ORIGINAL RESEARCH

Bioinformatic Exploration for Prognostic Significance of Sphingolipid Metabolism-Related Genes in Invasive Ductal Carcinoma Using the Cancer Genome Atlas Cohort

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Introduction: Sphingolipid metabolism is a highly controlled process that is involved in regulating bioactive lipid signaling pathways and serves important roles in several cellular processes in breast cancer. Invasive ductal carcinoma (IDC), which is characterized by the malignant proliferation of the ductal epithelium and stromal invasion, is the most common type of breast cancer. Recent advances in genetic research have accelerated the discovery of novel prognostic factors and therapeutic targets for the disease. The aim of the present study was to investigate the expression and prognostic significance of sphingolipid metabolismrelated genes in female IDC.

Methods: The present study used gene expression RNAseq data obtained from The Cancer Genome Atlas breast invasive carcinoma (TCGA BRCA) datasets.

Results: Sphingolipid metabolism-related genes exhibited dysregulated mRNA expression levels in IDC. The Student's t-test revealed that SMPDL3B, B4GALNT1, LPAR2, and LASS2 were significantly upregulated, while LASS3, LPAR1, B4GALT6, GAL3ST1, HPGD, ST8SIA1, UGT8, and S1PR1 were significantly downregulated in female IDC tissues compared with normal solid tissues. Kaplan-Meier survival analyses revealed that high SMPDL3B mRNA expression levels were associated with good prognosis in female IDC, suggesting that SMPDL3B plays a tumor suppressor role. To the best of our knowledge, the present study was the first to report that dysregulated expressions of SMPDL3B are significantly associated with age, estrogen receptor status, progesterone receptor status, and histological subtype.

Conclusion: Taken together, our study indicated that *SMPDL3B* may have a pathophysiological role and serve as a novel prognostic biomarker in IDC.

Keywords: SMPDL3B, sphingolipid metabolism, invasive ductal carcinoma, TCGA

Introduction

Breast cancer is the most common and life-threatening malignancy in females worldwide.¹ Breast carcinoma is the most prevalent malignant type and is classified as carcinoma in-situ and invasive breast cancer.² Invasive ductal carcinoma (IDC) is the most common type of invasive breast cancer, accounting for up to 80% of diagnosed breast cancer cases.³ There are clinical prognostic biomarkers for breast cancer, including size, histological grade, and estrogen receptor, progesterone receptor, and human epidermal growth factor receptor 2 status.⁴ In particular,

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molecular sub-classification systems such as receptors play an important role in clinical therapeutic strategy.⁵ However, despite the availability of therapeutic strategies for breast cancer based on molecular subtype, breast cancer is still not overcome.¹ Therefore, further studies are required to identify promising molecular biomarkers that can provide new treatment avenues.

Recent advances in genomic profiling using next generation sequencing have made it possible to identify the genetic characteristics of disease, particularly in cancer. Several large-scale cancer genome studies have been conducted, and The Cancer Genome Atlas (TCGA) is a research consortium that may be used to investigate genes in different cancer types.⁶ Moreover, TCGA may be used to investigate specific histological types of cancer, such as female IDC, by utilizing histological data and clinical parameters.

Sphingolipid metabolism is a highly-regulated intracellular process that controls the synthesis and degradation of bioactive lipids, including ceramide and sphingosine-1phosphate,⁷ which plays an important role in biological processes such as angiogenesis, ageing, cancer biology, degenerative diseases, diabetes, immune responses, and inflammation.8 Accumulating evidences revealed that dysregulated sphingolipid metabolism-related genes are implicated in human breast cancer. For example, LAG1 longevity assurance homolog 2 (LASS2) and LASS6 mRNA levels are increased in breast cancer tissues compared with matched normal tissues,⁹ and overexpressed LASS4 and LASS6 reduced cell proliferation.¹⁰ Furthermore, upregulation of sphingosine kinase 1 (SPHK1) was significantly associated with poor prognosis-¹¹ and metastasis.¹² Moreover, downregulation of 15hydroxyprostaglandin dehydrogenase (HPGD) has an unfavorable effect on the overall survival of patients with triple negative breast cancer.¹³ While the altered expression of various sphingolipid metabolism-related genes in breast cancer and their potentials as prognostic factors have been reported in the aforementioned studies, few studies have investigated sphingolipid metabolism-related genes in female IDC.

Therefore, the aim of the present study was to investigate the expression of sphingolipid metabolism-related genes in female IDC, as well as to evaluate their prognostic significance, using gene expression RNAseq data obtained from TGCA breast invasive carcinoma (TCGA BRCA) datasets.

