

Investigation on Probable Association Between *IL-13*, *IL-13RA1*, and *IL-13RA2* Genes Polymorphism and Pulmonary Tuberculosis

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Objective: Our study aimed to explore the association of *IL-13*, *IL-13RA1*, and *IL-13RA2* genes polymorphisms with PTB susceptibility and its clinical features.

Methods: Nine SNPs were genotyped by improved multiple ligase detection reaction (iMLDR) in 476 PTB patients and 473 controls. The association between these SNPs and PTB risk was analyzed using SPSS software and haplotype analysis was assessed using SHEsis software.

Results: The *IL-13RA1* rs2495636 GA genotype frequency in PTB patients was significantly decreased, and *IL-13RA2* rs5946039 A allele was related to the lower risk of PTB. In *IL-13* gene, rs20541 variant was found to be associated with PTB risk under recessive mode. Moreover, *IL-13RA1* rs141573089 C allele was significantly lower in PTB presenting with fever, drug resistance, and CC genotype was decreased in PTB presenting with leukopenia. *IL-13RA1* rs2495636 polymorphism was associated with drug resistance, pulmonary infection, and *IL-13RA2* rs3795175, rs638376 polymorphisms were related to drug resistance in PTB patients.

Conclusion: *IL-13* rs20541, *IL-13RA1* rs2495636, *IL-13RA2* rs5946039 polymorphisms might be contributed to the genetic background of PTB in Chinese population.

Keywords: pulmonary tuberculosis, *Mycobacterium tuberculosis*, interleukin-13, single nucleotide polymorphisms

Introduction

Tuberculosis (TB) is a common infectious disease caused by *Mycobacterium tuberculosis* (MTB) infection, which could not only spread to the respiratory system but also to other body systems, and is the leading infectious cause of death worldwide.¹ It is worth noting that there were estimated 9.9 million new incidents of TB patients globally in 2021.² Approximately one third of the world's population was infected with MTB, with only 10% of them developing active pulmonary TB (PTB),^{3,4} and this statistic remained largely unexplained. Increasing evidence suggested that genetic and environmental factors, as well as bacterial agents, play key roles in PTB development.^{5,6} Studying the significant roles of host genetic factors in TB disease and determining the specific host genes associated with PTB susceptibility were helpful to further understand the pathogenesis of PTB and formulate appropriate treatment strategies.

Additionally, immune pathways are important in controlling latent PTB infections, while the mechanisms underlying the progression from latent stage to the onset of active disease are still not fully understood.⁷ Complex interactions between lymphocytes, antigen-presenting cells, and secreted cytokines are involved in the immune response against PTB.³ Several inflammatory cytokines abnormally expressed in PTB could serve as biomarkers for disease severity and bacterial burden.⁸ Genetic variations in cytokine genes might alter the expression levels by affecting the recognition sites

of transcription factors, and their transcriptional activity.⁹ Hence, polymorphisms of cytokine genes were closely related to PTB susceptibility.^{10–12} For example, *IFN- γ -874* gene polymorphisms were reported to be related to PTB, and *interleukin (IL)-10* gene variations contributed to the progression of PTB.

IL-13 was a cytokine secreted by activated T-helper 2 (Th2) cells, which involved in the regulation of humoral immunity and the type II cytokine-mediated immune response, and had many biological functions, including promoting inflammatory immune response and inhibiting the production of inflammatory cytokines.^{13,14} Study had shown that IL-13 was involved in the development of PTB, and the plasma level of IL-13 was significantly increased in PTB patients.⁷ In human TB, the relative increased expression of IL-13, which associated with lung damage, indicated that a subversive Th2 component in the response to MTB might undermine protective immunity and contribute to reactivation and tissue pathology.¹⁵ Recently, another study analyzed the association between *IL-13* gene polymorphism and PTB susceptibility, while no meaningful result was found.¹⁶

IL-13 had two cognate receptors, known as IL-13RA1 and IL-13RA2, which specifically bind to IL-13 in vitro.^{17,18} The role of genetic variations in *IL-13RA1* and *IL-13RA2* in human disease had also been reported,^{19,20} however, similar studies had not been conducted in PTB patients. Therefore, to improve our understanding of the role of *IL-13* in the pathogenesis of PTB, we conducted this study to explore the possible association between *IL-13*, *IL-13RA1*, and *IL-13RA2* gene single nucleotide polymorphisms (SNPs) and susceptibility to PTB in a Chinese population.

