end fastq sequence libraries (raw read length: 38 x 2). Reads adapter trimmed filtered using Trimmomatic 0.38)3 were and quality ver. (http://www.usadellab.org/cms/?page=trimmomatic) setting the cutoff threshold for average base quality score)4)5 at 20 over a window of 3 bases. Reads shorter than 20 bases post-trimming were excluded. About 94% of the reads have both the mates passing the quality filters. Cleaned reads mapped to Human genome reference)6 sequence GRCh38.p12 with gencode v.28 annotation, using STAR version STAR_2.5.2b (63). Only samples)7 with about 99% of the cleaned reads aligned to the genome reference. Differential expression analysis was)8 performed using DESeq2 ver. 1.12.3 (https://bioconductor.org/packages/release/bioc/html/DESeq2.html).)9 Counts table containing mapped read counts for each gene, to be used as input for DESeq2 was generated using LO featureCounts tool of subread package (https://doi.org/10.1093/bioinformatics/btt656). Alternatively, we ran t-Ι1 tests comparing the normalized read counts for the set of replicates from High risk samples to those for the set L2 of replicates from Average risk samples. The normalized read counts were obtained from the DESeq2 run L3 L4 described above. The *p* values from the *t*-test were corrected for multiple testing using Benjamini-Hochberg method. ۱5

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17 **DNA methylation analysis**

Library was generated by using MethylCap Library Kit (Diagenode, Denville NJ, US) according to the ٢8 ٤9 manufacturer's protocols followed by single-end 75-bp sequencing on Illumina Nextseq4000. Internal controls 20 and duplicate samples were used to account for any batch effect and technical artifact. The data comprises of 146 paired end read libraries in FASTQ format. These libraries represent replicates for two samples - High risk 21 (68 libraries) and Average risk (78 libraries). The libraries were sequenced across multiple runs and the 22 combined read counts for each library were generated. Reads were adapter trimmed and quality filtered using 23 24 Trimmomatic 0.38 (http://www.usadellab.org/cms/?page=trimmomatic) with the cutoff threshold for average base quality score set at 20 over a window of 3 bases. Reads shorter than 20 bases post-trimming were 25 excluded. Approximately, 96% of the sequenced reads passed the quality filters to be considered "cleaned" 26 27 reads. This quality control reduced the number of samples to 57 high- and 55 average-risk. Cleaned reads were mapped to Human genome reference GRCh38.p12 using BWA ver. 0.7.15 (64). Insert sequences were imputed 28 from the concordantly mapped read pair alignments. More than 95% of the cleaned read pairs were 29 30 concordantly mapped. A previously described differential methylation analysis using either Zratio or ΔZ (65, 66) was applied to the current methyl-capture dataset with a slight improvisation on the validation of the 31 significance of differential methylation. For any given local bin of a given width on the genome, the method 32 compares across samples, variation in deduplicated insert coverage distribution quantified as the bin's z-score 33 with respect to a larger genome region containing the bin. For this analysis, we used local non-overlapping bins 34 with a fixed width of 250 bp with their z-scores computed relative to 25KB regions. Z-score is the number of 35 standard deviations by which the bin coverage varies from the larger region's mean coverage. A significant 36 37 difference in Zscores, calculated as either as ΔZ or Zratio between the samples would indicate potential differential methylation for that bin, as previously described (67),. The analysis identified 159,438 bins, each 38 250bp wide, to be potentially differentially methylated between High risk and Average risk samples with z-39 ratios or ΔZ significant at 5% FDR and p-values from t-test ≤ 0.05 . Based on positional overlap, these bins were 10 11 annotated using annotation from gencode v28.

12 Data analysis

Ingenuity Pathways Analysis (IPA, Qiagen, Redwood City, CA) was used for canonical pathway and 13 14 molecular network analyses (68). Publicly available transcriptomic data from primary and immortalized breast epithelial cells were obtained from GEO (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE108541) 15 (20). Analysis of The Cancer Genome Atlas (TCGA) was performed by interrogating both cBioPortal 16 (https://www.cbioportal.org/) and UALCAN (http://ualcan.path.uab.edu/) databases (69). Copy number 17 variations (CNV) analysis was obtained from the interrogation of the Molecular Taxonomy of Breast Cancer 18 19 International Consortium, METABRIC (17, 18). Oncoscore was used to rank genes according to their association with cancer, based on the available scientific literature (http://www.galseq.com/next-generation-50 sequencing/oncoscore-software/; accessed on 3/31/2021) (19). 51