Materials and Methods Gene Expression Datasets and Cluster

Analysis

TCGA BRCA gene expression RNAseq datasets (level 3, dataset ID: TCGA.BRCA.sampleMap/HiSeqV2) and clinical parameters (dataset IDs: TCGA.BRCA.sampleMap/ BRCA clinicalMatrix and survival/BRCA survival.txt) were downloaded from the UCSC Xena public database (https://xena.ucsc.edu). TCGA BRCA dataset consisted of 1218 samples, including 1097 primary tumor tissues, 7 metastatic tumor tissues, and 114 normal solid tissues (NST). NST were taken from normal tissues adjacent to the tumor. To analyze the RNAseq data of female IDC, female IDC datasets were sorted from TCGA BRCA using clinical parameters. The mRNA expression of the sphingolipid metabolism-related genes¹⁴ was identified from the female IDC dataset. In order to identify genes with ≥2-fold changes (2FC) in mRNA expression levels between IDC and NST, the difference between average values of the two groups was calculated, and genes with a value greater than 1 were selected. This study met the publication guidelines for using TCGA datasets (http://www.cancer.gov/aboutnci/organization/ccg/research/structrual-genomics/tcga/ using-tcga/citing-tcga). Cluster 3.0^{15} was used for cluster analysis, and samples with statistically similar gene expression were classified into groups. TreeView 1.6 (www.eisenlab.org/eisen) was used to visualize the resulting heat map. The mRNA expression levels of the heat maps were scaled (quantile normalization¹⁶ and mediancentered) within columns for visualization.

Survival Analysis

Survival data (death event and survival time) were available for 655 female IDC patients. For survival analysis, the mean gene expression value of the selected sphingolipid metabolism-related genes was used as a cutoff to divide the patients into high- and low-expression groups. Survival analysis was performed using the Kaplan–Meier method, and the log rank test was used to identify statistically significant differences between the two groups.

Statistical Analysis

Statistical analysis was performed using SPSS software (version 25.0; IBM SPSS, Armonk, NY, USA). The Kolmogorov–Smirnov test was performed to assess normality. Differences in mRNA expression levels between groups were analyzed using the Student's *t*-test. The

associations between clinicopathological parameters and the dysregulated sphingolipid metabolism-related genes were analyzed using Chi-square test or Fisher's exact test for categorical variables. Correlation analysis between inter-individual mRNA expression levels of the sphingolipid metabolism-related genes was performed using the Spearman correlation coefficient analysis for continuous variables. P<0.05 was considered to indicate a statistically significant difference.

Results

Sphingolipid Metabolism-Related Genes are Dysregulated in Female IDC Compared with NST in TCGA BRCA

The heat map revealed relative mRNA expression levels of various sphingolipid metabolism-related genes in female IDC tissues and NST (Figure 1). The Student's *t*-test revealed that a total of 36 sphingolipid metabolism-related



-7.4 0 10.4

Figure I Heat map showing the relative mRNA expression levels of the sphingolipid metabolism-related genes in female IDC tissues and NST obtained from TCGA BRCA cohort. In the data shown in matrix format, each row represents an individual gene and each column represents a single tissue. Each cell in the matrix represents the relative mRNA expression level of a gene feature in an individual tissue. The red and green in the cells reflects relatively high and low expression levels, respectively, as indicated by the scale bar. The samples are sorted into the NST group on the left and the IDC group on the right. Each cells is arranged in descending order of the mean difference between the scaled mRNA expression levels of each gene in the NST and IDC groups.

Abbreviations: TCGA BRCA, The Cancer Genome Atlas breast invasive carcinoma; IDC, invasive ductal carcinoma; NST, normal solid tissues.



Figure 2 (A) Heat map showing significantly altered mRNA expression of the sphingolipid metabolism-related genes in female IDC tissues compared with female NST obtained from TCGA BRCA cohort. In the data shown in matrix format, each row represents an individual gene and each column represents a single tissue. Each cell in the matrix represents the relative mRNA expression level of a gene feature in an individual tissue. The red and green in the cells represent relatively high and low expression levels, respectively, as indicated by the scale bar. The samples are sorted into the NST group on the left and the IDC group on the right. Each cells is arranged in descending order of the mean difference between the scaled mRNA expression levels of each gene in the NST and IDC groups. *Student's t-test, P<0.05 (NST versus IDC). (B) Relatively altered mRNA expression levels of various sphingolipid metabolism-related genes in IDC samples obtained from TCGA BRCA. **Student's t-test, P<0.001, $|2FC| \ge 1.0$. **Abbreviations:** TCGA BRCA, The Cancer Genome Atlas breast invasive carcinoma; IDC, invasive ductal carcinoma; NST, normal solid tissues.

