# **ORIGINAL ARTICLE**

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# $T_{peak}$ - $T_{end}$ , $T_{peak}$ - $T_{end}$ /QT ratio and $T_{peak}$ - $T_{end}$ dispersion for risk stratification in Brugada Syndrome: A systematic review and meta-analysis

Gary Tse MPH, PhD, FESC, FACC, FHRS, FRCP<sup>1,2,3</sup> | Mengqi Gong BS<sup>4</sup> | Christien Ka Hou Li<sup>1,2,3,5</sup> | Keith Sai Kit Leung BSc (Hons) LIBMS<sup>1,2,3,6</sup> | Stamatis Georgopoulos MD<sup>7</sup> | George Bazoukis MSc, MD<sup>7</sup> | Konstantinos P. Letsas MD, FESC, FEHRA<sup>7</sup> | Abhishek C. Sawant MD, MPH<sup>8</sup> | Giacomo Mugnai MD, PhD<sup>9</sup> | Martin C.S. Wong MPH, MD, FESC, FACC, FFPH<sup>10</sup> | Gan Xin Yan MD, PhD<sup>11,12</sup> | Pedro Brugada MD, PhD<sup>9</sup> | Gian-Battista Chierchia MD, PhD<sup>9</sup> | Carlo de Asmundis MD, PhD<sup>9</sup> | Adrian Baranchuk MD, FACC, FRCPC, FCCS<sup>13</sup> | Tong Liu MD, PhD<sup>4</sup> | International Health Informatics Study (IHIS) Network

<sup>1</sup>Department of Medicine and Therapeutics, Faculty of Medicine, Chinese University of Hong Kong, Hong Kong, China

<sup>2</sup>Li Ka Shing Institute of Health Sciences, Faculty of Medicine, Chinese University of Hong Kong, Hong Kong, China

<sup>3</sup>Shenzhen Research Institute, The Chinese University of Hong Kong, Shenzhen, China

<sup>4</sup>Tianjin Key Laboratory of Ionic-Molecular Function of Cardiovascular Disease, Department of Cardiology, Tianjin Institute of Cardiology, Second Hospital of Tianjin Medical University, Tianjin, China

<sup>5</sup>Faculty of Medicine, Newcastle University, Newcastle, UK

<sup>6</sup>Aston Medical School, Aston University, Birmingham, UK

<sup>7</sup>Second Department of Cardiology, Laboratory of Cardiac Electrophysiology, Evangelismos General Hospital of Athens, Athens, Greece

<sup>8</sup>Division of Cardiology, Department of Internal Medicine, State University of New York at Buffalo, Buffalo, New York

<sup>9</sup>Heart Rhythm Management Center, Postgraduate Program in Cardiac Electrophysiology and Pacing, Universitair Ziekenhuis Brussel-Vrije Universiteit Brussel, Brussels, Belgium

<sup>10</sup>JC School of Public Health and Primary Care, The Chinese University of Hong Kong, Hong Kong, China

<sup>11</sup>Lankenau Institute for Medical Research and Lankenau Medical Center, Wynnewood, Pennsylvania

<sup>12</sup>Beijing Anzhen Hospital, Capital Medical University, Beijing, China

<sup>13</sup>Division of Cardiology, Kingston General Hospital, Queen's University, Kingston, ON, Canada

#### Correspondence

Gary Tse, Department of Medicine and Therapeutics, Faculty of Medicine, Chinese University of Hong Kong, Hong Kong, China. Email: tseg@cuhk.edu.hk

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# Abstract

**Background:** Brugada syndrome is an ion channelopathy that predisposes affected subjects to ventricular tachycardia/fibrillation (VT/VF), potentially leading to sudden cardiac death (SCD).  $T_{peak}$ - $T_{end}$  intervals,  $(T_{peak}$ - $T_{end})/QT$  ratio and  $T_{peak}$ - $T_{end}$  dispersion have been proposed for risk stratification, but their predictive values in Brugada syndrome have been challenged recently.

**Methods:** A systematic review and meta-analysis was conducted to examine their values in predicting arrhythmic and mortality outcomes in Brugada Syndrome.

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**Results:** Nine studies involving 1740 subjects (mean age 45 years old, 80% male, mean follow-up duration was  $68 \pm 27$  months) were included. The mean  $T_{peak}$ - $T_{end}$  interval was 98.9 ms (95% CI: 90.5-107.2 ms) for patients with adverse events (ventricular arrhythmias or SCD) compared to 87.7 ms (95% CI: 80.5-94.9 ms) for those without such events, with a mean difference of 11.9 ms (95% CI: 3.6-20.2 ms, P = 0.005;  $l^2 = 86\%$ ). Higher ( $T_{peak}$ - $T_{end}$ )/QT ratios (mean difference = 0.019, 95% CI: 0.003-0.036, P = 0.024;  $l^2 = 74\%$ ) and  $T_{peak}$ - $T_{end}$  dispersion (mean difference = 7.8 ms, 95% CI: 2.1-13.4 ms, P = 0.007;  $l^2 = 80\%$ ) were observed for the event-positive group.

**Conclusion:**  $T_{peak}$ - $T_{end}$  interval,  $(T_{peak}$ - $T_{end})/QT$  ratio and  $T_{peak}$ - $T_{end}$  dispersion were higher in high-risk than low-risk Brugada subjects, and thus offer incremental value for risk stratification.