52 **Primary breast epithelial cells and immunofluorescence**

Primary breast epithelial cells were generated from cryopreserved breast tissue cores obtained from the 53 KTB as previously described (14, 20). Immunofluorescence staining was performed as previously described 54 55 (14). Briefly, 5,000 cells were cultures overnight into each well of an 8 well-chamber slide (BD Biosciences, San Jose, CA) and fixed with acetone: methanol (1:1) at -20°C for 10 min. After washing and blocking 56 (PBS1X, 5% normal goat serum, 0.1% TritonX-100) steps cells were incubated with primary either rabbit anti-57 vimentin (Cell Signaling, D21H3, 1:100) or mouse anti-E-cadherin (Cell Signaling, 14472, 1:50) overnight. 58 Upon three washes with PBS, cells were incubated with secondary antibodies (goat anti- mouse Alexa Fluor 59 568 or goat anti-rabbit Alexa Fluor 488; Thermo Fisher Scientific, 1:500) for 1 h at room temperature. After 50 three washes with PBS, the coverslide was mounted using DAKO fluorescent mounting medium (S3023 51 52 Agilent, Santa Clara, CA) and the staining was visualized using a fluorescent microscope (Eclipse TS100, Nikon Instruments inc, Melville, NY). 53

54 Quantitative real time polymerase chain reaction (qPCR)

Total RNA was extracted from cells using AllPrep DNA/RNA/miRNA kit (Qiagen). Reverse
 transcription was performed using SuperScriptTM IV VILOTM Master Mix (Invitrogen cat#: 11756050)

57 according to the manufacturer's instructions. qPCR was performed using the TaqManTM Universal PCR Master Mix (Applied Biosystems, cat# 4304437) and the following TaqMan Gene Expression Assays (Applied 58 59 Biosystems/Thermo Fisher Scientific, Grand Island, NY): ACTB (Hs9999903 m1), FAM83A 70 (Hs04994801 m1), and NEK2 (Hs00601227 m1). qPCR reactions were run on a StepOne Plus Real-Time PCR 71 System (Applied Biosystems/Thermo Fisher Scientific) and data analyzed using the StepOne Software v2.3 (Applied Biosystems). Relative quantification was calculated with reference to ACTB and analyzed using the 72 comparative C_T method. gPCR experiments were performed in triplicate. 73

74 Tissue microarray (TMA) immunohistochemistry (IHC) analysis

Normal breast tissues microarrays from 683 women were generated from paraffin-embedded blocks 75 obtained from the KTB at the Tissue procurement & Distribution core of the IU Simon Comprehensive Cancer 76 77 Center. Due to loss of material during TMA construction and processing, 58% (n=395) of these tissue biopsies were interpretable. TMA was analyzed with the following antibodies FAM83A (Protein Tech 20618-1-AP, 78 1:100), NEK2 (MyBioSource MBS9607934, 1:100), Ki67 (DAKO IR 626, ready-to-use), estrogen receptor 79 30 alpha (ERa) (clone:EP1, DAKO IR 084, ready-to-use), FOXA1 (Santa Cruz Biotechnology sc-6553, 1:100), and GATA3 (Santa Cruz Biotechnology sc-268, 1:50) (70). IHC was performed in a Clinical Laboratory 31 Improvement Amendments (CLIA)-certified histopathology laboratory and evaluated by 3 pathologists in a 32 blinded manner. Quantitative measurements generating positivity and H-score were done using the automated 33 Aperio Imaging system using an FDA-approved algorithm (71). 34

35 Statistical analysis

Comparisons between groups were done using either Student's *t*-test or nonparametric Mann-Whitney test on GraphPad Prism 9. Difference between groups is considered significant at *p*-values<0.05. Pearson's correlation analysis was performed to determine the strength and direction of the linear relationship between DNA methylation and transcription for given targets. Only correlations with a p<0.05 are shown. For transcriptome and methylome data, differential analysis was performed using DESeq2 and the previously described Z-score method (65, 66), respectively. *P*-values <0.05 are considered significanct and are corrected

Э2 for multiple testing using the Benjamini-Hochberg False Discovery Rate (FDR) algorithm. For the tissue microarrays analysis nonparametric Wilcoxon rank-sum tests were used for unpaired analyses, as positivity and ЭЗ Э4 H-scores were not normally distributed, whereas nonparametric Wilcoxon signed-rank tests were used for Э5 paired analyses. The statistical software SAS version 9.4 (SAS Institute Inc., Cary, NC) was used to complete the statistical analyses with p < 0.05 considered significant. Baseline demographic characteristics were Э6 summarized as median (range) for continuous variables and number and percentage for categorical variables. Э7 Comparisons between groups were done using Chi-square tests (or Fisher's Exact test, where appropriate) for 98)9 categorical variables, or Wilcoxon test for continuous variables.