Materials and Methods

Study Participants

In total, 476 PTB patients and 473 unrelated healthy individuals were included. All PTB patients were selected from Anhui Chest Hospital and were diagnosed using the following criteria: suspicious clinical symptoms, chest radiography, sputum and/or bronchoalveolar lavage fluid MTB culture, microscopy of acid fast bacilli (AFB), and the effects of anti-TB treatment. The patients with cancer, HIV positivity, hepatitis, and immune-compromised conditions were excluded, as well as those with poor compliance. Healthy individuals with the same ethnic background, no history of TB, cancer, and HIV were enrolled from the health examining center in the same area as controls. Moreover, all controls were examined by a senior physician to ensure that they were asymptomatic with negative sputum smear and culture, and normal chest radiograph.

This study was carried out after being approved by the ethics committee of Anhui Medical University (20200250) in complies with the Declaration of Helsinki. After obtaining informed consent, we collected peripheral blood samples and relevant data from the study subjects, including basic information such as age and sex, as well as some clinical data such as pulmonary infection, leukopenia, fever, drug resistance, drug-induced liver injury (DILI), sputum smear status.

DNA Extraction

Approximately 5 mL of peripheral blood was drawn from the medial cubital vein, placed in EDTA-containing tubes, and stored at -20°C until DNA extraction. The Flexi Gene-DNA Kit (Qiagen, Valencia, CA) was used to extract genomic DNA from the peripheral blood leukocytes with the standard procedures.

SNP Selection and Genotyping

The SNP selection methods mainly included literature searches and tag SNP selection. We systematically searched the previous studies on the relationship between *IL-13*, *IL-13RA1*, and *IL-13RA2* gene polymorphisms and human diseases to look for the SNPs associated with human disease susceptibility. Meanwhile, we obtained genotype data of *IL-13*, *IL-13RA1*, and *IL-13RA2* in CHB from the Ensembl Genome Browser 85 and CHBS_1000 g and used the pairwise option of the HaploView 4.0 software (Cambridge, MA, USA) to select the tag SNPs of these genes through linkage disequilibrium (LD) analysis with r^2 threshold >0.8 . The tag SNPs were selected with a minor allele frequency (MAF) ≥ 0.05 capturing all the common SNPs located in the chromosome locus transcribed into these genes and their flanking 2000 bp region.

Finally, we selected *IL-13* rs2066960, rs2069744, rs20541, *IL-13RA1* rs147857000, rs141573089, rs2495636, *IL-13RA2* rs3795175, rs638376, rs5946039 for genotyping.

Genetic polymorphisms were detected by an improved multiple ligase detection reaction (iMLDR) genotyping assay with the technical support of the Center for Genetic & Genomic Analysis, Genesky Biotechnologies (Inc., Shanghai). IMLDR is a commonly used, well-established genotyping method, which is widely used in our study and other studies, and its specific steps can be found in the previous studies.^{21,22} The detailed experimental steps were as follows: (1) 1 μ L DNA sample was extracted, and the quality of the sample was checked and the concentration was estimated by 1% agarose electrophoresis. Then, the DNA sample was diluted to 5–10 ng/ μ L according to the estimated concentration. (2) Multiplex PCR reaction was carried out with 20 μ L reaction system included 1x HotStarTaq bufer, 3.0 mM Mg²⁺, 0.3 mM dNTP, 1U HotStarTaq polymerase (Qiagen Inc.), 1 μ L sample DNA and 1 μ L multiple PCR primers. (3) Purification of multiple PCR products: 5U SAP enzyme and 2U Exonuclease I enzyme were added to 20 μ L PCR product, 37°C warm bath for 1 h, then 75°C inactivated for 15 min. (4) Ligating reaction system: 1 μ L 10x ligating buffer, 0.25 μ L high-temperature ligase, 0.4 μ L 5' ligating primer mixture (1 μ M), 0.4 μ L 3' ligating primer mixture (2 μ M), 2 μ L purified multiple PCR products, 6 μ L ddH₂O mixing. (5) The 0.5 μ L diluted product was mixed with 0.5 μ L Liz500 SIZE STANDARD, 9 μ L Hi-Di, denatured at 95°C for 5 min, then placed on the ABI3730XL sequencer. (6) The raw data collected on the ABI3730XL sequencer are analyzed by GeneMapper 4.1 (Applied Biosystems, USA). Only the subjects with all nine SNP successfully genotyped were included in the final analysis.

