

Polymorphisms in lncRNA *MIR2052HG* and susceptibility to breast cancer in Chinese population

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ABSTRACT

Background: Published studies based on pharmacokinetics have explored the relationship between the lncRNA *MIR2052HG* and the prognosis of breast cancer (BC) resistance and recurrence. However, the underlying association of *MIR2052HG* SNPs with BC development remains unclear.

Methods: Combining bioinformatics and databases, SNPs (Single Nucleotide Polymorphisms) in the *MIR2052HG* gene were screened, and SNPs in the lncRNA *MIR2052HG* were selected for genotyping among 504 Chinese Han patients and 505 healthy controls, which were frequency-matched for age (± 2 years). Logistic regression analysis was used to explore the association between *MIR2052HG* SNPs and the BC risk. Interactions between the *MIR2052HG* SNPs and reproductive factors were further evaluated using the multifactor dimensionality reduction (MDR) method. qRT-PCR was performed to detect *MIR2052HG* expression in individuals with different genotypes of rs34841297. The target miRNA, miR-4456 of *MIR2052HG* rs34841297 was predicted by websites and confirmed by performing dual luciferase gene reporter assays. CCK-8 and Transwell experiments were designed to explore the effects of miR-4456 on the proliferation, invasion and migration of BC cells.

Results: In this study, nine SNPs were screened. After adjusting for age, menarche age, menopausal status, number of pregnancies, history of abortions, breast feeding history and family history of BC, the results of the logistic regression analysis showed the rs34841297 A/- gene polymorphism was positively correlated with the incidence of BC. Compared with the AA genotype, patients with the A+/- genotype of rs34841297 at age<50 years, and menarche age<14 years, Premenopausal status, history of abortion, no history of breastfeeding and no family history of tumors in first-degree relatives had an increased risk of BC. MDR results revealed that individuals with rs34841297 - (homozygous deletion) of the A allele who were not menopausal and had no history of breastfeeding had a higher risk of BC. qRT-PCR results revealed that homozygous deletion (1.68 \pm 1.37) of the rs34841297 A- genotype resulted in higher *MIR2052HG* expression than the heterozygous deletion genotype (0.95 \pm 0.94) and wild AA genotype (0.26 \pm 0.12). Binding between *MIR2052HG* and miR-4456 was occurred when rs34841297 carried the AA genotype. Moreover, preliminary functional studies indicated that the overexpression of miR-4456 increased the proliferation, invasion and migration of BC cells.

Conclusion: Our study showed that the *MIR2052HG* gene polymorphism may be related to BC susceptibility, and the *MIR2052HG* rs34841297 A/- genotype may probably affect the proliferation, invasion and migration of BC cells by modulating the interactions with of miR-4456.

INTRODUCTION

According to the latest global cancer burden data released by the International Agency for Research on Cancer (IARC) of the World Health Organization in 2020 [1], the incidence of breast cancer (BC) in women exceeds that of lung cancer, accounting for 11.7% of all new tumors and becoming the most common cancer in the worldwide. Approximately 685,000 deaths from BC were reported, making it the fifth leading cause of cancer-related death in the world [1], 90% of which were caused by distant metastasis of primary tumor cells [2, 3]. Among women, the BC incidence and mortality rank first among 159 countries in the world, and the relative increase in cancer risk was the largest in low or medium countries/regions (95% increase and 64% increase from 2020, respectively). Although the mortality rate of BC had decreased in 1991-2017 [4], in the past ten years (2008-2017), the rate of decrease in mortality of females with BC has gradually slowed. The aforementioned survey data showed that the previous preventive or treatment measures were not effective, and the high morbidity and mortality rates have become a public health problem that seriously threatens women's health. Therefore, effective and cost-effective early detection, early diagnosis and individualized treatment of BC have become urgent problems that remain to be solved worldwide.

Current studies have reported many reproductive and environmental factors related to BC [5–7]. Genetic factors are also indispensable for increasing the risk of BC. *BRCA1* and *BRCA2* are currently recognized BC susceptibility genes and are widely measured as predictors of BC risk [8–10]. Recently, with the development of high-throughput sequencing technology, noncoding RNAs have been extensively studied, and lncRNAs, new BC biomarkers, are involved in some biological processes, such as cell proliferation, cell cycle, apoptosis, pluripotency differentiation and maintenance [11], resulting in the occurrence and development of some cancers including BC, liver cancer, and lung cancer, by promoting tumor proliferation, invasion, and metastasis [12–15].

The lncRNA *MIR2052HG*, also known as *FLJ39080* and *LOC441355*, is a long non-coding RNA located on chromosome 8. James N et al. [16] found that the risk of BC recurrence in individuals with homozygous mutant and heterozygous genotypes of the lncRNA *MIR2052HG*

was lower than that in the wild type homozygous individuals. In addition, *MIR2052HG* overexpression increases BC cell proliferation and promotes colony formation. Pharmacogenomics studies [17] have shown that as a functional polymorphic gene, *MIR2052HG* might affect the risk of BC recurrence in women treated with aromatase inhibitors. Single Nucleotide Polymorphisms (SNPs) are polymorphisms in DNA sequences caused by variations in single nucleotides at the genome level. According to genome-wide association studies (GWAS) [18], SNPs in lncRNAs are related to susceptibility to many diseases, and SNPs at the key regulatory position of lncRNAs may substantially disrupt their function. Wang L et al. [19] also found that the SNP rs3802201 in *MIR2052HG* is closely related to the recurrence. However, this study was mainly based on pharmacogenomics to explore the relationship between the lncRNA *MIR2052HG* and BC resistance and recurrence. Researchers have not clearly determined whether there is an association between the genetic variants of *MIR2052HG* and BC susceptibility exists.

Therefore, relying on the Han population in Henan, this project screened SNPs in the lncRNA *MIR2052HG* that affect the occurrence of BC and studied the possible molecular mechanism to discover new risk markers for BC. The results might facilitate the early identification and diagnosis of BC in high-risk populations to achieve the purpose of early prevention of BC.

RESULTS

Basic characteristics of study subjects

Based on a case-control study, the basic information of 504 patients and 505 healthy controls was presented in Table 1. The results showed that the age at menarche of the case group (14.21 ± 1.70) was higher than that of the control group (13.97 ± 1.75) ($P=0.030$), and ≥ 2 pregnancies ($OR:2.049$, 95% $CI: 1.592-2.637$, $P<0.001$) and the family history of BC ($OR:1.869$ 95% $CI: 1.116-3.130$, $P=0.017$) were risk factors for BC. However, a breastfeeding history ($OR:0.724$, 95% $CI: 0.535-0.980$, $P=0.037$) protected against BC.

Association of *MIR2052HG* SNPs with BC susceptibility

Different genotype models of nine *MIR2052HG* SNPs with BC susceptibility were presented in Table 2. The

Table 1. Basic characteristics of 504 breast cancer cases and 505 healthy controls.

Variable	Case (%)	Control (%)	<i>P</i> ^b
	n=504	n=505	
Age ($\bar{x} \pm s$)	48.00 \pm 9.85	48.15 \pm 9.61	0.806 ^a
Age at menarche ($\bar{x} \pm s$)	14.21 \pm 1.70	13.97 \pm 1.75	0.030 ^a
Age at menopause ($\bar{x} \pm s$)	48.60 \pm 3.90	48.72 \pm 3.70	0.760 ^a
Menopause statue			
Un-menopause	320 (63.5)	294 (58.2)	
Post-menopause	184 (36.5)	211(41.8)	0.086
Number of pregnancies			
<2	53(10.5)	94(18.6)	
\geq 2	451 (89.5)	411 (81.4)	<0.001
Number of abortions			
<2	340 (67.5)	345(68.3)	
\geq 2	164 (32.5)	160 (31.7)	0.771
Breast-feeding history			
No	121 (24.0)	94 (18.6)	
Yes	383 (76.0)	411(81.4)	0.037
Family history			
No	461 (91.5)	481(95.2)	
Yes	43 (8.5)	24 (4.8)	0.017
ER receptor			
Negative	149(30.3)		
Positive	342(69.7)		
PR receptor			
Negative	191(39.1)		
Positive	298(60.9)		
HER-2 receptor			
Negative	136(29.3)		
Positive	328(70.7)		

^at test.^b χ^2 test, Two-sided *P*<0.05 is statistically significant.

MIR2052HG rs3802201 CG+GG genotype, rs2553716 AC genotype and AC+CC genotype, rs2588297 GT genotype and GT+TT type, rs4259395 AG+GG genotype, rs10957736 TT genotype and CT+TT genotype and rs12546233 AC+CC genotype might reduce the risk of BC, while the rs34841297 deletion mutation increases the risk of breast cancer. Compared with the wild homozygous AA genotype, the SNP rs34841297 --(homozygous deletion) genotype (*OR*: 1.936, 95% *CI*: 1.208-3.123) and A+--genotype (*OR*: 1.704, 95% *CI*: 1.087-2.672) resulted in a higher risk in BC.

