Spatial and nonspatial aspects of TQ-ST segment mapping were studied with the solid angle theorem and randomly coded data from 15,000 electrograms of 160 anterior descending artery occlusions each of 100-s duration performed in 18 pigs. Factors analyzed included electrode location, ischemic area and shape, wall thickness, and increases in plasma potassium (K\(^+\)). Change from control in the TQ-ST recorded at 60 s (ΔTQ-ST) was measured at 22 ischemic (IS) and nonischemic (NIS) epicardial sites overlying right (RV) and left (LV) ventricles. In IS regions, ΔTQ-ST decreased according to LV > septum > RV and LV base > LV apex. In NIS regions, LV sites had negative (Neg) ΔTQ-ST which increased as LV IS border was approached. However, RV NIS had positive (Pos) ΔTQ-ST which again increased as RV IS border was approached. With large artery occlusion IS area increased 123±18%, ΔTQ-ST at IS sites decreased (−38.1±3.6%), and sum of ΔTQ-ST at IS sites increased by only 67.3±10.3%. In RV NIS Pos ΔTQ-ST became Neg. With increased K\(^+\), ΔTQ-ST decreased proportionately to log K\(^+\) (r = 0.97±0.01) at IS and NIS sites on the epicardium and precordium. TQ-ST at 60 s was obliterated when K\(^+\) = 8.7±0.2 mM. All findings were significant (P < 0.005) and agreed with the solid angle theorem. Thus, a transmembrane potential difference and current flow […]
Spatial and Nonspatial Influences on the TQ-ST Segment Deflection of Ischemia

THEORETICAL AND EXPERIMENTAL ANALYSIS IN THE PIG

ROGER P. HOLLAND and HAROLD BROOKS with the technical assistance of BARBARA LIDL

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ABSTRACT Spatial and nonspatial aspects of TQ-ST segment mapping were studied with the solid angle theorem and randomly coded data from 15,000 electrograms of 160 anterior descending artery occlusions each of 100-s duration performed in 18 pigs. Factors analyzed included electrode location, ischemic area and shape, wall thickness, and increases in plasma potassium (K⁺). Change from control in the TQ-ST recorded at 60 s (ΔTQ-ST) was measured at 22 ischemic (IS) and nonischemic (NIS) epicardial sites overlying right (RV) and left (LV) ventricles. In IS regions, ΔTQ-ST decreased according to LV > septum > RV and LV base > LV apex. In NIS regions, LV sites had negative (Neg) ΔTQ-ST which increased as LV IS border was approached. However, RV NIS had positive (Pos) ΔTQ-ST which again increased as RV IS border was approached. With large artery occlusion IS area increased 123±18%, ΔTQ-ST at IS sites decreased (−38.1±3.6%), and sum of ΔTQ-ST at IS sites increased by only 67.3±10.3%. In RV NIS Pos ΔTQ-ST became Neg. With increased K⁺, ΔTQ-ST decreased proportionately to log K⁺ (r = 0.97±0.01) at IS and NIS sites on the epicardium and precordium. TQ-ST at 60 s was obliterated when K⁺ = 8.7±0.2 mM. All findings were significant (P < 0.005) and agreed with the solid angle theorem. Thus, a transmembrane potential difference and current flow at the IS boundary alone are responsible for the TQ-ST. Nonspatial factors affect the magnitude of transmembrane potential difference, while spatial factors alter the position of the boundary to the electrode site.

INTRODUCTION

“No other part of the electrocardiogram is subject to so many theories, interpretations, and even more misinterpretations as the ST interval,” Shaefer and Haas (1). “The nature of these electrocardiographic changes is in no way mysterious,” Wilson et al. (2).