Statistical Analysis

All statistical analyses were two-sided and conducted using SPSS (version 23.0; Armonk, NY: IBM Corp, USA). Hardy-Weinberg equilibrium (HWE) of all SNPs was evaluated by Chi-square (χ^2) in normal controls. The differences in genotype, allele frequencies differences of all SNPs between PTB patients and normal controls were compared using the chi-square test (χ^2), and logistic regression analysis was performed to calculate odds ratios (OR), 95% confidence intervals (CI). Haplotype analysis was assessed using SHEsis software,²³ and the associations between all SNPs and PTB risk in two genetic models (dominant and recessive models) were also calculated. $P < 0.05$ was considered to be statistically significant. Bonferroni correction was used for multiple testing in SNP analysis, and $P < 0.0055$ (0.05/9) was considered to be statistically significant.

Results

Subject Characteristics

In this study, the mean ages of 476 PTB patients and 473 controls were 45.09 ± 17.79 years and 43.18 ± 13.68 years, respectively. The PTB group included 163 females and 313 males, while the control group consisted of 268 females and 205 males. In the PTB group, 129 (27.10%) patients had smear-positive sputum, the proportion of other clinical features were pulmonary infection (104, 21.84%), fever (83, 17.44%), drug resistance (73, 15.34%), DILI (68, 14.29%), hypoproteinemia (34, 7.14%), leukopenia (31, 6.51%).

Association of *IL-13*, *IL-13RA1*, and *IL-13RA2* Genes Polymorphisms with PTB Susceptibility

The genotype distributions and allele frequencies of all SNPs in *IL-13*, *IL-13RA1*, and *IL-13RA2* genes are shown in Table 1, and all the SNPs frequencies among in controls were conformed to HWE. The results demonstrated that *IL-13* rs20541 AA genotype frequency appeared to be increased in PTB patients in comparison to normal controls (AA vs GG: $P = 0.017$ after adjustment of sex and age), while the difference was not statistically significant after Bonferroni correction ($P > 0.0055$). In addition, rs20541 variant was found to be associated with PTB risk under recessive mode (AA vs GG+GA: $P = 0.001$ after adjustment of sex and age). However, we did not find any association of rs2066960 and rs2069744 polymorphisms with the predisposition to PTB.

The *IL-13RA1* rs2495636 GA genotype frequency in PTB patients was significantly decreased (GA vs GG: $P = 0.000$ after adjustment of sex and age), while the association between rs2495636 variant and PTB susceptibility under dominant

Table I Genotypes and Alleles Frequencies of *IL-13*, *IL-13RA1*, and *IL-13RA2* Genes in PTB Patients and Normal Controls