Stratified analysis of the association between *MIR2052HG* SNPs and breast cancer susceptibility

A stratified analysis was conducted to further explore the relationship between the nine SNPs in the

MIR2052HG gene and BC susceptibility. As shown in Table 3, compared with the rs34841297 AA genotype, the A+-- genotype of rs34841297 in patients age<50 years (*OR*: 1.998, 95% *CI*: 1.072-3.723) with an age at menarche<14 years (*OR*: 2.823, 95% *CI*: 1.316-6.005) who were non-menopausal (*OR*: 2.490, 95% *CI*: 1.288-4.815), had a history of abortion (*OR*: 2.045, 95% *CI*: 1.117-3.744), no history of breastfeeding (*OR*: 3.290, 95% *CI*: 1.183-3.070) and had family history of tumors in first-degree relatives (*OR*: 1.905, 95% *CI*: 1.183-3.070) had an increased risk of BC. However, the CG+GG genotype of rs3802201, the AC+CC genotype of rs2553716, the AG+GG genotype of rs4259395, the rs2588297 GT+TT genotype, the CT+TT genotype of rs10957736 and the AC+CC genotype of rs12546233 were all found to exert a protective effect on the BC risk.

Table 2. Association between nine SNPs and breast cancer susceptibility.

SNPs	Genetic model	Genotype	Case (n=504, %)	Control (n=505, %)	<i>P</i> ^a	Adjusted OR (95%CI)	<i>P</i> ^b	
rs3802201	Codominant	CC	257(51)	232(45.9)	0.0545	1	0.071	
		CG	218(43.3)	231(45.7)		0.776(0.590,1.022)		
		GG	29(5.8)	42(8.3)		0.642(0.375,1.098)		0.105
	Dominant	CC	257(51.0)	232(45.9)		1		
		CG+GG	247(49.0)	273(54.1)		0.756(0.580,0.986)		0.039
		CC+CG	475(94.2)	463(91.7)		1		
	Recessive	GG	29(5.8)	42(8.3)		0.725(0.431,1.219)		0.225
		CC+GG	286(56.7)	274(54.3)		1		
		CG	218(43.3)	231(45.7)		0.822(0.630,1.073)		0.148
rs2553716	Codominant	AA	261(51.8)	231(45.7)	0.1535	1	0.031	
		AC	211(41.9)	232(45.9)		0.739(0.560,0.973)		
		CC	32(6.3)	42(8.3)		0.728(0.432,1.228)		0.234
	Dominant	AA	261(51.8)	231(45.7)		1		
		AC+CC	243(48.2)	274(54.3)		0.737(0.565,0.931)		0.024
		AA+AC	472(93.7)	463(91.7)		1		
	Recessive	CC	32(6.3)	42(8.3)		0.841(0.507,1.394)		0.501
		AA+CC	293(58.1)	273(54.1)		1		
		AC	211(41.9)	232(45.9)		0.770(0.590,1.006)		0.055
rs4259395	Codominant	AA	186(36.9)	163(32.3)	0.041	1	0.086	
		AG	253(50.2)	264(52.3)		0.775(0.579,1.073)		
		GG	65(12.9)	78(15.4)		0.693(0.457,1.453)		0.086
	Dominant	AA	186(36.9)	163(32.3)		1		
		AG+GG	318(63.1)	542(67.7)		0.756(0.573,0.999)		0.049
		AA+AG	439(87.1)	427(84.6)		1		
	Recessive	GG	65(12.9)	78(15.4)		0.807(0.552,1.179)		0.268
		AA+GG	251(49.8)	241(47.7)		1		
		AG	253(50.2)	264(52.3)		0.862(0.662,1.123)		0.272
rs2588297	Codominant	GG	343(68.1)	291(57.6)	0.3605	1	0.000	
		GT	147(29.2)	195(38.6)		0.597(0.448,0.794)		
		TT	14(2.8)	19(3.8)		0.799(0.335,1.463)		0.343
	Dominant	GG	343(68.1)	291(57.6)		1		
		GT+TT	161(31.9)	214(42.4)		0.606(0.459,0.798)		0.000
		GG+GT	490(97.2)	486(96.2)		1		
	Recessive	TT	14(2.8)	19(3.8)		0.839(0.404,1.742)		0.637
		GG+TT	357(70.8)	310(61.4)		1		
		GT	147(29.2)	195(38.6)		0.608(0.458,0.807)		0.001
rs10957736	Codominant	CC	242(48.0)	211(41.8)	0.2226	1	0.124	
		CT	224(44.4)	235(46.5)		0.804(0.608,1.062)		
		TT	38(7.5)	59(11.7)		0.562(0.348,0.907)		0.018
	Dominant	CC	242(48.0)	211(41.8)		1		
		CT+TT	262(52.0)	294(58.2)		0.756(0.579,0.986)		0.039
		CC+CT	466(92.5)	446(88.3)		1		
rs10957736	Recessive	TT	38(7.5)	59(11.7)	0.627(0.397,0.991)	0.046		
		CC+TT	280(55.6)	270(53.5)	1			
		CT	224(44.4)	235(46.5)	0.889(0.682,1.160)	0.387		
	Codominant	TT	394(78.2)	396(78.4)	1			
		CT	106(21.0)	103(20.4)	1.009(0.727,1.399)	0.959		
		CC	4(0.8)	6(1.2)	0.580(0.151,2.236)	0.429		
rs269183	Dominant	TT	394(78.2)	396(78.4)	1			
		CT+CC	110(21.8)	109(21.6)	0.983(0.713,1.355)	0.917		
		TT+CT	500(99.2)	499(98.8)	1			

rs269198	Over-Dominant	CC	4(0.8)	6(1.2)		0.579(0.151,2.229)	0.427
		TT+CC	398(79.0)	402(79.6)		1	
		CT	106(21.0)	103(20.4)		1.015(0.733,1.408)	0.927
	Codominant	CC	380(75.4)	384(76.0)	0.733	1	
		CA	117(23.2)	114(22.6)		1.036(0.756,1.459)	0.826
	Dominant	AA	7(1.4)	7(1.4)		0.934(0.309,2.822)	0.903
		CC	380(75.4)	384(76.0)		1	
	Recessive	CA+AA	124(24.6)	121(24.0)		1.029(0.757,1.400)	0.853
		CC+CA	497(98.6)	498(98.6)		1	
	rs34841297	Over-Dominant	AA	7(1.4)	7(1.4)		0.926(0.307,2.793)
CC+AA			387(76.8)	391(77.4)		1	
CA			117(23.2)	114(22.6)		1.037(0.758,1.420)	0.820
Codominant		AA	39(7.7)	63(12.5)	0.3673	1	
		A-	223(44.2)	236(46.7)		1.502(0.938,2.405)	0.091
Dominant		--	242(48.0)	206(40.8)		1.936(1.208,3.123)	0.006
		AA	39(7.7)	63(12.5)		1	
Recessive		A+--	465(92.3)	442(87.5)		1.704(1.087,2.672)	0.020
		AA+A-	262(52.0)	299(59.2)		1	
rs12546233		Over-Dominant	--	242(48.0)	206(40.8)		1.388(1.062,1.812)
	AA+--		281(55.8)	269(53.3)		1	
	Codominant	A-	223(44.2)	236(46.7)		0.875(0.671,1.142)	0.326
		AA	276(54.8)	242(47.9)	0.6976	1	
	Dominant	AC	195(38.7)	220(43.6)		0.764(0.578,1.006)	0.055
		CC	33(6.5)	43(8.5)		0.664(0.394,1.119)	0.124
	Recessive	AA	276(54.8)	242(47.9)		1	
		AC+CC	228(45.2)	263(52.1)		0.747(0.573,0.973)	0.031
	Over-Dominant	AA+AC	471(93.5)	462(91.5)		1	
		CC	33(6.5)	43(8.5)		0.749(0.451,1.244)	0.264
		AA+CC	309(61.3)	285(56.4)		1	

^a*P* value of Hardy-Weinberg equilibrium in controls.

^b*P* value of logistic regression analysis with adjusted for age, age at menarche, menopausal status, number of pregnancies and abortions, breast-feeding status, and family history of BC in first degree relative.