10 LIST OF ABBREVIATIONS

BC: breast cancer; **KTB:** Susan G. Komen Tissue bank at IU Simon Comprehensive cancer center; **IHC**:

)2 immunohistochemistry; **IPA**: Ingenuity pathway analysis

DECLARATIONS

Ethics approval and consent to participate: Breast specimens were obtained from the Susan G. Komen Tissue Bank at the IU Simon Comprehensive Cancer Center (KTB) and donated upon informed consent by healthy women volunteers. Subjects were recruited under a protocol approved by the Indiana University Institutional Review Board (IRB protocols number 1011003097 and 1607623663)

Consent for publication: Not applicable

Availability of data and materials: Methylome and Transcriptome data are available in Gene Expression Omnibus (GEO) with GSE164694 (<u>https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE164694</u>) which includes the Sub Series GSE164640 (MeCap dataset) and GSE164641 (RNA-seq dataset). Transcriptomic data of primary and immortalized breast epithelial cells from Dr. Nakshatri's team (20) were obtained from GEO with accession number GSE108541 (<u>https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE108541</u>).

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Authors' contributions: N.M. conceived the idea and designed the experiments, analyzed, and interpreted the data, and was major contributor in writing the manuscript. N.M. and RG performed the experiments and generated the data. A.V. assisted with tissue processing. R.P, D.R., J.H., J.W., G.S., and S.A. analyzed the data. C.T. performed the immunostaining of the tissue microarray. P.R. provided the specimens used in this study. K.N. and H.N. provided input on the manuscript. A.M.S. participated in hypothesis generation, funded the work, and contributed to the manuscript preparation. All the authors read and approved the manuscript.

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ADDITIONAL FILES

- Additional File 1: Supplementary Tables. File format:.xls. It includes subjects demographics and raw data in
 form of tables.
- Additional File 2: Supplementary Figues. File format:.pdf. It includes additional data related to the main
 findings shown in the main figures.
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FIGURE LEGENDS

Figure 1: Transcriptome profiling of breast tissues from women at either high- or average risk of breast
 cancer. A) Schematics of the study design. Cancer-free breast tissue cores (N=146) were divided in either

)8 high-risk or average-risk group according to the Tyrer-Cuzick breast cancer risk evaluation score (20% used as threashold). The tissues were processed and analyzed for whole transcriptome and methylome profiling and)9 LO differentially expressed genes (DEG) and differentially methylated sites between high- and average-risk samples were identified. Thirty five women (10 high risk and 25 average risk) donated also a second biopsy ι1 L2 (D2) allowing to detect age-dependent aberrations. **B**) Pathway analysis of the transcripts differentially expressed (FDR<0.05) between average and high- risk breasts. C) Major molecular network of the differentially L3 expressed transcripts between the two experimental groups. Genes upregulated in high-risk breasts are in red, L4 ۱5 while downregulated genes are in green. D) FAM83A and NEK2 transcription level in breast tissues from women at either average- or high-risk of developing breast cancer. E) Upper panel: Representative image of the ۱6 immunofluorescence staining of primary breast epithelial cells with the epithelial marker, E-Cadherin (red), L7 L8 mesenchymal marker, Vimentin (green) as control, and nuclear staining, DAPI (blue). E-Cadherin and Vimentin staining of primary cells revealed that isolated primary cells are epithelial in nature. Lower panel: ٢9 FAM83A and NEK2 expression in primary epithelial cells isolated from either average-risk (n=4) and high-risk 20 21 breast (n=3). F) FAM83A and NEK2 expression in primary and h-TERT immortalized isogenic breast epithelial cells (n=7) from the GSE108541 dataset. G) Representative images of immunohistochemical staining for 22 FAM83A and NEK2 are shown at 20X magnification. Staining quantification is expressed as positivity and H-<u>23</u> 24 score. Data are shown as mean \pm standard error. #: FDR<0.005, *: p<0.05, **p<0.001, ***p<0.0001. Pvalue is 25 calculated using either unpaired nonparametric Mann-Whitney test or nonparametric Wilcoxon test.

