Research paper

DAGM: A novel modelling framework to assess the risk of HER2-negative breast cancer based on germline rare coding mutations

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\textbf{A B S T R A C T}

Background: Breast cancers can be divided into HER2-negative and HER2-positive subtypes according to different status of HER2 gene. Despite extensive studies connecting germline mutations with possible risk of HER2-negative breast cancer, the main category of breast cancer, it remains challenging to obtain accurate risk assessment and to understand the potential underlying mechanisms.

Methods: We developed a novel framework named Damage Assessment of Genomic Mutations (DAGM), which projects rare coding mutations and gene expressions into Activity Profiles of Signalling Pathways (APSPs).

Findings: We characterized and validated DAGM framework at multiple levels. Based on an input of germline rare coding mutations, we obtained the corresponding APSP spectrum to calculate the APSP risk score, which was capable of distinguishing HER2-negative from HER2-positive cases. These findings were validated using breast cancer data from TCGA (AUC = 0.7). DAGM revealed that HER2 signalling pathway was up-regulated in germline of HER2-negative patients, and those with high APSP risk scores had exhibited immune suppression. These findings were validated using RNA sequencing, phosphoproteome analysis, and CyTOF. Moreover, using germline mutations, DAGM could evaluate the risk for HER2-negative breast cancer, not only in women carrying BRCA1/2 mutations, but also in those without known disease-associated mutations.

Interpretation: The DAGM can facilitate the screening of subjects at high risk of HER2-negative breast cancer for primary prevention. This study also provides new insights into the potential mechanisms of developing HER2-negative breast cancer. The DAGM has the potential to be applied in the prevention, diagnosis, and treatment of HER2-negative breast cancer.

\begin{acknowledgement}
Abbreviations: AUC, area under the receiver operating characteristic curve; CCLE, Cancer Cell Line Encyclopedia; DAGM: damage assessment framework of genomic mutations; ER, Estrogen receptor; GDF: global driving force; GSEA, Gene Set Enrichment Analysis; GWAS: genome-wide association study; HER2, human epidermal growth factor 2; PBMC, peripheral blood mononuclear cells; PCC: Pearson correlation coefficient; PR, Progesterone receptor; PRS, polygenic risk score; RCM, rare coding mutations; SVM, Support Vector Machine; TCGA, The Cancer Genome Atlas; TNBC, Triple-negative breast cancer; WES, whole-exome sequencing
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Research in Context

Evidence before this study

The majority of hereditary breast cancers are caused by BRCA1/2 mutations, and the presence of these mutations is strongly associated with an increased risk of breast cancer. Meanwhile, BRCA1/2 gene mutations are rarely found in sporadic breast cancers and only account for a modest percentage of all breast cancer patients. Polygenic risk score (PRS), a widely-used approach for stratifying individuals according to their risk of a certain kind of complex disease, has been used to predict subjects at high risk for breast cancer. However, relying on single nucleotide polymorphisms (SNPs) from genome-wide association studies (GWAS) without including gene expressions or pathway activities means, PRS is not very suitable for cross-population prediction and describes disease risk in terms of genomic mutations without alluding to the underlying pathogenic mechanisms(s). Therefore, there is still an urgent need for a comprehensive, population-independent method to accurately assess the risk of breast cancer and to gain insights on potential mechanism(s).

Added value of this study

Analysing germline rare coding mutations (gRCMs) using DAGM framework results in the corresponding APSP (Activity Profiles of Signalling Pathways) and APSP risk score. Both APSP and APSP risk score can identify HER2-negative from HER2-positive breast cancers. These findings suggest HER2-negative breast cancer develops by a defined genomic evolutionary strategy. Furthermore, this study also revealed the up-regulation of HER2 signalling pathway in germlines of HER2-negative breast cancers and the immune suppression in subjects with high APSP risk score, shedding new light on the potential mechanisms of developing HER2-negative breast cancer. Moreover, our APSP risk score was able to relatively accurately evaluate the risk of developing HER2-negative breast cancer for each female, including not only BRCA1/2 carriers, but also non-carriers.

Implications of all the available evidence

The present study suggests that HER2 signalling pathway activity, as an aggressive factor, contributes to the development of different types of breast cancers, either via the combined effects of multiple germline mutations in HER2-negative germlines or via amplifying the gene itself in HER2-positive tumour cells. This provides a theoretical basis for the prevention, diagnosis, and treatment of breast cancers. At the same time, the study provides preliminary methods for assessing the relative risk of HER2-negative breast cancer for females with or without BRCA1/2 mutations. Finally, our findings provide a new perspective and theoretical basis for identifying high-risk female subjects, based on the high APSP risk score, for early screening and prevention of HER2-negative breast cancer.

1. Introduction

Breast cancer is the most common malignancy and the second leading cause of cancer death in women worldwide [1,2]. Breast cancer can be classified into two types: human epidermal growth factor 2 (HER2)-positive and HER2-negative. HER2-positive breast cancer is characterized by HER2 amplification in tumour tissues, whereas the HER2-negative breast cancer is not [3-7]. The majority (80%) of breast cancers cases are HER2-negative, including triple-negative breast cancer (TNBC), luminal A and luminal B (HER2-negative) [3-5]. Although germline mutations in genes such as BRCA1/2 are known to be associated with breast cancer, they are detected in only 5.3% of breast cancer patients and 11.2% of TNBC patients in China [8]. Signalling pathways such as STING pathway and immunosuppressive pathways have been linked to the development and progression of breast cancers [9-11]. However, the ability to accurately assess the risk and understand the underlying mechanisms of developing sporadic HER2-negative breast cancer remains elusive.

