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Gao and colleagues identify eight loci of breast cancer susceptibility in a genome-wide association study. By leveraging gene expression data from multiple tissues, they also discover 309 breast cancer susceptibility genes in a transcriptome-wide association study. Known genetic risk variants could act on these genes to cause breast cancer.





A joint transcriptome-wide association study across multiple tissues identifies candidate breast cancer susceptibility genes

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Summary

Genome-wide association studies (GWASs) have identified more than 200 genomic loci for breast cancer risk, but specific causal genes in most of these loci have not been identified. In fact, transcriptome-wide association studies (TWASs) of breast cancer performed using gene expression prediction models trained in breast tissue have yet to clearly identify most target genes. To identify candidate genes, we performed a GWAS analysis in a breast cancer dataset from UK Biobank (UKB) and combined the results with the GWAS results of the Breast Cancer Association Consortium (BCAC) by a meta-analysis. Using the summary statistics from the meta-analysis, we performed a joint TWAS analysis that combined TWAS signals from multiple tissues. We used expression prediction models trained in 11 tissues that are potentially relevant to breast cancer from the Genotype-Tissue Expression (GTEx) data. In the GWAS analysis, we identified eight loci distinct from those reported previously. In the TWAS analysis, we identified 309 genes at 108 genomic loci to be significantly associated with breast cancer at the Bonferroni threshold. Of these, 17 genes were located in eight regions that were at least 1 Mb away from published GWAS hits. The remaining TWAS-significant genes were located in 100 known genomic loci from previous GWASs of breast cancer. We found that 21 genes located in known GWAS loci remained statistically significant after conditioning on previous GWAS index variants. Our study provides insights into breast cancer genetics through mapping candidate target genes in a large proportion of known GWAS loci and discovering multiple new loci.

Introduction

Breast cancer is the most common malignancy among women in most countries around the world, accounting for one-quarter of all cancer cases in women. In the past 15 years, genome-wide association studies (GWASs) have identified over 200 loci significantly associated with breast cancer.²⁻⁴ Although some of these findings have yielded functional insights into breast cancer, 4 these genetic variants account for a relatively small proportion of heritability, suggesting that more genetic variants have yet to be identified. Because the vast majority of risk variants identified in GWASs are located in intergenic regions and are not nonsynonymous coding variants, the putative genes on which these risk variants act to cause breast cancer remain unclear for most GWAS-identified loci.

To further elucidate the role of genetic variants in complex traits, transcriptome-wide association studies (TWASs) have been conducted to quantify the relationship between a predicted level of genetically regulated gene expression and the phenotype of interest.^{5,6} TWASs of breast cancer have identified dozens of genes whose expression is significantly associated with breast cancer and its subtypes. 4,7-9 However, these genes account for only a small proportion of known GWAS loci of breast cancer. These TWASs were performed primarily by associating

a cis-regulated level of gene expression with breast cancer in single tissues (breast tissue or whole blood). Our recent study demonstrated that integrating information from multiple tissues in a TWAS could improve association detection. 10 In addition, existing TWASs used gene expression prediction models trained on data from older versions of the Genotype-Tissue Expression (GTEx) project, such as v.6 or v.7. The recent v.8 of GTEx has much larger sample sizes compared to the older versions, so the expression models trained in GTEx v.8 will be more accurate than those trained in older versions in prediction of expression levels. 11,12 By using GTEx v.8, one can explore heritability for expression more efficiently, and more genes can pass the filtering threshold and be used for TWAS analysis. 12 Therefore, the prediction models trained in GTEx v.8 have the potential to increase the power of TWAS in detecting susceptibility genes.

In this study, we aimed to identify candidate genes for breast cancer by performing joint TWAS analyses of breast cancer by combining TWAS information from multiple tissues. We applied our TWAS method to the summary statistics from a meta-analysis of data from 122,977 breast cancer cases and 105,974 controls in the Breast Cancer Association Consortium (BCAC)³ and 10,534 breast cancer cases and 185,116 controls in UK Biobank (UKB).¹³

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Subjects and methods

GWAS summary statistics on women of European ancestry from the BCAC

In our meta-analysis, we used the summary statistics data from the GWAS of breast cancer in 122,977 cases and 105,974 controls of European ancestry from the BCAC. The details of the BCAC have been described previously.^{3,14} Briefly, the BCAC included: (1) 61,282 female cases with breast cancer and 45,494 female controls of European ancestry that were genotyped using the OncoArray, including 570,000 SNPs; (2) 46,785 breast cancer cases and 42,892 controls of European ancestry from Collaborative Oncological Gene-environment Study (iCOGS) that were genotyped using a custom Illumina iSelect genotyping array containing \sim 211,155 variants; and (3) 11 other breast cancer GWASs (14,910 cases and 17,588 controls). Genotype data from iCOGS, OncoArray, and GWASs were imputed using the October 2014 release of the 1000 Genomes Project data as a reference. Genetic association results for breast cancer risk were combined using inverse-variance fixed-effect meta-analyses.³

GWAS analysis using data from UK Biobank

The UK Biobank project recruited approximately 500,000 participants, ages 40-69, between 2007 and 2010, across 22 study centers in the United Kingdom. The project collected detailed demographic, lifestyle, and disease histories at baseline, as well as disease occurrences through prospective follow-up and database linkages.¹³ Whole-genome genotyping was conducted using UK Biobank Axiom Arrays for 488,377 participants, and imputation was performed using the Haplotype Reference Consortium and 1000 Genomes phase 3 as reference panels to obtain >90 million genetic markers.¹³ In this study, we selected female individuals with both phenotypic and genotypic data available. Unrelated individuals with European ancestry were selected using principalcomponent analysis (PCA). We further filtered out samples with genotyping call rates <95%. After these exclusions, the analysis included 10,534 breast cancer cases (including 6,055 prevalent cases before enrollment and 4,479 incident cases during a median of 7 years follow-up) and 185,116 controls (Table S1). We performed GWAS analysis using logistic regression, comparing breast cancer cases with controls, adjusting for age at enrollment and top ten eigenvectors from PCA of genotypes with software package PLINK 2.0.¹⁵ As a sensitivity analysis, we performed a case-case GWAS analysis, comparing incident cases with prevalent cases. In the logistic regression models adjusting for age and top ten eigenvectors, we found that no variants were genome-wide significant (alpha = 5×10^{-8}), suggesting that it is reasonable to combine incident and prevalent cases in the primary analysis.