Figure 2: Methylome profiling of breast tissues from women at either high- or average risk of breast cancer. A) Chromosomal distribution of the DNA methylation aberrations observed in high-risk versus average-risk group. B) Heatmap of the 20 highest differentially methylated regions in high-risk breasts as compared with average-risk breasts at FDR<0.05. The overlapping gene name is indicated on the left. C) Genomic localization (intron, coding, promoter or UTR) of the DNA methylation aberrations including regions either hypo- or hyper-methylated in high-risk versus average-risk breasts. Data are shown as percentage of each genomic localizaton versus the total number of sites. D) Pathway analysis of the genes affected by DNA

- 33 methylation aberrations (FDR<0.05) in high-risk breasts as compared with breast from women at average risk
- for breast cancer. **E**) One of the molecular networks including the genes affected by DNA hypermethylation.

Figure 3: Correlation between degree of DNA methylation and gene expression. A) Pearson's correlation analysis between DNA methylation value and expression of the genes found differentially expressed between high- and average-risk breasts. B) Pearson's correlation analysis of the DNA methylation and expression of PHACTR1, hypermethylated in the breasts of high-risk women. *r* is the correlation coefficient and *p* is *p*value.

Figure 4: Age-related transcriptome and DNA methylation changes in healthy breast tissues. A) 39 10 Differentially expressed genes between the first (D1) and second (D2) donation time point in the breast tissues from average (blue bars) and high- (orange bars) risk women. Ratio between D2 and D1 is shown. B) Pearson's 11 correlation test between DNA methylation and transcription of GRIA4 and DNM3 in average- and high-risk 12 breasts at the two time points, D1 and D2, C) Number of genomic locations (intron, coding regions, promoter, 13 UTR) of the age-related DNA methylation events. N.A.: not available. D) Venn diagram of the DNA 14 methylation changes associated with age comparing our data set (D2/D1) with Horvath' epigenetic clock (353 15 CpGs) or Johnson's age-associated loci (787 CpGs) E) Differentially methylated regions between the first (D1) 16 and second (D2) donation time point in the breast tissues from average (blue bars) and high- (orange bars) risk 17 women. Ratio between D2 and D1 is shown. p<0.05; p<0.00118

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Table 1: Gene expression differences in high- versus average-risk breasts (FC>2; FDR<0.05)

_	Description	log2fc ^a	FDR	% genetic alterations ^b	Tumor/ Normal expression (<i>p</i> value) ^c	Copy Number Variation (CNV) ^d		Onco
Gene name						CNV=2 (%), p value	CNV=-2 (%), <i>p</i> value	score
MEPE	Matrix extracellular phosphoglycoprotein	2.28	2E-02	0.7	0.02 (n.s.)	12 (0.6), <i>n.s.</i>	2 (0.1), <0.001	15.6
OPRPN	Opiorphin prepropeptide	2.10	3E-03	1.3	N.A.	31 (1.4), <i>n.s.</i>	0 (0)	N.A.
CXCL13	C-X-C motif chemokine ligand 13	2.07	4E-03	1.3	6.6 (0,003)	26 (1.2), <i>n.s</i> .	0 (0)	33.7
APELA	Apelin receptor early endogenous ligand	1.87	8E-04	0.3	N.A.	N.A.	0 (0)	N.A.
CA6	Carbonic anhydrase 6	1.78	6E-04	0.8	0 (n.s.)	2 (0.1), <i>n.s.</i>	3 (0.1), <0.001	14.4
DIO2	lodothyronine deiodinase 2	1.60	2E-03	0.6	1.94 (n.s.)	13 (0.6), <i>n.s.</i>	0 (0)	7.7
FEZF2	FEZ family zinc finger 2	1.55	7E-03	0.7	0.04 (<0.001)	3 (0.1), <i>n.s.</i>	0 (0)	16.1
TNNT1	Troponin T1%2C slow skeletal type	1.52	9E-03	2.3	51.87 (n.s.)	36 (1.7), <i>n.s.</i>	0 (0)	12.3
MMP3	Matrix metallopeptidase 3	1.43	2E-02	1.8	5.66 (<0.001)	26 (1.2), <i>n.s</i> .	1 (0.04),<0.001	31.9
SERPINA12	Serpin family A member 12	1.42	2E-02	0.9	1.26 (<0.001)	12 (0.6), <i>n.s.</i>	1 (0), <0.001	11.9
C8B	Complement C8 beta chain	1.42	3E-02	1.8	0.014 (n.s.)	37 (1.7), <i>n.s.</i>	1 (0), <0.001	7.3
KCNJ13	Potassium voltage-gated channel subfamily J member 13	1.41	3E-03	0.6	0.16 (0.03)	2 (0.1), <i>n.s</i> .	1 (0), <0.001	9.0
CXCL6	C-X-C motif chemokine ligand 6	1.37	5E-03	2.2	0.10 (0.04)	43 (2), <i>n.s.</i>	0 (0), <i>n.s.</i>	31.0
SLC12A1	Solute carrier family 12 member 1	1.33	1E-02	0.9	0.48 (<0.001)	4 (0.2), <i>n.s.</i>	1 (0),<0.001	5.6
CYP24A1	Cytochrome P450 family 24 subfamily A member 1	1.33	3E-02	7.0	0.22 (n.s.)	164 (7.5),<0.001	1 (0), n.s.	30.2
ASB5	Ankyrin repeat and SOCS box containing 5	1.29	4E-03	1.3	0.01 (n.s.)	6 (0.3), n.s.	5 (0.2),<0.001	0.0
NPY2R	Neuropeptide Y receptor Y2	1.27	3E-02	1.0	0.003 (<0.001)	10 (0.5), n.s.	0 (0)	7.9
C2CD4A	C2 calcium dependent domain containing 4A	1.26	2E-02	0.6	0.9 (<0.001)	12 (0.6), n.s.	1 (0),<0.001	11.2
GABRR1	gamma-aminobutyric acid type A receptor rho1 subunit	1.26	3E-02	1.1	1.03 (0.03)	13 (0.6), n.s.	5 (0.2),<0.001	8.7
KIAA1210	KIAA1210	1.25	7E-03	1.6	0.43 (n.s.)	18 (0.8), n.s.	3 (0.1),<0.001	0.0