genes were significantly altered in female IDC tissues compared with NST. The results are presented as a heat map (P<0.05; Figure 2A). When a \geq 2FC in mRNA expression was used a cutoff, sphingomyelin phosphodiesterase acid like 3B (SMPDL3B), beta-1,4-N-acetyl-galactosaminyltransferase 1 (B4GALNTI), lysophosphatidic acid receptor 2 (LPAR2), and LAG1 longevity assurance homolog 2 (LASS2) were significantly upregulated in female IDC, whereas LAG1 longevity assurance homolog 3 (LASS3), lysophosphatidic acid receptor 1 (LPAR1), beta-1,4-galactosyltransferase 6 (B4GALT6), galactose-3-O-sulfotransferase 1 (GAL3ST1), HPGD, ST8 alpha-N-acetylneuraminide alpha-2,8-sialyltransferase 1 (ST8SIA1), UDP glycosyltransferase 8 (UGT8), sphingosine-1-phosphate receptor 1 (SIPR1) were significantly downregulated $(P<0.001, |2FC| \ge 1.0;$ Figure 2B), compared with NST. These results suggested that the aforementioned 12 dysregulated sphingolipid metabolism-related genes may play a crucial role in pathophysiology of female IDC.

KM Survival Analysis Identified SMPDL3B as a Prognostic Biomarker in Female IDC

Among the altered sphingolipid metabolism-related genes in female IDC, the KM survival analysis and Log rank test demonstrated that higher *SMPDL3B* mRNA expression levels were found to be associated with favorable overall survival in female IDC (Figure 3, P=0.003).

Altered mRNA Expression Levels of Sphingolipid Metabolism-Related Genes are Associated with Clinicopathological Parameters in Female IDC

To investigate the clinicopathological implications of the dysregulated sphingolipid metabolism-related genes, Chisquare test or Fisher's exact test were performed. The clinicopathological characteristics of female IDC patients of TCGA BRCA are presented in Tables 1 and 2. The *SMPDL3B* mRNA expression level was significantly associated with age, estrogen receptor (ER) status, progesterone receptor (PR) status, and histological subtypes according to immunohistochemistry (IHC) (P=0.002, P<0.001, P<0.001, and P<0.001, respectively, Table 1). The *B4GALNT1* mRNA expression level was significantly associated with T stage, ER status, PR status, and histological subtype according to IHC (P=0.004, P<0.001, P<0.001, and P<0.001, respectively, Table 1). The LPAR2 mRNA expression level was significantly associated with T stage, ER status, PR status, and histological subtype according to IHC (P<0.001, P<0.001, P<0.001, and P<0.001, respectively, Table 1). The LASS2 mRNA expression level was significantly associated with age, ER status, PR status, and histological subtype according to IHC (P=0.003, P<0.001, P<0.001, and P<0.001, respectively, Table 1). Moreover, the GAL3ST1 mRNA expression level was significantly associated with ER status, PR status, and histological subtype according to IHC (P=0.003, P=0.011, and P<0.001, respectively, Table 2). The LASS3 mRNA expression level was significantly associated with age and histological subtype according to IHC (P=0.036 and P=0.041, respectively, Table 2). The B4GALT6 mRNA expression level was significantly associated with ER status, PR status, epidermal growth factor receptor type 2 (HER2), and histological subtype according to IHC (P=0.006, P<0.001, P=0.003, and P<0.001, respectively, Table 2). The HPGD mRNA expression level was significantly associated with age, ER status, PR status, and histological subtype according to IHC (P=0.024, P<0.001, P=0.007, and P<0.001, respectively, Table 2). The UGT8 mRNA expression level was significantly associated with age, N stage, ER status, PR status, HER2 and histological subtype according to IHC (P=0.003, P=0.029, P<0.001, P<0.001, P=0.003, and P<0.001, respectively, Table 2). The ST8SIA1 mRNA expression level was significantly associated with age, T stage, ER status, PR status, and histological subtype according to IHC (P=0.019, P=0.008, P<0.001, P<0.001, and P<0.001, respectively, Table 2). The S1PR1 mRNA expression level was significantly associated with M stage and histological subtype according to IHC (P=0.005 and P<0.001, respectively, Table 2). The LPAR1 mRNA expression level was significantly associated with T stage, ER status, PR status, and histological subtype according to IHC (P<0.001, P<0.001, P<0.001, and P<0.001, respectively, Table 1). Interestingly, we found that lower expression levels of SMPDL3B were 2.676 times more frequent in ER-positive IDC than in ER-negative (odds ratio (OR), 2.676; 95% confidence interval (CI), 1.821 to 3.932; P<0.001). Moreover, we found that lower expression levels of SMPDL3B were 1.898 times more frequent in PR-positive IDC than in PR-negative (OR, 1.898; 95% CI, 1.357 to 2.655; P<0.001).