Table 3. Stratification analysis of the nine SNPs and BC susceptibility.

	rs3802201(CG+GG/CC) OR(95%CI)	<i>P</i> ^a	rs2553716(AC+CC/AA) OR(95%CI)	<i>P</i> ^a	rs4259395(AG+GG/AA) OR(95%CI)	<i>P</i> ^a
age						
<50	0.845(0.599,1.193)	0.339	0.822(0.583,1.160)	0.266	0.908(0.638,1.293)	0.749
≥50	0.718(0.466,1.106)	0.133	0.704(0.457,1.085)	0.112	0.622(0.395,0.979)	0.040
Age at menarche						
<14	0.817(0.532,1.253)	0.354	0.844(0.550,1.294)	0.436	0.813(0.515,1.285)	0.376
≥14	0.736(0.523,1.035)	0.078	0.686(0.488,0.965)	0.031	0.752(0.527,1.072)	0.115
Menopausal state						
Pre-menopausal	0.770(0.542,1.092)	0.142	0.750(0.528,1.064)	0.106	0.822(0.570,1.185)	0.293
Post-menopausal	0.760(0.499,1.158)	0.202	0.749(0.491,1.141)	0.178	0.690(0.930,2.257)	0.101
Age at menopause						
<45	0.945(0.224,3.987)	0.938	0.945(0.224,3.987)	0.938	0.784(0.172,3.571)	0.753
≥45	0.725(0.463,1.135)	0.160	0.709(0.453,1.110)	0.133	0.678(0.424,1.085)	0.105
No. of pregnancies						
<3	0.854(0.572,1.273)	0.438	0.861(0.577,1.283)	0.462	0.882(0.583,1.333)	0.551
≥3	0.759(0.536,1.074)	0.120	0.702(0.495,0.994)	0.046	0.688(0.473,1.003)	0.052
History of abortion						
no	0.667(0.429,1.036)	0.071	0.665(0.429,1.033)	0.069	0.715(0.448,1.142)	0.160

yes	0.834(0.598,1.164)	0.286	0.789(0.572,1.113)	0.183	0.721(0.507,1.023)	0.067
Breast-feeding						
no	1.773(0.971,3.237)	0.062	1.959(1.072,3.581)	0.029	0.670(0.357,1.260)	0.214
yes	1.230(0.913,1.657)	0.174	1.235(0.917,1.664)	0.165	0.779(0.570,1.064)	0.116
Family history						
no	0.717(0.544,0.944)	0.018	0.691(0.524,0.910)	0.009	0.738(0.554,0.985)	0.039
yes	1.594(0.542,4.691)	0.397	1.838(0.615,5.495)	0.276	0.965(0.291,3.199)	0.953
	rs2588297(GT+TT/GG)	<i>P</i>^a	rs10957736(CT+TT/CC)	<i>P</i>^a	rs269183(CT+CC/TT)	<i>P</i>^a
	OR(95%CI)		OR(95%CI)		OR(95%CI)	
age						
<50	0.671(0.468,0.963)	0.030	0.934(0.658,1.326)	0.703	1.098(0.730,1.649)	0.654
≥50	0.583(0.373,0.911)	0.018	0.605(0.392,0.932)	0.023	0.795(0.453,1.396)	0.425
Age at menarche						
<14	0.651(0.417,1.019)	0.060	0.915(0.592,1.414)	0.688	0.798(0.481,1.323)	0.381
≥14	0.580(0.407,0.828)	0.003	0.704(0.500,0.990)	0.043	1.203(0.785,1.845)	0.396
Menopausal state						
Pre-menopausal	0.583(0.403,0.844)	0.004	0.840(0.589,1.197)	0.334	1.068(0.707,1.614)	0.755
Post-menopausal	0.679(0.439,1.048)	0.080	0.700(0.459,1.066)	0.097	0.860(0.496,1.492)	0.592
Age at menopause						
<45	1.787(0.416,7.670)	0.435	1.667(0.404,6.880)	0.480	1.226(0.194,7.738)	0.828
≥45	0.599(0.376,0.953)	0.031	0.633(0.404,0.992)	0.046	0.837(0.462,1.516)	0.558
No. of pregnancies						
<3	0.597(0.397,0.912)	0.017	0.783(0.524,1.168)	0.230	1.038(0.633,1.702)	0.884
≥3	0.638(0.446,0.912)	0.014	0.708(0.498,1.005)	0.054	0.866(0.573,1.307)	0.492
Abortion						
no	0.468(0.296,0.739)	0.001	0.619(0.398,0.961)	0.032	1.181(0.698,1.998)	0.534
yes	0.717(0.508,1.013)	0.059	0.795(0.56,1.113)	0.181	0.800(0.528,1.212)	0.293
Breast-feeding						
no	0.515(0.271,0.978)	0.043	0.730(0.397,1.342)	0.311	0.990(0.492,1.994)	0.979
yes	0.623(0.457,0.850)	0.003	0.775(0.575,1.045)	0.094	0.989(0.687,1.425)	0.953
Family history						
no	0.551(0.412,0.737)	0.000	0.751(0.570,0.991)	0.043	1.028(0.736,1.437)	0.870
yes	1.877(0.635,5.550)	0.255	0.768(0.256,2.309)	0.639	0.559(0.165,1.896)	0.351
	rs269198(CA+AA/CC)	<i>P</i>^a	rs34841297(A+--/AA)	<i>P</i>^a	rs12546233(CC+AC/AA)	<i>P</i>^a
	OR(95%CI)		OR(95%CI)		OR(95%CI)	
age						
<50	1.015(0.909,1.132)	0.795	1.998(1.072,3.723)	0.029	0.809(0.572,1.145)	0.231
≥50	0.864(0.508,1.471)	0.591	1.479(0.737,2.969)	0.271	0.718(0.467,1.10)	0.131
Age at menarche						
<14	0.990(0.614,1.597)	0.967	2.823(1.316,6.055)	0.008	0.881(0.573,1.353)	0.562
≥14	1.133(0.752,1.709)	0.550	1.220(0.679,2.191)	0.506	0.699(0.496,0.984)	0.040
Menopausal state						
Pre-menopausal	1.145(0.767,1.709)	0.508	2.490(1.288,4.815)	0.007	0.720(0.506,1.024)	0.067
Post-menopausal	0.855(0.509,1.438)	0.555	1.063(0.546,2.070)	0.858	0.857(0.563,1.305)	0.472
Age at menopause						
<45	0.725(0.126,4.158)	0.718	0.333(0.027,4.106)	0.391	1.689(0.406,7.026)	0.471
≥45	0.896(0.514,1.562)	0.700	1.214(0.592,2.487)	0.597	0.799(0.510,1.250)	0.325
No. of pregnancies						
<3	1.035(0.646,1.658)	0.888	1.728(0.889,3.358)	0.107	0.688(0.460,1.030)	0.069
≥3	0.918(0.616,1.368)	0.674	1.573(0.861,2.876)	0.141	0.750(0.530,1.060)	0.103
Abortion						
no	1.164(0.707,1.918)	0.550	1.218(0.608,2.438)	0.578	0.517(0.333,0.804)	0.003
yes	0.865(0.580,1.289)	0.475	2.045(1.117,3.744)	0.020	0.889(0.637,1.240)	0.488
Breast-feeding						
no	1.124(0.570,2.216)	0.735	3.290(1.142,9.476)	0.027	0.650(0.357,1.182)	0.158

yes	0.995(0.702,1.410)	0.976	1.504(0.902,2.510)	0.118	0.785(0.583,1.057)	0.111
Family history						
no	1.063(0.771,1.464)	0.710	0.627(0.103,3.820)	0.613	0.737(0.560,0.971)	0.030
yes	0.661(0.199,2.192)	0.498	1.905(1.183,3.070)	0.008	0.943(0.323,2.754)	0.914

^a*P* values adjusted for age, menarche age, menopausal status, number of pregnancies, number of abortions, history of breast feeding, and family history of breast cancer in first-degree relatives in logistic regression analysis (except for stratification factors).

The distributions of hormone receptor status (ER, PR and HER-2) and molecular subtypes (triple-negative, Her-2 overexpression and luminal) of patients with BC presenting different SNP genotypes are shown in Supplementary Tables 4 and 5, respectively. The rs269183 T>C variant was associated with Her-2 receptor positive and Her-2 overexpressing BC. The rs269198 C>A variant reduces the risk of Her-2-overexpressing BC. The GG genotype of rs4259395 was related to the PR receptor-positive BC, and the TT genotype of rs2588297 was related to the Luminal-type BC. SNPs rs3802201 C>G and rs2553716 A>C were both associated with PR positivity and luminal BC. In addition, the rs12546233 AC+CC genotype was associated with triple-negative BC and the AA genotype was associated with Her-2-overexpression BC.

False positive report probability (FPRP)

FPRP analysis [20] was used to evaluate the reliability of the positive results for *MIR2052HG* SNPs associated with BC susceptibility. As presented in Supplementary Table 6, when the critical value of FPRP was set to 0.5 and prior probability was 0.25, the FPRP values of all positive results for SNPs rs2553716, rs269183, rs269198, and rs12546233 were lower than the critical value. A possible association between the SNPs and the risk of BC was observed, which was worthy of further research and verification.

Haplotype analysis

Haplotype analysis was used to test the combined effect of *MIR2052HG* SNPs (Supplementary Table 7). Haplotype C_{rs3802201}A_{rs2553716}A_{rs4259395}G_{rs2588297}C_{rs10957736}T_{rs269183}C_{rs269198} W_{rs34841297}A_{rs12546233} was the highest frequency haplotype, and individuals with this haplotype have an increased risk of BC (*OR*: 1.203, 95% *CI*: 1.001-1.445), while Haplotype G_{rs3802201}C_{rs2553716}G_{rs4259395}T_{rs2588297} T_{rs10957736} T_{rs269183}C_{rs269198} M_{rs34841297} A_{rs12546233} and haplotype G_{rs3802201}C_{rs2553716}G_{rs4259395}T_{rs2588297}T_{rs10957736}T_{rs269183}C_{rs269198}M_{rs34841297}C_{rs12546233} were associated with a low risk of BC (*OR*: 0.258, 95% *CI*: 0.073-0.908; *OR*: 0.698, 95% *CI*: 0.549-0.887, respectively).

Multifactor dimensionality reduction

MDR software (multifactor dimensionality reduction 3.0.2) was used to analyze the interaction of genes and reproductive factors. As shown in Table 4, there was an interaction with the rs34841297 - (homozygous deletion) of A genotype, non-menopausal status, and no history of breastfeeding was observed, furthermore, the interaction model revealed a higher risk of BC (*OR*: 1.771, 95% *CI*: 1.367-2.941, *P*<0.001).