To address these issues, extensive studies have been carried out in the past decade, leading to various methods such as Gene Set Enrichment Analysis (GSEA) [12] and Polygenic risk score (PRS) [13]. The PRS has been widely used to stratify individuals according to their risk for complex diseases including breast cancer [14-16]. The PRS relies on SNPs from genome-wide association studies (GWAS) but does not include gene expressions or pathway activities. Thus, it does not allude to the underlying pathogenic mechanism although describes the risk in terms of the association between genomic mutations and the disease. In predicting breast cancer risk, the performance of PRS models demonstrated an area under receiver-operator curve (AUC) of around 0.6 [14-17]. Due to the ancestry-dependence of GWAS, PRS is only suitable for risk stratification of individuals from the same cohort rather than across populations. Therefore, there is an urgent need for a comprehensive, population-independent method, which can integrate genome mutations, gene expressions, and pathway activities to accurately assess the risk of breast cancer and to gain insights into the pathogenesis.

2. Methods

2.1. Overview

To accurately assess the risk of breast cancer and to understand the underlying mechanisms and pathogenesis, we developed the new DAGM (Damage Assessment of Genomic Mutations) framework to mine genomic information including rare coding mutations, gene expression, and signalling pathway activities. Instead of looking for certain genes that possibly determine cell fate, we assume the unique fate of a cell is determined by a set of genes through regulating the activities of functional modules such as signalling pathways. Therefore, we first constructed a framework based on public data from the COSMIC cell line project. Then we validated its reliability and applied the framework to whole-exome data from breast cancer patients and controls. Eventually, we validated our findings by applying experimental data from RNA-Seq, phosphoproteome analysis, CyTOF analysis, and using breast cancer data from TCGA.
2.2. Study participants

This study recruited 721 subjects including 434 breast cancer patients and 287 controls. The average age of the controls was 81 years. Patients with breast cancer were recruited from two independently operated hospitals in Guangdong province in southern China (Table 1), including 316 patients in the Guangdong Provincial People’s Hospital (GZ cohort: 62 ERBB2-positive and 43 Luminal B (HER2-positive), 77 Luminal B (HER2-negative), 24 Luminal A, and 110 TNBC) and 118 patients from the Shantou Affiliated Hospital (ST cohort: 15 ERBB2-positive and 103 TNBC). A total of 246 AD cases and 173 age- and ethnicity-matched cognitively normal individuals that do not carry APOE are were sent for whole exome sequencing in a previous study [18].

2.3. Ethics

The study protocol was approved by the Research Ethics Committee at Guangdong General Hospital, Guangdong Academy of Medical Sciences. Written informed consent was obtained from all participants for the use of banked tissue (including white blood cells and buccal cells) and for the collection of pathological data and clinical follow-up data.

2.4. Whole-exome sequencing and variant calling

Peripheral blood mononuclear cell (PBMC) samples were collected from breast cancer patients. DNA was extracted using QIAamp DNA mini kit (Qiagen) according to the blood and body fluid protocol in the user manual. Paired-end multiplex sequencing of case samples was performed on a Illumina HiSeq X Ten sequencing platform to a median depth of 150-250X. The whole blood genomic DNA samples from the control cohort were enriched using the TruSeq Kit (Illumina, California, USA) and were sequenced on an Illumina HiSeq 2000 system to a median depth >60X. Paired-end raw sequence reads were mapped to the human reference genome (UCSC hg19) using the Burrows-Wheeler Aligner [19] with default settings. Variant calling was carried out using the Genome Analysis Toolkit (GATK) with the HaplotypeCaller module [20] according to GATK Best Practices. Briefly, the aligned BAM files were first marked for duplicate reads by Picard. Local realignment around indels and base quality score recalibration were performed using GATK. The processed BAM files were then used to call SNPs and indels. Variant filtering of SNPs and indels were performed separately by variant quality score recalibration using GATK. The filtered variants were then annotated by ANNOVAR [21].

2.5. Transcriptomics

A total of 3 \( \mu \)g of RNA per sample was used as the input material for RNA sample preparation. Sequencing libraries were generated using NEBNext UltraTM RNA Library Prep Kit for Illumina (NEB, USA) following the manufacturer’s instructions. Index codes were added to attribute the sequences to each sample. Sequencing was performed on an Illumina HiSeq platform. Raw data (raw reads) in FASTQ format were processed through our in-house Perl scripts to clean the data (clean reads), which removed reads containing adapters, reads containing poly-N, and low quality reads. All downstream analyses were based on high-quality clean data. Paired-end clean reads were aligned to the reference genome using Hisat2 v2.2.1 [22]. FeatureCounts v2.0.1 [23] was used to count the numbers of reads mapped to each gene. Only genes with at least one count per million in at least two samples were kept for the following analysis. Read counts were normalized using the TMM normalization method performed in the edgeR package v3.26.8 [24]. The RPKM values of the genes were calculated based on the length of the gene and read counts mapped to the gene.

