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# ORIGINAL RESEARCH Association between the COMT Vall 58Met polymorphism and risk of cancer: evidence from 99 case-control studies

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Abstract: Catechol-O-methyltransferase (COMT) plays a central role in DNA repair and estrogen-induced carcinogenesis. Many recent epidemiologic studies have investigated the association between the COMT Val158Met polymorphism and cancer risk, but the results are inconclusive. In this study, we performed a meta-analysis to investigate the association between cancer susceptibility and COMT Val158Met in different genetic models. Overall, no significant associations were found between COMT Val158Met polymorphism and cancer risk (homozygote model: odds ratio [OR] = 1.05, 95% confidence interval [CI] = [0.98, 1.13]; heterozygote model: OR =1.01, 95% CI = [0.98, 1.04]; dominant model: OR =1.02, 95% CI [0.97, 1.06], and recessive model: OR =1.03, 95% CI [0.97, 1.09]). In the subgroup analysis of cancer type, COMT Val158Met was significantly associated with increased risks of bladder cancer in recessive model, and esophageal cancer in homozygote model, heterozygote model, and dominant model. Subgroup analyses based on ethnicities, COMT Val158Met was significantly associated with increased risk of cancer in homozygote and recessive model among Asians. In addition, homozygote, recessive, and dominant models were significantly associated with increased cancer risk in the subgroup of allele-specific polymerase chain reaction genotyping. Significant associations were not observed when data were stratified by the source of the controls. In summary, this meta-analysis suggested that COMT Val158Met polymorphism might not be a risk factor for overall cancer risk, but it might be involved in cancer development at least in some ethnic groups (Asian) or some specific cancer types (bladder and esophageal cell cancer). Further evaluations of more preclinical and epidemiological studies are required.

Keywords: COMT, polymorphism, cancer, meta-analysis, susceptibility

#### Introduction

Cancer constitutes an enormous burden on the society in more and less economically developed countries alike.<sup>1,2</sup> Based on GLOBOCAN estimates, ~14.1 million new cancer cases and 8.2 million deaths occurred in 2012 worldwide.<sup>1</sup> According to the development trend, the new cases in 2030 will reach 22.2 million.<sup>2</sup> It is well known that the etiology and development of cancer are as a result of complex interactions between genetic and environmental factors.<sup>3</sup> Genes determine the susceptibility of individual to environment, and environmental factors often damage the DNA in turn. Recent studies have shown that host genetic factors are closely related to the pathophysiology of many human cancers.<sup>4</sup> The most common form of genetic variation, that is, single-nucleotide polymorphisms, is known to contribute individual susceptibility to cancer.<sup>5</sup> Therefore, it is anticipated that the identification of key gene polymorphisms associated with cancer risk is essential for predicting risk of individuals, and that it

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will greatly assist the global control and therapeutic strategies of this lethal disease.

The catechol-O-methyltransferase (COMT) gene is located on chromosome 22q11.2 and consists of six exons.6 It is an important enzyme involved in the inactivation of endogenous catecholamine and catechol estrogens. Catechol estrogens have been shown to have the ability to damage DNA and carcinogenetic potential.7 Therefore, the loss of or changes in COMT is supposed to contribute to genomic instability and tumor genesis. In line with these considerations, it has been hypothesized that COMT Val158Met might influence the development of all cancers. Up to now, many researches have indicated the link between COMT polymorphism and cancer susceptibility. Several polymorphisms have been identified, including the widely studied polymorphism Val158Met(rs4680).8 This change has been associated with a three- to four-fold decrease in the activity of COMT compared with the wild-type COMT-Val allele.9,10 It is biologically reasonable to hypothesize that women who carry mutant COMT-Met allele may have higher cancer risks.

In recent years, many studies have investigated the relationship between COMT Val158Met polymorphism in different races and different types of cancer, but the results were inconclusive or controversial.<sup>11-101</sup> The inconsistent conclusions may be due to a possible minor effect of the polymorphism on cancer or the small sample size in studies with inadequate statistical power of complex traits. Meta-analysis is a powerful statistical tool to pool different studies to overcome deficiencies such as small sample size and to provide more reliable results. Although some previous meta-analyses have reported the association between COMT Val158Met polymorphism and ovarian cancer (up to eight case-control studies included),<sup>102,103</sup> breast cancer (up to 56 case-control studies included),65,104-108 endometrial cancer (up to seven case-control studies included),<sup>103,109,110</sup> prostate cancer (up to six case-control studies included),111-113 and lung cancer (evidence from six case-control studies),<sup>114</sup> only specific cancer types or race populations were included, which led to their limitations. To update the results of previous meta-analyses and to provide a more precise assessment of the association between COMT Val158Met and cancer risk, we performed a comprehensive meta-analysis by including the most recent and relevant articles.

## Materials and methods Identification and eligibility of relevant studies

The meta-analysis was conducted following the criteria of Preferred Reporting Items for Systematic Reviews and

Meta Analyses. A comprehensive literature search was performed using the PubMed, Cochrane Library, Chinese National Knowledge Infrastructure, and EMBASE database for relevant articles published (the last search update was February 15, 2015) with keywords "COMT", "Catechol-O-methyltransferase", "Val158Met", "rs4680", "single nucleotide polymorphism", "polymorphism", "Variant", "Mutation", "Cancer", "tumor", "neoplasm", "malignancy", or "Carcinoma". In addition, studies were identified by a manual search of reviews and retrieved studies. Search results were restricted to human populations, and the articles were written in English or Chinese. We included all the casecontrol studies and cohort studies that have investigated the association between COMT Val158Met polymorphisms and cancer risk with genotyping data. All eligible studies were retrieved, and their bibliographies were checked for other relevant publications. When the same patient population was used in several publications, only the most recent, the largest or the most complete study was included.