Real-time PCR results

The *MIR2052HG* expression levels in individuals with the rs34841297 --, A- and AA genotypes were shown in Figure 1A. The relative expression in individuals with the homozygous deletion genotype (1.68±1.37) was significantly higher than that in individuals with the heterozygous deletion genotype (0.94±0.95) (*P*=0.011) and AA genotype (0.26±0.12) (*P*<0.001). In addition, the relative expression of *MIR2052HG* in individuals with a homozygous deletion of rs34841297 was significantly higher than that of individuals with the AA genotype (*P*=0.001).

Dual-luciferase reporter assays

A dual-luciferase reporter assay was performed in 293T cells to determine the biological association between rs34841297 and miR-4456. As shown in Figure 1B, the relative luciferase activity of the rs34841297 W-NC group was significantly higher than rs34841297 W-miR-4456 group (*P*<0.001), suggesting an interaction between the rs34841297 wild genotype plasmid vector and miR-4456. Meanwhile, the relative luciferase activity of the rs34841297 W-miR-4456 group was lower than the rs34841297 MUT-miR-4456 group (*P*<0.001), showing that the interaction between the plasmid vector of *MIR2052HG* and miR-4456 disappeared due to the deletion of the rs34841297 A allele.

The effect of miR-4456 combined with *MIR2052HG* on cell proliferation, invasion and migration

The relative expression of *MIR2052HG* and miR-4456 in MDA-MB-231 cells (2.02±0.34, 3.95±0.65) were

Table 4. Interaction results between *MIR2052HG* SNPs and reproductive factors.

Model	TBA ^a	CVC ^b	χ^2	P	OR(95%CI)
Pre-menopausal, no breast-feeding	0.4966	3/10	10.061	0.002	1.543(1.180,2.018)
Pre-menopausal, no breast-feeding, rs34841297	0.5313	7/10	18.883	<0.001	1.771(1.367,2.941)

^aTesting balance accuracy.

^bCross-validation consistency.

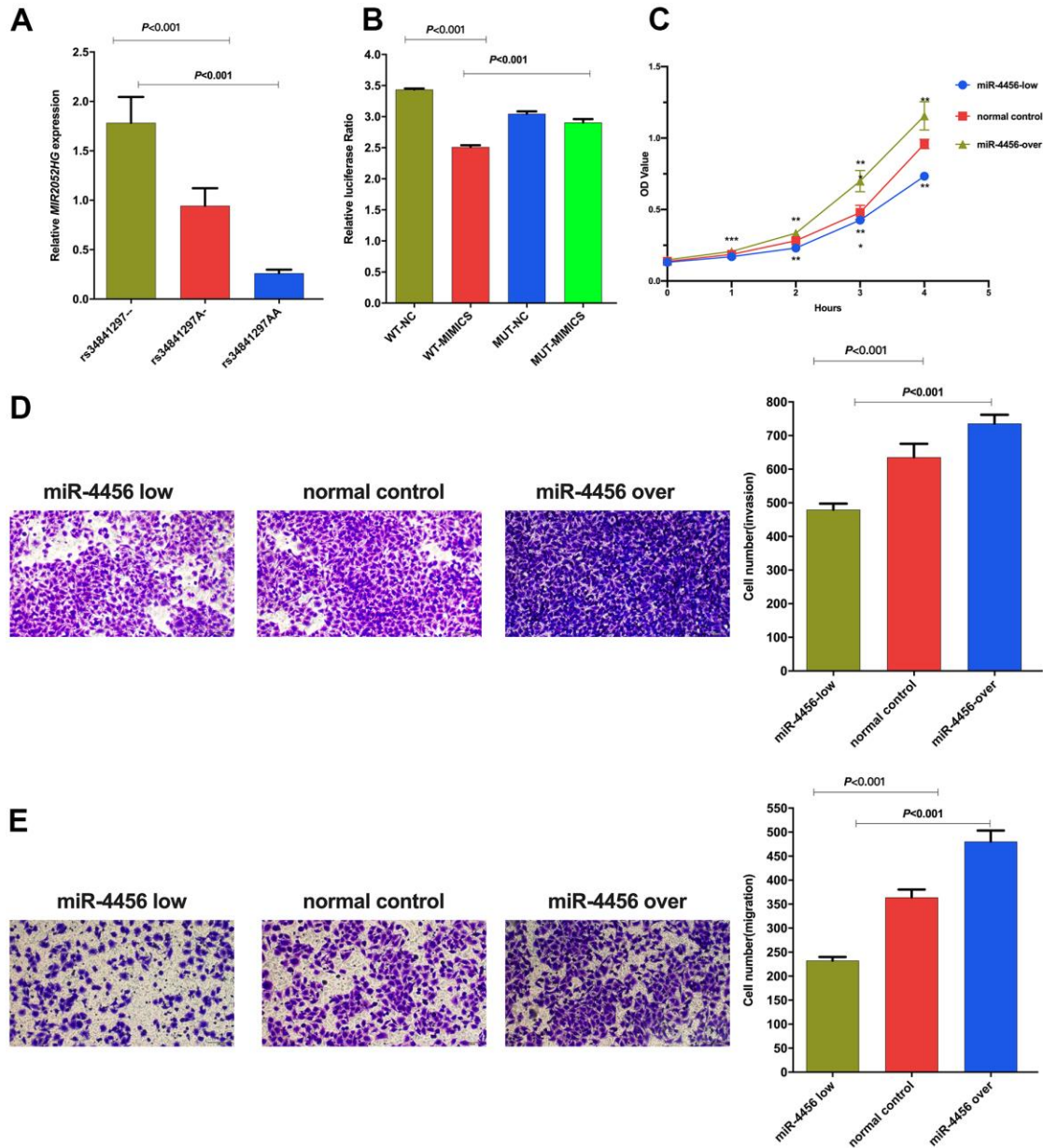


Figure 1. (A) Relative expression of lncRNA *MIR2052HG* in different genotypes of rs34841297. (B) The relative fluorescence value of different experimental groups in the dual luciferase report experiment. WT-NC: rs34841297 wild plasmids-Normal Control mimics, WT-MIMIC: rs34841297 wild plasmids-miR-4456 mimics, MUT-NC: rs34841297 mutant plasmids-Normal Control mimics, MUT-MIMIC: rs34841297 mutant plasmids-miR-4456 mimics. (C) The effect of miR-4456 combined with *MIR2052HG* on the proliferation of breast cancer cells (0-96h). *represents $P < 0.05$; **represents $P < 0.01$; ***represents $P < 0.001$. (D) Transwell experiment explores the effect of miR-4456 combined with *MIR2052HG* on the invasion ability of MDA-MB-231 cells. (E) Transwell experiment explores the effect of miR-4456 combined with *MIR2052HG* on the migration ability of MDA-MB-231 cells.

both higher than that in MCF10A cells (0.92 ± 0.08 , 1.17 ± 0.79), and the differences were statistically significant (Supplementary Figure 1).

qRT-PCR was performed to detect the relative expression of miR-4456 in BC cells (MDA-MB-231) stably transfected with three lentiviruses. The results of the CCK-8 test are shown in Figure 1C. In the MDA-MB-231 cells, the OD values for the miR-4456 low-expression group at 24 h ($P<0.001$), 48 h ($P=0.001$), 72 h ($P<0.001$) and 96 h ($P=0.002$) were lower than those of the NC group, and the OD values for the miR-4456 overexpression group at 24 h ($P<0.001$), 48 h ($P=0.005$), 72 h ($P<0.001$) and 96 h ($P=0.001$) were higher than those of the NC group. Then, the number of cells passing through the Matrigel-covered Transwell chamber was observed to determine the ability of miR-4456 to modulate the invasion of MDA-MB-231 cells. The results presented in Figure 1D showed that the number of invading MDA-MB-231 cells in the miR-4456 low-expression group was lower than that in the NC group ($P<0.001$), and the number of invading MDA-MB-231 cells in the miR-4456 overexpression group was higher than that in the NC group ($P=0.002$). Finally, Transwell experiments were performed to explore the migration ability of BC cells with different miR-4456 expression levels. As presented in Figure 1E, the number of migrating MDA-MB-231 cells in the miR-4456 low-expression group MDA-MB-231 was lower than that in the NC group ($P=0.004$), and the number of migrating MDA-MB-231 cells in the miR-4456 overexpression group MDA-MB-231 was higher than that in the NC group ($P=0.022$).