#### KEYWORDS

Brugada syndrome, risk stratification, sudden cardiac death, Tpeak-Tend, ventricular arrhythmia

# 1 | INTRODUCTION

Brugada syndrome is a used to describe the combination of specific ECG changes, the Brugada pattern, in addition to life threatening arrhythmias and sudden cardiac death (SCD).<sup>1</sup> Traditionally, it has been considered a congenital ion channelopathy linked to abnormalities in the cardiac sodium channel.<sup>2,3</sup> Recently, pathogenic mutations in other ion channels have been described. Mechanisms of arrhythmogenesis can be broadly divided into triggered activity and reentry. Of these, re-entry is thought to be the predominant mechanism underlying increased arrhythmogenicity in Brugada syndrome requiring an increased spatial dispersion of repolarization. Such reentrant activity may involve direct electrotonic activation during phase 2 of the cardiac action potential, as shown in pre-clinical studies using arterially perfused, canine wedge preparations,<sup>4</sup> or circustype/spiral wave activity around an anatomical or functional obstacle. Regardless of the precise underlying mechanism for re-entry, this transmural dispersion of repolarization can be quantified electrocardiographically by the interval from the peak to the end of the Twave (T<sub>peak</sub>-T<sub>end</sub> interval), (T<sub>peak</sub>-T<sub>end</sub>)/QT ratio and T<sub>peak</sub>-T<sub>end</sub> dispersion.5,6

However, not all studies have shown an association between higher  $T_{peak}$ - $T_{end}$  intervals,  $(T_{peak}$ - $T_{end})/QT$  ratio or  $T_{peak}$ - $T_{end}$  dispersion with an arrhythmogenic phenotype in Brugada Syndrome. Recently, Mugnai and colleagues conducted one of the largest retrospective studies to date, including a total of 448 patients with spontaneous or drug induced type 1 Brugada pattern.<sup>7</sup> They found no statistically significant difference in all three indices between asymptomatic subjects and patients with syncope and malignant arrhythmias. Morita and colleagues also found in 471 patients no difference in  $T_{peak}$ - $T_{end}$  intervals between patients with syncope or VT/VF and

those who were asymptomatic.<sup>8</sup> These findings contrast with a meta-analysis published previously by some members of our group, which extracted and pooled odds or hazard ratios for the relationship between  $T_{peak}$ - $T_{end}$  and arrhythmic and/or mortality outcomes in various clinical conditions, including Brugada Syndrome.<sup>9</sup> This demonstrated prolonged  $T_{peak}$ - $T_{end}$  interval was associated with an increased risk of ventricular arrhythmias and SCD in Brugada Syndrome.

However, our previous study did not determine the absolute mean values for  $T_{peak}$ - $T_{end}$ , nor was it possible to include the largest dataset from Mugnai and colleagues. Moreover, it did not investigate the utility of other indices such as  $(T_{peak}-T_{end})/QT$  ratio or  $T_{peak}-T_{end}$  dispersion. Therefore, we conducted a systematic review with meta-analysis into the relationships between  $T_{peak}$ - $T_{end}$  interval,  $(T_{peak}-T_{end})/QT$  ratio and  $T_{peak}-T_{end}$  dispersion and arrhythmic and/or mortality endpoints in Brugada Syndrome.

# 2 | METHODS

# 2.1 Search strategy, inclusion and exclusion criteria

This study was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISM) statement. PubMed and Embase were searched for studies that investigated the association between  $T_{peak}$ - $T_{end}$  or  $T_{peak}$ - $T_{end}$  /QT with arrhythmic or mortality endpoints in Brugada syndrome. The following search terms were used for both databases: ["Tpeak-Tend" or "Tpeak-end" or "Tp-e" AND Brugada]. The databases were searched until 1 May 2018 without language restrictions. The following inclusion criteria were used: (a) the study was a case-control, prospective or

retrospective cohort study in human subjects with a Brugada phenotype, (b)  $T_{peak}$ - $T_{end}$  intervals or  $(T_{peak}$ - $T_{end})$  /QT ratios were provided; (c) predefined adverse events (appropriate implantable cardioverterdefibrillator therapy [ICD], syncope, ventricular tachycardia/fibrillation [VT/VF], SCD, cardiovascular death [CVD], major adverse cardiac events [MACE]) or all-cause mortality were reported. In cases of incomplete data from the published studies, the original authors were contacted, but no replies were received.

The Newcastle-Ottawa Quality Assessment Scale (NOS) was used for quality assessment of the included studies.<sup>10</sup> The NOS system evaluated the categories of study participant selection, results comparability, and quality of the outcomes. Specifically, the following characteristics were assessed: (a) representativeness of the exposed cohort; (b) selection of the non-exposed cohort; (c) ascertainment of exposure; (d) demonstration that outcome of interest was not present at the start of study; (e) comparability of cohorts based on study design or analysis; (f) assessment of outcomes; (g) follow-up periods that were sufficiently long for outcomes to occur; and (h) adequacy of follow-up of cohorts. This scale varied from zero to nine stars, which indicated that studies were graded as poor quality if the score was <5, fair if the score was 5-7, and good if the score was >8. Studies with a score equal to or higher than six were included. The details of the NOS quality assessment are shown in Tables S1 and S2.

## 2.2 Data extraction and statistical analysis

Data from the different studies were entered in pre-specified spreadsheets in Microsoft Excel. All potentially relevant studies were retrieved as complete manuscripts, which were assessed fully to determine their compliance with the inclusion criteria. We extracted the following data from the included studies: (a) publication details: last name of first author, publication year and locations; (b) study design; (c) endpoint(s); (d) quality score; and (e) characteristics of the population including sample size, gender, age and number of subjects. Two reviewers (GT and MG) reviewed each included study independently. Disagreements were resolved by adjudication with input from a third reviewer (TL).