MMP10	Matrix metallopeptidase 10	1.23	2E-02	1.6	7.07 (<0.001)	26 (1.2), n.s.	1 (0), <0.001	38.2
FAM83A	Family with sequence similarity 83 member A	1.22	5E-03	16.0	1.23 (<0.001)	503(23.1),<0.001	0 (0)	74.5
LPO	Lactoperoxidase	1.21	1E-02	7.0	0.5 (<0.001)	168 (7.7),2E-24	1 (0), n.s.	11.5
CRISP2	Cysteine rich secretory protein 2	1.19	3E-02	1.5	0.06 (0.01)	31 (1.4), 3E-05	0 (0)	8.2
NMU	Neuromedin U	1.19	2E-02	0.8	3.6 (<0.001)	18 (0.8), n.s.	1 (0), <0.001	41.6
MAGEB4	MAGE family member B4	1.18	9E-03	0.8	8.6 (0.004)	10 (0.5), n.s.	2 (0.1),<0.001	55.9
MAG	Myelin associated glycoprotein	1.17	4E-02	2.3	5.3 (<0.001)	42 (1.9), 0.001	0 (0)	13.2
DAPL1	Death associated protein like 1	1.17	5E-03	0.7	0.09 (n.s.)	10 (0.5), n.s.	0 (0)	14.0
PRSS51	Serine protease 51	1.16	2E-02	1.6	N.A.	0 (0)	0 (0)	N.A.
PBK	PDZ binding kinase	1.14	4E-03	3.0	15.7 (<0.001)	20 (0.9), n.s.	15(0.7),<0.001	28.3
KRT77	Keratin 77	1.13	4E-02	0.8	0.04 (n.s.)	12 (0.6), n.s.	0 (0)	0.0
CALML3	Calmodulin like 3	1.12	3E-02	4.0	0.15 (n.s.)	108 (5),<0.001	0 (0)	37.7
ACBD7	Acyl-CoA binding domain containing 7	1.12	3E-03	2.3	1.13 (0.002)	78 (3.6),<0.001	0 (0)	0.0
UNC5D	Unc-5 netrin receptor D	1.11	2E-02	8.0	0.001 (n.s.)	152 (7), n.s.	6 (0.3),<0.001	44.8
ESCO2	Establishment of sister chromatid cohesion N-acetyltransferase 2	1.11	2E-03	3.0	8.02 (<0.001)	20 (0.9), n.s.	14(0.6),<0.001	25.1
BARX1	BARX homeobox 1	1.09	4E-02	5.0	1.54 (9E-08)	9 (0.4), n.s.	1 (0), <0.001	22.3
CTXND1	Cortexin domain containing 1	1.09	3E-02	0.0	N.A.	0 (0)	0 (0)	N.A.
SYT13	Synaptotagmin 13	1.08	4E-03	1.3	4.6 (<0.001)	36 (1.7), <0.001	1 (0), n.s.	38.8
PRAME	Preferentially expressed antigen in melanoma	1.06	2E-02	1.2	1.8 (<0.001)	21 (1), n.s.	1 (0), <0.001	82.6
SLC39A12	Solute carrier family 39 member 12	1.05	4E-03	2.4	0.18 (n.s.)	72 (3.3), <0.001	1 (0), n.s.	12.0
IGHV2-26	Immunoglobulin heavy variable2-26	1.04	4E-02	0.1	N.A.	0 (0)	0 (0)	N.A.
APLN	Apelin	1.04	7E-04	0.6	0.93 (n.s.)	16 (0.7), n.s.	2 (0.1),<0.001	13.8
IGHV3-30	Immunoglobulin heavy variable3-30	1.04	2E-02	0.1	N.A.	0 (0)	0 (0)	48.0
LPAR3	Lysophosphatidic acid receptor 3	1.04	8E-03	0.9	0.28 (n.s.)	13 (0.6), n.s.	0 (0)	12.9
ECEL1	Endothelin converting enzyme like1	1.03	2E-02	0.8	0.9 (n.s.)	1 (0), n.s.	1 (0), <0.001	N.A.
DCX	Doublecortin	1.03	6E-03	0.5	0.1 (0.02)	13 (0.6), n.s.	2 (0.1), <0.001	8.7
NEK2	NIMA related kinase 2	1.02	7E-03	12.0	25.78 (<0.001)	473 (21.8),<0.001	0 (0)	61.4