Figure 3 Survival analysis of the dysregulated sphingolipid metabolism-related genes in IDC samples obtained from TCGA BRCA cohort. Kaplan–Meier estimates of female patients with IDC according to the relative mRNA expression values of GAL3ST1, SMPDL3B, LASS3, LASS2, B4GALT6, HPGD, UGT8, ST8SIA1, S1PR1, LPAR1, B4GALNT1, and LPAR2.

Abbreviations: TCGA BRCA, The Cancer Genome Atlas breast invasive carcinoma; IDC, invasive ductal carcinoma.

	SMPDL3B (Number)			B4GALNTI (Number)			LPAR2	(Numb	er)	LASS2 (Number)		
	Low	High	Р	Low	High	Р	Low	High	Р	Low	High	Р
Age			0.002 ^a			0.120 ^b			0.931 ^b			0.003 ^a
< 50	70	117		93	94		97	90		108	79	
≥ 50	224	234		270	208		251	227		214	264	
T stage			0.738 ^a			0.004 ^a			<0.001 ^a			0.443 ^a
ті	86	89		115	60		124	51		88	87	
T2	183	208		200	191		182	209		191	200	
ТЗ	25	26		27	24		18	33		23	28	
T4	9	15		9	15		12	12		8	16	
N stage			0.255ª			0.630 ^a			0.140 ^a			0.753 ^a
N0	137	167		172	132		154	150		153	151	
NI	107	117		123	101		124	100		102	122	
N2	47	37		41	43		48	36		40	44	
N3	12	19		16	15		11	20		15	16	
M stage			0.105 ^b			0.582 ^b			0.781 ^b			0.580 ^b
M0	292	335		344	283		329	298		305	322	
МІ	9	4		6	7		6	7		5	8	
ER status			<0.001 ^a			<0.001 ^a			<0.001 ^a			<0.001 ^a
Negative	47	114		69	92		46	115		134	27	
Positive	246	223		280	189		283	186		174	295	
PR status			<0.001 ^a			<0.001ª			<0.001 ^a			<0.001ª
Negative	81	140		91	130		73	148		152	69	
Positive	213	194		257	150		256	151		154	253	
HER2			0.054 ^a			0.028 ^b			0.155 ^b			0.446 ^b
Negative	237	284		281	240		283	238		248	273	
Positive	56	44		66	34		46	54		52	48	
Subtype according to IHC			<0.001 ^a			<0.001 ^a			<0.001 ^a			<0.001 ^a
Luminal A	145	148		182	111		195	98		104	189	
Luminal B	98	62		80	80		91	69		51	109	
HER2-enriched	29	33		41	21		20	42		40	22	
TN	27	94		38	83		29	92		108	13	

Table I	Upregulated	mRNA	Expression	Levels	of	Sphingolipid	Metabolism-Related	Genes	in	Relation	to	Clinicopathological
Parametei	rs of IDC											

Notes: Luminal A: ER+ or PR+, and HER2-; Luminal B: ER+ or PR+, and HER2+; HER2-enriched: ER-, PR-, and HER2+; TN: ER-, PR-, and HER-^aPearson's Chi-square Test. ^bFisher's Exact Test.

Abbreviations: IDC, invasive ductal carcinoma; ER, estrogen receptor; PR, progesterone receptor; HER2, epidermal growth factor receptor type 2; IHC, immunohistochemistry; TN, triple-negative.

Altered mRNA Expression Levels of Sphingolipid Metabolism-Related Genes Have Inter-Individual Correlations in Female IDC

Spearman correlation coefficient analysis was performed to explore the correlation among the significantly altered sphingolipid metabolisms-related genes in female IDC tissues. Prior to the correlation analysis, the mRNA expression levels of the 12 significantly dysregulated sphingolipid metabolism-related genes in female IDC samples were retrieved from TCGA BRCA cohort. The resulting analysis uncovered 43 significant correlations among the mRNA expression levels of specific sphingo-lipid metabolism-related genes in female IDC samples (Table 3).