Adverse events were defined as ventricular arrhythmias (VT/VF), SCD, cardiovascular death, MACE or all-cause mortality. If more than one mortality endpoint was described, then SCD was preferentially used for analysis, followed by cardiovascular and all-cause mortality in this order. Mean differences between event-positive and event-negative groups, with 95% confidence intervals (CIs) for  $T_{peak}$ - $T_{end}$  interval, ( $T_{peak}$ - $T_{end}$ )/QT ratio and  $T_{peak}$ - $T_{end}$  dispersion were extracted and subsequently combined to generate a pooled estimate.

Heterogeneity between studies was quantified using The Cochran's Q value and the  $l^2$  statistic from the standard chi-square test, which describes the percentage of the variability in effect estimates resulting from heterogeneity.  $l^2 > 50\%$  was considered to reflect significant statistical heterogeneity. A fixed effects model was used if  $l^2 < 50\%$ . The random-effect model using the inverse variance heterogeneity method was used when  $l^2 > 50\%$ . To locate the

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origin of the heterogeneity, sensitivity analysis by excluding one study at a time, and subgroup analyses based on different disease conditions and different endpoints were performed. Funnel plots, Begg and Mazumdar rank correlation test and Egger's test were used to detect publication bias.

# 3 | RESULTS

Figure 1 shows a flow diagram detailing the above search terms with inclusion and exclusion criteria. A total of 29 and 57 entries were retrieved from PubMed and Embase, respectively. Nine studies met the inclusion criteria and were included in our final meta-analysis.<sup>6,7,11-17</sup> In this meta-analysis, a total of 1740 subjects with Brugada Syndrome were included (mean age 45 years old, 80% male). The mean follow-up duration was  $68 \pm 27$  months. Of the entire cohort, 40% had a spontaneous Type 1 pattern and 19% were positive for SCN5a mutation. The baseline characteristics of these studies and of the study populations are shown in Table 1.

# 3.1 | T<sub>peak</sub>-T<sub>end</sub>

For determining  $T_{end}$ , the tangent method and the return of the voltage to baseline method were used.  $T_{peak}$ - $T_{end}$  intervals from different leads and the maximum of these measurements have been presented by most studies. Regarding maximum  $T_{peak}$ - $T_{end}$  intervals, the mean value for the event-positive group was 98.9 ms (95% CI: 90.5-107.2 ms) (Figure 2A) and event-negative group was 87.7 ms (95% CI: 80.5-94.9 ms) (Figure 2B). Five studies reported longer values in the event-positive compared to event-negative groups, whereas four studies reported no significant difference (Figure 2C).  $T_{peak}$ - $T_{end}$  intervals were 11.9 ms longer (95% CI: 3.6-20.2 ms, P = 0.005) in event-positive patients than in event-negative patients. The Cochran's Q value was greater than the degrees of freedom (56 vs 8), indicating that the true effect size was different between studies.

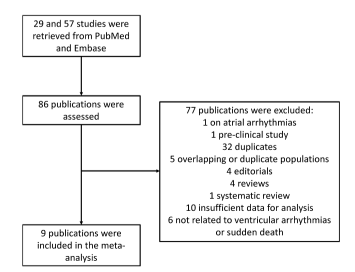


FIGURE 1 Flow diagram of the study selection process

FirstTpeak-Tend measurement: yearyearSamplemeasurement: method and leadsMorita471Tangent method; V1, V2, V3, V5Mugnai448End of the T- wave; V1 to V6Zo17143Tangent method; v1 to V6Zumhagen143Tangent method; v1 to V6Zumhagen78V1 to V6Zumhagen78V1 to V6Zumhagen78Tangent method; v1 to V6Zumhagen78Tangent method; v1 to V6Zumhagen78Tangent method; v1 to V6Zumhagen78Tangent method; v1 to V6	10	Age         No. of           (SD)         males (%)           47 (19)         447 (95)           45 (16)         273 (61)	No. of Sp. type 1 patients (%)	No. of SCN5a positive			No. of patients with adverse events /without adverse	Follow-up duration	:	
i 471 . De 143 . Ben 78 .		<ul> <li>9) 447 (95)</li> <li>6) 273 (61)</li> </ul>		patients (%)	Endpoints	Comparisons	events/%/% per year	(months)	Quality score	Ref.
448           143           78           325			118 (25)	27 (15)	Syncope or VT/VF	Syncope/VT/VF vs asymptomatic	145/326/31/4.09	91	7	16
143			96 (21)	55 (22)	Spontaneous VF or SCD	AT/SD vs asymptomatic	43/290/13/1.67	63	6	7
78 325		2) 140 (98)	84 (59)	I	VF	VF vs no VF	35/108/24/1.9	105	7	17
325		4) 57 (73)	22 (28)	17 (22)	Spontaneous VT/VF	VT/VF/aborted SCD vs asymptomatic/ syncope	22/54/29/-	I	Ŷ	14
		47 (13) 260 (80)	143 (44)	43 (13)	Spontaneous VT/VF	AT/SD vs asymptomatic	26/226/10/2.50	48	7	11
Letsas         23         End of the T-           2010         wave; V2, V6	e T- 43 (15) 2, V6	5) 19 (83)	10 (43)	I	Inducible VT/ VF	Inducible VT vs no inducible VT	17/6/74/16.15	55	6	12
Junttila 200 End of the T- 2008 wave; V2, II		40 (16) 143 (72)	200 (100)	25 (50)	Syncope, VT/ VF, SCD	Syncope/VT/VF/ aborted SCD vs asymptomatic	66/134/33/-	I	7	15
Wang23End of the T-2007wave; Max fromV1 to V6	е Т- 45 (8) ax from	) 23 (100)	I	I	Spontaneous VT/VF	Syncope//T//F/ inducible VT vs asymptomatic	11/9/55/5.12	43	00	13
Castro 29 Tangent method, Hevia Max from V1 to 2006 V6	nethod, 41 (12) n V1 to	2) 25 (86)	15 (52)	I	Spontaneous VT/VF	Presyncope/syncope/ aborted SCD vs asymptomatic	12/17/41/3.81	43	8	Ŷ