CWH43	Cell wall biogenesis 43 C-terminal homolog	1.02	3E-02	1.0	0.5 (<0.001)	6 (0.3), n.s.	0 (0)	12.9
PRSS21	Serine protease 21	1.01	3E-02	5.0	0.2 (n.s.)	154 (7.1),5E-102	0 (0)	46.3
FOXI3	Forkhead box I3	1.01	2E-02	0.3	0.01 (<0.001)	10 (0.5), n.s.	0 (0)	8.5
FCER2	Fc fragment of IgE receptor II	-0.98	1E-03	1.3	0.07 (0.04)	11 (0.5), n.s.	2 (0.1),<0.001	17.1
DACH2	Dachshund family transcription factor 2	-1.01	1E-02	0.8	0.3 (n.s.)	17 (0.8), n.s.	9 (0.4),<0.001	25.3
LILRB5	Leukocyte immunoglobulin like receptor B5	-1.02	8E-04	2.1	0.15 (<0.001)	39 (1.8), n.s.	0 (0)	0.0
SBK3	SH3 domain binding kinase family member 3	-1.03	7E-03	2.3	N.A.	48 (2.2), n.s.	0 (0)	0.0
TRDN	Triadin	-1.03	3E-02	2.3	0.02 (n.s.)	41 (1.9), n.s.	1 (0),< 0.001	1.0
NXF3	nuclear RNA export factor 3	-1.04	3E-03	0.6	0.9 (n.s.)	8 (0.4), n.s.	4 (0.2),< 0.001	32.2
LILRA6	leukocyte immunoglobulin like receptor A6	-1.05	2E-03	2.1	1 (n.s.)	39 (1.8), n.s.	1 (0), n.s.	0
SYNDIG1L	synapse differentiation inducing 1 like	-1.07	9E-03	0.5	N.A.	8 (0.4), n.s.	1 (0),< 0.001	0
ARPP21	cAMP regulated phosphoprotein 21	-1.13	2E-02	1.1	0.42 (n.s.)	11 (0.5), n.s.	1 (0),< 0.001	24.04
SLC22A12	solute carrier family 22 member 12	-1.13	2E-02	1.1	0.9 (<0.001)	20 (0.9), n.s.	0 (0)	8.9
CCL24	C-C motif chemokine ligand 24	-1.17	1E-02	0.7	0.98 (<0.001)	21 (1), n.s.	0 (0)	16.2
TPSD1	tryptase delta 1	-1.17	2E-02	5.0	0.55 (0.04)	170 (7.8),<0.001	0 (0)	0
PROK2	prokineticin 2	-1.19	2E-02	0.7	0.24 (0.01)	5 (0.2), n.s.	1 (0), 0.001	18.8
HBG2	hemoglobin subunit gamma 2	-1.59	4E-02	1.0	0.2 (n.s.)	19 (0.9), n.s.	0 (0)	11.3
FGF8	fibroblast growth factor 8	-1.68	3E-04	0.3	0.88 (0.005)	2 (0.1), n.s.	1 (0),< 0.001	14.3
SULT1C2	sulfotransferase family1C member2	-1.74	2E-03	0.5	1.6 (0.02)	9 (0.4), n.s.	0 (0)	21.8
MS4A6E	membrane spanning 4-domains A6E	-2.24	4E-02	0.9	N.A.	23 (1.1),n.s.	0 (0)	0

N.A. : not available; a: log fold change; b: breast cancer data from cBioportal; c: from UALCAN portal; d: data retrieved from the METABRIC, number of samples with either CNV= 2 or -2 (%), p value.