DISCUSSION

In the present study, six SNPs rs3802201 (C>G), rs2553716 (A>C), rs4259395 (A>G), rs2588297 (G>T), rs10957736 (C>T) and rs12546233 (A>C), reduced the risk of BC. Among them, the rs3802201 (C>G) mutation reduces the risk of breast cancer, which corresponds with the study by Wang L et al. [19]. However, the rs34841297 gene polymorphism was positively correlated with the incidence of BC, and rs34841297 A gene deletion might increase the risk of BC. Furthermore, the associations between nine *MIR2052HG* SNPs and the BC receptor status (ER, PR, Her-2) were analyzed. Estrogen receptor (ER) can regulate normal breast epithelial cells and breast gland proliferation of cancer cells [21, 22]. Progesterone receptor (PR) was a member of the nuclear receptor superfamily of transcription factors that has the biological function of promoting functional recovery and reducing the volume of BC lesions [23]. Human epidermal growth factor receptor 2 (Her-2), as a marker for predicting the prognosis of BC, was regarded as the

key to evaluating the efficacy of targeted drugs [16]. Our results indicated that rs3802201, rs2553716 and rs4259395 may exert a protective effect on BC by affecting PR receptor status. The CT+CC genotype of rs269183 was related to Her-2 receptor status and may affect the prognosis of BC. MDR model results showed that with rs34841297, homozygous deletion of the A gene, a non-menopausal status and no history of breastfeeding resulted in a higher risk of BC, which might lead to an increased risk of BC.

MIR2052HG downregulation expression was reported to reduce ER α -positive BC cell growth [24]. Genetic variations in *MIR2052HG* were associated with the BC-free interval in the MA.27 trial (ClinicalTrials.gov number NCT00066573), and the variant SNPs were associated with increased *MIR2052HG* expression due to increased ER α binding to EREs [17, 25]. What's more, researchers have discovered that SNPs could mediate the occurrence of cancer by affecting the secondary structure and expression of lncRNAs and the biological effects mediated by the interactions between lncRNAs and miRNAs [26–28]. Here, we performed qRT-PCR to explore the total expression of *MIR2052HG* expression in individuals with different genotypes of rs34841297. A dual-luciferase reporter gene experiment was conducted to identify whether the rs34841297 A/- deletion mutation affects the binding ability of *MIR2052HG* to miR-4456. An interaction was observed between *MIR2052HG* and miR-4456 mimics when rs34841297 carried wild-type A allele, and the interaction disappeared with the deletion of the A allele, consistent with the results predicted by the LncRNASNP2 and DINAN websites. This result suggested that the rs34841297 polymorphism might affect the *MIR2052HG* and miR-4456 interaction.

MiRNAs are small noncoding RNAs of approximately 22 nucleotides in length that perform posttranscriptional regulatory functions by binding to specific sites on the target transcript [29]. miR-4456 is a tiny noncoding RNA located on chromosome 5 that putatively influences oxytocin signaling [30]. To date, no studies have reported the relationship between miR-4456 and cancer. Our study proposed the hypothesis that the effect of the lncRNA *MIR2052HG* on BC occurrence and development is based on the interaction of *MIR2052HG* and miR-4456 mediated by rs34841297. CCK-8, Transwell and scratch wound experiments were performed using MDA-MB-231 cells. Compared with the normal control group, the proliferation ability of MDA-MB-231 cells with high miR-4456 expression was significantly increased. The Transwell assays using MDA-MB-231 cells found that compared with the normal control group, the ability of cells with high miR-4456 expression to migrate through the chamber and

invade the Matrigel-coated Transwell chamber was significantly increased. Based on these experiments, we preliminarily suggested that the lncRNA *MIR2052HG* might mediate the binding of miR-4456 through rs34841297 to affect the proliferation, invasion and metastasis of BC.

This study is the first to explore the association between genetic variants in the lncRNA *MIR2052HG* and BC susceptibility. This study has several advantages. First, the patients included in our study were all newly diagnosed and the controls were selected according to the frequency matching, which might reduce the selection bias in the study. Second, SNPscan high-throughput typing technology was used to SNP typing, making the results more accurate and credible than traditional restriction fragment length polymorphism (PCR-RFLP) typing technology. Finally, genotyping of all SNPs was performed on 10% of randomly selected samples for sequencing verification. Additionally, all cell function experiments were repeated more than three times, which improved the authenticity and reliability of the study results. Nevertheless, this study still has some limitations. All the subjects included in this study were of the Chinese Han population, and further studies of other populations should be performed to verify our results. In addition, the mechanism by which *MIR2052HG* SNPs modulate BC must be further explored *in vivo*.

CONCLUSIONS

In conclusion, the study reveals the association between the *MIR2052HG* gene polymorphism and the occurrence of BC. The *MIR2052HG* rs34841297 A/--variant may affect the binding of miR-4456 to *MIR2052HG* and subsequently alter proliferation, invasion and migration of BC cells by regulating the expression of miR-4456 expression, which provides a baseline information for screening high-risk population populations for BC and formulating individualized preventive measures.

MATERIALS AND METHODS

Research sample

Based on a case-control study design, 504 new BC samples that were pathologically confirmed were collected from March 2016 to December 2018. All patients were recruited from the top three hospitals in Henan Province. Through frequency matching to patients by age (± 2 years), 505 healthy control samples were obtained from the biological sample bank of Henan Key Laboratory of Oncology Epidemiology. The basic information and clinical characteristics of the patients

included age, age at menarche, menopausal status, menopausal age, number of pregnancies, number of abortions, history of breastfeeding and family history of BC. Moreover, information on patients' hormone receptor status including estrogen receptor (ER), progesterone receptor (PR) and human epidermal growth factor receptor-2 (HER-2) was also acquired. The study was approved by the Ethics Review Committee of the Ethics Committee of Medical and Health Research of Zhengzhou University, Zhengzhou, Henan, China.

DNA extraction, SNP selection and genotyping

Sample DNA was extracted using a DNA Extraction Kit (Shanghai Laifeng Biotechnology Co., Ltd.) according to the manufacturer's instructions and stored at -80°C until use. The *MIR2052HG* SNPs were screened using Ensembl38, Ensembl37, NCBI 1000 Genomes and Haploview software, with a minor allele frequency (MAF) >0.05 in Chinese Han population. Finally, 9 SNPs in functional regulatory regions SNPs and tag SNPs were determined, and the basic information was presented in Supplementary Table 1. RNAfold was used to predict the secondary structure of the significant SNPs. LncRNASNP2 and DIANA were used to predict the binding capacity of miRNAs that SNPs might affect (Supplementary Table 2). Combined with the Δ Energy, correlation with BC, and site prediction intersection results, miR-4456 was selected for further functional research. All SNPs were genotyped with SNPscanTM multiple SNP typing kit.

Quantitative real-time PCR (qRT-PCR)

Plasma RNA was extracted from 72 randomly selected healthy controls with TRIzol reagent, DNA was removed and RNA was reverse transcribed into cDNA. The relative expression of *MIR2052HG* in individuals carrying the rs34841297 polymorphism was detected using qRT-PCR with SYBR-green among individuals with different genotypes of rs34841297, and GAPDH served as the endogenous control. All samples were analyzed in triplicate, and the relative expression was calculated using the method of $2^{-\Delta\text{Ct}}$ method. The sequences of primers used in this study were listed in Supplementary Table 3.

Dual-luciferase report assay

According to LncRNASNP2, carried with wild-type *MIR2052HG*-rs34841297 might gain a binding site for miR-4456, and the dual-luciferase assay verified the biological association between rs34841297 and miR-4456. Following the construction of the *MIR2052HG* rs34841297 wild-type and mutant pmirGLO plasmids, cotransfected HEK 293T cells with miR-4456 mimic or

normal control (NC) mimics by using the riboFECT™ CP kit. Firefly fluorescence activity and Renilla fluorescence activity in each group were detected 72 hours after transfection, and relative luciferase activity was calculated based on firefly/Renilla fluorescence.

Cytological function experiment

qRT-PCR experiments first detected the relative expression of *MIR2052HG* and miR-4456 in MDA-MB-231 cells and MCF10A cells. The relative expression was calculated using the method of 2- $\Delta\Delta C_t$ method. The sequences of *MIR2052HG* primers used in this study were listed in Supplementary Table 3. And the sequences of miR-4456 and U6 internal reference primers are shown in Supplementary Table 8.

BC cells (MDA-MB-231) stably transfected with low-expression, overexpression and negative control (NC) lentiviral vectors carrying miR-4456 were established. Then, the stably transfected cell line was used for CCK-8 experiments and Transwell experiments to detect the proliferation, invasion and migration ability of BC cells with different miR-4456 expression levels.

Statistical analysis

Unconditional logistic regression analysis was used to explore BC-related *MIR2052HG* SNPs and adjusted for age, menarche age, menopausal status, number of pregnancies, number of abortions, breastfeeding history, and family history. SHEsis online software was applied to conduct the haplotype analysis of *MIR2052HG*. Multifactor dimensionality reduction (MDR) was used to analyze the interaction between genes and the environment. Independent *t* tests were applied to compare the relative expression of the lncRNA *MIR2052HG* with different rs34841297 genotypes, and the false-positive report probability (FPRP) [20] analysis was conducted to verify the positive results and the cut-off value to ensure the reliability and accuracy of the positive results. A *t* test was used to compare the OD values from the CCK8 experiment among groups with low expression of miR-4456 and the high expression of miR-4456 and the NC group. Independent *t* tests were used to analyze the accurate counts of stained cells in the two groups obtained in Transwell experiments. Differences in invasion and migration capabilities were calculated using *t* tests. All data analysis and cell number statistical analyses in this study were performed using SPSS 21.0 (t-test, χ^2 test and unconditional logistic regression model analysis), ImageJ (calculate scratch healing area, Transwell migration and invasion cell count), and GraphPad Prism 7.0 (plot the calculation results). Two-sided $P < 0.05$ was statistically significant.