2.6. Phosphoproteome analysis

Samples were minced individually in liquid nitrogen. The enrichment was carried out using PHOS-Select iron affinity gel (Sigma, P9740) following the manufacturer’s instructions. Shotgun proteomics analyses were performed using an EASY-nLC™ 1200 UHPLC system (Thermo Fisher) coupled with an Orbitrap Q Exactive HF-X mass spectrometer (Thermo Fisher) operating in a data-dependent acquisition (DDA) mode. The resulting spectra from each fraction were searched separately against the UniProt database [25]. For protein identification, proteins with at least one unique peptide were identified at an FDR less than 1.0% at the peptide and protein level, respectively.

2.7. CyTOF

Antibodies were either purchased pre-conjugated (Fluidigm, DVS Sciences) or purchased and conjugated in-house using MaxPar X8 Polymer Kits (Fluidigm) according to the manufacturer’s instructions (Suppl. Table 1). Scans were acquired on a Helios 2.0 (Fluidigm) at an event rate of 300 events/s. After normalizing and randomizing values to near zero using the Helios software, FCS files were then generated.

Table 1

Study population characteristics.

| Characteristic                     | Luminal A | Luminal B HER2- | Luminal B HER2+ | ERBB2-positive | TNBC |
|-----------------------------------|-----------|-----------------|-----------------|----------------|------|
| Age, Mean (range, years)          | 50 (38-71)| 47 (22-80)      | 48 (25-63)      | 51 (22-72)     | 50 (23-82) |
| Histological grade, n (%)         | 1 (42)    | 1 (77.1)        | 0 (43)          | 0 (77)         | 5 (213) |
| 2                                 | 14 (24)   | 36 (77.1)       | 23 (43.5)       | 28 (77.36)     | 78 (213) |
| 3                                 | 7 (24)    | 31 (77.1)       | 16 (43.72)      | 40 (77.51)     | 109 (213) |
| Unknown                           | 2 (24)    | 9 (77.117)      | 4 (43.9)        | 9 (77.117)     | 21 (213) |
| Turnover size, n (%)              | ≤2 cm     | 6 (24.5)        | 24 (77.1)       | 10 (43.33)     | 51 (213) |
| >2 cm                             | 15 (24)   | 50 (77.164)     | 31 (43.71)      | 57 (77.740)    | 157 (213) |
| Unknown                           | 3 (24)    | 37 (77.139)     | 2 (43.47)       | 9 (77.47)      | 5 (213) |
| Nodal status, n (%)               | ≤10       | 2 (24)          | 11 (77.143)     | 8 (43.186)     | 5 (213) |
| >10                               | 10 (24)   | 23 (77.129)     | 13 (43.302)     | 23 (77.299)    | 47 (213) |
| unknown                           | 1 (24)    | 3 (77.139)      | 1 (43.23)       | 1 (77.13)      | 10 (213) |

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for the analysis. Mass cytometry data was de-barcoded using a dou-
blet filtering scheme with mass-tagged barcodes, and then manually
gated to retain live, singlet, and valid immune cells. Data generated
from different batches were normalized through the bead normaliza-
tion method. All cell events in each individual sample were pooled
and included in the analysis.

2.8. The Damage Assessment of Genomic Mutations (DAGM) framework

We developed a novel framework called Damage Assessment of
Genomic Mutations (DAGM), which integrates genome-wide informa-
tion of rare coding mutations and gene expression from the COSMIC cell
line project to calculate the activity profile of signalling pathways
(APSPs). DAGM consists of three sequential steps (Fig. 1, see supplemen-
tary method for details). The first step takes the mutated genes and gene
expressions as the inputs to determine the driving force of each
mutated gene on the expression of all genes as the output. The global
driving force (GDF) is then calculated and the global driving genes are
identified in the cell. The second step evaluates the activity pro-
file of signalling pathways (APSP) in three sequential steps (see supplemen-
tary method for full details). The first step takes the mutated genes and
gene expressions as the input and calculates the effect of the rare
mutations on gene expression changes within the cell lines, which
will generate a n x n matrix. In this matrix each row represents a
mutated gene, and each column denotes the expression of a gene,
and the value in the cell was de
cided. The second step calculates the combined effects of all muta-
tions that one individual carries. The third step is to evaluate the activity
profile of signalling pathways (APSPs). To identify the global driver
genes regardless of different extrinsic/intrinsic cellular conditions, we
assessed the driving force based on the genome-wide data of rare cod-
ing mutations and gene expression in 970 cancer cell lines (COSMIC Cell
Line project), considering cancer cell lines exhibit high tumour purity
and low expression heterogeneity. To validate this framework, we used
another dataset from Cancer Cell Line Encyclopaedia (CCLE) [26] to build
a ‘driving force’ matrix.

Similar to the eQTL method that uses gene expression as an inde-
dependent trait to determine its linkage site in the genome, we can cor-
correlate the gene expression in cell lines to ‘traits’ and associate them
with a variety of different genes with rare coding mutations (RCMs).
However, rather than examining how all genes with rare mutations
change the gene expression traits, DAGM screens out genes defined as
global driver genes, whose mutations determinationistically cause
almost the same alterations in the global gene expression across dif-
f erent cellular or tissue contexts. For example, the well-known driver
gene TP53 drives similar altered gene expressions in tumour cells
from different origins. We are continuously collecting RCM data and
gene expression traits from cancer cell lines or primary cells to
improve the database and to provide more accurate global driver
genes. Based on a sample with only RCM data, DAGM was able to
determine candidate genes based on the RCM distribution, in which
the effects of driver genes were combined to give the altered gene
expression, representing the mutations as an overall impact on gene
expression. The DAGM can then determine the activity profiles of sig-
nalling pathways (APSP) in three sequential steps (see supplemen-
tary method for full details). The first step takes the mutated genes and
gene expressions as the input and calculates the effect of the rare
mutations on gene expression changes within the cell lines, which
will generate a n x n matrix. In this matrix each row represents a
mutated gene, and each column denotes the expression of a gene,
and the value in the cell was determined.