Gene expression prediction models

Gene expression prediction models were built with the genotype and RNA-seq data in 49 tissues of European ancestry from the GTEx project (v.8). ¹² Specifically, building prediction models for a gene includes the following steps. (1) Across all tissues, *cis*-expression quantitative trait loci (*cis*-eQTLs) were discovered with a false discovery rate of 5% per tissue. Only genes with *cis*-eQTLs were selected. (2) Fine mapping was performed in each tissue in the corresponding *cis*-gene region by the DAP-G method ^{16,17} to select variants with minor allele frequency >0.01 and posterior inclusion probabilities (PIPs) >0.01 and to select genes with at least one credible set of PIP >0.1 (where the cred-

ible-set PIP is the sum of PIPs of variants in the set). Then, in each credible set, only the variant with the highest PIP was kept. For the 49 tissues, a union of selected variants across 49 tissues was obtained, and linkage disequilibrium (LD) pruning was applied to the union of variants to remove redundant variants. (3) The multivariate adaptive shrinkage (MASH) method was applied to the marginal eQTL effects across the 49 tissues at the union of variants to jointly estimate effects of eQTLs, allowing sparse effects (that is, with many zero effects) and accounting for correlation among non-zero effects in different tissues. 18 (4) The predicted expression of the gene in each tissue was calculated as the linear combination of genotypes multiplying by their estimated effect sizes. In this study, we used the prediction models for 11 tissues, including female tissues (breast, ovary, uterus, and vagina), tissues that resemble connective and fat tissues in the breast (subcutaneous adipose, visceral adipose, and cultured fibroblasts), tissues related to immune cells (spleen, EBV-transformed lymphocytes, and whole blood), and liver. These tissues are potentially relevant to breast cancer development or carcinogen metabolism.

Summary statistic-based imputation

For variants included in the GTEx prediction models but not in the GWAS summary statistics, we imputed Z scores with the method ImpG-Summary. The ImpG-Summary method assumes that, under null hypothesis, the vector \mathbf{Z} of Z scores at all SNPs in a locus is approximately distributed as a Gaussian distribution, $\mathbf{Z} \sim N(\mathbf{0}, \mathbf{\Sigma})$, with $\mathbf{\Sigma}$ being the correlation matrix among all pairs of SNPs induced by LD. We estimated posterior mean of Z scores at unobserved SNPs. We used the GWAS summary statistics and correlation matrix estimated from the genotype data in the GTEx samples as the input for the ImpG-Summary method.

Joint TWAS across multiple tissues

The joint TWAS analysis includes two steps: (1) performing a traditional TWAS analysis in each of the 11 tissues with the software S-PrediXcan²⁰ to obtain the p values p_k (k=1,...,11) and (2) constructing test statistics using the aggregated Cauchy association test (ACAT) method²¹ that combined p values for each gene from the single-tissue TWAS analyses across the 11 tissues. Specifically, the ACAT test statistic is $T_{ACAT} = \sum_{k=1}^{11} w_k \tan{((0.5 - p_k)\pi)}$, where w_k is a nonnegative weight. We used $w_k = 1/11$. The p value of the ACAT test statistic is approximated by $\frac{1}{2}$ – (arctan T)/ π . We noticed that, for some genes, expression prediction models were only available for K of the 11 tissues (K < 11), then the ACAT test statistic was calculated using the S-PrediXcan p value p_k from the K tissues, with a weight $w_k = 1/K$.

Conditional joint TWAS

To test if the signals at the 309 genes detected by the TWAS are independent of a set of published GWAS index SNPs and newly identified index SNPs in our GWAS analysis, we performed a TWAS conditional on these index SNPs that were genome-wide significant (p < 5×10^{-8}). At each gene, we considered two sets of SNPs: the target set of SNPs used for predicting gene expression and the conditioning set of significant index SNPs from GWASs within ± 2 Mb of the transcription start or stop sites of the gene. For the target set of SNPs, we calculated adjusted effects (beta) on breast cancer risk and their variances, after conditioning on the index SNPs using the conditional and joint multiple-SNP (COJO) analysis method of Yang et al.²² We then ran

S-PrediXcan²⁰ on these conditional summary statistics in single tissues and performed the joint TWAS analysis that combines p values from the single-tissue analyses using the ACAT method.²¹

Colocalization analysis

For genes identified in the TWAS, we calculated regional colocalization probabilities (RCPs) using the method ENLOC. 17 ENLOC divides the genome into roughly independent LD blocks using the approach described in Berisa and Pickrell.²³ For a gene located in a specific LD block, ENLOC calculates the colocalization probability of causal GWAS hits and causal eQTLs in the LD block. Variant-level fine-mapping analysis was done to reveal possible causal eQTLs and GWAS hits. To calculate RCP for a gene in an LD block, we used the GTEx (v.8) eQTLs for the gene and the meta-analysis GWAS summary statistics in the LD block. Because ENLOC can calculate RCP only for single tissues, we calculated RCP in each of the 11 tissues and then calculated maximum RCP (Max RCP) among the 11 tissues. An RCP at a gene greater than a threshold (such as 0.5) provides supportive information that the gene identified by the TWAS has a high probability of colocalizing with a nearby GWAS variants, which strengthens the association signal.

Gene-based fine-mapping

We performed a gene-based statistical fine-mapping over the genetrait association signals from our TWAS using the software FOCUS (fine-mapping of causal gene sets). ²⁴ For an LD block, we identified a credible set of genes that contain the causal genes at a predefined confidence level of 90%. We also computed the marginal PIP in the target tissue (breast tissue) for each gene in the region to be causal given the observed TWAS statistics. FOCUS accounts for the correlation structure induced by LD and prediction weights used in the TWAS and controls for certain pleiotropic effects. We used 11 tissues and related expression prediction weights from the GTEx v.8 and assigned the breast tissue as the target tissue. When the expression prediction model for a gene in the breast tissue was unavailable (18% of the genes), we randomly selected an alternative tissue with a prediction model as a proxy for the gene.