Informed consent

Informed consent was obtained from all individual participants included in the study.

Ethical approval

All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Ethical responsibilities of authors

The manuscript has not been submitted to more than one journal for simultaneous consideration. The manuscript has not been published previously (partly or in full).

AUTHOR CONTRIBUTIONS

CH S and LP X conceived of the study, participated in its design and helped to draft the manuscript. H Y carried out the molecular genetic studies, performed the statistical analysis, participated in the cell experiment and drafted the manuscript. QY S, FF C, XR J, YL W, KD X and YL Z revised the manuscript. All authors read and approved the final manuscript.

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CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

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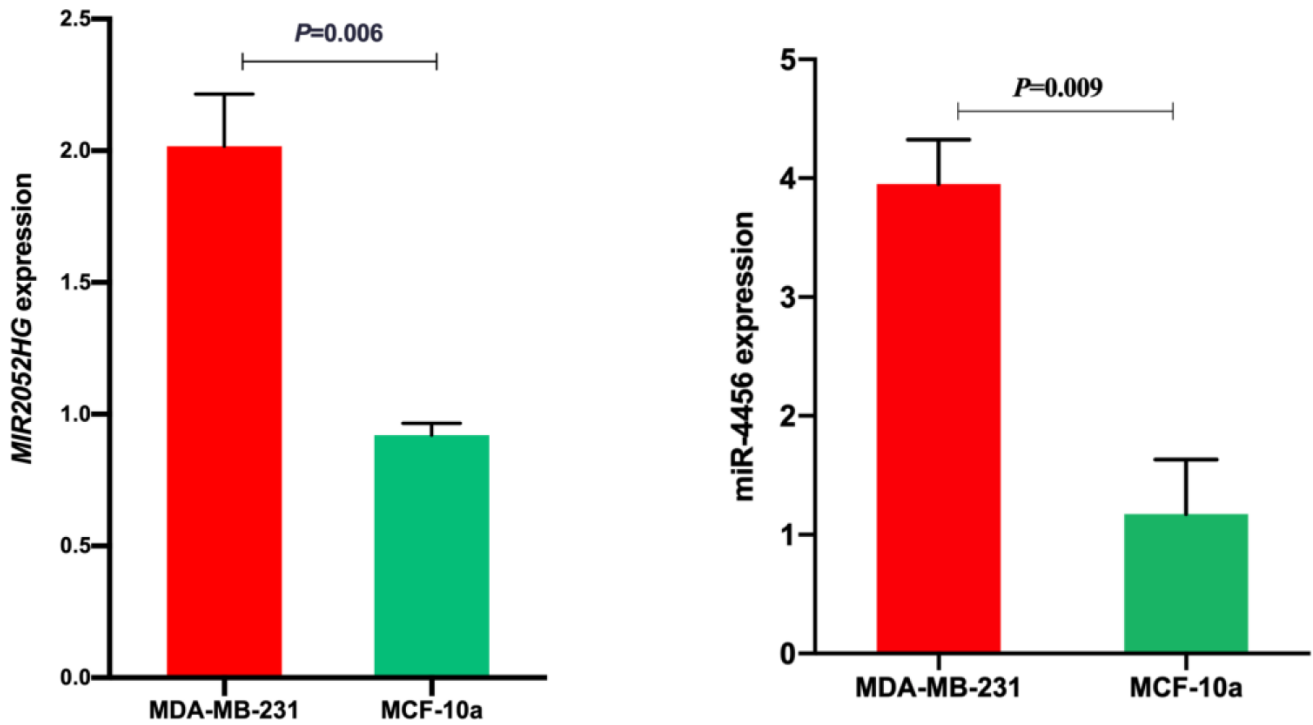
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SUPPLEMENTARY MATERIALS

Supplementary Figure



Supplementary Figure 1. Relative expression of *MIR2052HG* and miR-4456 in MDA-MB-231 and MCF-10A cells.

Supplementary Tables

Supplementary Table 1. Basic information of nine *MIR2052HG* SNPs.

SNP	Position	Ref Allele	%	Alt Allele	%
rs3802201	8:75678754	C	0.705	G	0.295
rs2553716	8:75696873	A	0.705	C	0.295
rs4259395	8:75702373	A	0.602	G	0.398
rs2588297	8:75732970	G	0.797	T	0.203
rs10957736	8:75750323	C	0.679	T	0.321
rs269183	8:75758791	T	0.890	C	0.110
rs269198	8:74702379	C	0.874	A	0.126
rs34841297	8:74752460	-	0.676	A	0.324
rs12546233	8:74758311%	A	0.721	C	0.279

Supplementary Table 2. Biological function prediction of *MIR2052HG* functional SNPs.

LncRNA	SNP	Ref/alt	Δ Energy (kCal/Mol)	miRNA	Energy (kCal/Mol)	Effect
<i>MIR2052HG</i>	rs12546233	A/C	3	has-miR-4659b-3p	-0.20	loss
				has-mir-4659a-3p	-0.20	loss
				has-miR-141-5p	-3.70	gain
				has-miR-3126-3p	-1.60	gain
				has-miR-3686	-0.40	gain
	rs269198	G/T	0.3	has-miR-452-3p	-1.00	gain
				has-miR-1207-3p	0.00	loss
				has-miR-2115-5p	-4.10	loss
				has-miR-1537-5p	-3.60	loss
				hsa-miR-185-3p	-4.90	gain
	rs34841297	-/A	2.4	hsa-miR-4498	-3.20	gain
				hsa-miR-5001-5p	-4.70	gain
				hsa-miR-4492	-4.00	gain
				hsa-miR-1298-3p	0.00	gain
				hsa-miR-762	-7.20	gain
				has-miR-4456	-0.70	loss
				has-miR-6842-3p	-3.50	loss

Supplementary Table 3. Sequence of *MIR2052HG* primers of qRT-PCR.

Gene	Sequence of primers ^a
<i>MIR2052HG</i>	F: ATCAGCGAGATTCCGTGGG R: GAAACTGCCTCATCAGACATAAAAAG
<i>GAPDH</i>	F: CGGAGTCAACGGATTTGGTTCGTAT R: AGCCTTCTCCATGGTGGTGAAGAC

^aqRT-PCR method was carried out in plasma to detect the relative *MIR2052HG* expression in AA genotype, A- genotype and -- genotype of rs34841297. Meanwhile the qRT-PCR was also performed to explore the relative *MIR2052HG* expression in MDA-MB-231cells and MCF10A cells.

Supplementary Table 4. The Associations between *MIR2052HG* SNPs and ER, PR and HER-2 status of breast cancer patients.