#### Assessment of study quality

The quality of the included studies was assessed by the Newcastle–Ottawa Scale (NOS; <u>http://www.ohri.ca/programs/</u> <u>clinical epidemiology/oxford.asp</u>),<sup>115</sup> including selection of groups, comparability of groups, and ascertainment of exposure. The NOS score ranges from 0 to 10 stars. Studies with NOS score > five stars were included in the final analysis.

#### Inclusion criteria

All studies were included if they met the following criteria: 1) only the case-control studies or cohort studies were considered, 2) studies that investigated the COMT Val158Met polymorphism and the risk of cancer susceptibility were included, and 3) the genotype distribution of the polymorphism in cases and controls was described in details, and the results were expressed as odds ratio (OR) and corresponding 95% confidence interval (95% CI). Major reasons for exclusion of studies were as follows: 1) not for cancer research, 2) only case population, 3) duplicate of previous publication, and 4) review articles, editorials, case reports, studies with preliminary results not on COMT Val158Met polymorphism or outcome, and investigations of the role of COMT expression related to disease. Ethics approval for the study was granted by the local institute, the People's Hospital of Three Gorges University Ethics Committee.

#### Data extraction

Using a standardized form, data from published studies were extracted independently by two reviewers to evaluate

their eligibility for inclusion by first screening the title and abstract of each identified reference and then establishing the eligibility of the included papers based on the full text when necessary. For each included study, the following information was collected: first author, year of publication, region, study design, sample size, source of control, genotyping method, allele or genotype frequencies, and evidence of Hardy–Weinberg equilibrium (HWE). Any discrepancy between the two reviewers was resolved by discussion and consultation with a third reviewer.

#### Statistical analysis

ORs and their 95% CIs were used to determine the strength of association between the COMT Val158Met polymorphism and cancer risk. The significance of the pooled OR was determined using the Z test, and P < 0.05was considered statistically significant. Homozygote model (AA vs GG), heterozygote model (GA vs GG), dominant model (GA + AA vs GG), and recessive model (AA vs GG + GA) were investigated. Subgroup analysis was performed by ethnicity, cancer type (if one cancer type contained less than two studies, it was defined as "other"), source of controls, and hospital or population controls. Effective modification by a subgroup was assessed by testing the interaction between genotypes and stratification variables by using logistic regression analyses (random-effects estimator). HWE was tested using the chi-square test among controls, and P < 0.05 was considered a significant departure from HWE. If the *P*-value for heterogeneity was >0.05 and  $I^2 < 50\%$ , indicating an absence of heterogeneity among studies, the fixed-effects model (the Mantel-Haenszel method) was used.<sup>116</sup> In contrast, if either the P-value for heterogeneity was  $\leq 0.05$  or  $I^2$  was  $\geq 50\%$ , indicating heterogeneity among the studies, the more appropriate random-effects model (the DerSimonian and Laird method) was used.<sup>117</sup> Sensitivity analyses were performed to assess the stability of the results. Begg's funnel plots were used to diagnose potential publication bias, and P < 0.05 was used to indicate possible publication bias.<sup>118</sup> All analyses were performed using RevMan 5.3 (updated in March 2012 by the Cochrane Collaboration). P-values were based on two-sided tests.

#### Results

# Literature search and meta-analysis databases

Following the searching strategy, 337 potentially relevant studies were retrieved. After title and abstract screening, nine of them were ruled out because of repeated data. A total of 202 irrelevance articles were excluded. In addition, after the

full texts of the remaining 182 articles were read, 90 articles were excluded for the following reasons: article was a review (n=27), articles had insufficient data (n=13), articles were not related to cancer (n=34), and articles were not related to COMT (n=16). A total of 92 publications with full text were selected and were subjected to further examination. Because seven studies included more than one ethnicity, genotype method, control source, or tumor type and were performed by the same author, we treated them separately in this metaanalysis. Of those, 99 case-control studies with 43,085 cancer cases and 57,882 control subjects were included in our metaanalysis. A flow chart showing the detailed steps of study selection is shown in Figure 1. All studies were case-control studies with the following tumor-type distribution: three were conducted for bladder cancer, two for renal cancer, nine for endometrial cancer, eight for ovarian cancer, 62 for breast cancer, six for lung cancer, three for liver cancer, two for colon cancer, two for esophageal cell cancer, one for thyroid cancer and non-Hodgkin lymphoma, and one for testicular germ cell tumor. Fifty studies investigated the risks in Caucasian populations, 35 studies investigated Asian populations, ten studies investigated mixed populations, and the remaining studies were conducted in African populations. Five main genotyping methods were used such as polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP), TaqMan, sequencing, matrix-assisted laser desorption ionization time of flight mass spectrometry (MALDI-TOF), and allele-specific PCR (AS-PCR). By source of controls, 50 studies were population based, 45 studies were hospital based, and four studies were not clear. The distribution of the genotypes in the control subjects was in agreement with HWE, except for eight studies.<sup>34,37,70,72,80,88,95,119</sup> The quality assessment showed that the quality scores ranged from 5 to 9 with a median score of 6, suggesting that all studies were of high quality. The main characteristics of the eligible studies are listed in Table 1.

#### Quantitative synthesis

Overall, no significant associations between COMT Val158Met and cancer risk were found using homozygote model (OR =1.05, 95% CI [0.98, 1.13]), heterozygote model (OR =1.01, 95% CI [0.98, 1.04]), dominant model (OR =1.02, 95% CI [0.97, 1.06]), or recessive model (OR =1.03, 95% CI [0.97, 1.09]).

Significant heterogeneity was observed among the 99 studies on COMT Val158Met polymorphism. To explore the source of heterogeneity, we performed stratified analyses on ethnicity, cancer type, source of controls, and genotyping method. In the subgroup analysis on cancer type, COMT



Figure I Flow chart of publication selection.