Gene set enrichment and functional annotation

For the set of significant genes identified by our TWAS, we conducted enrichment of protein-coding and long non-coding RNA (lncRNA) genes against gene sets from multiple biological pathways, functional categories, and databases using the FUMA package.²⁵ Specifically, we used the GENE2FUNC module of FUMA and specified 33,527 protein-coding and lncRNA genes as the background genes for enrichment testing. Multiple testing correction was performed per data source of tested gene sets (e.g., canonical pathways, GWAS catalog categories) using Benjamini-Hochberg false discovery rate adjustment. We reported pathways/categories with adjusted p values ≤0.05 and at least two genes that overlapped with the gene set of interest.

Results

GWAS in BCAC and UKB

We performed a GWAS analysis in a breast cancer dataset from UKB (genomic control $\lambda = 1.02$) and then combined the UKB GWAS results with the previously published GWAS results of the BCAC data³ by a meta-analysis using

the software METAL²⁶ (Figure S1). We identified eight GWAS loci that were not reported by previous studies (Table 1; Figure S2). In six loci, the sentinel variants are located at least 2 Mb away from any of the risk variants identified by previous GWASs, and in two loci, the sentinel variants are located at least 500 kb away (rs9833726 and rs62483813), but none of these index variants are in LD with previous GWAS signals. Each of the index variants showed the same association direction in BCAC and UKB GWASs. No significant heterogeneity was observed in the meta-analysis at any sentinel variants. All the sentinel variants are common and are located in the introns of nearby genes.

Joint TWAS combining information across multiple

We used expression prediction models trained in 11 tissues of European ancestry (with sample sizes ranging from 129 to 670 and a median of 227) from the GTEx v.8 data using the MASH method. ^{11,12,18} In total, 19,274 genes across the 11 tissues with prediction models, including 14,613 genes expressed in breast tissue, were tested in our TWAS analysis. Using the meta-analysis summary statistics, we performed a single-tissue TWAS analysis in each tissue with S-PrediXcan²⁰ and then a joint TWAS analysis using ACAT²¹ to combine p values of single-tissue TWASs across the 11 tissues for each gene.

The results of the joint TWAS analysis are summarized in the Manhattan plots against the variant-based GWAS analysis results (Figure S1). Of the 19,274 genes tested in our joint TWAS analysis, we identified 299 genes whose predicted expression was associated with breast cancer risk at the Bonferroni-corrected threshold of p $< 2.59 \times 10^{-6}$ (Table S2). Only 141 genes were identified when TWAS analysis used only breast tissue, i.e., a conventional single-tissue TWAS approach²⁰ (Table S3). Of these 141 genes, 131 genes were also identified in the joint, multi-tissue TWAS. The remaining 10 genes identified only in the breast-tissue TWAS analysis were also marginally significant in the joint TWAS (p < 1.62×10^{-5}), so we described the 309 genes from either TWAS in further analysis. Table S4 shows the detailed single-tissue TWAS results for the 309 genes in the analysis of two databases (BCAC and UKB) pooled and separately. We found that Z scores across tissues were moderately concordant on average, with an intraclass correlation coefficient of 0.561, but the agreement between tissues as well as the strongest association signals varied across genes. These findings suggest that the multi-tissue joint TWAS could provide additional information compared to a traditional singletarget-tissue TWAS and address the possibility that the target tissue(s) could vary for different genes. We also found that TWAS results using the BCAC and UKB databases were very consistent, with a Pearson r = 0.871(Figure S3).

Of the 309 genes identified in our TWAS, 108 genes have been reported in previous TWASs (Tables S2 and S5), and

rsid	Position (hg38)	Locus	Nearest gene ^b	Alleles ^c	EAF	Data source	OR (95% CI)	p value
rs707475	7857016	1p36.23	UTS2	A/G	0.393	UKB	0.97 (0.94–1.00)	2.05E-02
						BCAC	0.97 (0.95-0.98)	1.29E-07
						Meta	0.97 (0.95-0.98)	7.17E-09
rs60504827	168127440	1q24.2	GPR161	T/C	0.126	UKB	0.96 (0.92–1.00)	4.03E-02
						BCAC	0.95 (0.93-0.97)	1.76E-07
						Meta	0.95 (0.93-0.97)	1.90E-08
rs9833726	86154659	3p12.1	LOC102723364	T/G	0.135	UKB	0.93 (0.89-0.97)	8.21E-04
			CADM2			BCAC	0.96 (0.94-0.98)	5.98E-06
						Meta	0.95 (0.94-0.97)	4.02E-08
rs35016840	150318622	4q31.3	LRBA	T/C	0.650	UKB	1.03 (1.00-1.06)	3.55E-02
						BCAC	1.03 (1.02–1.05)	3.73E-07
						Meta	1.03 (1.02–1.05)	4.25E-08
rs62483813	102478605	7q22.1	POLR2J	T/C	0.369	UKB	1.04 (1.01–1.07)	7.79E-03
						BCAC	1.04 (1.02–1.05)	1.89E-07
						Meta	1.04 (1.02–1.05)	5.34E-09
rs77457752	13942941	9p23	LINC00583	A/G	0.127	UKB	0.91 (0.87-0.96)	1.41E-04
						BCAC	0.95 (0.94-0.97)	7.95E-07
						Meta	0.95 (0.93-0.97)	1.70E-09
s3235	13907609	10p13	FRMD4A	A/G	0.656	UKB	1.04 (1.01–1.07)	8.68E-03
						BCAC	1.03 (1.02–1.05)	9.34E-07
						Meta	1.03 (1.02–1.05)	3.37E-08
s71063528	113863643	11q23.2	USP28	(A) ₁₇ /delA	0.275	UKB	0.96 (0.93-0.99)	1.29E-02
						BCAC	0.96 (0.95-0.98)	7.91E-07
						Meta	0.96 (0.95-0.98)	3.15E-08