Genotype	ER		<i>P</i> ^a	OR(95%CI)	PR		<i>P</i> ^a	OR(95%CI)	HER-2		<i>P</i> ^a	OR(95%CI)
	Negative (n=149)	Positive (n=342)			Negative (n=191)	Positive (n=298)			Negative (n=138)	Positive (n=329)		
rs3802201												
CC	80	169		1	106	142		1	66	168		1
CG	63	150	0.430	1.178(0.785,1.768)	77	135	0.109	1.371(0.932,2.018)	60	144	0.877	0.967(0.633,1.478)
GG	6	23	0.090	2.303(0.877,6.050)	8	21	0.048	2.424(1.006,5.837)	12	17	0.311	0.656(0.290,1.484)
CG+GG	69	173	0.248	1.263(0.850,1.877)	85	156	0.049	1.458(1.001,2.124)	72	161	0.688	0.920(0.611,1.385)
rs2553716												
AA	82	170		1	107	144		1	68	169		1
AC	58	149	0.212	1.300(0.851,1.962)	74	132	0.093	1.397(0.946,2.061)	54	144	0.595	1.124(0.730,1.730)
CC	9	23	0.333	1.514(0.653,3.510)	10	22	0.103	1.966(0.872,4.433)	16	16	0.066	0.485(0.224,1.049)
AC+CC	67	172	0.165	1.326(0.891,1.974)	84	154	0.050	1.457(1.000,2.124)	70	160	0.959	0.989(0.656,1.491)
rs4259395												
AA	56	124		1	74	106		1	53	116		1
AG	75	172	0.581	1.128(0.736,1.728)	98	147	0.633	1.103(0.737,1.650)	63	171	0.226	1.320(0.842,2.068)
GG	18	46	0.364	1.350(0.707,2.580)	19	45	0.046	1.905(1.012,3.585)	22	42	0.933	0.973(0.517,1.833)
AG+GG	93	218	0.455	1.169(0.776,1.760)	117	192	0.302	1.226(0.833,1.805)	85	213	0.335	1.233(0.805,1.888)
rs2588297												
GG	107	226		1	134	198		1	92	223		1
GT	40	104	0.213	1.327(0.850,2.070)	54	89	0.445	1.176(0.776,1.783)	43	95	0.906	0.973(0.622,1.524)
TT	2	12	0.112	3.484(0.476,16.247)	3	11	0.081	3.249(0.867,12.183)	3	11	0.401	1.772(0.467,6.727)
GT+TT	42	116	0.112	1.424(0.921,2.200)	57	100	0.239	1.276(0.851,1.913)	46	106	0.920	1.023(0.661,1.583)
rs10957736												
CC	68	166		1	90	144		1	70	152		1
CT	72	148	0.633	0.905(0.601,1.363)	86	132	0.874	0.969(0.656,1.431)	61	147	0.361	1.220(0.796,1.869)
TT	9	28	0.398	1.423(0.627,3.229)	15	22	0.798	1.099(0.533,2.269)	7	30	0.098	2.132(0.869,5.230)
CT+TT	81	176	0.856	0.964 (0.649,1.431)	101	154	0.949	0.988(0.680,1.436)	68	177	0.192	1.315(0.871,1.986)
rs269183												
TT	110	274		1	145	238		1	116	249		1
CT	38	65	0.155	0.713(0.448,1.136)	44	58	0.378	0.816(0.519,1.282)	22	76	0.057	1.683(0.985,2.875)
CC	1	3	0.839	1.271(0.126,12.867)	2	2	0.696	0.667(0.088,5.062)	0	4	NC	NC
CT+CC	39	68	0.175	0.728(0.460,1.152)	46	60	0.353	0.810(0.519,1.264)	22	80	0.039	1.755(1.030,2.991)
rs269198												
CC	105	265		1	139	230		1	109	243		1
CA	42	72	0.130	0.707(0.451,1.108)	48	65	0.414	0.834(0.539,1.290)	29	79	0.298	1.300(0.793,2.131)
AA	2	5	0.967	1.036(0.192,5.581)	4	3	0.368	0.494(0.106,2.297)	0	7	NC	NC
CA+AA	44	77	0.147	0.722(0.465,1.121)	52	68	0.326	0.808(0.528,1.237)	29	86	0.175	1.403(0.860,2.290)
rs34841297												
AA	10	28		1	15	23		1	7	31		1
A-	68	151	0.583	0.799(0.360,1.776)	85	132	0.728	0.879(0.424,1.820)	62	146	0.199	0.554(0.225,1.366)
--	71	163	0.508	0.766(0.348,1.686)	91	143	0.746	0.888(0.434,1.820)	69	152	0.106	0.478(0.196,1.169)
A+--	139	314	0.528	0.781(0.363,1.681)	176	275	0.728	0.884(0.441,1.772)	131	298	0.132	0.512(0.214,1.225)
rs12546233												
AA	80	187		1	101	166		1	79	174		1
AC	64	128	0.590	0.894(0.593,1.346)	79	111	0.411	0.848(0.572,1.256)	52	130	0.315	1.248(0.810,1.920)
CC	5	27	0.078	2.459(0.904,6.694)	11	21	0.471	1.335(0.609,2.928)	7	25	0.253	1.695(0.686,4.183)
AC+CC	69	155	0.955	1.012(0.681,1.502)	90	132	0.613	0.908(0.624,1.321)	59	155	0.211	1.302(0.861,1.969)

^a*P* values adjusted for age, menarche age, menopausal status, number of pregnancies, number of abortions, history of breast feeding, and family history of breast cancer in first-degree relatives in logistic regression analysis.

Supplementary Table 5. The Associations between *MIR2052HG* SNPs and luminal breast cancer, Her-2 over-expression breast cancer and triple-negative breast cancer.

Genotype	Triple-negative		<i>P</i> ^a	OR(95%CI)	Her-2 over-expression		<i>P</i> ^a	OR(95%CI)	Luminal		<i>P</i> ^a	OR(95%CI)
	no (n=418)	yes (n=49)			no (n=390)	yes (n=77)			no (n=195)	yes (n=272)		
rs3802201												
CC	211	23		1	192	42		1	107	127		1
CG	182	22	0.906	1.039(0.550,1.965)	171	33	0.521	0.847(0.509,1.407)	80	124	0.116	1.370(0.925,2.028)
GG	25	4	0.869	1.105(0.336,3.640)	27	2	0.105	0.291(0.065,1.296)	8	21	0.026	2.730(1.127,6.609)
CG+GG	207	26	0.881	1.048(0.566,1.939)	198	35	0.305	0.770(0.467,1.269)	88	145	0.045	1.478(1.009,2.146)
rs2553716												
AA	213	24		1	195	42		1	109	128		1
AC	180	18	0.523	0.805(0.414,1.565)	165	33	0.691	0.902(0.543,1.500)	76	122	0.069	1.443(0.972,2.144)
CC	25	7	0.204	1.895(0.706,5.085)	30	2	0.089	0.274(0.062,1.215)	10	22	0.054	2.233(0.985,5.061)
AC+CC	205	25	0.871	0.950(0.512,1.761)	195	35	0.395	0.805(0.488,1.327)	86	144	0.030	1.526(1.041,2.237)
rs4259395												
AA	151	18		1	141	28		1	74	95		1
AG	211	23	0.587	0.829(0.422,1.630)	194	40	0.956	0.985(0.574,1.689)	101	133	0.643	1.102(0.730,1.666)
GG	56	8	0.896	0.940(0.369,2.390)	55	9	0.459	0.731(0.319,1.676)	20	44	0.031	2.006(1.065,3.778)
AG+GG	267	31	0.629	0.854(0.450,1.619)	249	49	0.777	0.928(0.552,1.559)	121	177	0.281	1.243(0.837,1.846)
rs2588297												
GG	280	35		1	260	55		1	137	178		1
GT	124	14	0.434	0.760(0.382,1.512)	118	20	0.404	0.786(0.446,1.383)	55	83	0.340	1.228(0.806,1.871)
TT	14	0	1.000	0.000	12	2	0.585	0.650(0.138,3.057)	3	11	0.052	3.716(0.987,13.996)
GT+TT	138	14	0.266	0.677(0.341,1.346)	130	22	0.354	0.772(0.446,1.335)	58	94	0.158	1.343(0.892,2.024)
rs10957736												
CC	195	27		1	190	32		1	91	131		1
CT	187	21	0.220	0.673(0.357,1.267)	171	37	0.334	1.297(0.765,2.199)	89	119	0.852	0.963(0.647,1.433)
TT	36	1	0.092	0.170(0.022,1.339)	29	8	0.432	1.429(0.587,3.434)	15	22	0.561	1.242(0.598,2.578)
CT+TT	223	22	0.098	0.592(0.318,1.101)	200	45	0.283	1.319(0.796,2.184)	104	141	0.991	1.002(0.685,1.466)
rs269183												
TT	323	42		1	312	53		1	147	218		1
CT	91	7	0.139	0.524(0.223,1.233)	75	23	0.039	1.802(1.031,3.150)	46	52	0.334	0.797(0.504,1.262)
CC	4	0	1.000	0.000	3	1	0.742	1.477(0.145,15.059)	2	2	0.781	0.749(0.098,5.740)
CT+CC	95	7	0.112	0.510(0.217,1.198)	78	24	0.039	1.787(1.031,3.097)	48	54	0.321	0.795(0.506,1.250)
rs269198												
CC	313	39		1	301	51		1	141	211		1
CA	98	10	0.377	0.712(0.335,1.514)	84	24	0.056	1.710(0.987,2.962)	50	58	0.355	0.811(0.520,1.265)
AA	7	0	1.000	0.000	5	2	0.456	1.908(0.349,10.423)	4	3	0.448	0.551(0.118,2.573)
CA+AA	105	10	0.310	0.677(0.319,1.437)	89	26	0.046	1.723(1.009,2.941)	54	61	0.290	0.791(0.513,1.221)
rs34841297												
AA	37	1		1	30	8		1	16	22		1
A-	190	18	0.266	3.263(0.405,26.271)	171	37	0.883	0.936(0.386,2.266)	87	121	0.782	0.903(0.437,1.866)
--	191	30	0.070	6.725(0.855,5.887)	189	32	0.468	0.721(0.298,1.744)	92	129	0.724	0.879(0.429,1.799)
A+--	381	48	0.127	4.908(0.636,37.867)	360	69	0.637	0.816(0.351,1.896)	179	250	0.741	0.890(0.445,1.779)
rs12546233												
AA	221	32		1	218	35		1	102	151		1
AC	166	16	0.080	0.554(0.285,1.074)	144	38	0.037	1.738(1.033,2.921)	82	100	0.339	0.823(0.552,1.228)
CC	31	1	0.115	0.190(0.024,1.495)	28	4	0.716	0.812(0.265,2.493)	11	21	0.306	1.511(0.686,3.329)
AC+CC	197	17	0.034	0.497(0.260,0.949)	172	42	0.081	1.563(0.946,2.580)	93	121	0.603	0.904(0.618,1.323)

^a*P* values adjusted for age, menarche age, menopausal status, number of pregnancies, number of abortions, history of breast feeding, and family history of breast cancer in first-degree relatives in logistic regression analysis.

Supplementary Table 6. False positive report probability analysis.