SNP	Analyze Model	PTB (N = 476) n (%)	Control (N = 473) n (%)	P value	OR (95% CI)
IL-13					
rs2066960	Genotypes				
	AA	59(12.39)	69(14.59)	0.811	0.951(0.630,1.437) ^a
	CA	226(47.48)	206(43.55)	0.181	1.214(0.914,1.613) ^a
	CC	191(40.13)	198(41.86)		Reference
	Alleles				
	A	344(36.13)	344(36.36)	0.917	0.994(0.882,1.120)
	C	608(63.87)	602(63.64)		Reference
	Dominant model				
	CC	191(40.13)	198(41.86)	0.314	0.871(0.667,1.139) ^a
	CA+AA	285(59.87)	275(58.14)		Reference
rs2069744	Recessive model				
	AA	59(12.39)	69(14.59)	0.438	0.859(0.585,1.262) ^a
	CC+CA	417(87.61)	404(85.41)		Reference
	Genotypes				
rs20541	TT	8(1.68)	4(0.85)	0.370	1.764(0.510,6.105) ^a
	CT	82(17.23)	81(17.12)	0.942	1.013(0.715,1.435) ^a
	CC	386(81.09)	388(82.03)		Reference
	Alleles				
	T	98(10.29)	89(9.41)	0.517	1.094(0.833,1.437)
	C	854(89.71)	857(90.59)		Reference
	Dominant model				
	CC	386(81.09)	388(82.03)	0.774	0.952(0.679,1.335) ^a
	CT+TT	90(18.91)	85(17.97)		Reference
	Recessive model				
rs20541	TT	8(1.68)	4(0.85)	0.372	1.760(0.509,6.082) ^a
	CC+CT	468(98.32)	469(99.15)		Reference
	Genotypes				
	AA	64(13.45)	32(6.77)	0.017	1.784(1.017,2.874) ^a
	GA	188(39.49)	218(46.09)	0.135	0.809(0.612,1.068) ^a
	GG	224(47.06)	223(47.14)		Reference
	Alleles				
	A	316(33.19)	282(29.81)	0.113	1.002(0.888,1.129)
	G	636(66.81)	664(70.19)		Reference
	Dominant model				
GG	224(47.06)	223(47.14)	0.625	1.068(0.820,1.390) ^a	
GA+AA	252(52.94)	250(52.86)		Reference	
rs20541	Recessive model				
	AA	64(13.45)	32(6.77)	0.004	1.972(1.247,3.118) ^a
	GG+GA	412(86.55)	441(93.23)		Reference
IL-13RA1					
rs147857000	Genotypes				
	AA	8(1.68)	5(1.06)	0.685	1.270(0.400,4.029) ^a
	GA	14(2.94)	23(4.86)	0.823	0.924(0.459,1.858) ^a
	GG	454(95.38)	445(94.08)		Reference
	Alleles				
	A	30(3.15)	33(3.49)	0.682	0.900(0.544,1.488)
G	922(96.85)	913(96.51)		Reference	

(Continued)

Table I (Continued).