Techniques currently used to assess ischemic/infarcted myocardium include hemodynamic monitoring, creatine phosphokinase (CPK) curve analysis, myocardial radionuclide imaging, and precordial TQ-ST segment mapping. The latter has enjoyed considerable popularity due primarily to its low cost, noninvasive approach, and simply applied rules of interpretation. Controversy has arisen recently, however, for despite all the time, effort, and resources invested in TQ-ST segment mapping studies, basic appreciation of the complex manner by which the TQ-ST segment deflection relates to the underlying ischemic region has been lacking. Inevitably, confusion and disagreement over the specificity and quantitative value of this electrocardiographic measure of

1 Abbreviations used in this paper: CPK, creatine phosphokinase; ECG, electrocardiogram; [K⁺]o, plasma potassium; LAD, left anterior descending coronary artery; LV, left ventricle; RV, right ventricle; SA, average solid angle; ΣSA; summed solid angle; SUM(+), SUM(−), sum of all positive and negative TQ-ST deflections, respectively; TQ-STo, preocclusion TQ-ST segment deflection; ΔTQ-ST, individual TQ-ST segment deflection changes; ATQ-ST, average change in the TQ-ST segment; ΣΔTQ-ST, sum of individual TQ-ST segment deflection changes from the ischemic region; Vm, transmembrane potential of the ischemic region; Vm+n, transmembrane potential of the normal region; δVm, transmembrane potential difference; Ω, solid angle subtended at recording site by the ischemic boundary.
myocardial injury was to be expected (3–11). Recognizing the current renewed interest in this area and the profound clinical value of being able to quantify ischemic damage, we have in this study examined in a theoretical and experimental manner some basic spatial and nonspatial factors influencing the magnitude and polarity of TQ-ST deflection during myocardial ischemia.

**METHODS**

**Nomenclature**

Many studies have shown that the injury deflection, conventionally referred to as the “ST segment deflection,” has a diastolic as well as a systolic component. For this reason we have here, as in other studies (12, 13), referred to this deflection as the “TQ-ST segment deflection.” So-called “ST segment elevation,” then, is actually the summation of TQ depression and ST elevation (14–17). Due to capacitive coupling of most electrocardiogram (ECG) recording apparatus, the fact that the TQ segment is often not at zero potential (isoelectric) may not be obvious. The relative contribution of the diastolic (TQ segment) and systolic (ST segment) components to the total TQ-ST segment deflection has not yet been clearly established (14–17).

**Mathematical formulations from the solid angle theorem**

The solid angle theorem was used in this study to mathematically describe the behavior of the TQ-ST segment during ischemia. According to the theorem (Fig. 1) the magnitude and polarity of the TQ-ST segment deflection (e) recorded at an electrode site P is equal to the product of three parameters. Ω is the solid angle subtended at P by the ischemic boundary; V_m and V_m and K denote the transmembrane potentials of the normal and ischemic region during either diastole (TQ segment) or systole (ST segment); and K is a term correcting for differences in intra- and extracellular conductivity and the occupancy of much of the heart muscle by interstitial tissue and space. See text for additional discussion. Adapted from Holland and Arnsdorf (19).

**Experiments in the intact heart**

**Animal model.** Experiments were performed in 18 open-chest, domestic, neutered male pigs weighing from 35 to 60 kg. The pig was chosen because of similarities between the gross coronary artery architecture (22) and collateral circulation (23) of this animal and man. In addition, the high degree of reproducibility of the coronary architecture (22) among individual animals, as well as the less extensive distribution of collateral vessels in the pig ventricle as compared to the dog ventricle, insures that one can have a certain length of a coronary artery (24) results in a visually (cyanotic discoloration) and electrically identifiable area of ischemic damage. The degree of reproducibility is documented in the Results section of this paper.