EAF, effect allele frequency; OR, odds ratio; CI, confidence interval; GWAS, genome-wide association study; UKB, UK Biobank; BCAC, Breast Cancer Association Consortium; meta, meta-analysis.

multiple genes have been implicated in previous GWASs, such as FGFR2, TOX3, and ESR1. The 309 genes identified in our TWAS are distributed among 108 genomic loci (Figure 1). Based on NHGRI-EBI GWAS Catalog²⁷ and literature review, we curated 226 GWAS loci of breast cancer susceptibility (Table S6). Including the eight GWAS loci discovered in the current study (Table 1), there are a total of 234 GWAS breast cancer susceptibility loci (Figure 1). Our TWAS identified 292 significant genes that are located in 100 known GWAS loci. The remaining 17 genes are located in eight TWAS loci that are at least 1.4 Mb away from any risk variant identified in previous GWASs and are not in LD with risk variants (Table 2). Of the 17 genes found in the eight TWAS loci, 10 genes in six loci were significant in the breast-tissue-based TWAS at the Bonferroni threshold. For example, we found MAP2K4 in the 17p12 locus was significant in both the multi-tissue joint TWAS

and the breast-tissue-based TWAS, although there was no reported GWAS signal in this locus (Figure 2). Notably, the 1q24.2 locus was identified both in our GWAS and in our TWAS, and the GWAS index variant rs60504827 is located in an intron of *GPR161* (Figure 2).

Conditional joint TWAS on known GWAS index variants

To determine whether the associations for the genes identified by the joint TWAS were independent of GWAS association signals, we performed conditional analyses adjusting for nearby GWAS index risk variants. We found 21 genes located in 15 known GWAS loci that were conditionally significant (Table 3). This suggests that additional genetic variants, which are neither genome-wide significant nor in LD with GWAS-significant variants, may account for the association between expression of these genes and breast cancer risk at these loci.

^aThe heterogeneity tests comparing effects in BCAC and UKB were not significant in any of the loci

^bVariants are located in the introns of nearby genes

^cEffect allele/reference allele

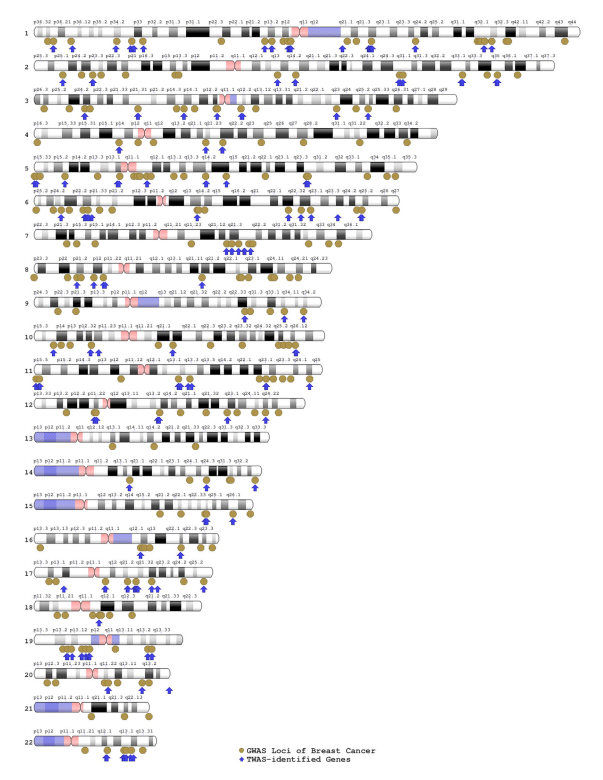


Figure 1. Ideogram of the 309 TWAS-identified genes in the context of known breast cancer GWAS loci

Colocalization analysis and gene-based fine-mapping

In colocalization analysis using ENLOC, ¹⁷ we found 45 of 309 genes with RCP values greater than 0.5 using breast tissue and 75 genes with maximum RCP values greater than 0.5 (Table S2). The colocalization RCP value for breast tissue was inversely correlated with the p value from breast-tissue TWAS (Spearman r = -0.287), suggesting that genes

with stronger TWAS significance are more likely to colocalize with GWAS causal variants.

As the LD among variants can induce significant genetrait associations for non-causal genes, we conducted gene-level fine-mapping analysis using the package FOCUS.²⁴ We found that 141 genes are in credible sets that contain causal genes at the confidence level of 90%;

Table 2. The 17 genes identified by TWAS located at 8 genomic loci at least 1 Mb away from previous GWAS hits

Locusa	Gene symbol	Position (hg38)	Joint ACAT p value	Breast p value	PIPb	In credible set	Max RCP ^c	RCP in breast
1q21.1, L1	Н3-2	chr1: 143,894,544–143,905,977	1.13E-10	9.13E-01	-	_	0.000	0.000
	FAM72C	chr1: 143,955,287-143,971,986	2.77 E-12	_	0.419	Yes	0.000	0.000
1q24.2	GPR161	chr1: 168,079,542–168,137,667	1.04 E-06	2.30E-01	0.000	No	0.513	0.000
6q24.1	TXLNB	chr6: 139,240,061–139,291,998	8.21 E-06	2.24E-06	0.761	Yes	0.681	0.681
7q22.1, L1	SPDYE3	chr7: 100,307,702–100,322,196	1.91 E-07	-	0.253	Yes	0.127	0.040
	PILRB	chr7: 100,352,176–100,367,831	2.04 E-07	1.71E-07	0.412	Yes	0.505	0.505
	PILRA	chr7: 100,367,530-100,400,096	2.69 E-07	3.47E-07	0.016	No	0.386	0.386
	ZCWPW1	chr7: 100,400,826-100,428,992	6.72 E-07	8.53E-06	0.005	No	0.261	0.202
	MEPCE	chr7: 100,428,322-100,434,126	1.74 E-07	_	0.219	Yes	0.259	0.057
	C7orf61	chr7: 100,456,620-100,464,260	7.19 E-07	2.92E-07	0.079	Yes	0.155	0.088
	TSC22D4	chr7: 100,463,359-100,479,232	1.08 E-06	1.38E-06	0.001	No	0.228	0.228
	NYAP1	chr7: 100,483,927-100,494,802	1.15 E-06	5.41E-02	0.000	No	0.086	0.000
10p12.1	YME1L1	chr10: 27,110,111-27,155,266	4.99 E-06	4.99E-07	0.920	Yes	0.249	0.084
17p12	MAP2K4	chr17: 12,020,829-12,143,830	9.59 E-07	7.28E-07	0.893	Yes	0.630	0.630
17q23.1	RPS6KB1	chr17: 59,893,046-59,950,574	3.97 E-06	1.80E-06	0.823	Yes	0.144	0.095
20q13.33	RGS19	chr20: 64,073,181-64,079,988	9.27 E-07	1.76E-06	0.166	Yes	0.922	0.862
_	OPRL1	chr20: 64,080,082-64,100,643	4.37 E-07	2.81E-07	0.792	Yes	0.925	0.791