**TABLE 1** Characteristics of the nine studies included in this meta-analysis

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 $l^2$  took a value of 86%, suggesting the presence of substantial heterogeneity. A funnel plot plotting standard errors against differences in means is shown in Figure S1. Begg and Mazumdar rank correlation analysis demonstrated that Kendall's Tau took a value of 0.3 with P = 0.30, which suggests no significant publication bias. Egger's test demonstrated no significant asymmetry (intercept 2.4, t-value 1.2; P = 0.25). To identify the source of the heterogeneity, sensitivity analysis was performed by removing one study at a time, but this did not significantly influence the mean difference (Figure S2), suggesting that no single study was responsible for the heterogeneity observed in this meta-analysis. Subgroup analysis based on the method of T<sub>end</sub> determination was performed. For the tangent method, the  $T_{peak}$ - $T_{end}$  mean difference was 15.5 ms (95% CI: 3.9-27.2 ms; P = 0.009) and  $l^2$  remained high at 90%. For full recovery of voltage to baseline, the mean difference was 6.0 ms (95% CI: 0.7-11.4 ms; P = 0.006) and  $l^2$  remained high at 76%. Therefore, different methods of  $T_{\mbox{\scriptsize end}}$  determination did not introduce significant heterogeneity to the pooled effect estimate.

# 3.2 | (T<sub>peak</sub>-T<sub>end</sub>)/QT ratio

Regarding maximum (Tpeak-Tend)/QT ratio, the mean value for the event-positive group was 0.221 (95% CI: 0.208-0.234) (Figure 3A) and event-negative group was 0.210 (95% CI: 0.205-0.214) (Figure 3B). Two studies reported higher values in Brugada subjects with positive events compared to those without such events, whereas four studies demonstrated no significance between the groups (Figure 3C). Pooling of the mean values demonstrated significantly higher (T<sub>peak</sub>-T<sub>end</sub>)/QT ratios in the event-positive group than in the event-negative group (mean difference = 0.019, 95% CI: 0.003-0.036, P = 0.024). The Cochran's O value was greater than the degrees of freedom (19 vs 5), indicating that the true effect size was different between studies. I<sup>2</sup> took a value of 74%, suggesting significant heterogeneity. A funnel plot plotting standard errors against differences in means is shown in Figure S3. Begg and Mazumdar rank correlation analysis demonstrated that Kendall's Tau took a value of 0.07 with P = 1, which suggested no significant publication bias. Egger's test demonstrated no significant asymmetry (intercept 3.5, t-value 1.1; P = 0.31). To identify the source of the heterogeneity, sensitivity analysis was performed by removing one study at a time, but this did not significantly influence the mean difference (Figure S4), suggesting that no single study was responsible for the heterogeneity observed in this meta-analysis. Subgroup analysis based on the method of T<sub>end</sub> determination was performed. For the tangent method, the mean difference of (T<sub>peak</sub>-T<sub>end</sub>)/QT ratio was 0.03 (95% CI: 0.01-0.05; P < 0.05) and  $I^2$  was lowered to 55%. For full recovery of voltage to baseline, the mean difference was only 0.004 (95% CI: -0.03 to 0.03 ms; P = 0.81) and  $I^2$ remained high at 74%. Therefore, different method of  $T_{end}$  determination appeared to contribute partially to the heterogeneity of the pooled effect estimate. Moreover, statistical significance was achieved when the tangent method was used, but was lost when the return to baseline method was used, which may suggest the former approach may be more sensitive.