**Surgery.** After induction of anesthesia with a small intravenous injection of thiopental, the animals were anesthetized with an intravenous infusion of a warmed solution of alpha-chloralose (60 mg/kg). During the study, supplementary doses of chloralose were given to maintain a relatively uniform state of anesthesia. The hemodynamic actions of chloralose are minimal and transient in duration (25). Respiration was maintained by a volume respirator (Harvard Apparatus Co., Inc., Millis, Mass.), regulated to maintain an arterial pH of 7.45±0.05 throughout the experiment. The pump was connected to a tracheostomy tube and supplemental oxygen was administered to maintain arterial oxygen saturation at 95%. The heart was exposed by a mid-sternal thoracotomy, a pericardiotomy was performed, and a pericardial cradle was created to support the exposed heart. **Hemodynamic measurements.** Heart rate was kept constant at 125 beats/min. This was accomplished by placing a bipolar electrode in the wall of the right auricular appendage and stimulating with a Grass stimulator (Grass Instrument Co., Quincy, Mass.) and an optical isolation unit. Stimulus duration was 5 ms. Animals whose resting
In this study, pressure determinations were made through 6-inch Teflon 16T gauge catheters connected directly to Statham P23 Dh pressure transducers (Statham Instruments Div., Gould, Inc., Oxnard, Calif.) without intervening tubing. Systemic pressure was obtained by placing the catheter in the left carotid artery and left ventricular pressure by placing it in the left ventricular cavity at the apex. The left ventricular pressure was differentiated electronically to obtain the time derivative of left ventricular pressure (LV dP/dt). Left ventricular end-diastolic pressure measurements were made from high-gain tracings via an additional amplifier (1 mm Hg/division).

In those animals in which the pattern of TQ-ST segment deflections was studied at different levels of plasma potassium ([K+]o), [K+]o was increased from control values by a slow intravenous infusion of KCl with an infusion pump. [K+]o levels were obtained from heparinized carotid arterial samples taken immediately after the beginning of each coronary occlusion and were measured with a flame photometer (Instrumentation Laboratory, Inc., Lexington, Mass.).

Ischemic area. The anterior left ventricular free wall supplied by branches of the left anterior descending (LAD) artery was selected for study. Reproducible areas of ischemic involvement in all animals were obtained by placing a silk ligature (OO) around the LAD artery at a point approximately one-third to one-half the distance from the apical termination of the LAD to its origin from the left main coronary artery. Reversible occlusion was obtained with a polyethylene collar. In those experiments in which large and small areas of ischemic involvement were compared, the larger area was produced by occluding the anterior descending artery at a point approximately 1.0 cm from its origin with the left main coronary artery. This insured that the large area completely enclosed the small area. Occlusions were limited to 100-s duration. Recent studies have demonstrated that although ischemic TQ-ST segment deflections return towards normal after release of a brief occlusion (5-10 min), return of contractile activity is delayed and may exhibit a permanent deficit (26-28). Thus, to insure complete functional (biochemical, electrical, mechanical, etc.) recovery of the ischemic segment, it would appear necessary to limit the occlusions to very short periods of time, instead of the 15-20-min occlusions of most studies (5, 29). Although the TQ-ST segment deflection has not obtained a steady-state value by this time (100 s), there is no indication from earlier studies that steady-state values are reached at any time within the 1st h after occlusion. Finally, the incidence of conduction abnormalities (QRS complex, loss of the S wave, ectopic beats, etc.) which may either obliterate, mimic, or alter the TQ-ST segment deflection characteristic of ischemia increases with the duration of the occlusion (5, 30). In the pig, such

Influences on the TQ-ST Segment Deflection
abnormalities are clearly in evidence beyond the first 100 s of the occlusion (30).

The area of ischemic involvement was estimated in all animals by the following methods: (a) The area of cyanotic discoloration after the first occlusion was drawn on to a previously sketched anterolateral view of the porcine heart which included the distribution of the LAD artery and its primary branches (31); (b) At postmortem, the length of LAD artery occluded was perfused with methylene blue dye and the stained epicardial surface along with various anatomic arterial landmarks was measured with a pair of calipers. The two areas (cyanotic and stained) were then calculated by planimetry and the caliper values; (c) The mass of stained tissue (ischemic weight) after the above dye infusion was then cut out and weighed (24). Agreement in the two methods of estimating ischemic area is documented in the Results section of this article.

Electrical measurements. Epicardial electrical potentials were recorded from the heart's surface with 22 atrumatic, firmly attaching electrodes designed in this laboratory. The electrodes are a modification of the original design (32) and are constructed from polished brass screws having a surface area of 14 mm² (radius = 2.2 mm). The electrodes were spaced equidistant from one another over the anterior surface of the left and right ventricles in the distribution of the LAD artery (Fig. 2, left).