a"L1" and "L2" denote the first and second locus defined by LD block in the same cytoband, respectively

for these genes, the median PIP was 0.792 (Table S2). As the fine-mapping analysis in FOCUS mainly used breast tissue as the target tissue (82%), genes identified in the breast-tissue TWAS were more likely in the credible sets (94 genes).

As candidate causal genes, we selected 114 genes if they are in the credible sets with gene PIPs greater than 0.15 and located in regions in which the null model is not a possible outcome (Table S7). These 114 genes are located in 83 loci. For most loci (n=61), fine-mapping identified only one causal gene candidate, eliminating many TWAS-identified genes; for example, CHEK2 (PIP = 1.0) was identified as the possible causal gene in locus 22q12.1-q12.2 (out of six TWAS-identified genes). For fewer loci, multiple candidate genes were identified after fine-mapping analysis; for example, GSTM1, GSTM2, and GSTM4, three members of the glutathione S-transferase multigene family, were suggested to be possible causal genes in locus 1p13.3 (Table S7; Figure 2).

Gene set enrichment and functional annotation

Of the 309 TWAS-identified genes, 272 are protein-coding genes, 34 are lncRNA genes, and 3 are pseudogenes. We tested the enrichment of this set of protein-coding and lncRNA genes against background gene sets from multiple databases using the FUMA software package. We found that these TWAS-identified genes were significantly enriched in several biological pathways, such as the Trail

signaling pathway, Fas signaling pathway, apoptosis pathway, biosynthesis, and cell cycle regulation; all of these pathways are important in cancer development or a hallmark of cancer, which further warrants efforts into studying how the genes identified in our TWAS may contribute to breast cancer etiology (Table S8). Interestingly, we found significant enrichment in the genes underlying several breast cancer risk factors, including body fat distribution, mammographic density, alcohol use, and body height,²⁸ suggesting that the TWAS-identified genes may indirectly contribute to breast cancer susceptibility through their impacts on known lifestyle/environmental risk factors. We also found strong enrichment for other diseases, such as inflammatory bowel disease, diabetes, and other cancers (including melanoma, chronic myeloid leukemia, and cancers of the esophagus, pancreas, bladder, and prostate), suggesting that some of breast cancer genes have pleiotropic effects. These results were consistent with the notion that there are shared genetic components between various cancer sites²⁹ and suggest that further research into collating TWAS results across cancers may be beneficial to understanding their shared genetic etiologies. Lastly, differential gene expression analysis in GTEx showed that the TWAS-identified genes had strong tissue specificity, although our joint TWAS weighted each tissue similarly; the most up-expressed tissues of these genes were uterus, ovary, vagina, and breast (Figure S4).

^bPosterior inclusion probability (PIP) calculated by FOCUS

^cMaximum marginal posterior inclusion probability (RCP) in all tissues calculated in colocalization analysis

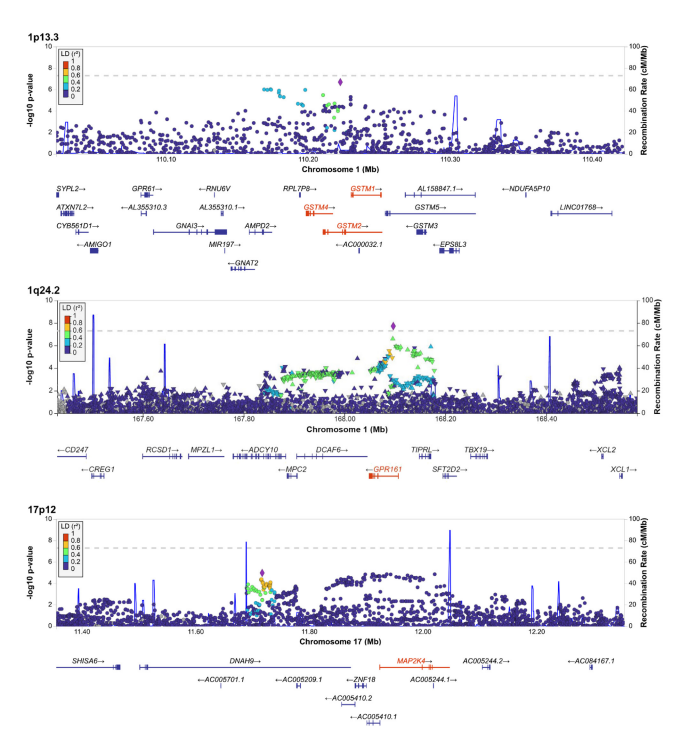


Figure 2. Manhattan plots of exemplar genes in three loci identified by the joint TWAS analysis The genes highlighted in red are significant genes identified in the joint transcriptome-wide association study.