# 3.3 | T<sub>peak</sub>-T<sub>end</sub> dispersion

Regarding maximum  $T_{peak}$ - $T_{end}$  dispersion, the mean value for the event-positive group was 40.8 ms (95% CI: 26.9-54.8 ms) (Figure 4A) and event-negative group was 29.7 ms (95% CI: 24.5-34.8 ms) (Figure 4B). Regarding T<sub>peak</sub>-T<sub>end</sub> dispersion, two studies reported longer values in event-positive group compared to event-negative groups, whereas three studies found no significant difference (Figure 4C). Overall, pooling of the data showed that T<sub>peak</sub>-T<sub>end</sub> dispersion was significantly higher in the event-positive than in the event-negative groups (mean difference = 7.8 ms, 95% CI: 2.1 to 13.4 ms, P = 0.007). The Cochran's Q value was greater than the degrees of freedom (20 vs 4), indicating that the true effect size was different between studies.  $I^2$ took a value of 80%, suggesting significant heterogeneity. A funnel plot plotting standard errors against differences in means is shown in Figure S5. Begg and Mazumdar rank correlation analysis demonstrated that Kendall's Tau took a value of -2 with P = 0.62, which suggests no significant publication bias. Egger's test demonstrated no significant asymmetry (intercept -5.4, *t*-value 0.8; P = 0.48). To identify the source of the heterogeneity, sensitivity analysis was performed by removing one study at a time, but this did not significantly influence the mean difference between event-positive and event-negative groups (Figure S6), suggesting that no single study was responsible for the heterogeneity observed in this meta-analysis. Subgroup analysis based on the method of  $T_{\rm end}$  determination was performed. For the tangent method, the mean difference of  $T_{\text{peak}}\text{-}T_{\text{end}}$  dispersion was 16.2 ms (95% CI: 7.9-24.5 ms; P < 0.0001) and  $I^2$  was 65%. For full recovery of voltage to baseline, the mean difference was 0.4 ms (95% CI: -7.3 to 8.2 ms; P = 0.91) and I<sup>2</sup> was reduced to 19%. Therefore, different method of  $T_{\text{end}}$  determination contributed heterogeneity to the pooled effect estimate. Moreover, statistical significance was achieved when the tangent method was used, but was lost when the return to baseline method was used, which may suggest the former approach may be more sensitive.

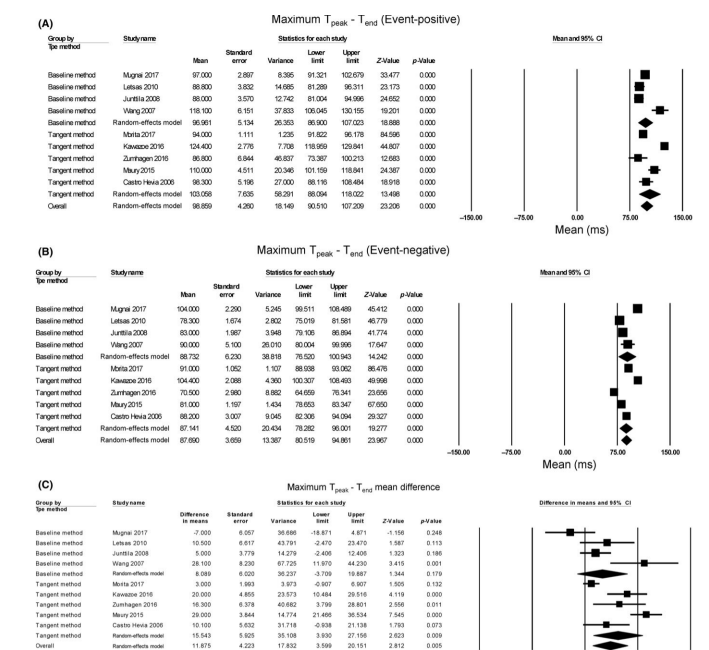
# 3.4 Comparisons between patients with and without SCN5A mutations

SCN5A is the commonest ion channel gene that is mutated in Brugada syndrome.<sup>2,3</sup> Separate meta-analyses were conducted to compare the different  $T_{peak}$ - $T_{end}$  parameters between patients with and without SCN5A mutations. Two of the included studies provided sufficient information for such analyses.<sup>7,14</sup> No significant difference in  $T_{peak}$ - $T_{end}$  (mean difference = 8.2 ms, 95% CI: -6.7 to 23.2 ms, P = 0.28;  $l^2 = 59\%$ ; Figure S7),  $T_{peak}$ - $T_{end}/QT$  ratio (mean difference = -0.006 ms, 95% CI: -0.023 to 0.011 ms, P = 0.47;  $l^2 = 24\%$ ; Figure S8) or  $T_{peak}$ - $T_{end}$  dispersion (mean difference = 5.2 ms, 95% CI: -2.9 to 13.2 ms, P = 0.21;  $l^2 = 31\%$ ; Figure S9) was observed between patients with and without SCN5A mutations.

50.00

Higher in event-positive

group



**FIGURE 2** Forest plot demonstrating  $T_{peak}-T_{end}$  intervals obtained from event-positive (A) and event-negative (B) groups and the mean difference between both groups (C) in Brugada Syndrome

# 4 DISCUSSION

The main findings of our meta-analysis, which included 1597 Brugada subjects, are (a)  $T_{peak}-T_{end}$  intervals, (b)  $(T_{peak}-T_{end})/QT$  ratio and (c)  $T_{peak}-T_{end}$  dispersion are higher in Brugada subjects with adverse cardiac events (ventricular tachy-arrhythmias and SCD) when compared to Brugada subjects free from such events.

The presence of pre-existing electrophysiological heterogeneities is important for mediating the normal, unidirectional spread of action potentials in the heart.<sup>18,19</sup> These are attributed to differences in repolarization times of the different cell types, which are responsible for generation of the T-wave on the electrocardiogram (ECG).<sup>20,21</sup> However, exacerbation of such differences has been associated with ventricular tachy-arrhythmias in different conditions, thereby generating a pro-arrhythmic phenotype. These include congenital ion channelopathies such as long QT syndrome and Brugada syndrome<sup>22-24</sup> and acquired cardiac diseases such as myocardial infarction.<sup>25,26</sup> These heterogeneities can occur locally or across the

-25.00

Lower in event-positive

group

Mean difference (ms)

-50.00

(A)

(B)