Table 2: The 20 most differentially methylated regions between the high- and average-risk breast tissues

Genomic Locus	Overlapping	Gene Name	Description	$\Delta Z^{\#}$	FDR
	Gene Feature		•		
Chr8:120,669,501	Intron	SNTB1	syntrophin beta 1	2.4	7E-53
Chr18:6,803,751	Intron	ARHGAP28	Rho GTPase activating protein 28	2.0	3E-35
Chr6:12,944,751	Intron	PHACTR1	phosphatase and actin regulator 1	1.9	1E-31
Chr21:30,743,501	promoter	KRTAP21- 4P	keratin associated protein 21-4 2C pseudogene	1.9	6E-31
Chr2:115,663,751	Intron	DPP10	dipeptidyl peptidase like 10	1.8	2E-30
Chr4:87,111,001	Intron	AFF1	AF4/FMR2 family member 1	1.8	2E-29
Chr3:33,638,251	Intron	CLASP2	cytoplasmic linker associated protein 2	1.8	2E-29
Chr14:106,498,501	Intron	LINC01881	long intergenic non-protein coding RNA1881	1.8	2E-29
Chr6:129,416,001	Intron	LAMA2	laminin subunit alpha 2	1.8	4E-29
Chr14:31,750,251	Intron	NUBPL	nucleotide binding protein like	1.8	2E-28
Chr8:37,842,001	Coding	ADGRA2	adhesion G protein-coupled receptor A2	-1.2	1E-12
Chr1:44,724,501	Coding	C1orf228	chromosome 1 open reading frame 228	-1.2	9E-13
ChrX:46,575,001	Coding	CHST7	carbohydrate sulfotransferase 7	-1.2	5E-13
Chr14:104,729,501	Coding	ADSSL1	adenylosuccinate synthase like 1	-1.3	1E-15
Chr2:202,774,251	Intron/promoter	ICA1L	islet cell autoantigen 1 like	-1.3	1E-15
Chr1:155,190,001	Coding	MUC1	mucin 1, 2C cell surface associated	-1.4	2E-17
Chr20:3,751,751	Coding	HSPA12B	heat shock protein family A (Hsp70) member 12B	-1.4	8E-18
Chr11:58,141,001	Intron	OR9Q1	olfactory receptor family 9 subfamily Q member 1	-1.6	4E-22
Chr1:45,803,751	Coding	MAST2	microtubule associated serine/threonine kinase 2	-1.6	3E-23
Chr7:636,001	Intron	PRKAR1B	protein kinase cAMP- dependent type I regulatory subunit beta	-1.7	1E-24

#: High- versus average-risk value

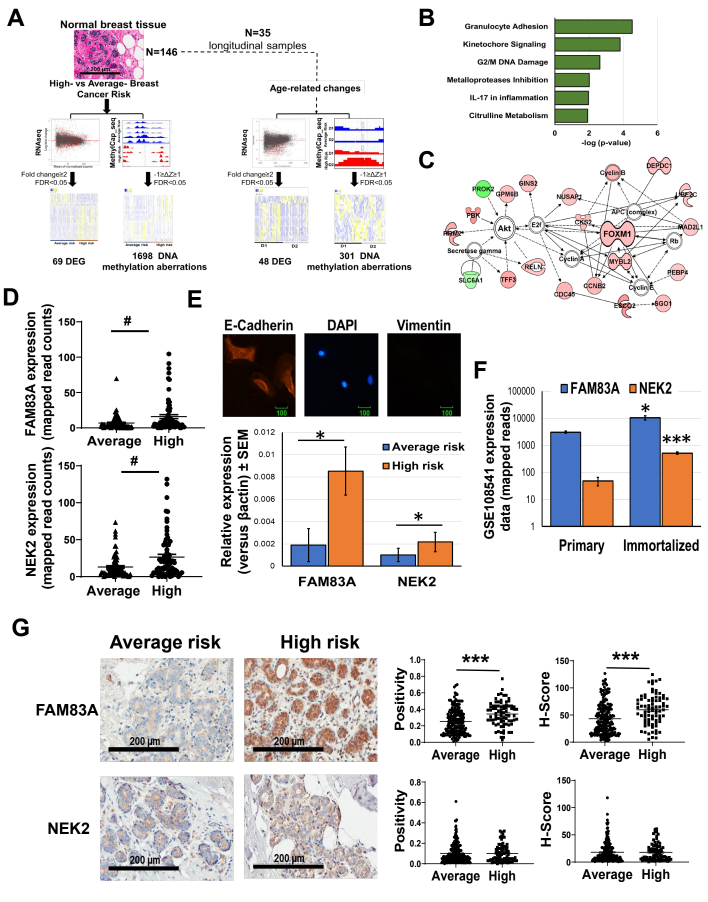
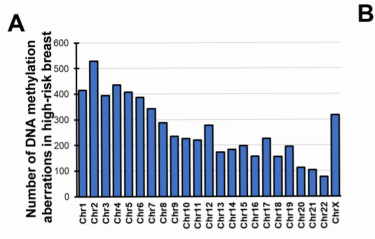
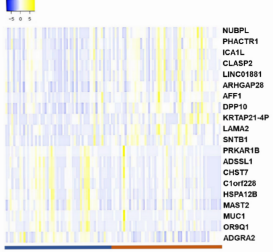