Enrichment of genetic variants in prediction models

Fachal et al.³⁰ identified a set of 11,289 credible causal variants (CCVs) for breast cancer risk in a fine-mapping study of 150 breast cancer risk regions. We performed a hypergeometric test to examine whether genetic variants included in the expression prediction models for the 292 significant genes identified by our TWAS in 100 known GWAS loci were enriched in the set of CCVs. In 100 known GWAS loci, there were 1,160,671 genetic variants with mi-

nor allele frequency >0.01 in the GTEx v.8 data. Among these variants, 7,537 were shared with the set of CCVs identified by Fachal et al. 30 The prediction models for the 292 genes used 1,702 unique genetic variants; of these variants, 114 variants were CCVs (Table S9). Therefore, there was an approximately 10-fold over-representation of CCVs among our genetic variants of prediction models (hypergeometric p value 3.17×10^{-75}). Although this analysis cannot take into account LD among genetic variants,

Locus	Gene symbol	Position (hg38)	Joint ACAT p value	Conditional p value ^a	Closest index SNP ^b	Dist (kb) ^c	PIP ^d	In Credible set	Max RCP ^e
1p13.3	GSTM1	chr1: 109,687,814-109,709,039	2.91E-08	1.10E-06	rs5776993	7.5	0.491	Yes	0.996
2q35	DIRC3	chr2: 217,284,019–217,756,593	1.67E-16	2.31E-10	rs6436017	102.1	1.000	Yes	0.001
3p24.1, L1	NEK10	chr3: 27,106,484–27,369,460	<5.0E-17	1.77E-07	rs4973768	5.1	1.000	Yes	0.209
5p15.33, L1	AHRR	chr5: 321,714-438,291	1.22E-07	4.78E-07	rs62641919	0	0.000	No	0.379
	EXOC3	chr5: 443,175–471,937	2.14E-06	1.87E-06	rs62641919	98.2	0.317	Yes	0.065
6q22.33	RSPO3	chr6: 127,118,671–127,199,481	2.21E-06	2.57E-06	rs2180341	80	0.055	No	0.324
8q21.13	HNF4G	chr8: 75,407,914–75,566,834	1.14E-12	2.18E-07	rs72658071	14.4	1.000	Yes	0.069
11p15.5, L2	LSP1	chr11: 1,850,904–1,892,267	<5.0E-17	7.24E-07	rs576603	0.6	0.000	No	0.478
11q13.1	OVOL1	chr11: 65,787,063-65,797,214	2.08E-12	2.23E-06	rs3903072	18.4	0.982	Yes	0.165
11q13.3	RP11-554A11.8	chr11: 69,147,228-69,171,564	1.58E-08	6.02E-09	rs72932540	0	0.004	No	0.000
12p11.22, L1	CCDC91	chr12: 28,133,249-28,581,511	2.22E-16	2.90E-11	rs7297051	111.4	0.005	No	0.000
12p11.22, L2	OVCH1	chr12: 29,412,474–29,497,686	1.43E-06	1.63E-06	rs1027113	425.1	0.000	No	0.000
16q12.1-q12.2	TOX3	chr16: 52,436,417–52,547,802	<5.0E-17	<5.0E-17	rs3803662	4.6	1.000	Yes	0.333
19q13.32	GIPR	chr19: 45,668,221-45,683,722	2.31E-08	8.44E-07	rs61373376	0	0.001	No	0.353
20q11.23	PHF20	chr20: 35,771,974–35,950,370	8.79E-07	2.33E-06	rs112208395	0	0.330	Yes	0.322
	CNBD2	chr20: 35,954,564–36,030,700	2.38E-06	3.66E-06	rs112208395	21.5	0.251	Yes	0.038
22q12.1–q12.2	TTC28	chr22: 27,978,014–28,679,840	1.14E-08	9.40E-12	rs62235681	4.9	0.000	No	0.000
	CHEK2	chr22: 28,687,743–28,742,422	9.02E-13	4.39E-15	rs62235681	3.0	1.000	Yes	0.001
	HSCB	chr22: 28,742,039–28,757,515	1.36E-12	6.49E-15	rs17879961	16.9	0.000	No	0.000
	CCDC117	chr22: 28,772,674–28,789,301	1.46E-07	1.62E-09	rs17879961	47.6	0.000	No	0.000
	XBP1	chr22: 28,794,555–28,800,597	6.35E-09	1.26E-10	rs17879961	69.5	0.003	No	0.000

^aConditional ACAT p value after adjusting for adjacent index SNPs
^bSNPs that were identified to be significant in previous genome-wide association studies
^cDistance from the gene to closest index SNP (kb)
^dPosterior inclusion probability (PIP) calculated by the FOCUS method
^eMaximum marginal posterior inclusion probability (RCP) in all tissues calculated in colocalization analysis

the strong enrichment is unlikely to be due to confounders.

Fachal et al.³⁰ also classified GWAS signals as strong and moderate signals. We examined whether CCVs in regions with strong GWAS signals are more likely to be genetic variants of our prediction models, compared with CCVs in regions with moderate GWAS signals. Of the 5,117 CCVs in regions with strong GWAS signals, 83 (1.6%) were genetic variants in our gene expression prediction models. In contrast, 30 of 5,973 (0.5%) CCVs in regions with moderate GWAS signals were genetic variants in our gene expression prediction models, so there was a >3-fold enrichment (p = 4.81×10^{-9}). In short, genetic variants used in gene expression models are more likely to be variants identified from fine-mapping of breast cancer GWAS, and there is a stronger enrichment for regions with strong GWAS signals.

Discussion

In this study, we performed a breast cancer TWAS analysis that leveraged the genetically predicted gene expression levels across multiple tissues. We identified 309 significant genes at the Bonferroni threshold, including 17 genes located in eight loci not reported in previous studies and 292 genes located in 100 known GWAS loci. In about 43% of known breast cancer GWAS loci, our study was able to identify possible susceptibility gene(s). We also found 21 genes in known GWAS loci that were independent of previously reported GWAS risk variants, suggesting potentially additional breast cancer susceptibility signals. Generally, our study findings are consistent with previous TWASs; of the 368 genes reported in previous TWASs, ^{4,7–9,31–36} 108 genes were replicated in our study.