Mean and 95% CI

# Maximum Tpeak - Tend/QT ratio (Event-positive)

Group by	Studyname			Statisti	cs for each	study				
Tpe method		Mean	Standard error	Variance	Lower limit	Upper limit	Z-Value	p-Value		
Baseline method	Mugnai 2017	0.200	0.009	0.000	0.182	0.218	21.858	0.000		
Baseline method	Letsas 2010	0.230	0.007	0.000	0.216	0.244	31.610	0.000		
Baseline method	Random-effects model	0.216	0.015	0.000	0.186	0.245	14.376	0.000		
Tangent method	Morita 2017	0.240	0.013	0.000	0.214	0.266	18.062	0.000		
Tangent method	Zumhagen 2016	0.240	0.023	0.001	0.194	0.286	10.234	0.000		
Tangent method	Maury 2015	0.210	0.008	0.000	0.195	0.225	26.770	0.000		
Tangent method	Castro Hevia 2006	0.220	0.012	0.000	0.197	0.243	19.053	0.000		
Tangent method	Random-effects model	0.222	0.008	0.000	0.207	0.237	29.090	0.000		
Overall	Random-effects model	0.221	0.007	0.000	0.208	0.234	32.446	0.000		
									-0.30	

# 0.15 Mean

#### Maximum T<sub>peak</sub> - T<sub>end</sub>/QT ratio (Event-negative)

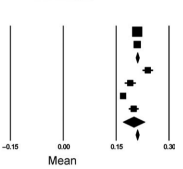
Group by	Studyname			Statisti	cs for each	study			
Tpe method		Mean	Standard error	Variance	Lower limit	Upper limit	Z-Value	<i>p</i> -Value	
Baseline method	Mugnai 2017	0.210	0.003	0.000	0.204	0.216	71.523	0.000	
Baseline method	Letsas 2010	0.210	0.004	0.000	0.202	0.218	51.439	0.000	
Baseline method	Random-effects model	0.210	0.002	0.000	0.205	0.215	88.100	0.000	
Tangent method	Morita 2017	0.240	0.008	0.000	0.225	0.255	30.952	0.000	
Tangent method	Zumhagen 2016	0.190	0.008	0.000	0.174	0.206	23.270	0.000	
Tangent method	Maury 2015	0.170	0.003	0.000	0.163	0.177	51.113	0.000	
Tangent method	Castro Hevia 2006	0.200	0.007	0.000	0.186	0.214	27.487	0.000	
Tangent method	Random-effects model	0.200	0.016	0.000	0.169	0.230	12.707	0.000	
Overall	Random-effects model	0.210	0.002	0.000	0.205	0.214	89.009	0.000	
									-0

Mean and 95% CI

0.00

0.30

0.15



Maximum T<sub>peak</sub> - T<sub>end</sub>/QT ratio mean difference (C) Statistics for each study Difference in means and 95% CI Studyname Group by Difference in means Standard Lower Upper Variance Z-Value p-Value Baseline method Mugnai 2017 -0.010 0.008 0.000 -0.026 0.006 -1.191 0.234 0.020 0.000 1.581 0.114 Baseline method Letsas 2010 0.013 -0.005 0.045 Random-effects mod Baseline method 0.004 0.015 0.000 -0.026 0.033 0.235 0.814 Tangent method Morita 2017 0.000 0.015 0.000 -0.029 0.029 0.000 1.000 Tangent method Zumhagen 2016 0.050 0.020 0.000 0.088 2.550 0.011 0.012 Tangent method Maury 2015 0.040 0.010 0.000 0.020 0.060 3.935 0.000 Tangent method 0.020 0.013 0.000 0.045 1.541 0.123 Castro Hevia 2006 -0.005 Random-effects mode Tangent method 0.027 0.010 0.000 0.007 0.047 2.592 0.010 Random-effects mode Overall 0.019 0.008 0 000 0.003 0.036 2 265 0.024 -0.15 Mean difference

Lower in event-positive Higher in event-positive group group

FIGURE 3 Forest plot demonstrating T<sub>peak</sub>-T<sub>end</sub>/QT ratios obtained from event-positive (A) and event-negative (B) groups and the mean difference between both groups (C) in Brugada Syndrome

myocardial wall,<sup>27</sup> potentially causing arrhythmias by inducing unidirectional conduction block and therefore circus-type or spiral wave re-entry.<sup>28,29</sup> Moreover, a greater epicardial-endocardial repolarization time difference may increase the propensity of phase 2 re-entry, which is hypothesized to generate extrasystolic activity in Brugada syndrome.<sup>30</sup> This occurs when sites with an action potential dome to sites which a dome morphology, leading to direct depolarization of the downstream sites.<sup>31</sup> Once an extrasystole is generated, together with a favorable re-entrant substrate, ventricular tachycardia and fibrillation can result.32

A number of electrocardiographic indices have been proposed for stratification of arrhythmic or mortality risk.<sup>33,34</sup> Of these, Yan and Antzelevitch were the first to propose the use of the difference between the peak and the end of the T-wave (the  $T_{peak}$ - $T_{end}$  interval) as a measure of transmural dispersion of repolarization.<sup>20,35-37</sup> Subsequent clinical studies have demonstrated that, confirmed recently in a systematic review and meta-analysis from our group,<sup>9</sup> that  $T_{peak}$ - $T_{\text{end}}$  prolongation significantly elevated the risk of ventricular tachyarrhythmias and/or SCD in heart failure, ischemic heart disease, Brugada syndrome, hypertension, and the general population. Recently, Mugnai and colleagues in a total of 448 subjects found no significant differences  $T_{peak}$ - $T_{end}$  intervals,  $(T_{peak}$ - $T_{end})/QT$  ratio or  $T_{peak}$ - $T_{end}$  dispersion between patients with VT/VF requiring anti-tachycardia pacing or with sudden death, and those who were asymptomatic.<sup>7</sup> Similarly, in a separate population of 471 subjects, Morita and colleagues found no significance difference in Tpeak-Tend intervals (A)