**Note:** A total of 99 studies were included in this meta-analysis and systematically reviewed after a comprehensive study selection. **Abbreviation:** COMT, catechol-0-methyltransferase.

Val158Met was significantly associated with an increased risk of bladder cancer in recessive model (OR =1.30, 95% CI [1.02, 1.66]), esophageal cell cancer in homozygote model (OR =1.77, 95% CI [1.07, 2.93]), heterozygote model (OR =1.40, 95% CI [1.01, 1.92]), and dominant model (OR =1.46, 95% CI [1.08, 1.98]). However, studies on renal, endometrial, lung, liver, ovarian, colon, and other cancer types have suggested null association (OR =0.70-1.46; Table 2). These studies were further stratified on the basis of ethnicities, and the results showed that COMT Val158-Met polymorphism may be a risk factor for cancer in Asian populations in the homozygote model (OR =1.25, 95% CI [1.03, 1.51]) and recessive model (OR =1.20, 95% CI [1.01, 1.43]). We failed to detect any association between the COMT Val158Met polymorphism and African, Caucasian, and mixed populations. In addition, homozygote models (OR =3.46, 95% CI [2.07, 5.80]), recessive models (OR =3.32, 95% CI [2.02, 5.44]), and dominant models (OR =1.54, 95% CI [1.12, 2.11]) were significantly associated

2794 submit your manuscript | www.dovepress.com Dovepress with increased cancer risk in the subgroup of AS-PCR genotyping method, but no significant associations were observed when PCR-RFLP, TaqMan, sequencing, MALDI-TOF, and other genotyping method were used. No significant associations were detected when the studies were stratified on the basis of the source of control subjects.

#### Test of heterogeneity and sensitivity

Heterogeneity among studies was observed in the overall comparisons as well as in the subgroup analyses. The source of heterogeneity was investigated by cancer ethnicity (European, Asian, African, and mixed; P=0.483), cancer types (bladder, breast, renal, endometrial, lung, liver, ovarian, colon, and other cancer types; P=0.684), control source (population based, hospital based, and family based; P=0.659), and genotyping method (AS-PCR, PCR-RFLP, TaqMan, sequencing, MALDI-TOF, and other genotyping method; P=0.647) using meta-regression, but no covariables were found to contribute to the heterogeneity.