The need for near-simultaneous recordings of the electrograms from 22 sites required that an automated rapid electromechanical switching circuit be designed (32). Basically the system consists of a pulse generator and two stepping relays. Upon receiving a pulse the stepping relay is advanced and connects a particular electrode site to the ECG amplifier for a period of 0.8 s. The heart rate of 125 beats/min used in this study permits approximately 1 1/4 heart beats to be recorded at each site. By using a pair of calibrated bioelectric amplifiers (No. 8811A, Hewlett-Packard Co., Palo Alto, Calif.) with a frequency response of DC to 10 kHz, all 22 electrode sites could be sampled within the space of 10 s. The signals were simultaneously recorded on FM magnetic tape (7% ips) and displayed on a high speed chart recorder along with the hemodynamic signals. The FM tape recorder had a frequency response of DC to 2.5 kHz, a signal-to-noise ratio of 44 dB, total harmonic distortion of 2%, and fluctuer of 0.4%. The resulting signals were then analyzed at a gain of 1.0 mV/div and an equivalent paper speed of 50 mm/s with a frequency response of DC to 400 Hz. With this system TQ-ST segment deflections after acute coronary occlusion were characterized in the following manner. TQ-ST deflections were measured from all sites immediately before occlusion (control) and at exactly 60 s after coronary occlusion. The total TQ-ST deflection was measured from that portion of the TQ segment occurring immediately before the inscription of the QRS complex to the systolic endpoint, a point on the ST segment occurring 100 ms after onset of the QRS complex (33). Track width was approximately 1/8 of a division and TQ-ST values were measured to 1/16 of a division (0.1 mV). The control values (TQ-ST₅ₐ) were obtained from the mean of multiple (2–4) complexes recorded during the 30 s immediately preceding the occlusion. The difference between this value and the magnitude of the deflection at 60 s after occlusion was labeled ΔTQ-ST. To obtain values for TQ-ST at 60 s from sites which were sampled at times other than 60 s (e.g., 56 and 66 s), linear interpolation was employed.

TQ-ST segment deflection changes at multiple ischemic and nonischemic sites were categorized solely on the basis of the position of the various recording sites to the ischemic region. This was accomplished by dividing the anteroseptal region of the heart into 10 regions as illustrated in Fig. 2 (right). All ischemic sites were required to be completely enclosed by the region of ischemic (cyanotic) myocardium.

Border sites were required to overlap both normal and cyanotic tissue and thus the width of the border was defined by the electrode diameter (4.4 mm). Septal ischemic sites were within 0.5 cm of the main trunk of the LAD. Left ventricular ischemic sites were further subdivided into sites lying in thin, apical portions of the ischemic region and those in the thicker-walled areas nearer the base of the heart. This was again accomplished in an unbiased fashion by drawing an imaginary line from the site of the occlusion to the apex of heart (see Fig. 2). All electrode locations were placed in 1 of the 10 categories before measurement of their electrogram tracings.

Data analysis. To avoid any prejudicial measurement of the TQ-ST segment deflections, all electrogram tracings were randomly coded and evaluated without knowledge as to electrode locations. As a result of the requirement for multiple measurements of the TQ-ST segment deflection an average of 100 determinations were made during each occlusion and thus in the course of this study in excess of 15,000 electrogram tracings were evaluated. Statistical analysis was confined to paired and unpaired t-tests and linear regression analysis (34) carried out with the help of a Wang 700 desk calculator (Wang Laboratories, Inc., Lowell, Mass.).