Genotype	Stratification factors	OR (95%CI)	P	Prior probability					
				0.25	0.1	0.01	0.001		
rs3802201	CC/CG+GG	All sample	BC	0.756(0.580,0.986)	0.039	0.124	0.299	0.824	0.979
		No family history	BC	0.717(0.544,0.944)	0.018	0.071	0.186	0.716	0.962
	CC/GG	All cases	PR	2.424(1.006,5.837)	0.048	0.505	0.753	0.971	0.997
	CC/CG+GG	All cases	PR	1.458(1.001,2.124)	0.049	0.210	0.444	0.898	0.989
	CC/GG	All cases	luminal	2.730(1.127,6.609)	0.026	0.458	0.717	0.965	0.996
rs2553716	CC/CG+GG	All cases	luminal	1.478(1.009,2.146)	0.045	0.184	0.404	0.882	0.987
	AA/AC	All sample	BC	0.739(0.560,0.973)	0.031	0.108	0.267	0.801	0.976
	AA/ AC+CC	All sample	BC	0.737(0.565,0.931)	0.024	0.038	0.105	0.565	0.929
	AA+CC/AC	All sample	BC	0.770(0.590,1.006)	0.055	0.163	0.368	0.865	0.985
	AA/ AC+CC	Age at menarche≥14	BC	0.686(0.488,0.965)	0.031	0.139	0.326	0.842	0.982
		Gravidity≥3	BC	0.702(0.495,0.994)	0.046	0.184	0.403	0.881	0.987
		No breast feeding	BC	1.959(1.072,3.581)	0.029	0.310	0.574	0.937	0.993
		No family history	BC	0.691(0.524,0.910)	0.009	0.041	0.113	0.584	0.934
	AA/ AC+CC	All cases	PR	1.457(1.000,2.124)	0.050	0.212	0.447	0.899	0.989
	AA/AC	All cases	luminal	2.233(0.985,5.061)	0.054	0.489	0.742	0.969	0.997
AA/ AC+CC	All cases	luminal	1.526(1.041,2.237)	0.030	0.164	0.370	0.866	0.985	
rs4259395	AA/AG+GG	All sample	BC	0.756(0.573,0.999)	0.049	0.154	0.353	0.857	0.984
		Age≥50	BC	0.622(0.395,0.979)	0.040	0.240	0.486	0.912	0.991
		No family history	BC	0.738(0.554,0.985)	0.039	0.135	0.318	0.837	0.981
	AA/GG	All cases	PR	1.905(1.012,3.585)	0.046	0.374	0.642	0.952	0.995
	AA/AG	All cases	luminal	2.006(1.065,3.778)	0.031	0.337	0.604	0.944	0.994
rs2588297	GG/GT	All sample	BC	0.597(0.448,0.794)	0.000	0.005	0.016	0.148	0.636
	GG/ GT+TT	All sample	BC	0.606(0.459,0.798)	0.000	0.004	0.013	0.126	0.592
	GG+TT/GT	All sample	BC	0.608(0.458,0.807)	0.001	0.007	0.019	0.178	0.686
	GG/ GT+TT	Age<50	BC	0.671(0.468,0.963)	0.030	0.151	0.348	0.854	0.983
		Age≥50	BC	0.583(0.373,0.911)	0.018	0.161	0.366	0.864	0.985
		Age at menarche≥14	BC	0.580(0.407,0.828)	0.003	0.035	0.099	0.547	0.924
		Un-menopause	BC	0.583(0.403,0.844)	0.004	0.051	0.138	0.638	0.947
		Gravidity<3	BC	0.597(0.397,0.912)	0.017	0.144	0.335	0.847	0.982
		Gravidity≥3	BC	0.638(0.446,0.912)	0.014	0.092	0.233	0.770	0.971
		Age at menopause≥45	BC	0.599(0.376,0.953)	0.031	0.219	0.458	0.903	0.989
		No history of abortion	BC	0.468(0.296,0.739)	0.001	0.050	0.135	0.633	0.946
		No breast feeding	BC	0.515(0.271,0.978)	0.043	0.372	0.640	0.951	0.995
		Have breast feeding	BC	0.623(0.457,0.850)	0.003	0.025	0.071	0.456	0.894
		No family history	BC	0.551(0.412,0.737)	0.000	0.002	0.005	0.055	0.372
	rs10957736	GG/TT	All case	luminal	3.716(0.987,13.996)	0.052	0.636	0.840	0.983
CC/TT		All sample	BC	0.562(0.348,0.907)	0.018	0.185	0.405	0.882	0.987
CC/ CT+TT		All sample	BC	0.756(0.579,0.986)	0.039	0.124	0.299	0.824	0.979
CC+CT/TT		All sample	BC	0.627(0.397,0.991)	0.046	0.257	0.509	0.919	0.991
CC/ CT+TT		Age≥50	BC	0.605(0.392,0.932)	0.023	0.171	0.382	0.872	0.986
		Age at menarche≥14	BC	0.704(0.500,0.990)	0.043	0.174	0.387	0.874	0.986
		Age at menopause≥45	BC	0.633(0.404,0.992)	0.046	0.252	0.502	0.917	0.991
		No history of abortion	BC	0.619(0.398,0.961)	0.032	0.209	0.442	0.897	0.989
		No family history	BC	0.751(0.570,0.991)	0.043	0.139	0.326	0.842	0.982
rs269183		TT/ CT+CC	All cases	HER-2	1.755(1.030,2.991)	0.039	0.291	0.552	0.931
	TT/CT	All cases	Her-2 over-expression	1.802(1.031,3.150)	0.039	0.309	0.573	0.937	0.993
	TT/ CT+CC	All cases	Her-2 over-expression	1.787(1.031,3.097)	0.039	0.303	0.566	0.935	0.993
rs269198	CC/CA+AA	All cases	Her-2 over-expression	1.723(1.009,2.941)	0.046	0.312	0.576	0.937	0.993
rs34841297	AA/--	All sample	BC	1.936(1.208,3.123)	0.006	0.121	0.292	0.819	0.979
	AA/ A+--	All sample	BC	1.704(1.087,2.672)	0.020	0.173	0.386	0.874	0.986

	AA+A-/-	All sample	BC	1.388(1.062,1.812)	0.016	0.063	0.167	0.688	0.957
	AA/ A+/-	Age<50	BC	1.998(1.072,3.723)	0.029	0.324	0.590	0.941	0.994
		Age at menarche<14	BC	2.823(1.316,6.055)	0.008	0.306	0.570	0.936	0.993
		Un-menopause	BC	2.490(1.288,4.815)	0.007	0.233	0.477	0.910	0.990
		Abortion history	BC	2.045(1.117,3.744)	0.020	0.280	0.538	0.928	0.992
		No breast feeding	BC	3.290(1.142,9.476)	0.027	0.530	0.772	0.974	0.997
		had family history	BC	1.905(1.183,3.070)	0.008	0.130	0.309	0.831	0.980
rs12546233	AA/AC	All sample	BC	0.764(0.578,1.006)	0.055	0.166	0.373	0.868	0.985
	AA/ AC+CC	All sample	BC	0.747(0.573,0.973)	0.031	0.103	0.256	0.791	0.974
		Age at menarche≥14	BC	0.699(0.496,0.984)	0.040	0.166	0.373	0.867	0.985
		No history of abortion	BC	0.517(0.333,0.804)	0.003	0.073	0.191	0.723	0.963
		No family history	BC	0.737(0.560,0.971)	0.030	0.106	0.262	0.796	0.975
		All cases	Triple-Negative	0.497(0.260,0.949)	0.034	0.354	0.622	0.948	0.955

Supplementary Table 7. Haplotype analysis of nine SNPs in *MIR2052HG*.

Gene	Haplotype ^a	Cases (%)	Controls (%)	χ^2	P	OR (95%CI)
<i>MIR2052HG</i>	C A A G C T C M ^b A	579.61(57.5)	545.74(54.0)	3.879	0.049	1.203 (1.001,1.445)
	C A A G T C A W C	14.72(1.5)	20.16(2.0)	0.787	0.375	0.737 (0.374,1.451)
	C A A T T T C W C	10.79(1.1)	6.42(0.6)	1.181	0.277	1.712 (0.642,4.565)
	C A G G T C A W A	32.98(3.3)	30.66(3.0)	0.126	0.722	1.095 (0.664,1.805)
	C A G G T C A W C	54.55(5.4)	55.25(5.5)	0.000	0.993	1.002 (0.681,1.473)
	G C G G C T C M A	90.02(8.9)	82.97(8.2)	0.440	0.507	1.112 (0.813,1.520)
	G C G T C T C M A	15.05(1.5)	10.51(1.0)	0.886	0.346	1.461 (0.661,3.228)
	G C G T T T C W A	3.06(0.3)	11.93(1.2)	5.160	0.023	0.258 (0.073,0.908)
	G C G T T T C W C	138.71(13.8)	189.63(18.8)	8.678	0.003	0.698 (0.549,0.887)

^aThe sequence of SNP locus is rs3802201; rs2553716; rs4259395; rs2588297; rs10957736; rs269183; rs269198; rs34841297 and rs12546233

^bM represents A genotype, W represents deletion

Supplementary Table 8. Sequence of miR-4456 primers of qRT-PCR in MDA-MB-231 and MCF-10A cells.

Gene	Sequence of primers (5'-3')
miR-4456	F: ATATATCGCGCTGGTGGCTT R: AGTGCAGGGTCCGAGGTATT RT: GTCGTATCCAGTGCAGGGTCCGAGGTATTCGCACTGGATACGACAAAAGG GCACTGGATACGACCAAGCC
U6	F: CGCAAATTCGTGAAGCGTTC R: GCAGGGTCCGAGGTATTC RT: GTCGTATCCAGTGCAGGGTCCGAGGTATTCG