InitialImitedLavigne et allLavigne et all1997USACaucasianMillikan et all21998USACaucasianMillikan et all21998USACaucasianThompson et all3Thompson et all31999People's RepublicAfricanMillikan et all21998USACaucasianGoodman et all31999People's RepublicAsianGoodman et all62000USACaucasianGoodman et all62001USACaucasianGoodman et all72001USACaucasianMirrunen et all82001USACaucasianMirrunen et all82001USACaucasianMirrunen et all82001SwedenCaucasianMirrunen et all82001USACaucasianMirrunen et all82001SwedenCaucasianMirrunen et all82003USACaucasianMirrunen et all82003USACaucasianMirrunen et all82003USACaucasianMirrunen et all82003USACaucasianMirrunen et all82003USACaucasianMirrunen et all82003USACaucasianMirrunen et all82003USACaucasianMirrune et all82003USACaucasianMirrune et all82003USACaucasianMirrune et all82003USACaucasianMirrune et all82003USACaucasian <t< th=""><th></th><th>type source source Breast HB Breast HB Breast PB Breast PB Breast PB Breast HB Covarian HB Covarian HB Breast HB Breast HB Breast HB Breast PB Breast PB Breast PB Breast PB PB</th><th>method PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP</th><th>35 <b>AA</b></th><th>BG 1</th><th>ម្ល</th><th>¥,</th><th><b>AG</b> 56</th><th><b>GG</b></th><th></th><th>score</th></t<>		type source source Breast HB Breast HB Breast PB Breast PB Breast PB Breast HB Covarian HB Covarian HB Breast HB Breast HB Breast HB Breast PB Breast PB Breast PB Breast PB	method PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP	35 <b>AA</b>	BG 1	ម្ល	¥,	<b>AG</b> 56	<b>GG</b>		score
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1998 USA   1998 USA   1998 USA   1999 People's Republic   2000 Germany   2001 USA   2001 Japan   2001 Finland   2001 Finland   2002 Urkey   2003 USA   2003 USA   2003 USA   2003 USA   2003 USA   2003 USA   2004 UK   2003 USA   2004 UK   2003 USA   2004 UK   2004 UK   2004 UK   2004 UK   2004 UK   2004 UK   2004 U	ian ian an a		PCR-RFLP PCR-RFLP PCR-RFLP PCR-RFLP	ç	57	21	- 1		i	0.862	9
1998 USA   1999 People's Republic   1999 People's Republic   2000 Germany   2001 USA   2001 Finland   2001 Finland   2002 USA   2003 USA   2004 UK   2003 USA   2004 UK   2003 USA   2004 UK   2004 UK   2004 UK   2004 UK   2004 UK   2004 UK   2004	ian ian an a	5 5	PCR-RFLP PCR-RFLP PCR-RFLP	29	901	130	34	118	Ξ	0.838	8
1998 USA   1999 People's Republic   2000 Germany   2001 USA   2001 Finland   2001 Finland   2001 Korea   2002 Urkey   2003 Italy   2003 Italy   2003 Italy   2003 USA   2003 USA   2003 USA   2003 Vechen   2003 UK   2003 UK   2004	ian		PCR-RFLP PCR-RFLP	102	184	103	105	188	86	0.916	œ
1999 People's Republic of China 2000 Germany 2001 USA 2001 USA 2001 USA 2001 USA 2001 Japan 2001 Finland 2001 Korea 2003 USA 2003 Italy 2003 Italy 2003 USA 2003 Of China 2004 USA 2004 UK 2004 UK 2004 UK	ian ian ian	5 5	PCR-RFLP	53	159	69	72	139	78	0.522	7
of China 2000 Germany 2001 USA 2001 USA 2001 USA 2001 USA 2001 Japan 2001 Finland 2001 Korea 2003 USA 2003 Italy 2003 Italy 2003 USA 2003 USA 2003 Italy 2004 UK 2004 UK 2004 UK 2004 UK 2004 UK	ilan ilan ilan	5 5		13	37	68	4	55	99	0.612	S
2000 Germany 2001 USA 2001 USA 2001 USA 2001 USA 2001 Japan 2001 Finland 2001 Korea 2002 USA 2003 Turkey 2003 Italy 2003 Italy 2003 Italy 2004 USA 2004 UK 2004 UK 2004 UK 2004 UK	ilan ilan ilan ilan										
2001 USA 2001 USA 2001 USA 2001 Japan 2001 Finland 2001 Korea 2002 Turkey 2003 Italy 2003 Italy 2003 Sweden 2004 USA 2004 UK 2004 UK 2004 USA 2004 UK	ilan ilan ilan	E	PCK-KFLF	27	54	27	29	52	25	0.905	7
2001 USA 2001 Japan 2001 Japan 2001 Sweden 2001 Finland 2001 Korea 2002 USA 2003 USA 2003 Italy 2003 Italy 2003 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA	ilan ilan ilan		PCR-RFLP	16	57	52	61	57	68	0.827	œ
rom and Wingren <sup>19</sup> 2001 Japan 2001 Sweden 2001 Finland 2001 Korea 2002 USA 2003 USA 2003 Italy 2003 Italy 2003 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA	sian sian sian sian		PCR-RFLP	35	57	20	31	55	27	0.788	œ
strom and Wingren <sup>19</sup> 2001 Sweden 2001 Finland 2001 Korea 2002 USA 2003 USA 2003 Italy 2003 Italy 2003 China 2003 China 2003 USA 2004 USA 2004 UK 2004 UK 2004 UK	sian sian sian sian		PCR-RFLP	81	72	60	23	63	79	0.079	9
2001 Finland 2001 Korea 2002 USA 2002 Turkey 2003 USA 2003 Raly 2003 Raly 2003 China 2003 China 2003 USA 2004 USA 2004 UK 2004 UK 2004 USA 2004 UK	sian sian sian		PCR-RFLP	46	64	16	43	61	13	0.209	ъ
2001 Korea 2002 USA 2003 USA 2003 USA 2003 Italy 2003 Raly 2003 China 2003 Sweden 2003 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA	sian sian sian		PCR-RFLP	128	238	115	143	237	001	0.921	S
2002 USA 2002 Turkey 2003 USA 2003 Italy 2003 People's Republic 2003 USA 2004 USA 2004 UK 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA 2004 USA	sian sian	st HB	PCR-RFLP	m	79	8	16	46	101	0.004	9
2002 Turkey 2003 USA 2003 Italy 2003 People's Republic 2003 Sweden 2004 USA 2004 UK 2004 Austria 2004 USA 2004 UK	sian sian	Ovarian PB	PCR-RFLP	48	103	59	54	611	52	0.861	9
2003 USA 2003 Italy 2003 Raly 2003 People's Republic 2003 USA 2004 USA 2004 UK 2004 Austria 2004 USA 2004 USA 2004 USA	sian	ist HB	PCR-RFLP	4	42	28	13	55	35	0.227	7
2003 Italy 2003 People's Republic of China 2003 Sweden 2003 USA 2004 UK 2004 UK 2004 Austria 2004 USA 2004 USA	sian	ist PB	PCR-RFLP	12	24	31	38	78	29	0.335	9
2003 People's Republic of China 2003 Sweden 2003 USA 2004 UK 2004 UK 2004 Austria 2004 USA 2004 USA		г HB	PCR-RFLP	15	56	16	23	51	16	P>0.05	9
of China 2003 Sweden 2003 USA 2004 USA 2004 UK 2004 Austria 2004 USA 2004 USA		ist HB	PCR-RFLP	26	103	121	13	105	132	0.174	œ
2003 Sweden 2003 USA 2004 USA 2004 UK 2004 Austria 2004 USA 2004 USA											
2003 USA 2004 USA 2004 UK 2004 Austria 2004 USA 2004 USA	casian Breast	ist PB	DASH	442	767	281	433	662	245	0.772	9
2004 USA 2004 UK 2004 Austria 2004 France 2004 USA	n Breast	ist PB	TaqMan	48	213	328	51	229	282	0.646	9
2004 UK 2004 Austria 2004 France 2004 USA		ist FB		73	156	84	60	44	58	0.108	9
2004 Austria 2004 France 2004 USA	Caucasian Breast	ist PB	TaqMan	845	1,360	645	534	926	448	0.232	œ
2004 France 2004 USA 2004 Turkov	Caucasian Breast	st PB	Sequencing	98	192	101	478	835	385	0.577	œ
al <sup>33</sup> 2004 USA 2004 Turlovi	Caucasian Blad	Bladder HB	PCR-RFLP	43	96	62	43	114	57	P>0.05	7
2004 Turkey		Endometrial HB	PCR-RFLP	55	105	55	172	308	161	0.874	7
	Caucasian Breast	st PB	PCR-RFLP	28	69	33	16	146	62	0	9
2004 People's Republic	n Liver	ΗB	PCR-RFLP	30	21	m	49	31	9	AA	7
of China											
1 <sup>36</sup> 2004 Russia	Caucasian End	etrial	PCR-RFLP	30	65	29	44	73	23	0.996	9
Cheng et al <sup>37</sup> 2005 People's Republic Asian	n Breast	st HB	NR	35	197	237	58	262	420	0.006	9
of China											
38 2005 USA		etrial	PCR-RFLP	8	174	67	123	207	90	0.953	9
al <sup>39</sup> 2005 Austria	Caucasian Colon		PCR-RFLP	0	58ª	8	0	519ª	203	AN	9
Lin et al <sup>40</sup> 2005 People's Republic Asian	n Breast	st PB	PCR-RFLP	ъ	31	51	81	133	190	0.393	9
Lin et al <sup>41</sup> 2005 People's Republic Asian of China	n Breast	ist PB	PCR-RFLP	6	35	58	23	138	205	0.972	9
Le Marchand et al <sup>42</sup> 2005 USA Mixed	ed Breast	st PB	PCR-RFLP	196	624	519	206	614	550	0.109	7
2005 USA	ian	st PB	TaqMan	77	124	49	1,104	1,943	903	0.391	œ
2005 USA		- -	PCR-RFLP	119	224	011	147	269	127	0.903	7
2005 USA			PCR-RFLP	0	I 7ª	61	0	30ª	23	0.059	9