RESULTS

Theoretical analysis

In Fig. 3 the effect of different wall thicknesses and areas of ischemic involvement on the solid angle subtended at precordial and epicardial locations is illustrated. Although increases in ischemic size increase the magnitude of the solid angle at precordial locations, and decrease it at epicardial sites (12), at either location increases in wall thickness increase the solid angle. This finding immediately suggested that for equivalent areas of ischemic involvement, TQ-ST deflections recorded at thick-walled portions

![Figure 3](attachment:figure3.png)

**Figure 3** Effect of changes in wall thickness on the magnitude of the solid angle at precordial and epicardial electrode locations. Curves were derived with the electrode centrally positioned over the ischemic region. The outer wall radius used in this analysis was 3.5 cm. At either location increases in wall thickness increase the solid angle for a given area of ischemic involvement.
(base) of the ischemic left ventricle should exceed those recorded at the thinner-walled apex. Similarly, the TQ-ST deflection at left ventricular sites should exceed the magnitude of the deflections from right ventricular ischemic sites.

The solid angle values calculated above were for electrodes centrally positioned over a circular area of ischemic involvement. However, in mapping studies, electrodes overlie all regions of ischemic and nonischemic tissue. The effect of moving the electrode from the center of the ischemic area ($\beta/P = 0.0$), towards the ischemic boundary ($\beta/P = 1.0$), and beyond ($\beta/P > 1.0$) was computed and is shown in Fig. 4. The solid angle at precordial sites is maximal when the electrodes directly overlie the ischemic region ($\beta/P = 0.0$) and decreases at sites distant from the center ($\beta/P > 0.0$). The solid angle does not become negative until electrodes are positioned at a considerable distance from the ischemic boundary ($\beta/P \gg 1.0$). The solid angle at epicardial sites, in comparison, increases in positivity as the boundary is approached. At the boundary a discontinuity in the curve occurs and beyond this point, in the nonischemic regions, the epicardial solid angle is negative, rapidly decreasing in magnitude as the electrode moves further away from the ischemic boundary. Although the solid angle can assume values of either positive or negative polarity in the immediate vicinity of the boundary, on the average the solid angle values should have a positive polarity with a magnitude of approximately 40% that recorded by electrodes directly overlying the center of the ischemic region.

Having determined how the magnitude and polarity of the solid angle varies depending upon its location within the ischemic region, the summed solid angle ($\Sigma$SA) for different areas of ischemic involvement may be computed. $\Sigma$SA is equal to the sum of the individual solid angles subtended at each electrode site overlying the ischemic region. In Fig. 5 $\Sigma$SA and the average solid angle $\bar{SA}$ were plotted as a function of ischemic area. At the precordium both $\bar{SA}$ and $\Sigma$SA increase with ischemic area. The curve relating $\bar{SA}$ to ischemic area is remarkably linear over the range of areas from 5 to 50 cm². At the epicardium the situation is

Influences on the TQ-ST Segment Deflection
The involvement, and electrodes overlying the precordium percentage in depends. Although \( W - C^0 \) in doubling is ischemic area, \( ISA \). ischemic area, epicardial area (\( ISA \) +100\%) or epicardium. Although \( ISA \) increases with increases in ischemic area at either position, a given percentage change in ischemic area results in substantially smaller changes in \( ISA \) at the epicardium. A purely linear relationship between \( ISA \) and ischemic area is expected only when \( SA \) is constant for all ischemic areas. This situation never exists but is approximated at precordial sites and large ischemic areas whereupon changes in ischemic area are accompanied by nearly equivalent changes in \( ISA \).

different. Although \( \bar{SA} \) decreases with increases in ischemic area, \( \Sigma SA \) increases, but in a nonlinear manner. At this location a given percentage change in ischemic area results in a substantially smaller change in \( \Sigma SA \). In the range of areas from 10 to 50 cm\(^2\), a doubling in ischemic area is met by a 22\% decrease in \( \bar{SA} \) and only a 60\% increase in \( \Sigma SA \) at epicardial sites; at precordial sites the change in \( \Sigma SA \) (+153\%) is greater than the change in area (+100%).

**Experimental verification**

**Spatial influences.** In Table I anatomic and hemodynamic characteristics of the experimental model and the ischemic region are tabulated. Occlusion of a specific length (38.1±1.1 mm) of the anterior descending coronary artery in animals of equivalent weight class resulted in the production of standard areas of ischemic involvement (13.2±0.8 cm\(^2\)). The area of tissue becoming ischemic (cyanotic) closely approximated that area normally perfused by the occluded artery (stained).