The number of TWAS-significant genes identified in our study is similar to the number identified in all previous studies combined, possibly because of several notable differences in methodologies. First, we used GWAS data from a large number of breast cancer cases (n = 133,511) and controls (n = 291,090) from BCAC and UKB. This large sample size provided high statistical power in the association analysis and helped to identify eight GWAS loci that were not reported previously. Second, we aggregated TWAS signals across 11 tissues. This multi-tissue approach resulted in more genes being identified compared to the TWAS using breast tissue alone. This suggests that while breast tissue is an important tissue to be utilized when conducting breast cancer TWASs, other tissues can contribute additional information for gene discovery. For example, our multi-tissue approach identified FGFR2, a gene with strong evidence in breast cancer etiology, but this gene has not been identified in previous TWASs and would have been missed if we had utilized only the TWAS exclusive to breast tissue. Expression of FGFR2 in fibroblasts, ovary, vagina, and liver were associated with breast cancer risk (Table S4). Third, we used expression prediction models trained in GTEx v.8 with the MASH method based

on fine-mapping that selected possible causal eQTLs as predictors for each gene. The expression models trained in GTEx v.8 can be more accurate than those trained in older versions of GTEx for three reasons: (1) The sample sizes for tissues in GTEx v.8 are larger than those in older versions of GTEx (for example, we used 329 samples from GTEx v.8 to build prediction models of breast tissue in European ancestry individuals, while Wu et al.⁸ used 67 samples from v.6); (2) selecting possibly causal eQTL through fine-mapping can reduce the probability that non-causal eQTLs were used in the prediction models¹²; and (3) MASH accounts for eQTL correlation across tissues and provides more accurate estimates of beta coefficients of eQTLs used as final weights in the prediction models. By using the prediction models trained in GTEx (v.8) data, we were able to perform this joint TWAS analysis on 19,274 genes with prediction models of good performance (i.e., with eQTL signals). In contrast, two previously published large TWASs that relied on breast tissues in older versions of GTEx and traditional methods could evaluate a smaller subset of genes. ^{7,8} Wu et al. ⁸ evaluated 8,597 genes in their TWAS and commented that several highly implicated breast cancer susceptibility genes, such as ESR1, TERT, and MRPS30, could not be investigated because of poor performance of prediction models. Our study was able to identify these three genes as significant at the Bonferroni threshold. Similarly, Feng et al. investigated 901 genes in their TWAS.

We identified 17 genes in eight loci that are at least 1 Mb away from any risk variants and not in LD with risk variants reported in previous GWASs. This finding suggests that transcriptome-based association studies are able to discover cancer susceptibility signals, extending the capacity of variant-based association studies. These genes and loci are plausibly important in breast cancer susceptibility, based on evidence from previous studies in other cancers or known cancer pathways. For example, MAP2K4 in the 17p12 locus has not been implicated in breast cancer susceptibility. We found that the predicted expression of MAP2K4 in multiple tissues, including breast tissue, was positively associated with breast cancer risk. Both colocalization analysis (RCP = 0.63) and fine-mapping analysis (PIP = 0.893 in breast tissue) suggested that MAP2K4 is a possible causal breast cancer gene, and the effect was driven by multiple weak variants. MAP2K4 (also known as MKK4) is a member of the MAPK family, which act as integration points for multiple biochemical signals and are involved in a wide variety of cellular processes such as proliferation, differentiation, transcription regulation, and development. MAP2K4 has been found to be a metastasis suppressor gene in ovarian carcinoma.³⁷ Furthermore, MAP2K4 was identified as a driver gene mutated in both early and metastatic breast cancer. 38,39 Taken together, it is possible that MAP2K4 was a breast cancer susceptibility gene in the 17p12 locus.

Most of our TWAS-identified genes are located in known GWAS susceptibility loci. Interestingly, we are able to

identify possible susceptibility genes in a large proportion of the known breast cancer GWAS loci. In these scenarios, our TWAS revealed possible target genes that GWAS-identified risk variants act on to cause breast cancer. Interestingly, genetic variants used in our expression prediction models and TWAS analysis were highly enriched in the set of CCVs identified in a previous fine-mapping study of breast cancer, ³⁰ and the enrichment was even stronger for loci with strong GWAS signals. Using eQTL analyses, Guo et al. ⁴⁰ inferred 101 target genes in known breast cancer GWAS loci. We re-discovered 51 of these 101 genes in our TWAS.

One interesting GWAS locus is 1p13.3, a region containing >20 genes within 400 kb (Figure 2). Although none of the SNPs in this locus reached GWAS significance in BCAC, this locus was recently reported to be associated with breast cancer in a cross-ancestry study. 41 Of the genes in this region, it is unclear which ones are breast cancer susceptibility genes simply based on GWAS signals. Our TWAS found that the predicted expression of GSTM1, GSTM2, and GSTM4 in multiple tissues, including breast tissue, was inversely associated with breast cancer risk. After adjusting for GWAS index SNPs in conditional analysis, the three genes were no longer significant, suggesting that GWAS risk SNPs may be responsible for the observed TWAS signals. Both colocalization analysis (RCP >0.99) and finemapping analysis suggested that all three genes are possible breast cancer candidate genes. GSTM1, GSTM2, and GSTM4 encode members of the glutathione S-transferase multigene family, which can detoxify xenobiotics, including carcinogenic compounds, and thus were proposed as cancer susceptibility genes. 42 The GSTM1 null genotype has been associated with risk of several cancers. 43-49 Therefore, there exists evidence that all three genes are possible cancer suppressors responsible for GWAS signals in 1p13.3 through carcinogen metabolism.

Another interesting example is *GPR161* at 1q24.2 (Figure 2). We discovered this locus in our meta-analysis GWAS, and its expression in adipose and fibroblasts was inversely associated with breast cancer risk in our TWAS. This gene is overexpressed in triple-negative breast cancer, promotes cell proliferation, stimulates migration and invasion, and disrupts E-cadherin localization *in vitro*. Overexpression of *GPR161* in fibroblasts has also been experimentally shown to increase cAMP signaling, ultimately resulting in decreased signaling of the Sonic hedgehog (Shh) pathway, which plays an essential role in embryonic development and tumorigenesis. Germline *GPR161* mutations have been associated with pediatric medulloblastoma. Taken together, there is some evidence that *GPR161* may play a role in breast cancer etiology.