Table I (Continued)														
Authors	Year	Country	Ethnicity	Cancer	Control	Genotype	Gend	Genotype (cases)	ases)	Geno	Genotype (controls)	ntrols)	HWE	NOS
			mixed	type	source	method	AA	AG	U U	AA	AG	U U		score
Skibola et al <sup>45</sup>	2005	NSA	Caucasian	NHL	PB	TaqMan	77	153	75	163	323	193	P>0.05	7
Wen et al <sup>46</sup>	2005	People's Republic	Asian	Breast	B	PCR-RFLP	83	425	612	93	470	628	0.698	7
Chang et al <sup>47</sup>	2006	People's Republic	Asian	Breast	ВН	PCR-RFLP	6	77	103	30	159	132	0.068	7
ı		of China												
Gallicchio et al <sup>48</sup>	2006	NSA	Caucasian	Breast	PB	TaqMan	24	41	16	371	608	272	0.44	6
Gaudet et al <sup>49</sup>	2006	NSA	Caucasian	Breast	BB	MALDI-TOF	240	521	287	266	549	277	0.853	œ
Gaudet et al <sup>49</sup>	2006	Poland	Caucasian	Breast	PB	TaqMan	439	993	551	539	1,123	617	0.525	œ
Onay et al <sup>50</sup>	2006	Canada	Caucasian	Breast	BB	TaqMan	94	202	102	96	196	80	0.283	œ
Song et al <sup>51</sup>	2006	People's Republic	Asian	Breast	NR	PCR-RFLP	m	4	66	=	36	65	0.09	2
:		of China												
Tao et al <sup>52</sup>	2006	People's Republic	Asian	Endometrial	ΗB	TaqMan	85	383	563	67	425	534	0.683	9
		of China												
Akisik and Dalay <sup>53</sup>	2007	Turkey	Caucasian	Breast	NR	PCR-RFLP	26	59	29	21	53	34	0.966	9
Fan et al <sup>99</sup>	2007	People's Republic	Asian	Breast	면	PCR-RFLP	29	75	96	ъ	44	51	0.25	9
		of China												
Gemignani et al <sup>54</sup>	2007	European	Caucasian	Lung	면	PCR-RFLP	59	<u>+</u>	83	75	I 46	81	0.569	7
Holt et al <sup>55</sup>	2007	NSA	Caucasian	Ovarian	PB	TaqMan	79	129	72	137	209	104	0.948	œ
Holt et al <sup>55</sup>	2007	NSA	African	Ovarian	PB	TaqMan	0	61	4	16	58	52	0.2	œ
Hu et al <sup>57</sup>	2007	People's Republic	Asian	Breast	면	Sequencing	=	36	65	m	41	99	0.252	9
		of China												
Liu et al <sup>119</sup>	2007	People's Republic	Asian	Endometrial	면	PCR-RFLP	ъ	33	42	m	46	35	0.01	9
		of China												
Ralph et al <sup>56</sup>	2007	NSA	Caucasian	Breast	몀	TaqMan	405	825	396	006	1,631	755	0.758	7
Szyllo et al <sup>s8</sup>	2007	Poland	Caucasian	Endometrial	HB	PCR-RFLP	24	8	46	39	011	48	0.253	6
Takata et al <sup>59</sup>	2007	NSA	Mixed	Breast	PB	PCR-RFLP	89	257	229	47	108	95	0.104	8
Tanaka et al <sup>60</sup>	2007	Japan	Asian	Renal	PB	Sequencing	0	54	59	=	61	85	AA	œ
Zhao et al <sup>61</sup>	2007	People's Republic	Asian	Endometrial	HB	PCR-RFLP	16	77	39	œ	50	52	0.779	9
		of China												
Delort et al <sup>62</sup>	2008	France	Caucasian	Ovarian	PB	TaqMan	8	22	=	283	480	237	0.916	7
Hirata et al <sup>63</sup>	2008	NSA	Caucasian	Endometrial	PB	PCR-RFLP	37	8	32	27	90	48	0.277	œ
Justenhoven et al <sup>64</sup>	2008	Germany	Caucasian	Breast	PB	MALDI-TOF	I 45	298	163	147	305	170	0.654	8
Onay et al <sup>65</sup>	2008	Canada	Caucasian	Breast	PB	TaqMan	273	642	302	201	353	160	0.832	8
Onay et al <sup>65</sup>	2008	Finland	Caucasian	Breast	PB	TaqMan	206	361	4	168	267	114	0.676	7
Yuan et al <sup>66</sup>	2008	People's Republic	Asian	Liver	몀	PCR-RFLP	8	144	258	32	157	286	P>0.05	9
		of China												
Zhu <sup>100</sup>	2008	People's Republic	Asian	Esophageal	ΈB	PCR-RFLP	16	51	23	0	37	30	P>0.05	5
	0000	of China					0	ç		G				G
Zienolddiny et al%	2000	Norway	Caucasian	Lung .	ድ የ	Sequencing	32	79	163	20	09 į	707	0.182	× 0
Cote et al <sup>w</sup>	2000	USA	African	Lung .	ድ	TaqMan	0 .	46 201	2.6 1	4	4/	59	0.332	× 0
Cote et al <sup>66</sup>	2009	USA -	Caucasian	Lung	82 5	PCR-RFLP	102	205	8/	4	197	7.6	0.696	∞ •
Fontana et al <sup>69</sup>	2009	France	Caucasian	Bladder	HB	TaqMan	4	28	6	0	24	=	AA A	9
He et al <sup>/1</sup>	2009	NSA	Caucasian	Breast	원	TaqMan	334	607	1/7	446	83/	400	0.85	7