2.9. Classification

Binary classifiers were built using the support vector machine
(SVM) and logistic regression implemented in the caret R package
[27]. Support vector machine (SVM), aims to create a decision bound-
ary (known as the hyperplane) between two classes that enables the
prediction of labels from one or more feature vectors, is a powerful
supervised learning method for building a classifier. The radial basis
function (RBF) kernel that is commonly used in SVM classification was
used in this study [28]. Logistic regression is an extremely robust and
flexible method for dichotomous classification prediction. It is simple
and efficient for binary and linear classification problems and is widely
used in biological studies [29]. All data were randomly split into the
training dataset (60% or 75%) and the testing dataset (40% or 25%),
while preserving the overall class distribution of the data. The five-fold
cross-validation procedure was used to optimize the hyperparameters
of a classifier on the training dataset. The criteria of the area under the
receiver operating characteristics curve (AUC) were used to measure
the performance of the classifiers on the testing dataset.

2.10. Activity profiles of signalling pathway (APSP) risk score

Three pathway panels including growth factor, pro-tumour, and
immune function pathways were significantly downregulated in
HER2-negative breast cancers and were selected to calculate the APSP
risk score. The APSP risk score was calculated as the weighted mean of
the APSPs of the three selected pathway panels in each person.

![Workflow of the DAGM framework.](image)

The DAGM framework mainly composed of three steps. In the first step (grey colour), driving force (DF) and global driving force (GDF) are established for future analysis based on the rare coding mutations and gene expression from the COSMIC Cell Lines project. When obtaining the germline rare coding mutations of a subject, we calculated the combined
effect of all these mutations (Step 2). In the last step, the activity profile of signalling pathways (APSPs) is evaluated for this subject.
Table 2
Signalling pathways for DAGM framework.

| Index | Pathway                      | Pathway Panel          |
|-------|------------------------------|------------------------|
| 1     | CDK5                         | Development            |
| 2     | Hedgehog                    | Development            |
| 3     | NOTCH                       | Development            |
| 4     | Hypoxia                     | Response to stress     |
| 5     | ROS                         | Response to stress     |
| 6     | Unfolded protein response   | Response to stress     |
| 7     | UBV induced MAPK             | Response to stress     |
| 8     | UV response                  | Response to stress     |
| 9     | Adipogenesis                | Energy & metabolism    |
| 10    | AMPK                        | Energy & metabolism    |
| 11    | mTORC1                      | Energy & metabolism    |
| 12    | mTOR                        | Energy & metabolism    |
| 13    | Beta Cells                  | Energy & metabolism    |
| 14    | PI3K-AKT-mTOR               | Energy & metabolism    |
| 15    | PI3K-AKT                    | Energy & metabolism    |
| 16    | PPARA-RXRA                  | Energy & metabolism    |
| 17    | PRA                         | Energy & metabolism    |
| 18    | VDR-RXR                     | Energy & metabolism    |
| 19    | 4-1BB                       | Immune system          |
| 20    | BCR                         | Immune system          |
| 21    | CD40                        | Immune system          |
| 22    | IL2-Stat5                   | Immune system          |
| 23    | INF alpha                   | Immune system          |
| 24    | INF gamma                   | Immune system          |
| 25    | Jak-Stat                    | Immune system          |
| 26    | LBC                         | Immune system          |
| 27    | NF-kappaB                   | Immune system          |
| 28    | Toll like receptor          | Immune system          |
| 29    | AML                         | Pro-tumour             |
| 30    | Angiogenesis                | Pro-tumour             |
| 31    | Angiopoietin                | Pro-tumour             |
| 32    | Colorectal cancer Metastasis| Pro-tumour             |
| 33    | Estrogen-dependent BCA      | Pro-tumour             |
| 34    | Gisieblastoma               | Pro-tumour             |
| 35    | Gioma invasiveness          | Pro-tumour             |
| 36    | Gioma                       | Pro-tumour             |
| 37    | NSCLC                       | Pro-tumour             |
| 38    | Pancreatic adenocarcinoma   | Pro-tumour             |
| 39    | Renal cell carcinoma        | Pro-tumour             |
| 40    | Renin angiotensin           | Pro-tumour             |
| 41    | Apoptosis                   | Anti-tumour            |
| 42    | Ceramide                    | Anti-tumour            |
| 43    | TP53                        | Anti-tumour            |
| 44    | PTEN                        | Anti-tumour            |
| 45    | DNA synthetic checkpoint    | Cell cycle & proliferation |
| 46    | DNA damage checkpoint       | Cell cycle & proliferation |
| 47    | Erk/MapK                    | Cell cycle & proliferation |
| 48    | KRAS                        | Cell cycle & proliferation |
| 49    | Telomerase                  | Cell cycle & proliferation |
| 50    | EGFR                        | Growth factors         |
| 51    | Her2                        | Growth factors         |
| 52    | Erbb4                       | Growth factors         |
| 53    | HGF                         | Growth factors         |
| 54    | IGF-1                       | Growth factors         |
| 55    | PDGF                        | Growth factors         |
| 56    | TGF-beta                    | Growth factors         |
| 57    | VEGF                        | Growth factors         |
| 58    | Wnt calcium                 | Growth factors         |
| 59    | Wnt beta-catenin            | Growth factors         |
| 60    | Wnt calcium                 | Growth factors         |

weights were $-1/n$ and $1/n$ for the HER2 signalling pathway and for other pathways, respectively.