Determining causality of TWAS-identified genes remains challenging because these genes may be associated with disease phenotypes through their correlation with disease-causal gene(s) in the same LD region. Based on genebased fine-mapping, we proposed 114 genes in 83 loci to have a high probability of being causal genes. Still, these

genes need to be investigated in future functional experiments. Guo et al. 40 used luciferase reporter assays to study functional target genes, and they found a significant difference between alternative and reference alleles in promoter activity for five genes (*DCLRE1B*, *SSBP4*, *MRPS30*, *ATG10*, and *PAX9*) but failed to show functional activity for *ARRDC3*. These findings are consistent with ours: we found that all five genes were TWAS significant, and three (*DCLRE1B*, *SSBP4*, and *PAX9*) are in our proposed list of causal genes (Table S7). Consistent with Guo et al., 40 we did not find *ARRDC3* to be TWAS significant.

The current study has several limitations. First, although the joint TWAS identified more genes than single-tissue TWAS, it may generate more false-positive hits because it utilizes other tissues that may not be causal to breast cancer risk, 54 and it uses a large number of prediction models from multiple tissues. This may increase the chance of poor/unreliable prediction models being used for downstream association analysis and may result in identification of non-causal genes. We tried to control the type I error using a stringent Bonferroni alpha level and conducted genebased fine-mapping analysis to suggest causal genes. Furthermore, the target tissue for cancer development might not be distinct, and gene expression across multiple tissues could be partially correlated. 10,11 We also observed moderate consistency between the results of single-tissue TWASs. For instance, breast tissue is presumably the target tissue for breast cancer, but gene expression in liver might better reflect carcinogen metabolism. Fortunately, the ACAT method used in our joint TWAS analysis calculates a weighted average of p values from multiple tissues and is relatively conservative in identifying significant genes. Strikingly, the top tissues in which the joint TWAS-identified genes were upregulated were all female tissues (breast, uterus, ovary, and vagina), suggesting that our joint TWAS method was able to automatically prioritize target tissues. One focus of our future methods research is to develop more efficient methods to combine TWAS signals across tissues by effectively accounting for the correlation of the signals across tissues or giving high weights to potential target tissues.

Second, the current study analyzed only data from individuals with European ancestry and focused on overall breast cancer risk. Breast cancer is a heterogeneous disease consisting of several molecular subtypes. The genetic architecture of estrogen receptor (ER)-negative breast cancer may be different from the ER-positive subtype. In the BCAC consortium, 76% of patients had ER-positive breast cancer, 3 so the current study may mainly identify genes for susceptibility of ER-positive breast cancer. Future TWASs that focus on ER-negative breast cancer or in other racial/ethnic populations are highly desirable.

Third, the current study examined only overall expression of genes but did not consider the effect of RNA splicing on disease etiology. Li et al.⁵⁵ reported that RNA splicing is another primary link between genetic variation and complex diseases. Therefore, TWASs evaluating

associations of genetically predicted splicing with breast cancer have great promise for identifying novel putative candidate disease genes. We are currently working on a splicing-based TWAS of breast cancer. Last, our study focused on *cis*-eQTL effects when constructing the expression prediction models. We did not consider *trans*-eQTL due to limited sample sizes in the GTEx data, but this is an interesting topic for future studies.

In conclusion, our joint TWAS identified more than 300 breast cancer genes for further functional investigation. Our approach has discovered susceptibility loci not reported previously and mapped out candidate genes in multiple known susceptibility loci. Future studies in diverse populations and with a focus on homogeneous phenotypes of breast cancer using innovative TWAS methodology are warranted. There is potential to map out most candidate genes in GWAS loci of breast cancer, the most common malignancy affecting women across the world.

Data and code availability

In this study, we used only existing datasets that are publicly available (see web resources). The code pipeline and results for our joint TWAS analysis are available at https://zenodo.org/record/7814694#.ZDaspXbMK5d (DOI: 10.5281/zenodo.7814694). For specific method code, we made minor modifications to S-PrediXcan to combine results with ACAT (https://github.com/shugamoe/MetaXcan/tree/catch_up). We also made minor modifications to FOCUS to accommodate PrediXcan GTEx v.8 MASHR models (https://github.com/shugamoe/focus).

Supplemental information

Supplemental information can be found online at https://doi.org/10.1016/j.ajhg.2023.04.005.

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Author contributions

G.G., H.K.I., and D.H. conceived the study and contributed to the study design. P.N.F., J.M., A.N.B., G.G., and D.H. performed statis-

tical analyses. G.G., P.N.F., J.M., A.N.B., J.L.L., O.I.O. , H.K.I, and D.H. wrote and revised the manuscript.

Declaration of interests

O.I.O. reported receiving grants from Tempus (scientific advisory board) during the study, being cofounder of CancerIQ, serving as a member of the board of directors for 54gene, and receiving grants from Color Genomics (research support) and from Roche (clinical trial support for IIT) outside the submitted work.

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Web resources

BCAC summary statistics, https://bcac.ccge.medschl.cam.ac.uk/bcacdata/oncoarray/oncoarray-and-combined-summary-result COJO (GCTA), https://yanglab.westlake.edu.cn/software/gcta/Enloc, https://github.com/xqwen/integrative FOCUS, https://github.com/bogdanlab/focus FUMA, http://fuma.ctglab.nl GTEx Portal, https://gtexportal.org/home/Metal, http://csg.sph.umich.edu/abecasis/Metal/PLINK 2.0, https://www.cog-genomics.org/plink/2.0/PrediXcan GTEx v.8 MASHR models, https://predictdb.org/S-PrediXcan, https://github.com/hakyimlab/MetaXcan and https://github.com/hakyimlab/summary-gwas-imputation UK Biobank, http://ukbiobank.ac.uk

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