SNP	Analyze Model	PTB (N = 476) n (%)	Control (N = 473) n (%)	P value	OR (95% CI)
rs141573089	Dominant model				
	GG	454(95.38)	445(94.08)	0.985	0.994(0.548,1.802) ^a
	GA+AA	22(4.62)	28(5.92)		Reference
	Recessive model				
	AA	8(1.68)	5(1.06)	0.683	1.272(0.401,4.038) ^a
	GG+GA	468(98.32)	468(98.94)		Reference
	Genotypes				
	CC	60(12.61)	55(11.63)	0.424	0.847(0.563,1.273) ^a
	CA	57(11.97)	81(17.12)	0.793	0.949(0.639,1.407) ^a
	AA	359(75.42)	337(71.25)		Reference
rs2495636	Alleles				
	C	177(18.59)	191(20.19)	0.379	0.921(0.766,1.106)
	A	775(81.41)	755(79.81)		Reference
	Dominant model				
	AA	359(75.42)	337(71.25)	0.481	1.113(0.826,1.501) ^a
	CA+CC	117(24.58)	136(28.75)		Reference
	Recessive model				
	CC	60(12.61)	55(11.63)	0.438	0.852(0.568,1.227) ^a
	AA+CA	416(87.39)	418(88.37)		Reference
	Genotypes				
AA	184(38.66)	148(31.29)	0.702	0.943(0.700,1.271) ^a	
GA	68(14.28)	151(31.92)	0.000	0.482(0.331,0.699) ^a	
GG	224(47.06)	174(36.79)		Reference	
Alleles					
A	436(45.80)	447(47.25)	0.526	0.969(0.880,1.067)	
G	516(54.20)	499(52.75)		Reference	
Dominant model					
GG	224(47.06)	174(36.79)	0.034	1.336(1.021,1.747) ^a	
GA+AA	252(52.94)	299(63.21)		Reference	
Recessive model					
AA	184(38.66)	148(31.29)	0.272	1.169(0.885,1.545) ^a	
GG+GA	292(61.34)	325(68.71)		Reference	
IL-13RA2					
rs3795175	Genotypes				
	TT	62(13.03)	38(8.03)	0.177	1.354(0.873,2.100) ^a
	CT	45(9.45)	78(16.49)	0.228	0.774(0.511,1.173) ^a
	CC	369(77.52)	357(75.48)		Reference
	Alleles				
	T	169(17.75)	154(16.28)	0.393	1.090(0.894,1.331) ^a
	C	783(82.25)	792(83.72)		Reference
	Dominant model				
	CC	369(77.52)	357(75.48)	0.966	0.993(0.727,1.356) ^a
	CT+TT	107(22.48)	116(24.52)		Reference
Recessive model					
TT	62(13.03)	38(8.03)	0.139	1.391(0.898,2.154) ^a	
CC+CT	414(86.97)	435(91.97)		Reference	
rs638376	Genotypes				
	CC	40(8.41)	41(8.67)	0.296	0.779(0.487,1.245) ^a
	CT	41(8.61)	77(16.28)	0.064	0.669(0.438,1.023) ^a

(Continued)

Table 1 (Continued).

SNP	Analyze Model	PTB (N = 476) n (%)	Control (N = 473) n (%)	P value	OR (95% CI)
rs5946039	TT Alleles	395(82.98)	355(75.05)		Reference
	C	121(12.71)	159(16.81)	0.012	0.756(0.608,0.941)
	T	831(87.29)	787(83.19)		Reference
	Dominant model				
	TT	395(82.98)	355(75.05)	0.045	1.397(1.007,1.939) ^a
	CT+CC	81(17.02)	118(24.95)		Reference
	Recessive model				
	CC	40(8.40)	41(8.67)	0.379	0.810(0.507,1.295) ^a
	TT+CT	436(91.60)	432(91.33)		Reference
	Genotypes				
	AA	18(3.78)	27(5.71)	0.025	0.488(0.260,0.914) ^a
	AT	28(5.88)	49(10.36)	0.280	0.757(0.458,1.254) ^a
	TT	430(90.34)	397(83.93)		Reference
	Alleles				
	A	64(6.72)	103(10.89)	0.001	0.617(0.458,0.832)
	T	888(93.28)	843(89.11)		Reference
	Dominant model				
	TT	430(90.34)	397(83.93)	0.027	1.572(1.052,2.350) ^a
AT+AA	46(9.66)	76(16.07)		Reference	
Recessive model					
AA	18(3.78)	27(5.71)	0.028	0.494(0.263,0.925) ^a	
TT+AT	458(96.22)	446(94.29)		Reference	

Notes: ^aCalculated by non-conditional logistic regression model adjusted for age and sex. P value was corrected by Bonferroni correction (0.05/9), $P < 0.0055$ was considered as statistical significance.

Abbreviations: SNP, single nucleotide polymorphisms; PTB, pulmonary tuberculosis; OR, odds ratios, CI, confidence intervals.

mode was not reached statistically insignificant after Bonferroni correction (GG vs GA+AA: $P > 0.0055$). Regarding *IL-13RA2* gene variations, we found that rs5946039 A allele was related to the lower risk of PTB (A vs T: $P = 0.001$ after adjustment of sex and age). The rs638376 CT genotype, C allele, rs5946039 AA genotype frequencies seemed to be associated with PTB susceptibility; however, no statistically significant association was found after Bonferroni correction ($P > 0.0055$).