(B)

Group by	Studyname	Statistics for each study									
Tpe method		Mean	Standard error	Variance	Lower limit	Upper limit	Z-Value	p-Value			
Baseline method	Mugnai 2017	42.000	2.897	8.395	36.321	47.679	14.495	0.000			
Baseline method	Letsas 2010	15.100	2.231	4.979	10.727	19.473	6.767	0.000			
Baseline method	Random-effects model	28.486	13.450	180.898	2.125	54.848	2.118	0.034			
Tangent method	Kawazoe 2016	58.800	4.953	24.528	49.093	68.507	11.873	0.000			
Tangent method	Maury 2015	47.000	5.295	28.038	36.622	57.378	8.876	0.000			
Tangent method	Castro Hevia 2006	31.700	3.868	14.963	24.118	39.282	8.195	0.000			
Tangent method	Random-effects model	45.608	8.385	70.311	29.173	62.043	5.439	0.000			
Overall	Random-effects model	40.816	7.116	50.632	26.869	54.762	5.736	0.000			

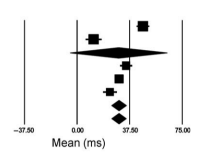
# Maximum $T_{peak}$ - $T_{end}$ dispersion (Event-negative)

Group by	Studyname	yname Statistics for each study							
Tpe method		Mean	Standard error	Variance	Lower limit	Upper limit	Z-Value	<i>p</i> -Value	
Baseline method	Mugnai 2017	47.000	2.349	5.517	42.396	51.604	20.010	0.000	
Baseline method	Letsas 2010	11.800	2.939	8.640	6.039	17.561	4.014	0.000	
Baseline method	Random-effects model	29.444	17.600	309.758	-5.051	63.940	1.673	0.094	
Tangent method	Kawazoe 2016	35.000	2.252	5.070	30.587	39.413	15.544	0.000	
Tangent method	Maury 2015	30.000	1.131	1.279	27.784	32.216	26.529	0.000	
Tangent method	Castro Hevia 2006	23.500	2.571	6.609	18.461	28.539	9.141	0.000	
Tangent method	Random-effects model	29.655	2.677	7.168	24.407	34.902	11.076	0.000	
Overall	Random-effects model	29.650	2.647	7.006	24,462	34.838	11.202	0.000	

Mean and 95% CI

0.00 Mean (ms)

Mean and 95% CI



Froup by permethod	Studyname			Statistics	for each stu	dy			
pe metroa		Difference in means	Standard error	Variance	Lower	Upper limit	Z-Value	p-Value	
aseline method	Mugnai 2017	-5.000	6.207	38.528	-17.166	7.166	-0.806	0.421	
aseline method	Letsas 2010	3.300	4.162	17.325	-4.858	11.458	0.793	0.428	
aseline method	Random-effects model	0.427	3.949	15.591	-7.312	8.166	0.108	0.914	
angent method	Kawazoe 2016	23.800	4.853	23.551	14.288	33.312	4.904	0.000	
angent method	Maury 2015	17.000	3.779	14.281	9.593	24.407	4.498	0.000	
angent method	Castro Hevia 2006	8.200	4.457	19.865	-0.536	16.936	1.840	0.066	
angent method	Random-effects model	16.204	4.242	17.996	7.890	24.519	3.820	0.000	
werall	Random-effects model	7.751	2.890	8.354	2.086	13.416	2.682	0.007	

-75.00

-37.50

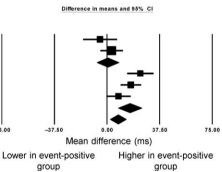


FIGURE 4 Forest plot demonstrating T<sub>peak</sub>-T<sub>end</sub> dispersion obtained from event-positive (A) and event-negative (B) groups and the mean difference between both groups (C) in Brugada Syndrome

between patients with syncope or VT/VF and asymptomatic patients.<sup>16</sup> Publication of these two studies prompted us to conduct this meta-analysis, which confirms the value of  $T_{peak}$ - $T_{end}$  interval,  $(T_{\text{peak}}\text{-}T_{\text{end}})/QT$  ratio and  $T_{\text{peak}}\text{-}T_{\text{end}}$  dispersion, in distinguishing highrisk patients from low-risk patients.