Reding et al <sup>73</sup> Sangrajrang et al <sup>74</sup> Shrubsole et al <sup>75</sup>	2009 2009 2009	USA Thailand People's Republic	Caucasian Asian Asian	breast Breast Breast	2 9 2	i aqirlari TaqMan PCR-RFLP	42 0	427 233 497ª	290 296 596	0 30 0	190 554ª	266 266 615	0.61 NA	0 ~ ~
Yadav et al <sup>76</sup> Zhou <sup>98</sup>	2009 2009	or Crima India People's Republic of China	Asian Asian	Breast Colon	8 8	PCR-RFLP SNPlex	28 23	82 121	44 208	29 38	85 262	52 327	0.57 P>0.05	~ ~
Delort et al <sup>77</sup> Ferlin et al <sup>78</sup> MARIE-GENICA Consortium on Genetic	2010 2010 2010	France Italy Germany	Caucasian Caucasian Caucasian	Breast TGCT Breast	888	TaqMan PCR-RFLP MALDI-TOF	254 0 844	455 200 1,569	201 34 731	283 2 1,569	480 182 2,669	237 34 1,243	0.23 P>0.05 0.094	8 7 8
Susceptibility for Menopausal Hormone Therapy Related Breast Cancer Risk <sup>70</sup> Jakubowska et al <sup>72</sup> Li et al <sup>97</sup>	2010 2010	Poland People's Republic	Caucasian Asian	Breast Endometrial	위 위	PCR-RFLP PCR-RFLP	6 84	164 26	17 17	8 54	168 35	68 71	0.01	ωı
Martínez et al <sup>ioi</sup> Moreno-Galvan et al <sup>79</sup>	2013 2010	Mexico Mexico	Caucasian Caucasian	Breast Breast	9 원	PCR-RFLP PCR-RFLP	32 12	66 42	52 37	23 14	59 42	68 38	0.085 0.669	6 7
Peterson et a <sup>180</sup> Syamala et a <sup>181</sup> Svamala et a <sup>181</sup>	2010 2010 2010	USA India India	Caucasian Asian Asian	Breast Breast Breast	28 99 89	TaqMan PCR-RFLP PCR-RFLP	420 41 28	794 104 64	370 74 48	403 65 65	665 164 164	348 138 138	0.026 0.183 0.183	8 9 9
Wang et al <sup>82</sup> Xu et al <sup>96</sup>	2010 2010	People's Republic of China People's Republic	Asian Asian	Breast Breast	8 8	AS-PCR AS-PCR	34 38	62 42	80 60	4 0	66 44	96 68	0.58 0.45	~ ~
Cerne et al <sup>83</sup> Cribb et a <sup>184</sup> Huang et al <sup>85</sup>	2011 2011 2011	of China Slovenia Canada People's Republic	Caucasian Caucasian Asian	Breast Breast Esophageal	<u></u>	TaqMan PCR-RFLP PCR-RFLP	144 51 25	263 108 95	123 48 90	67 155 30	136 326 146	67 140 180	0.903 0.208 NA	6 7 7
Lajin et a <sup>186</sup> Naushad et a <sup>187</sup> dos Santos et a <sup>188</sup> Wang et al <sup>89</sup>	2013 2011 2011 2011	of China Syria India Brazil People's Republic	Mixed Asian Mixed Asian	Breast Breast Breast Breast	8888	PCR-RFLP PCR-RFLP PCR-RFLP Sequencing	31 66 68	70 154 41ª 145	34 122 21 187	30 36 36	54 107 26ª 156	28 120 36 208	0.887 0.201 - 0.389	N 9 N N
Heck et al <sup>%</sup> Lim et al <sup>91</sup> Wolpert et al <sup>92</sup>	2012 2012 2012	of China USA Singapore Egypt	Mixed Asian Mixed	Renal Lung Bladder	뿜 뿜 뽑	Sequencing PCR-RFLP TaqMan	0 39 160	632ª 220 245	242 284 110	0 63 95	I,496ª 353 180	557 549 114	0.36 0.539 P>0.05	8 ~ 8
Zhang et al <sup>93</sup> Ghisari et al <sup>94</sup> Son et al <sup>95</sup>	2013 2014 2015	People's Republic of China Denmark Korea	Asian Caucasian Asian	Lung Breast Breast	뿜 뽑 뿜	Sequencing TaqMan Assay	0	69 11 423 <sup>a</sup>	120 7 427	0 4 19	78 53 212ª	103 19 178	0.454 P>0.05 0.008	8 9 7
Note: "Number of patients with the AA + GA genotype in the case and control groups. Abbreviations: HWE. Hardy-Weinberg equilibrium; NOS, Newcastle-Ottawa Scale; HB, hospital based; PCR-RFLP, polymerase chain reaction-restriction fragment length polymorphism; PB, population based; DASH, dynamic allele-specific hybridization; FB, family based; NA, not available; MALDI-TOF, matrix-assisted laser desorption ionization time of flight mass spectrometry; NHL, non-Hodgkin lymphoma; TGCT, testicular germ cell tumor; AS-PCR, allele-	genotype in ibrium; NO t available;	the case and control gro S, Newcastle-Ottawa Sco MALDI-TOF, matrix-assi	ups. ale; HB, hospita sted laser deso	l based; PCR-RFL rption ionization	P, polymeras time of flight	e chain reaction-resti mass spectrometry;	iction fra NHL, no	gment len n-Hodgkir	gth polyn Iymphor	iorphism; 1a; TGCT	PB, popula: , testicular	cion based; germ cell	DASH, dyna tumor: AS-P	mic allele CR. allele