Association of *IL-13*, *IL-13RA1*, and *IL-13RA2* Genes SNPs with Clinical Features Among PTB Patients

A case-only analysis was performed to investigate the potential relationship between *IL-13*, *IL-13RA1*, and *IL-13RA2* genes polymorphism and several common clinical features of PTB patients (Table S1). We noted that there was no significant relationship between *IL-13* gene rs2066960, rs2069744, rs20541 and any clinical features in PTB patients.

In *IL-13RA1* gene, the rs141573089 C allele frequency was significantly lower in PTB patients with fever, drug resistance when compared to the patients without these features, respectively ($P = 0.009$, $P = 0.032$, respectively), and the rs141573089 CC genotype frequency was significantly decreased in PTB patients with leukopenia ($P = 0.015$) (Table 2). Furthermore, the AA genotype and A allele frequencies of rs2495636 in PTB patients with drug resistance were both significantly higher than that in PTB patients without drug resistance ($P = 0.049$, $P = 0.001$, respectively), and the rs2495636 A allele frequency was also increased in PTB patients with pulmonary infection ($P = 0.041$).

For the *IL-13RA2* gene, when compared to PTB patients without drug resistance, the increased frequency of rs3795175 T allele, as well as the decreased frequency of rs638376 C allele, was found in PTB patients with drug resistance ($P = 0.029$, $P = 0.009$, respectively).

Table 2 The Positive Findings of Associations Between *IL-13*, *IL-13RA1*, and *IL-13RA2* Genes Polymorphisms and Several Clinical Features of PTB Patients

SNP	Allele	Clinical Features	Group	Genotypes n (%)			P value	Alleles n (%)		P value
	(M/m)			MM	Mm	mm		M	m	
rs141573089	A/C	Fever	+	70	7	6	0.108	147	19	0.009
			-	285	50	53		620	156	
rs141573089	A/C	Drug resistance	+	62	4	7	0.090	128	18	0.032
			-	294	53	53		641	159	
rs141573089	A/C	Leukopenia	+	20	9	2	0.015	49	13	0.625
			-	338	48	58		724	164	
rs2495636	G/A	Drug resistance	+	26	10	37	0.049	62	84	0.001
			-	198	58	144		454	346	
rs2495636	G/A	Pulmonary infection	+	42	16	46	0.283	100	108	0.041
			-	182	52	137		416	326	
rs3795175	C/T	Drug resistance	+	51	9	13	0.223	111	35	0.029
			-	316	36	48		668	132	
rs638376	T/C	Drug resistance	+	67	3	3	0.090	137	9	0.009
			-	325	38	37		688	112	

Note: Part of the study subjects of data missing.

Abbreviations: ±, with/without; M, major alleles; m, minor alleles; SNP, single nucleotide polymorphisms.

Haplotype Analysis

Five main haplotypes (ACA, ACG, CCA, CCG, CTA) for *IL-13*, four main haplotypes (AAG, GAA, GAG, GCG) for *IL-13RA1*, four main haplotypes (CCA, CCT, CTT, TTT) for *IL-13RA2* were detected using SHEsis software. These haplotype frequency distributions between PTB patients and controls are summarized in Table 3.

Table 3 Haplotype Analysis of *IL-13*, *IL-13RA1*, *IL-13RA2* Genes in PTB Patients and Controls