In Table II TQ-ST segment deflection changes at multiple ischemic and nonischemic sites have been tabulated from a total of 77 occlusions performed in 16 animals. As was suggested by the theoretical findings, different TQ-ST potential populations can be successfully categorized simply on the basis of spatial relationships existing between the various electrode sites and the geometry of the ischemic region. In the ischemic region TQ-ST deflections were always observed to increase in magnitude as a thicker-walled ischemic boundary was approached. Thus the magnitudes of the individual ischemic populations decreased accordingly: left ventricle (LV) > SEPT > right ventricle (RV) and LV base > LV apex. The relative magnitudes of the deflections at these anatomic locations exhibited even smaller variability when deflection populations were compared in each animal (Fraction column). In this case the LV and RV ischemic sites were respectively 1.47±0.06 and 0.65±0.04 the magnitude recorded at the septal locations, while the sites closer to the LV base were 1.25±0.06 (\( P < 0.001 \)) those nearer the apex.

Although there was no significant difference in the means of the positive potential deflections recorded at the RV and LV ischemic borders, a distinction between these two populations may be noted. At the

**TABLE I**

**Characteristics of the Experimental Model**

<table>
<thead>
<tr>
<th>Anatomic</th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Animal weight, kg</td>
<td>46.6±1.7</td>
</tr>
<tr>
<td>Heart weight, g</td>
<td>184.3±6.1</td>
</tr>
<tr>
<td>Heart/animal weight, %</td>
<td>4.2±0.2</td>
</tr>
<tr>
<td>Ventricle weight, g</td>
<td>152.9±4.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Hemodynamic</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP, mm Hg</td>
<td>102.5±5.4</td>
</tr>
<tr>
<td>LVP, mm Hg</td>
<td>109.1±5.2</td>
</tr>
<tr>
<td>LVEDP, mm Hg</td>
<td>6.4±0.4</td>
</tr>
<tr>
<td>LV dP/dt, mm Hg/s</td>
<td>1,649±106</td>
</tr>
</tbody>
</table>

**Abbreviations:** MAP, mean arterial pressure; LVP, left ventricular peak systolic pressure; LVEDP, left ventricular end-diastolic pressure; LV dP/dt, time derivative of left ventricular pressure.

Values shown are means±SEM obtained from experiments performed in 17 animals. Heart rate was kept constant at 125 beats/min.

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right ischemic border the potential deflections were always positive, while at the left negative deflections were measured in a few instances.

At electrode sites overlying nonischemic portions of the LV reciprocal negative deflections were recorded, their magnitudes decreasing at locations more distant from the LV ischemic boundary. These reciprocal deflections averaged $-0.25\pm0.03$ (near) and $-0.09\pm0.01$ (distant) the magnitude at the ischemic septal sites.

In contrast, at nonischemic positions on the RV the polarity of the TQ-ST deflections was positive despite the fact that the sites from where they were recorded neither overlay cyanotic tissue nor were in the apparent distribution of the occluded artery. The magnitude of these deflections maintained its positivity but diminished in intensity at locations more distant from the right ventricular ischemic boundary.

In Table III the effects of small and large coronary artery occlusion on the area of ischemic involvement and the magnitude and polarity of the deflections recorded from sites overlying ischemic and nonischemic portions of the heart are tabulated. With the porcine heart model, production and subsequent measurement of a standardized area of ischemic involvement was easily accomplished. The ischemic area after large artery occlusion was approximately twice that observed after the smaller occlusion. The percentage increase in ischemic weight, however, was greater than this due presumably to the involvement of the thicker, more basally located portions of the anterior wall only during large artery occlusion.