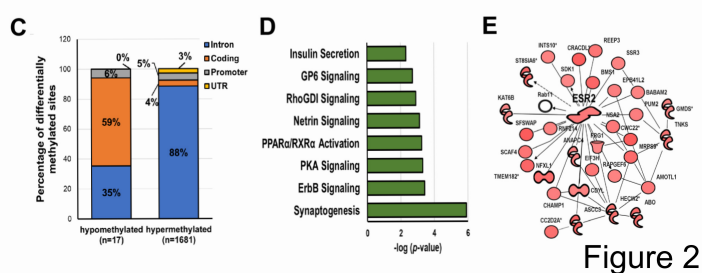
Figure 1

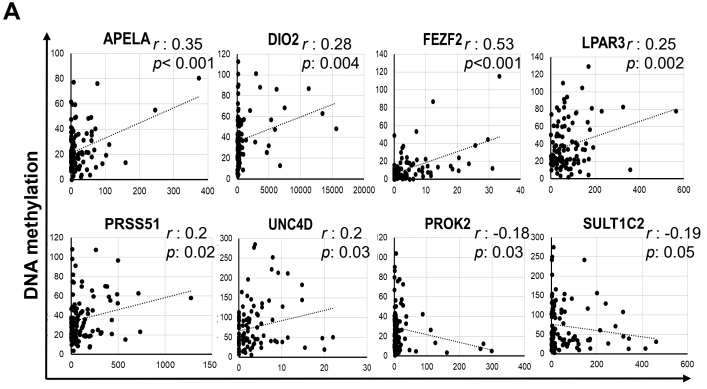




Average risk







Transcription (mapped reads count)

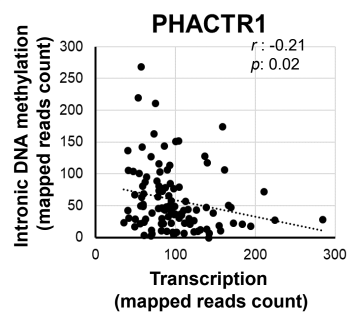
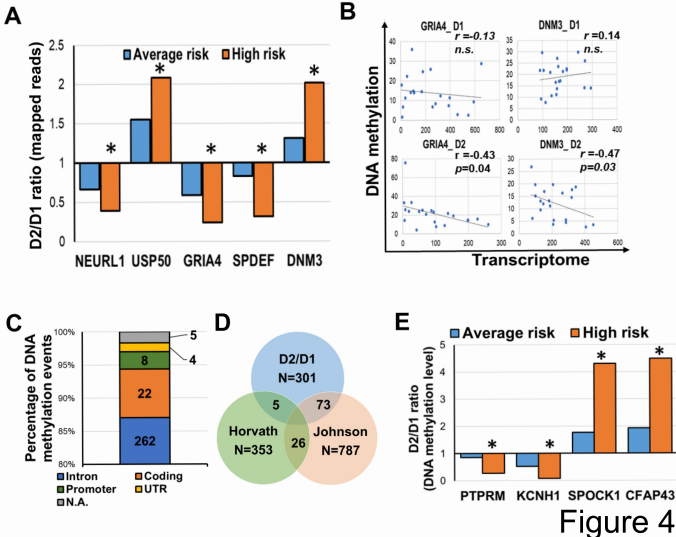


Figure 3

В



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