Haplotype	Case [n(%)]	Control [n(%)]	P value	OR (95% CI)
<i>IL-13</i> rs2066960-rs2069744-rs20541				
ACA	39.45(4.1)	30.88(3.3)	0.310	1.282(0.793,2.071)
ACG	299.06(31.4)	306.37(32.4)	0.652	0.956(0.788,1.161)
CCA	181.08(19.0)	163.45(17.3)	0.323	1.125(0.891,1.422)
CCG	334.42(35.1)	356.30(37.7)	0.252	0.896(0.743,1.081)
CTA	89.99(9.5)	81.30(8.6)	0.513	1.111(0.811,1.521)
<i>IL-13RA1</i> rs147857000-rs141573089-rs2495636				
AAG	29.99(3.2)	32.99(3.5)	0.682	0.900(0.544,1.488)
GAA	435.99(45.8)	446.99(47.3)	0.526	0.943(0.788,1.130)
GAG	309.01(32.5)	275.01(29.1)	0.110	1.173(0.965,1.425)
GCG	177.00(18.6)	191.00(20.2)	0.379	0.903(0.719,1.134)
<i>IL-13RA2</i> rs3795175-rs638376-rs5946039				
CCA	63.99(6.7)	102.99(10.9)	0.001	0.591(0.427,0.819)
CCT	54.71(5.7)	55.99(5.9)	0.884	0.972(0.662,1.427)
CTT	664.30(69.8)	633.02(66.9)	0.156	1.151(0.948,1.397)
TTT	166.70(17.5)	153.98(16.3)	0.459	1.095(0.861,1.393)

Notes: Global χ^2 is 848.632, df=7 (frequency<0.03 in both control and case has been dropped). Fisher's p value is <0.001.

Abbreviations: OR, odds ratios; CI, confidence intervals.

The results suggested that the frequency of *IL-13RA2* CCA haplotype was significantly lower in PTB patients than normal controls, while there was no significant difference regarding the *IL-13*, *IL-13RA1* haplotype frequencies between PTB patients and controls.

Discussion

Studies had increasingly focused on the genetic factors that affected the PTB risk, and many new genetic markers had been identified in previous studies.^{24,25} Additionally, since the susceptibility and severity of PTB were determined by complex immune interactions, discussing the role of immune-modulatory gene variants in PTB development was essential. Considering that T cell-derived IL-13 participated in promoting inflammation, regulating cytokine networks, and determining the outcome of PTB patients,²⁶ we analyzed the association between three SNPs in *IL-13* gene and susceptibility to PTB. Meanwhile, our study also examined the association between several SNPs of two IL-13 receptors (*IL-13RA1*, *IL-13RA2*) and PTB susceptibility. Finally, we found the *IL-13* rs20541 variant was associated with PTB susceptibility, and significant associations were also observed for *IL-13RA1* rs2495636, *IL-13RA2* rs5946039 variants with PTB.

Abnormal IL-13 level in PTB patients suggested that IL-13 was closely related to the pathogenesis of PTB. Heitmann et al demonstrated that IL-13/IL-4Ra mediated mechanisms were involved in PTB-associated tissue pathology by regulating the expression level of arginase-1 in an experimental PTB model.¹⁵ SNPs in *IL-13* were also associated with various human diseases, including renal cell carcinoma, colorectal cancer, breast cancer, and bladder cancer.^{27–29} Hence, the potential association between *IL-13* gene variation and PTB susceptibility also attracted more attention. Sun et al explored the association of *IL-13* gene polymorphisms (rs2066960, rs1295686, rs20541, rs2069757 and rs2243248) with PTB susceptibility in a Western Chinese Han population and did not find any associations.¹⁶ Consistent with these results, we observed that the allele and genotype frequencies of rs20541 were not associated with the increased risk of PTB. Meanwhile, genetic model analysis showed that rs20541 was associated with an increased PTB susceptibility in recessive mode, suggesting that rs20541 variant might be involved in the risk of PTB. We selected two other SNPs (rs2066960 and rs2069744) of *IL-13* for genotyping, but no correlation was found. Similarly, the authors found that *IL-13* rs2066960, rs2069744 had no relationship with the risk of renal cell carcinoma in another study.³⁰ Currently, research on the association between *IL-13* and PTB was very limited, and our results contributed to further understanding of the role of *IL-13* gene variation in the pathogenesis of PTB. It was important to note that increased sample sizes and additional SNP might need to be explored.