Variables	No of	Homozygote mo	del	Heterozygote mo	del	<b>Recessive model</b>		Dominant model	
	studies	OR (95% CI)	<b>1</b> <sup>2</sup> %	OR (95% CI)	<b>1</b> <sup>2</sup> %	OR (95% CI)	<b>1</b> 2%	OR (95% CI)	<b>1</b> <sup>2</sup> %
Total	99	1.05 (0.98, 1.13)	56	1.01 (0.97, 1.05)	29	1.03 (0.97, 1.09)	51	1.02 (0.97, 1.06)	44
Cancer type									
Bladder	3	1.38 (0.86, 2.21)	45	1.12 (0.71, 1.77)	57	1.30 (1.02, 1.66)	0	1.20 (0.74, 1.94)	65
Renal	2	1.31 (0.52, 3.28)	-	1.28 (0.78, 2.09)	-	1.18 (0.48, 2.86)	_	1.02 (0.83, 1.25)	12
Breast	62	1.04 (0.96, 1.13)	58	1.01 (0.96, 1.05)	21	1.03 (0.96, 1.10)	57	1.01 (0.96, 1.06)	40
Endometrial	9	0.99 (0.73, 1.35)	55	0.90 (0.73, 1.11)	52	1.03 (0.84, 1.26)	29	0.91 (0.73, 1.13)	61
Lung	6	1.09 (0.68, 1.75)	76	1.11 (0.96, 1.28)	2	1.04 (0.67, 1.57)	74	1.09 (0.87, 1.36)	60
Liver	3	0.68 (0.42, 1.09)	0	1.03 (0.80, 1.34)	0	0.70 (0.48, 1.03)	0	0.96 (0.75, 1.23)	0
Ovarian	8	1.05 (0.75, 1.47)	52	1.01 (0.80, 1.28)	33	1.02 (0.84, 1.24)	20	1.00 (0.79, 1.27)	43
Colon	2	0.95 (0.55, 1.64)	-	0.73 (0.55, 0.96)	-	1.08 (0.64, 1.85)	_	0.92 (0.56, 1.50)	63
Esophageal	2	1.77 (1.07, 2.93)	0	1.40 (1.01, 1.92)	0	1.46 (0.92, 2.34)	0	1.46 (1.08, 1.98)	0
Other	2	0.96 (0.29, 3.16)	24	1.18 (0.90, 1.56)	0	0.87 (0.29, 2.62)	21	1.18 (0.91, 1.54)	0
Ethnicities									
African	4	1.46 (0.43, 4.99)	83	1.23 (0.61, 2.49)	75	1.17 (0.53, 2.56)	69	1.09 (0.60, 1.98)	73
Caucasian	50	0.98 (0.91, 1.05)	43	1.00 (0.96, 1.05)	88	0.97 (0.92, 1.03)	38	0.99 (0.95, 1.04)	16
Asian	35	1.25 (1.03, 1.51)	62	1.04 (0.94, 1.14)	53	1.20 (1.01, 1.43)	60	1.06 (0.97, 1.15)	59
Mixed	10	0.96 (0.78, 1.20)	49	1.00 (0.86, 1.17)	38	0.99 (0.87, 1.13)	5	1.03 (0.88, 1.20)	58
Controls source									
PB	50	1.03 (0.94, 1.13)	63	0.99 (0.94, 1.04)	24	1.06 (0.95, 1.17)	58	1.01 (0.95, 1.07)	49
HB	45	1.09 (0.96, 1.24)	48	1.04 (0.96, 1.12)	36	1.02 (0.94, 1.09)	43	1.04 (0.96, 1.11)	41
Other	4	0.95 (0.59, 1.54)	48	1.00 (0.78, 1.27)	4	1.00 (0.69, 1.46)	38	0.99 (0.78, 1.26)	7
Genotyping meth	od								
PCR-RFLP	58	1.02 (0.91, 1.15)	49	1.01 (0.94, 1.09)	36	1.01 (0.92, 1.11)	42	1.02 (0.95, 1.09)	44
TaqMan	24	1.03 (0.94, 1.13)	46	1.02 (0.96, 1.08)	15	1.00 (0.93, 1.07)	35	1.02 (0.95, 1.08)	34
Sequencing	6	1.55 (0.79, 3.03)	85	0.98 (0.84, 1.14)	I.	1.55 (0.84, 2.86)	84	1.09 (0.84, 1.41)	67
MALDI-TOF	3	0.92 (0.83, 1.02)	0	0.98 (0.90, 1.08)	0	0.93 (0.85, 1.01)	0	0.96 (0.88, 1.05)	0
AS-PCR	2	3.46 (2.07, 5.80)	0	1.11 (0.78, 1.57)	0	3.32 (2.02, 5.44)	0	1.54 (1.12, 2.11)	0
Other	6	0.91 (0.77, 1.08)	0	0.94 (0.72, 1.24)	76	0.92 (0.80, 1.05)	0	0.93 (0.81, 1.08)	57

Notes: The bold values indicate that the results are statistically significant.