2.11. Statistical analysis

The heatmap R package was used to plot the clustered heatmaps using Ward’s criteria [30]. The ggpubr R package was used to draw the scatter plots, boxplots, bar plots, histograms, and linear regression lines with 95% confidence interval. The plotROC R package was used to draw ROC. Pearson correlation coefficients (PCC) and P-values were also labelled on the plots. The difference between two groups was analysed by Student’s t test. A P-value < 0.05 was considered statistically significant. To test the significance of the differences of APSP between different groups of individuals, P-values were calculated by permutation (1,000,000 times). P-values were adjusted for multiple testing since multiple hypotheses were tested simultaneously. The false discovery rate (FDR) was computed using Benjamini and Hochberg procedure [31] to correct for multiple hypothesis testing since FDR is often more appropriate and useful in high-throughput biological experiments [32]. FDRs passing cut-off of 5% were accepted as significant. All statistical analyses and plots were conducted using in-house scripts developed in Python, Perl, and R.

2.12. Data sharing statement

The raw exome sequencing data reported in this paper have been deposited in the Genome Sequence Archive (GSA) in National Genomics Data Centre, Beijing Institute of Genomics, Chinese Academy of Sciences, under accession number HRA000285, which is publicly accessible at https://bigd.big.ac.cn/gsa. All other data and materials are available upon request.

3. Results

3.1. Characterization and validation of the DAGM framework

We developed a novel framework called Damage Assessment of Genomic Mutations (DAGM), which consists of three sequential steps. To validate this framework, we used another dataset from Cancer Cell Line Encyclopaedia (CCLE) [26] to build a ‘driving force’ matrix. The GDFs calculated from the two different datasets were significantly correlated (coefficient of determination: $R^2 = 0.99$, P-value < 1.0 × 10⁻¹⁶, Suppl. Fig. 1a), suggesting the first step of DAGM is robust across different datasets. Next, we selected the top 10 genes with highest GDF in the DAGM and searched public databases of cancer driver genes. We found six of these genes, KRAS, TP53, MYC, BCL2, BRAF, and RPL22, were listed in COSMIC Cancer Gene Census [33], whereas all 10 were listed in DriverDBv3 [34], which confirmed the potential ability of using GDF to identify essential genes for tumourigenesis, including cancer driver genes.

To test whether the APSP calculated from DAGM reflects signaling pathway activities, we applied DAGM to somatic mutations from tumour samples from 16 ERBB2-positive patients and 42 TNBC patients (Suppl. Fig. 1b). Interestingly, although HER2 amplification was not included in the input mutation information, the resultant APSP showed significantly higher HER2 signalling pathway activity (T-test, P-value = 0.00095) in ERBB2-positive tumours compared to TNBC (Suppl. Fig. 1c). This result was consistent with the upregulation of the HER2 signalling pathway in HER2-positive breast cancer samples, which was also demonstrated by fluorescence in situ hybridization (FISH) [3].

3.2. APSP spectrum based on germline mutations distinguishes between HER2-negative and HER2-positive breast cancers

As is well known, it is of great importance to distinguish HER2-negative from HER2-positive breast cancer, but this currently relies on HER2 FISH of tumour tissues. To test whether the APSP from DAGM can be used to identify breast cancer subtypes, we collected germline rare coding variants from 721 subjects, which included 434 breast cancer patients of different subtypes and 287 cancer-free female subjects in the control group. The DAGM analysis using these input germline mutations (Suppl. Fig. 2) revealed the APSP spectrums for HER2-negative patients were remarkably lower than that for HER2-positive patients or controls (Fig. 2a). Further hierarchical clustering using Euclidean distance showed that Luminal A, Luminal B (HER2-nagetive), and TNBC were grouped together, whereas ERBB2-positive and Luminal B (HER2-positive) were clustered into another
group (Fig. 2b), suggesting the germline APSP spectrum can distinguish HER2-negative breast cancer from HER2-positive breast cancer subtypes. To quantitatively evaluate the performance of APSP in identifying breast cancer subtypes, we built a binary classifier based on the APSP of all 60 pathways using the support vector machine (SVM) with a radial basis function kernel with five-fold cross validation. The results showed an average value of the area under the receiver operating characteristic curve (AUC) of 0.77 for the classifiers between Her2-negative and Her2-positive patients (Fig. 2c), and 0.76 between Her2-negative and controls (Fig. 2d). Similar AUCs were obtained from different classifiers that were built using logistic regression with five-fold cross validation (Suppl. Table 2). Therefore, the APSP spectrum based on germline mutations can distinguish HER2-negative from HER2-positive breast cancer subtypes.