Image <th< th=""><th></th><th colspan="2">GAL3ST1 (Number)</th><th colspan="3">LASS3 (Number)</th><th>B4GA</th><th>.T6 (Nur</th><th>nber)</th><th colspan="3">HPGD (Number)</th></th<>		GAL3ST1 (Number)		LASS3 (Number)			B4GA	.T6 (Nur	nber)	HPGD (Number)			
Age < 50		Low	High	Þ	Low	High	P	Low	High	Þ	Low	High	Þ
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Age			0.256 ^b			0.036 ^b			0.490 ^b			0.024 ^b
≥ 50 273205 $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ $<$ <t< td=""><td>< 50</td><td>116</td><td>71</td><td></td><td>96</td><td>91</td><td></td><td>96</td><td>91</td><td></td><td>87</td><td>100</td><td></td></t<>	< 50	116	71		96	91		96	91		87	100	
T stage T T T T T T T T T T T T T T T T T T T	≥ 50	273	205		289	189		260	218		270	208	
Ti 108 6.7 104 71 102 73 103 81 94 T2 226 163 20 12 229 162 20 191 20 10 200 101 T4 15 9 10	T stage			0.829ª			0.567 ^a			0.351 ^ª			0.119 ^a
T2 226 165 162 229 162 200 191 200 171 200 171 T4 30 21 25 26 26 26 23 23 23 29 22 24 N stage 9 124 124 124 124 124 124 124 159 159 159 159 159 159 159 159 159 159 159 155 159 155 159 155 155 155 155 152 152 159 155	ті	108	67		104	71		102	73		81	94	
T3 30 21 10 25 26 10 28 23 10	T2	226	165		229	162		200	191		220	171	
T4 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15 9 15	Т3	30	21		25	26		28	23		29	22	
Name No NO NO NO NO NO NO NO NO NO NO NO NO NO	T4	15	9		13	П		15	9		15	9	
N0 180 124 181 123 154 150 150 159 145 N1 132 92 11 160 124 102	N stage			0.869ª			0.228ª			0.195ª			0.215 ^a
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	N0	180	124		181	123		154	150		159	145	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	NI	132	92		118	106		128	96		122	102	
N3 20 I 21 10 21 10 21 10 22 9 M stage M0 M1 371 256 0.779 ^b 361 266 0.573 ^b 336 291 0.400 ^b 386 291 0.400 ^b 386 291 0.400 ^b 386 291 0.001 ^b 38 291 0.002 ^b 234 234 234 234 234 234 234 234 234 235 234 234 235 234 234 234 234 234 234 234 234 234 234 234 234 234	N2	47	37		51	33		44	40		42	42	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	N3	20	П		21	10		21	10		22	9	
M0 M1371 7256 6 266 9 361 9 266 4 336 9 291 9 336 4 291 8 336 8 291 8 <td>M stage</td> <td></td> <td></td> <td>0.779^b</td> <td></td> <td></td> <td>0.573^b</td> <td></td> <td></td> <td>0.400^b</td> <td></td> <td></td> <td>0.780^b</td>	M stage			0.779 ^b			0.573 ^b			0.400 ^b			0.780 ^b
M1 7 6 1 9 4 9 4 1 8 5 1 ER status Negative Positive 77 84 0.00 ⁵ 98 63 204 1 63 1 16 57 234 16 1 57 235 1 1 57 235 1 1 57 235 1 1 57 2 1 1 57 2 1 1 57 2 1 1 5 1	M0	371	256		361	266		336	291		336	291	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	мі	7	6		9	4		9	4		8	5	
Negative Positive778484848663728972891457235PR status Negative Positive1131080.011b121100 3.12^{b} 98123164135860.007bNegative Positive2521550.01b1211000.740b98123164135860.007bHER2 Negative Positive3022190.149b3002210.740b2692520.001b2872340.155bSubtype according to HPC Luminal A HER2-enriched 	ER status			0.003 ^b			0.356 ^b			0.006 ^b			<0.001 ^a
Positive 288 181 265 204 270 199 234 235 PR status Negative Positive 113 252 108 .011 ^b 121 100 88 123 135 86 100 243 164 135 86 167	Negative	77	84		98	63		72	89		14	57	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Positive	288	181		265	204		270	199		234	235	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	PR status			0.011 ^b			0.312 ^b			<0.001ª			0.007 ^b