Abbreviations: COMT, catechol-0-methyltransferase; OR, odds ratio; Cl, confidence interval; PB, population based; HB, hospital based; PCR-RFLP, polymerase chain reaction-restriction fragment length polymorphism; MALDI-TOF, matrix-assisted laser desorption ionization time of flight mass spectrometry; AS-PCR, allele-specific PCR; *I*<sup>2</sup>, variation in OR attributable to heterogeneity.

Sensitivity analysis was conducted to verify the effect of each study on the overall OR by repeating the meta-analysis, but one study was omitted each time. When sensitivity analyses were performed without HWE violating studies, all the results were not materially altered. The results showed that the pooled



Figure 2 Begg's funnel plot of the meta-analysis of cancer risk and COMT Val I58Met polymorphism (AA + AG vs GG).

Note: Begg's funnel plot with pseudo 95% confidence limits.

Abbreviations: COMT, catechol-0-methyltransferase; OR, odds ratio; SE, standard error.

ORs of these three polymorphisms were not materially altered by the contribution of any individual study, thus confirming that the results of this meta-analysis were statistically robust.

#### **Publication bias**

Begg's funnel plot and Egger's test were performed to evaluate the publication bias of the studies. The shape of the funnel plots showed that the dots were almost symmetrically distributed and were predominantly in 95% confidence limits (dominant model, Figure 2). The results of Egger's test statistically confirmed the absence of publication bias in the dominant model (t=1.68, P=0.096).

#### Discussion

In the past several years, interest in the genetic susceptibility to cancers has drawn increased attention to the studies on polymorphisms of genes involved in tumor genesis. Genome-wide association study, also known as whole genome association study, is widely used in the study of genetic epidemiology. At present, >1,369 susceptibility loci associated with cancer risk have been identified by

genome-wide association study, but none of these studies had reported significant associations between cancer susceptibility and COMT Val158Met polymorphisms. We searched the manufacturers' websites (http://www.affymetrix.com/index.affx and http://www.illumina.com)120 and the relevant PubMed databases (Probe, Database of Genotypes and Phenotypes, and Gene Expression Omnibus DataSets) and found that the COMT Val158Met polymorphism was not included in the platforms commonly used in genome-wide association studies. But since the identification of COMT Val158Met polymorphism, the role of COMT Val158Met in cancers risk has been reported in an increasing number of studies, but the results remained controversial. Some recent meta-analyses studies reported such an association only for single cancer or specific populations. Importantly, several published studies were not included in the previous metaanalysis, and additional original studies with larger sample sizes have been published since then. Hence, the association between the COMT Val158Met polymorphism and the risk of cancer remains unknown. Therefore, meta-analysis can provide a quantitative summary of the available data supporting the association between COMT Val158Met and cancer risk. Compared with some previous meta-analyses, strengths of our meta-analysis include the large sample size and high statistical power of the analysis based on substantial number of cases and controls from differential studies, which minimized selection bias and led to relatively stable risk estimation.

In the current meta-analysis, 99 case-control studies with 43,085 cancer cases and 57,882 control subjects were considered. The results indicated no significant association between COMT Val158Met polymorphism and overall cancer risk in any genetic comparison model tested. In further subgroup analysis by cancer type, COMT Val158Met was significantly associated with an increased risk of bladder cancer and esophageal cancer in some specific genetic models. However, studies on renal, endometrial, lung, liver, ovarian, colon cancers, and other cancer types have suggested null associations. In line with most previous meta-analyses for single cancer, Zhang et al,<sup>111</sup> Du et al<sup>102</sup> and Mao et al<sup>121</sup> have reported that the COMT Val158Met polymorphism may not contribute to the risk of prostate cancer, ovarian cancer, or breast cancer in any of the assessed genetic model. In the subgroup analysis by ethnicity, no significant associations were found in African, Caucasian, and mixed populations. However, the significant association between the COMT Val158Met polymorphism and cancer risk remains to be determined in Asians. The discrepancy in ethnicity could be attributed to the evident difference in the minor allele frequency of Val158Met polymorphism in Asians and Caucasians in our meta-analysis. This genetic polymorphism variance with ethnicity was consistent with those described in a previous study.<sup>8</sup> In addition, stratified analyses by genotyping techniques indicated that studies involving AS-PCR likely acquired significant results in the overall comparison. However, this result should be carefully interpreted because of a relatively small sample size. Moreover, this result should be confirmed by further analysis of additional published studies.

Several limitations should be acknowledged in this meta-analysis. First, only studies in English or Chinese were included in this meta-analysis, which might cause publication bias. Second, the pooled results were based on unadjusted estimates because not all studies had provided adjusted ORs. Even in cases where adjusted ORs were found, they were not adjusted by the same confounders. Hence, a precise analysis should be performed. Third, several factors such as gene-gene or gene-environment interaction may influence gene-disease factor, and the lack of individual data from the included studies limited further evaluation of other potential interactions, as in other genes and environment factors. Finally, cancer is a multifactorial disease resulting from complex interactions among many genetic and environmental factors. Therefore, a single gene or single environmental factor is unlikely to explain cancer susceptibility.

#### Conclusion

In conclusion, the present meta-analysis suggested that COMT Val158Met polymorphism might not be a risk factor for overall cancer risk, but it might be involved in cancer development at least in some ethnic groups (Asian) or some specific cancer types (bladder and esophageal cancer). Further large-scale and well-designed studies regarding different ethnicities are required to confirm the results of our meta-analysis.

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### Disclosure

The authors report no conflicts of interest in this work.

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