IL-13 mediated its effects through interacting with its cognate receptors (*IL-13RA1*, *IL-13RA2*) on a variety of biological processes, and the influence of several SNPs in *IL-13RA1*, *IL-13RA2* genes had also been studied in some diseases, such as asthma, atopic dermatitis, and systemic sclerosis.^{19,31,32} The results by Namkung et al identified an association between *IL-13RA1* rs2265753, rs2254672 and atopic dermatitis phenotype.¹⁹ In another study, no significant association was found between *IL-13RA1* +1398 genotype frequencies in asthma, allergic rhinitis patients when compared to in controls.³² Granel et al suggested that *IL-13RA2* rs638376 was related to the risk of systemic sclerosis.³¹ Whereas no studies had been conducted to explore the association between *IL-13RA1*, *IL-13RA2* gene variation and PTB susceptibility, our study was the first to analyze the association between several SNPs in *IL-13RA1*, *IL-13RA2* genes and PTB susceptibility. We found that *IL-13RA1* rs2495636 GA genotype was significantly decreased in PTB patients, suggesting that this SNP might contribute to susceptibility to PTB. Moreover, the results demonstrated that the reduced frequency of rs5946039 A allele existed in PTB patients. In haplotype analysis, *IL-13RA2* CCA haplotype was also proved to be associated with PTB risk. We hypothesized that *IL-13RA1*, *IL-13RA2* gene polymorphism might be involved in the pathogenesis of PTB by modulating IL-13 function, and repetitive studies were necessary to verify our experimental results. Furthermore, the functional role of *IL-13RA1*, *IL-13RA2* genes polymorphism in the development of PTB is worth further investigation.

As a complex disease, PTB usually has multiple clinical manifestations and complications, such as drug resistance, pulmonary infection, fever, hypoproteinemia, and leukopenia, which may greatly impact treatment and rehabilitation. Many variations were found to be associated with clinical features.^{33,34} Song et al found that *lncRNA RP11-37B2.1* rs218916 and rs160441 were associated with thrombocytopenia in PTB patients.³³ Our recent study found a significant association between *CYP27A1* rs933994 T allele frequency and drug resistance in PTB patients.⁴ Similarly, *IL-13RA1* gene rs141573089 polymorphism was found to be related to fever, drug resistance, and leukopenia, while rs2495636 polymorphisms were

closely associated with pulmonary infection and drug resistance in PTB patients. These results indicated that the variants of *IL-13RA1*, *IL-13RA2* contributed to host response to drug treatment and played an active role in the development of appropriate treatment choices for PTB patients. However, the mechanism underlying how they affect the drug treatment response was unclear; hence, more rigorous studies at the molecular genetic level were needed.

Some limitations should be noted in our study. First, this study was conducted in a single center; therefore, the sample size might not be sufficient. Second, the potential influence of some confounding factors, such as environmental factors, treatment regimen, and so on, was not excluded. Finally, we only studied several SNPs within *IL-13*, *IL-13RA1*, *IL-13RA2*, other functional SNPs of these genes warrant further investigation. Further studies with large-scale samples and other ethnic backgrounds were warranted to determine the precise role of *IL-13*, *IL-13RA1*, and *IL-13RA2* gene variations in PTB.

Conclusion

Our study demonstrated that *IL-13* rs20541, *IL-13RA1* rs2495636, *IL-13RA2* rs5946039 polymorphisms might be associated with PTB susceptibility in Chinese population. Moreover, several SNPs in *IL-13RA1* and *IL-13RA2* genes were related to some clinical features among PTB patients, including leukopenia, pulmonary infection, and drug resistance.

Abbreviations

TB, tuberculosis; IL, interleukin; MAF, minor allele frequency; IMLDR, improved multiple ligase detection reaction; HWE, Hardy-Weinberg equilibrium; SNPs, single nucleotide polymorphisms; MTB, Mycobacterium tuberculosis; LD, linkage disequilibrium.

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Author Contributions

All authors made a significant contribution to study design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically; agreed to submit to the current journal; gave final approval for the version to be published; and agreed to be accountable for all aspects of the work.

Hong-Miao Li and Fei Tang should be considered as co-first authors.

Disclosure

The authors declare that they have no conflict of interest.

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