The change in ischemic area had a predictable influence on the magnitude and polarity of the TQ-ST deflections recorded from all areas. Sites originally overlying the smaller ischemic area (LV apex) maintained their positive polarity but decreased in magnitude as the LV ischemic boundary was removed to a more distant location nearer the base of the heart. New sites closer to the base of the heart, which after the small artery occlusion were in nonischemic or border regions, now registered positive deflections which exceeded in magnitude those deflections recorded at more apical portions of the ventricle. Similar though less dramatic changes were observed in the RV ischemic regions, for after the large artery occlusion the RV boundary extended itself upwards towards the pulmonary outflow tract but still maintained a position parallel to and a fixed distance away from the septum (Fig. 6).

Although the area of ischemic involvement more than doubled after large artery occlusion, the sum of the individual TQ-ST segment deflections from the ischemic region, $\Sigma$TQ-ST, increased by only 67.3 $\pm10.3\%$. This was predictable on the basis of the theoretical findings and is a reasonable expectation when one considers that although the number of ischemic sites increased to the same degree as the ischemic area, the average potential deflection recorded at these sites decreased by 20%. Indeed, if the thickness of the LV ischemic boundary had not increased after large artery occlusion, the average ischemic TQ-ST deflection might have experienced an even greater decrease (and $\Sigma$TQ-ST a smaller increase) and thus might have yielded results in even closer agreement with the theoretical predictions.

In the nonischemic portions of the RV, the normal positive deflections recorded with a small area of ischemia declined in magnitude and in many instances even became negative after the large artery occlusion ($P < 0.005$). In contrast, however, the normally negative deflections recorded from the nonischemic portions of the LV appeared to increase in magnitude after large artery occlusion. These results suggested that the position of both right and left ischemic ventricular boundaries helped to determine both the polarity and magnitude of the deflections recorded at all sites.

This hypothesis was tested further as is illustrated

| Table II |
|------------------|------------------|
| **TQ-ST Segment Deflections at Multiple Ischemic** |
| **Nonischemic Sites** |
| Magnitude | Fraction (septum) |
| mV |  |
| LV |  |
| Apex | 4.25$\pm0.29$ | 1.32$\pm0.06$ |
| Base | 5.26$\pm0.36$ | 1.64$\pm0.07$ |
| Difference | 1.00$\pm0.25^*$ | 0.32$\pm0.07^*$ |
| Right ventricle | 3.21$\pm0.18$ | 1.00 |
| Right ventricle | 2.03$\pm0.15$ | 0.65$\pm0.04$ |
| RV |  |
| LV | 1.66$\pm0.32$ | 0.56$\pm0.11$ |
| Right ventricle | 1.14$\pm0.15$ | 0.38$\pm0.08$ |
| Right ventricle | 0.52$\pm0.45$ | 0.18$\pm0.16$ |

Values shown are means$\pm$SEM obtained from experiments in 17 animals. Average number of occlusions performed in each animal was 4.5$\pm0.5$. $^*$P < 0.002; paired t test.
### Table III