Positive252155124016724316412022051.55HER2 Negative Positive302219 0.149^b 300221 0.740^b 269252 0.03^b 287234 0.155^b Negative Positive302219 0.01^a 40 0.04^b 269252 0.00^a 287234 0.155^b Subtype according to IHC Luminal A Luminal B HER2-enriched TN190103 0.01^a 158135 0.041^a 1170123 0.001^a 136 157 0.001^a MER2-enriched TN3923 0.01^a 1681351641170123 0.01^a 136157 0.01^a MER2-enriched TN96642121 0.041^a 15153032 0.01^a MER2-enriched TN9772 0.03^b 6853 0.041^a 1680 0.001^a 0.00^a 32 0.001^a MER2-enriched TN101869100Mig p Low $High$ p Low $High$ p 0.09^b 0.00^b	Negative	113	108		121	100		98	123		135	86	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Positive	252	155		240	167		243	164		202	205	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	HER2			0.149 ^b			0.740 ^b			0.003 ^b			0.155 ^b
Positive6634604068324753Subtype according to IHC Luminal A190103158135170123136157	Negative	302	219		300	221		269	252		287	234	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Positive	66	34		60	40		68	32		47	53	
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	Subtype according to IHC			<0.001 ^a			0.041 ^ª			<0.001 ^ª			<0.001 ^a
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	Luminal A	190	103		158	135		170	123		136	157	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Luminal B	96	64		106	54		82	78		87	73	
TN4972685341808932UGT8 UmberST8SI (Number)SIPRI (Number)SIPRI (Number)LPARI (Number)LowHigh p LowHigh p LowHigh p Age8998 < 50 10186939489989295T stage<	HER2-enriched	39	23		41	21		47	15		30	32	
UGT8 (Number)ST8SI/ (Number)SIPRI (Number)LPARI (Number)LowHigh p LowHigh p LowHigh p Age0.003b0.019b0.058b1 ≤ 50 10186939489989295 ≥ 50 319159287191267211200278T stage0.127a9877799654121T2239152215176222196188203	TN	49	72		68	53		41	80		89	32	
LowHigh p LowHigh p LowHigh p LowHigh p Age0.003b0.003b0.019b0.019b0.058b0.058b0.058b0.099b < 50 101869394267211200278 ≥ 50 3191590.127a1910.008a0.008a0.089a0.089a0.089aT stage121549877799654121T2239152215176222196188203		UGT8 (Number)		ST8SIA1 (Number)			SIPRI (Number)			LPARI (Number)			
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$		Low	High	Þ	Low	High	Þ	Low	High	Þ	Low	High	Þ
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Age			0.003 ^b			0.019 ^b			0.058 ^b			0.099 ^b
≥ 50 319 159 287 191 267 211 200 278 T stage 0.127 ^a 0.008 ^a 0.089 ^a <	< 50	101	86		93	94		89	98		92	95	
T stage 0.127 ^a 0.008 ^a 0.089 ^a < <0.001 ^a T I 121 54 98 77 79 96 54 121 T2 239 152 215 176 222 196 188 203	≥ 50	319	159		287	191		267	211		200	278	
TI 121 54 98 77 79 96 54 121 T2 239 152 215 176 222 196 188 203	T stage			0.127 ^a			0.008 ^a			0.089 ^a			<0.001 ^a
T2 239 152 215 176 222 196 188 203	ті	121	54		98	77		79	96		54	121	
	T2	239	152		215	176		222	196		188	203	
T3 29 22 34 17 30 21 31 20	Т3	29	22		34	17		30	21		31	20	
T4 18 6 21 3 16 8 7 17	T4	18	6		21	3		16	8		7	17	
N stage 0.029 ^a 0.809 ^a 0.287 ^a 0.724 ^a	N stage			0.029 ^a			0.809 ^a			0.287 ^a			0.724 ^a
NO 176 128 171 133 170 134 127 177	N0	176	128		171	133		170	134		127	177	