In the Mugnai study, the largest study to date, the percentage of patients with adverse events were the lowest at 13%.<sup>7</sup> Male gender, a spontaneous Type 1 Brugada pattern and SCN5a mutation positive status were significantly associated with ventricular arrhythmias.38 Therefore, the lower percentage of patients with adverse events can be explained by the lower percentage of Type 1 Brugada patients (21% vs 28%-100% in the remaining studies) and lower percentage male patients (61% vs 72%-100%) despite similar percentage with SCN5a positive status (22% vs 13%-50%). While these differences in patient characteristics affect the likelihood of adverse events occurring, they should not explain the lack of difference in T<sub>peak</sub>-T<sub>end</sub> intervals between event-positive and event-negative groups in the Morita study<sup>16</sup> or the Mugnai study. Interestingly, Mugnai and colleagues found a non-statistically significant lower T<sub>peak</sub>-T<sub>end</sub> intervals in event-positive groups. Of the remaining six studies, five studies had reported significantly higher  $T_{\text{peak}}$ - $T_{\text{end}}$  intervals and one study reported no difference.<sup>15</sup> A recent epidemiological study reported a U-shaped relationship between T<sub>peak</sub>-T<sub>end</sub> intervals and increased mortality.<sup>39</sup> Autonomic modulation, which is part of Coumel's triad for arrhythmogenesis,<sup>40</sup> is known to modulate the re-entrant substrate. Increased activity of the parasympathetic nervous system may reduce T<sub>peak</sub>-T<sub>end</sub> intervals, which may also be pro-arrhythmic.<sup>41</sup> By contrast, exercise, during which sympathetic activity is increased, can exacerbate pre-existing heterogeneities, such as producing conduction slowing<sup>42</sup> and increasing the dispersion of repolarization.<sup>43</sup>

In our previous meta-analysis pooling together studies that reported odds ratios or hazard ratios, the average cut-off for  $T_{peak}$ -T<sub>end</sub> was 95.8 ms across different clinical conditions.<sup>9</sup> The present meta-analysis pooling mean values for event-positive and -negative groups clearly indicates that the 100 ms cut-off is too high for Brugada syndrome. Our data would support a lower cut-off value between 88 and 99 ms to be used. This cut-off will also be methoddependent for determining T<sub>end</sub> in the case of the T<sub>peak</sub>-T<sub>end</sub> intervals. Previously, it was shown that in a cohort of high-risk Brugada subjects, only 10 of 16 studies reported a T<sub>neak</sub>-T<sub>end</sub> longer than 100 ms, supporting our notion that this cut-off value may be too high.<sup>44</sup> Moreover, different studies measured T<sub>peak</sub>-T<sub>end</sub> from different leads. Some had measured it from all 12 leads and taken the mean values while others have done so for V1 to V3 only. While there is no consensus as to which leads are most appropriate for measurement, obtaining it from all 12 leads is likely to be less useful clinically due to the time-consuming nature. To simplify T<sub>neak</sub>-T<sub>end</sub> determination, we would thus propose measuring it from the right precordial leads given BrS is primarily a right ventricular disorder.

While it may appear that the difference in  $T_{peak}$ - $T_{end}$  between high-risk and low-risk Brugada patients was only small, at around 12 ms, it should be emphasized that increased transmural dispersion of repolarization is only one mechanism by which re-entrant arrhythmogenesis is generated. Other mechanisms, such as reduced conduction velocity, increased dispersion of conduction<sup>45</sup> or dynamic substrates such as steep action potential restitution,<sup>46</sup> in which normal  $T_{peak}$ - $T_{end}$  interval,  $T_{peak}$ - $T_{end}/QT$  ratio or  $T_{peak}$ - $T_{end}$  dispersion may be observed, also contribute to arrhythmogenesis in Brugada syndrome. Therefore, better risk stratification scores will need to incorporate a combination of repolarization and conduction indices. Moreover, some of these dynamic changes may not be detectable on the ECG and may require additional tests such as non-invasive ECG imaging (ECGi),<sup>43</sup> or only becomes detectable only under stressful conditions such as exercise.<sup>43</sup>

#### 4.1 | Limitations

The following limitations of this meta-analysis should be noted. First, there is marked heterogeneity between the included studies. The method of T<sub>peak</sub>-T<sub>end</sub> determination across the studies was split even between the tangent method and full recovery of the voltage to baseline. Subgroup analysis based on the method used did not reduce the heterogeneity observed. Therefore, measurement method was unlikely to have significantly contributed to the heterogeneity observed. Moreover, the Letsas 2010 study<sup>12</sup> used a different endpoint of inducible VT compared to the remaining studies, but its exclusion did not significant affect the mean  $T_{\text{peak}}\text{-}T_{\text{end}}$  values for event-positive group, event-negative group, and mean difference between these groups. Second, retrospective studies may have more bias than prospective studies. Finally, it should be acknowledged that there is overlap between event-postiive and event-negative groups irrespective of the method of measuring  $T_{end}$ . This would suggest as a single measurement,  $T_{\text{peak}}\text{-}T_{\text{end}}$  is unlikely to be useful in its own 595

right. Indeed, accurate risk stratification will require a composite scoring system assessing not only dispersion of repolarization, but that of conduction, clinical symptoms, family history, the type of Brugada pattern, genetic background, electrical and drug provocation testing as well as electrophysiological mapping.<sup>38,41,45,47-49</sup>

# 5 | CONCLUSIONS

 $T_{peak}-T_{end}$  interval,  $T_{peak}-T_{end}/QT$  ratio and  $T_{peak}-T_{end}$  dispersion were higher in high-risk than low-risk Brugada subjects, and thus offer incremental value for risk stratification.

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#### CONFLICTS OF INTERESTS

Authors declare no Conflict of Interests for this article.

# ORCID

Gary Tse D http://orcid.org/0000-0001-5510-1253 George Bazoukis D http://orcid.org/0000-0003-1009-9772 Giacomo Mugnai D http://orcid.org/0000-0003-4733-9418 Carlo de Asmundis D http://orcid.org/0000-0001-9351-0760 Adrian Baranchuk D http://orcid.org/0000-0002-3042-6569

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# SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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