3.3. APSP reveals up-regulation of the HER2 signalling pathway in the germlines of HER2-negative breast cancer

Although BRCA1/2 mutations are currently used in breast cancer screening [35-37], carriers of these mutations represent only a small number of HER2-negative breast cancer patients. To find potential signalling features that are characteristic of HER2-negative breast cancer, we further examined the APSPs in more depth. Interestingly, most cellular signalling pathways were drastically down-regulated in HER2-negative patients compared to control subjects, whereas almost no significant changes in HER2-positive patients were observed (Fig. 3a). Next, we used TCGA data to crosscheck the activity differences in the 60 pathways between HER2-positive and HER2-negative patients, which showed significant correlations with our results (Pearson's correlation coefficient = 0.93, P-value = 3.8e-27, Suppl. Fig. 3a). Furthermore, the Z-scores of the APSPs of 60 signalling pathways were strongly correlated between HER2-negative subtypes (Luminal B (HER2-negative) and TNBC) (Pearson's correlation coefficient = 0.86, P-value = 1.14e-18, Fig. 3b) but not for HER2-positive subtypes (Luminal B (HER2-positive) vs. ERBB2-positive) (Pearson's correlation coefficient = 0.56, P-value = 3.92e-6, Fig. 3c). These findings suggest that HER2-negative breast cancer can be characterized by germline APSP features.

Among the significantly altered pathways according to APSP, we found significant up-regulation of the HER2 signalling pathway in germlines of HER2-negative breast cancer patients, even though HER2 gene was not amplified in the tumour tissues. To validate this finding, we performed RNA-Seq and phosphoproteome analysis using breast cancer and paracancerous (normal breast) tissues from both ERBB2-positive and TNBC patients. The expression of signature genes of the HER2 signalling pathway in paracancerous tissues were systematically lower in TNBC patients than in ERBB2-positive patients (Fig. 3d, left panel), despite high expression of these genes in TNBC tumour tissues (Fig. 3d, right panel). These observations indicated upregulation of the HER2 signalling pathway in TNBC paracancerous tissues and supported the conclusion inferred from APSP. Furthermore, the results from phosphoproteome analysis confirmed activation of the HER2 signalling pathway in paracancerous tissues of TNBC compared to ERBB2-positive patients (Fig. 3e). Therefore, the upregulation of HER2 signalling pathway in germlines revealed by APSP are a specific feature of HER2-negative patients, as validated by RNA-Seq and phosphoproteome data.

3.4. APSP risk score based on germline mutations is able to identify HER2-negative breast cancer

We demonstrated APSP spectrum could distinguish HER2-negative breast cancer from controls and HER2-positive breast cancer. Next, we classified the APSPs of the 60 signalling pathways into eight panels according to their functions. The pathways associated with the immune system, growth factor and pro-tumour pathways were significantly down-regulated in HER2-negative breast cancers (Fig. 4a), indicating these three panels are characteristic of HER2-negative breast cancer. Based on these three panels, we calculated the APSP risk score for each individual and found the risk scores were significantly higher in HER2-negative breast cancer patients than in controls (T-test, P-value = 3.96e-27) or in HER2-positive breast cancer.

Fig. 2. Germline APSP spectrum distinguishes HER2-negative breast cancer patients from HER2-positive patients. (a) Heatmap of the germline APSPs of 60 pathways for 721 subjects. Each row represents one pathway and each column represents one subject. (b) Hierarchical clustering dendrogram. HER2-negative subtype (Luminal B (HER2-negative), Luminal A, and TNBC) and HER2-positive subtype (ERBB2-positive and Luminal B (HER2-positive)) were well separated by hierarchical clustering with Euclidean distance as the distance measure. (c) Receiver operating characteristic (ROC) curve for distinguishing HER2-negative breast cancer patients from HER2-positive patients by APSPs. Area under the curve (AUC) was 0.77. (d) Receiver operating characteristic (ROC) curve for distinguishing HER2-negative breast cancer patients from controls by APSPs. Area under the curve (AUC) was 0.76.
Fig. 3. Upregulated HER2 signalling pathway in germlines of Her2-negative breast cancer patients.

(a) Difference of activity profiles of signalling pathways (APSPs) between breast cancer patients and controls. To test the significance of the difference, P-values were calculated by permutation (1,000,000 times). The FDR corrected P-values were also calculated to correct the multiple comparisons. The asterisk denotes FDR corrected P-value < 0.05. Most pathways activities were significantly down-regulated in HER2-negative breast cancer germlines (upper panel), whereas the HER2 signalling pathway activity was significantly up-regulated (rectangle, upper panel). However, only a few pathway activities were significantly altered in HER2-positive breast cancer patients (lower panel). (b) Correlation of APSPs between TNBC and Luminal B (HER2-negative). For HER2-negative breast cancer, germline pathway APSPs between TNBC and Luminal B (HER2-negative) subtype were strongly positively correlated. The black triangle represents the HER2 signalling pathway. (c) Correlation of APSP between ERBB2-positive and Luminal B (HER2-negative). For HER2-positive breast cancer, germline pathway APSPs between ERBB2-positive and Luminal B (HER2-positive) subtype were weakly positively correlated. (d) RNA-Seq validation of the upregulated HER2 signalling pathway. We performed RNA-Seq on both cancerous and paracancerous tissues from TNBC and ERBB2-positive breast cancer patients, the expression level of HER2 pathway signature genes in paracancerous tissue were significantly lower in the TNBC compared to ERBB2-positive patients, which validated the upregulated HER2 signalling pathway in HER2-negative breast cancer germlines. (e) The phosphoproteome validation the upregulated of the HER2 signalling pathway. Phosphoproteome analysis of both cancerous and paracancerous tissues from TNBC and ERBB2-positive breast cancer patients validated the upregulated HER2 signalling pathway in HER2-negative breast cancer germlines.
patients (T-test, $P$-value = 4.88e-22) (Fig. 4b). A similar difference in the APSP risk scores was observed using the TCGA data ($P$-value = 5.4e-11, Suppl. Fig. 3b), suggesting the APSP risk score can be used to identify HER2-negative breast cancer. Given that women carrying BRCA1/2 mutations have significantly increased risk for breast cancer [35-37], we compared the APSP risk scores for HER2-negative patients with or without BRCA1/2 mutations. We found all HER2-negative patients had higher APSP risk scores than control subjects regardless of BRCA1/2 mutations (Fig. 4c T-test, $P$-value=9.5e-06 and $P$-value=3.1e-24 for BRCA1/2 carriers and non-carriers, respectively). These findings imply that APSP risk score can be used to recognize not only HER2-negative patients with BRCA1/2 mutations, but also those without any known biomarkers.