Table 2 Downregulated mRNA Expression Levels of Sphingolipid Metabolism-Related Genes in Relation to ClinicopathologicalParameters of IDC

(Continued)

	GAL3STI (Number)			LASS3 (Number)			B4GAI	.T6 (Nur	nber)	HPGD (Number)		
	Low	High	Þ	Low	High	Þ	Low	High	Þ	Low	High	Þ
NI	152	72		128	96		116	108		102	122	
N2	61	23		50	34		42	42		40	44	
N3	19	12		20	П		21	10		13	18	
M stage			1.000 ^b			0.170 ^b			0.005 ^b			0.574 ^b
M0	397	230		357	270		336	291		272	355	
мі	8	5		10	3		12	I		7	6	
ER status			<0.001 ^a			<0.001 ^a			0.067 ^b			<0.001ª
Negative	41	120		28	133		95	66		101	60	
Positive	358	111		331	138		236	233		175	294	
PR status			<0.001 ^a			<0.001 ^a			0.211 ^b			<0.001 ^a
Negative	84	137		75	146		124	97		131	90	
Positive	315	92		284	123		207	200		144	263	
HER2			0.003 ^b			0.187 ^b			0.744 ^b			0.443 ^b
Negative	316	205		290	231		282	239		223	298	
Positive	76	24		63	37		56	44		47	53	
Subtype according to IHC			<0.001 ^a			<0.001 ^a			<0.001 ^a			<0.001 ^a
Luminal A	229	64		207	86		121	172		82	211	
Luminal B	122	38		121	39		108	52		82	78	
HER2-enriched	43	19		33	29		36	26		32	30	
TN	9	122		7	114		79	42		84	37	

Table 2 (Continued).

Notes: Luminal A: ER+ or PR+, and HER2-; Luminal B: ER+ or PR+, and HER2+; HER2-enriched: ER-, PR-, and HER2+; TN: ER-, PR-, and HER-^aPearson's Chi-square Test. ^bFisher's Exact Test.

Abbreviations: IDC, invasive ductal carcinoma; ER, estrogen receptor; PR, progesterone receptor; HER2, epidermal growth factor receptor type 2; IHC, immunohis-tochemistry; TN, triple-negative.

Discussion

Sphingolipid metabolism regulates diverse biological processes, including breast cancer pathophysiology.¹⁷ IDC of the breast cancer, arises from epithelial cells in the inner lining of the milk ducts, and is the most common type of invasive breast cancer that microscopically penetrates through the epithelium of the ducts into the stroma in breast tissue.^{2,3} As the importance of precision medicine becomes apparent, the number of studies using genomic profiles to identify novel prognostic and therapeutic targets is steadily increasing.^{18,19} Although methods for precisely targeting or imaging such as peptide-based SPECT radiopharmaceuticals have been studied using the discovered target for diagnosis or treatment of IDC,²⁰⁻²² its prognosis is still unfavorable. In the present study, gene expression RNAseq data from TCGA BRCA dataset were used to evaluate whether sphingolipid metabolism-related genes are significantly dysregulated, and whether they may serve as potential prognostic indicators in female IDC.

The present study investigated altered genes with $\geq 2FC$ in mRNA expression levels in IDC compared with NST. SMPDL3B, B4GALNT1, LPAR2, and LASS2 were significantly upregulated and the other genes (LASS3, LPAR1, B4GALT6, GAL3ST1, HPGD, ST8SIA1, UGT8, and S1PR1) were significantly downregulated (Figure 2). Among the dysregulated sphingolipid metabolism-related genes, only SMPDL3B had prognostic significance in female IDC (Figure 3). SMPDL3B is one of the lipid raft enzymes that regulates lipid composition and fluidity in the plasma membrane of macrophages.²³ Interestingly, SMPDL3B modulates podocyte migration and apoptosis, and activates integrin in podocytes.²⁴ Moreover SMPDL3B inversely regulates ceramide-1-phosphate (C1P) levels by interacting with ceramide kinase (CERK) in human podocytes.²⁵ In addition, it has been reported that C1P is involved in enhancement of cancer cell growth. migration, and survival.²⁶ However, CERK was downregulated in IDC (Figure 1), suggesting SMPDL3B and C1P

Samples	Correlations Between	Spearman Correlation	P-value*	
	Components	Coefficient Value		
IDC tissues from TCGA BRCA cohort (n = 665)	GAL3ST1 and SMPDL3B	0.108	0.006	
	GAL3ST1 and LASS3	0.078	0.046	
	GAL3ST1 and LASS2	-0.116	0.003	
	GAL3ST1 and HPGD	-0.126	0.001	
	GAL3ST1 and ST8SIA1	0.143	<0.001	
	GAL3ST1 and S1PR1	-0.083	0.032	
	GAL3ST1 and LPAR2	0.109	0.005	
	SMPDL3B and LASS2	-0.081	0.036	
	SMPDL3B and B4GALT6	-0.099	0.011	
	SMPDL3B and HPGD	-0.172	<0.001	
	SMPDL3B and UGT8	0.175	<0.001	
	SMPDL3B and ST8SIA1	0.145	<0.001	
	SMPDL3B and SIPRI	-0.128	0.001	
	SMPDL3B and LPARI	-0.133	0.001	
	SMPDL3B and LPAR2	0.324	<0.001	
	LASS3 and LASS2	-0.081	0.037	
	LASS3 and B4GALT6	0.101	0.009	
	LASS3 and UGT8	0.168	<0.001	
	LASS3 and SIPRI	0.099	0.010	
	LASS3 and LPAR1	-0.092	0.018	
	LASS2 and UGT8	-0.359	<0.001	
	LASS2 and ST8SIA1	-0.396	<0.001	
	LASS2 and B4GALNT1	-0.091	0.019	
	LASS2 and LPAR2	-0.127	0.001	
	B4GALT6 and UGT8	0.223	<0.001	
	B4GALT6 and ST8SIA1	0.086	0.026	
	B4GALT6 and SIPRI	-0.083	0.032	
	B4GALT6 and B4GALNTI	0.274	<0.001	
	B4GALT6 and LPAR2	-0.141	<0.001	
	HPGD and UGT8	-0.082	0.034	
	HPGD and SIPRI	0.241	<0.001	
	HPGD and LPAR1	0.177	<0.001	
	HPGD and LPAR2	-0.192	<0.001	
	UGT8 and ST8SIA1	0.456	<0.001	
	UGT8 and LPAR1	-0.081	0.036	
	UGT8 and B4GALNT1	0.178	<0.001	
	UGT8 and LPAR2	0.163	<0.001	
	ST8SIA1 and B4GALNT1	0.150	<0001	
	ST8SIA1 and LPAR2	0.105	0.007	
	SIPRI and LPARI	0.204	<0.001	
	SIPRI and B4GALNTI	-0.235	<0.001	
	SIPRI and LPAR2	-0.154	<0.001	
	LPAR1 and LPAR2	-0.287	<0.001	

Table 3 Spearman Correlation Analysis Between Inter-Individual Components of Sphingolipid Metabolism-Related Genes

Note: *Spearman correlation coefficient analysis.

are regulated differently in IDC and podocytes. Notably, a recent report demonstrated that increased *SMPDL3B* was associated improved prognosis in prostate cancer.²⁷ The present study demonstrated that increased mRNA

expression levels of *SMPDL3B* is significantly associated with improved prognosis (Figure 3). These results suggested that *SMPDL3B* may function as a tumor suppressor gene.

In the present study, we found that SMPDL3B and LPAR2 were upregulated (Figure 2B), and a significant positive correlation was identified between the genes (Table 3). Additionally, higher LPAR2 mRNA expression levels tended to show favorable overall survival in female IDC (Figure 3). However, LPAR2 showed aggressive characteristics of gastric cancer cells, such as inducing cell migration²⁸ and LPAR2-KO had significantly suppressed the invasion of ovarian cancer cells.²⁹ Although more studies on the mechanism of each gene in IDC are needed, the aforementioned results suggested that SMPDL3B and LPAR2 have different regulatory mechanisms and functions. On the other hand, the downregulated sphingolipid metabolism-related genes such as HPGD, S1PR1, and LPAR1, which are inversely correlated with the SMPDL3B, have been identified (Table 3). Additionally, higher HPGD mRNA expression levels tended to indicate a favorable prognosis (Figure 3). However, it was previously shown that HPGD was significantly upregulated and higher HPGD is significantly associated with a poor overall survival and the increased risk of disease relapse in breast cancer.³⁰ In order to determine the dysregulation and function of HPGD in breast cancer, it is necessary to analyze it by breast cancer subtype.

Recent studies reported significant correlations among sphingolipid metabolism-related genes, such as ceramide synthases. For example, significant correlations were determined between *LASS2* and *LASS4/LASS6*, and between *LASS4* and *LASS6* in breast cancer.⁹ Moreover, significant correlations between *LASS2* and *LASS4*, and between *LASS5* and *LASS4/LASS6* were identified in various colorectal cancer cohorts.³¹ Interestingly, in the present study, only the correlation between *LASS2* and *LASS3* was significant in female IDC (Table 3).

Although further studies are required to elucidate the underlying mechanisms involved in the inter-individual correlation analysis in female IDC, our results suggested that networks of sphingolipid metabolism-related genes may be implicated in the pathophysiology of the disease.

Conclusions

To the best of our knowledge, the present study was first to identify the significantly dysregulated sphingolipid metabolism-related gene, *SMPDL3B* as a novel prognostic biomarker for female IDC using TCGA BRCA datasets. Our results suggested that *SMPDL3B* plays an important role in the pathophysiology of female IDC, and may serve as a

potential prognostic biomarker as well as promising therapeutic target for the disease.

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Disclosure

The authors report no conflicts of interest in this work.

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