To quantitatively evaluate the use of APSP risk score in identifying HER2-negative breast cancer, we built a binary classifier based on the APSP risk score using the logistic regression with five-fold cross validation. The results showed an AUC of 0.79 for the classifiers between HER2-negative and HER2-positive cases (Fig. 4d), and 0.74 for the classifier between HER2-negative and controls (Fig. 4e). Similar AUCs were obtained from different classifiers that were built using support vector machine (SVM) with a radial basis function kernel with five-fold cross validation (Suppl. Table 2). Using data from TCGA, the logistic regression binary classifier could distinguish between HER2-
negative and HER2-positive breast cancer patients with an average AUC value of 0.70 (Suppl. Fig. 3c). Taken together, germline APSP risk score can be used to identify HER2-negative breast cancer patients.

3.5. Enhanced immune suppression in subjects with high APSP risk scores

The APSP risk score was determined using three significantly downregulated pathway panels, including the immune system panel. Therefore, we reasoned that subjects with higher APSP risk scores would exhibit different altered immune states. To this end, we performed single-cell mass cytometry to count the number of different types of T cell types. Strikingly, we observed significantly increased exhausted CD8+ T lymphocytes in those with a high APSP risk score compared to those with a low risk score group (Fig. 5a, T-test, P-value = 0.0028), indicating an association between APSP risk score and immune suppression. Consistently, we also found significantly increased levels of exhausted CD8+ T lymphocytes in TNBC patients compared to controls (Fig. 5b, T-test, P-value = 0.00038), indicating enhanced immune suppression in TNBC patients with high APSP risk scores. These results revealed that subjects with higher APSP risk score also have suppressed immunity, which possibly contributes to an increased risk of HER2-negative breast cancer.

**Fig. 5.** Enhanced immune suppression and risk stratification of HER2-negative breast cancer.

Mass cytometry analysis revealed percentage of exhausted CD8+ T cells in female subjects of low and high APSP risk scores. (b) Mass cytometry analysis revealed percentage of exhausted CD8+ T cells in TNBC patients and control subjects. (c) Stratification of HER2-negative patients (red dots) as well as control subjects (green dots) into six groups, G1 to G6, according to the different standard deviations of APSP risk scores from their mean value. G1 and G6 included subject with APSP risk score two standard deviations lower and higher than the mean value, respectively. G2 and G5 included subject with APSP risk score between one and two standard deviations lower and higher than the mean value, respectively. G3 and G4 included subject with APSP risk score within one standard deviation lower and higher than the mean value, respectively. (d) Odds ratios for HER2 negative breast cancer display exponential distribution when the subjects were stratified into six groups (G1 to G6) as in (c). The grey circles represent the sample sizes of each group.
3.6. APSP risk score based on germline mutations stratifies risk of HER2-negative breast cancer

As the APSP risk score could identify HER2-negative breast cancer subtype, we wondered if it could be used to quantitatively used to evaluate the risk of HER2-negative breast cancer. We divided the subjects into six groups (G1 to G6) and calculated their corresponding odds ratio for HER2-negative breast cancer. The odds ratio increased with increasing APSP risk score from G1 to G6 (Fig. 5c) and followed exponential distributions (Fig. 5d). There was only one patient in the highest risk group G6 and no controls. The odds ratio in group G5 reached 7.7, indicating these individuals have a high risk for HER2-negative breast cancer, whereas the odds ratio in G1 was 0.08, indicating these individuals have a much lower risk for HER2-negative breast cancer. In summary, DAGM can identify HER2-negative breast cancer based on germline mutations and the calculated APSP risk score can be used to stratify the risk.

4. Discussion

HER2-negative breast cancer is the main category of breast cancer, the most common malignancy worldwide and the second leading cause of cancer death in women. It remains challenging to evaluate the risk for HER2-negative breast cancer in healthy female, despite extensive studies [14-17,38]. Here, we developed a new framework, DAGM (Damage Assessment of Genomic Mutations). Different to other approaches [12,14-17,38,39], DAGM integrates gene mutations and gene expressions data to assess the risk and potential pathogenesis of a disease. DAGM can use this information to calculate the activity profiles of signalling pathways (APSPs) and then derive the APSP risk score to assess the disease risk in an individual subject. Using germline mutations, DAGM was able to distinguish HER2-negative breast cancer patients from HER2-positive patients. The derived germline APSP risk score was able to predict the risk of developing HER2-negative breast cancer, not only in those with BRCA1/2 mutations, but also in those without any known disease-associated mutations. Furthermore, DAGM revealed that the HER2 signalling pathway was upregulated in the germline of HER2-negative patients, and those with high APSP risk scores also had elevated immune suppression. The findings were validated by RNA-Seq, phosphoproteome analysis, and single-cell mass cytometry.

To test whether the APSP can distinguish different breast cancer subtypes or recognize HER2-negative breast cancer patients from healthy subjects, we collected germline rare coding variants from breast cancer patients and cancer-free control subjects. When planning the research design, two candidate cohorts of healthy female subjects without breast cancer could be used as controls: the 81-year-old group that we chose and the alternative group of around 50-year-old. As this study aims to determine the potential contribution of germline rare coding mutations on the risk of HER2-negative breast cancer, we need gender-matched healthy subjects whose germline genome does not include breast cancer-associated information. As is already known, around one eighth of the general female population is diagnosed with breast cancer in their lifetime. Likewise, more than 10% of ~50-year-old “healthy” females in the alternative cohort will be diagnosed with breast cancer in their future lifetime, which makes them unsuitable for a best choice as healthy controls containing no disease-associated germline mutations. Meanwhile, the female cohort with an average age of 81 could be regarded as “winners” free from HER2-negative breast cancer and will likely “never” be diagnosed with breast cancer. This cohort will be more qualified as healthy control subjects, given that the current study is focused on the contribution of germline rare coding mutations rather than somatic mutations on the risk of HER2-negative breast cancer. Furthermore, germline mutations are inherited from parents, present in virtually every cell in the body throughout a person’s whole life, and are not expected to change with increasing age. To validate this point, we counted the number of germline rare coding mutations carried by the subjects. The mean number of germline rare coding mutations were 310.96 (95% CI: 309.15, 312.77) in the around 50-year-old cases and 317.58 (95% CI: 302.76, 332.41) in the around 81-year-old controls. The difference between the two means was not statistically significant (Student’s T-Test, P-value=0.08), confirming that age difference between disease cases and controls should not affect the risk assessment results.

Notably, we observed an upregulation of the HER2 signalling pathway in the germlines of HER2-negative breast cancer patients, suggesting that HER2 signalling pathway activity is an important characteristic of HER2-negative breast cancer that can distinguish it from HER2-positive breast cancer. As the germline HER2 signalling pathway was activated in a receptor-independent manner, we presented different models for the activation of the HER2 signalling pathway and its contribution to the pathogenesis of the two breast cancer subtypes (Suppl. Fig. 4). In HER2-positive breast cancer patients, the HER2 signalling pathway is activated in tumour tissue due to the amplification of the HER2 genes [40-42], but remains unchanged in the germlines. Meanwhile, in HER2-negative breast cancer patients, the HER2 signalling pathway is upregulated by the germline mutations, but there is no amplification of HER2 genes in the tumour tissue. According to a previous study, the upregulation of HER2 signalling pathway may lead to immune suppression via AKT1-mediated disruption of STING signalling [9], which supports our DAGM findings.

Although the APSP risk score from DAGM and PRS both provide disease risk assessment based on the DNA mutations in the germline genome, they are different in the following respects (Suppl. Table 3). First, DAGM uses rare coding variants (minor allele frequency, MAF<1%) from whole-exome sequencing to derive the APSP risk score, whereas PRS generally uses common variants (MAF>5%) from large genome-wide association studies (GWAS), of which nearly 90% lie within non-coding regions of the genome [43]. As GWAS results are ancestry-dependent, it is difficult for PRS to be applied across populations, whereas DAGM was validated using data from a different population (Suppl. Fig. 3). Second, DAGM projects DNA mutations and gene expression onto signalling pathway activities, whereas PRS assesses only SNPs. As a result, DAGM could reveal not only genetic mutations, but also functional signalling pathways, which are biologically relevant to disease. In the future, DAGM could be improved by adding more cell lines and signalling pathways data. Third, PRS is suitable for the risk assessment of complex diseases such as breast cancer with an AUC ranging from 0.63 to 0.69 [14,17], whereas DAGM was used to assess breast cancer risk with an AUC ranging from 0.74 to 0.79. A broader investigation of the application of DAGM in different cancers besides breast cancer is currently underway.

Overall, the DAGM framework provides an effective risk assessment for HER2-negative breast cancer using information encoded in the germline genome, which can facilitate screening of those at high risk of HER2-negative breast cancer for primary prevention. The results from DAGM also provide new insight into the pathogenesis mechanism of breast cancer.

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Author contributions

GN and YF conceived the experiments. GN, KW, MY, YF, YN, and ZYW designed the experiments. MY, XLL, ZC, MC, JX, CY, MH, FJ, LJ, WL, JL, CL, ZL, and HG performed the clinical experiments. YF, GN, QZ, JG, SH, CZ, ZZ, XJ, HC, GT, YCH, YQS, and MQZ analysed the data. GN and YF developed the methodology. YF, MY, ZF, YN, KW, and GN wrote the paper. All authors discussed the results and contributed to the final manuscript.

Data sharing statement

The raw exome sequencing data have been deposited in the Genome Sequence Archive (GSA) in National Genomics Data Centre, Beijing Institute of Genomics, Chinese Academy of Sciences, under accession number CRA000285, which is publicly accessible at https://bigd.big.ac.cn/gsa. All other data and materials are available upon request to the corresponding authors.

Declaration of Competing Interest

The authors declare that they have no competing interests.

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Supplementary materials

Supplementary material associated with this article can be found in the online version at doi:10.1016/j.ebiom.2021.103446.

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