**Effect of Changes in Ischemic Area on the TQ-ST Segment Deflection**

<table>
<thead>
<tr>
<th></th>
<th>Small</th>
<th>Large</th>
<th>Difference</th>
<th>Change</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>%</td>
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<tr>
<td>Anatomic</td>
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<tr>
<td>LAO, mm</td>
<td>39.2±1.3</td>
<td>79.8±1.8</td>
<td>40.6±2.5*</td>
<td>105.4±9.5*</td>
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<tr>
<td>IA, cm²</td>
<td>13.0±0.6</td>
<td>28.5±1.8</td>
<td>15.6±1.9*</td>
<td>123.3±17.9*</td>
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<td>IW, g</td>
<td>19.6±1.0</td>
<td>45.3±1.6</td>
<td>25.6±1.0*</td>
<td>133.4±9.6*</td>
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<tr>
<td>#IS</td>
<td>8.1±0.3</td>
<td>16.6±0.5</td>
<td>8.5±0.5*</td>
<td>106.3±9.2*</td>
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<tr>
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</tr>
<tr>
<td>Electrical</td>
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<tr>
<td>Ischemic region</td>
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<td></td>
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<tr>
<td>LV + SEPT (ΔTQ-ST)</td>
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</tr>
<tr>
<td>Original</td>
<td>3.77±0.21</td>
<td>2.28±0.11</td>
<td>-1.49±0.22*</td>
<td>-38.5±3.6*</td>
</tr>
<tr>
<td>New</td>
<td>3.86±0.26§</td>
<td>3.01±0.14</td>
<td>-0.76±0.15*</td>
<td>-19.4±3.0*</td>
</tr>
<tr>
<td>Total</td>
<td>3.77±0.21</td>
<td>3.01±0.14</td>
<td>-0.76±0.15*</td>
<td>-19.4±3.0*</td>
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<tr>
<td>RV (ΔTQ-ST)</td>
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<tr>
<td>Original</td>
<td>1.73±0.15</td>
<td>1.32±0.17</td>
<td>-0.41±0.12†</td>
<td>-24.9±8.2†</td>
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<tr>
<td>New</td>
<td>2.62±0.28§</td>
<td>1.95±0.20</td>
<td>+0.23±0.23</td>
<td>+19.1±15.7</td>
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<tr>
<td>Total</td>
<td>1.73±0.15</td>
<td>1.95±0.20</td>
<td>+0.23±0.23</td>
<td>+19.1±15.7</td>
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<tr>
<td>ΣΔTQ-ST</td>
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<td></td>
</tr>
<tr>
<td>Original</td>
<td>28.5±2.1</td>
<td>17.4±1.1</td>
<td>-11.1±1.7*</td>
<td>-38.1±3.6*</td>
</tr>
<tr>
<td>New</td>
<td>29.8±2.9§</td>
<td>29.8±2.9§</td>
<td>0.0±0.0§</td>
<td>0.0±0.0§</td>
</tr>
<tr>
<td>Total</td>
<td>28.5±2.1</td>
<td>47.2±3.3</td>
<td>+18.7±2.5*</td>
<td>67.3±10.3*</td>
</tr>
<tr>
<td>Nonischemic region</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LV + SEPT (ΔTQ-ST)</td>
<td>-0.59±0.12</td>
<td>-1.14±0.50</td>
<td>-0.55±0.35</td>
<td>-93±59</td>
</tr>
<tr>
<td>RV (ΔTQ-ST)</td>
<td>+0.44±0.08</td>
<td>-0.25±0.12</td>
<td>-0.69±0.15*</td>
<td>-160±26*</td>
</tr>
</tbody>
</table>

**Abbreviations:** Difference, large–small; LAO, length of artery occluded; IA, area of cyanotic discoloration; IW, ischemic weight; #IS, number of electrode sites completely overlying cyanotic region; LV, left ventricle; SEPT, septal sites; RV, right ventricle; ΣΔTQ-ST, sum of ΔTQ-ST at all ischemic sites; Original, sites overlying small ischemic area; New, additional sites becoming ischemic after large artery occlusion.

Values shown are means±SEM obtained from experiments performed in eight animals. The average number of small and large artery occlusions performed in each animal was 5.0 and 2.1, respectively. ΔTQ-ST at nonischemic LV + SEPT sites after large occlusions were obtained in only three animals.

* $P < 0.005$; paired $t$ test (difference and percent change).
† $P < 0.02$, paired $t$ test (difference and percent change).
§ $P < 0.005$; paired $t$ test (original–new).

In Fig. 6, in this experiment multiple occlusions of the small coronary artery in an animal were made and the average TQ-ST (ΔTQ-ST) changes at all sites measured at 30, 60, and 90 s. During every other occlusion at precisely 65 s after the small artery occlusion, the larger artery was then occluded. Again despite increases in the amount of ischemic damage, the deflection magnitude at sites originally overlying the smaller area of ischemia began to decline, being significantly lower 25 s later than if the smaller occlusion had been maintained the full 90 s. In this example SUM(+) and SUM(−) refer to the sum, respectively, of all positive and negative ΔTQ-ST deflections measured from all 22 sites. With a sudden increase in ischemic at 65 s, sites originally overlying nonischemic regions of the LV now overlay ischemic regions and began to develop positive ΔTQ-ST. Thus these sites were no longer categorized in SUM(−) and this parameter dramatically decreased in value. SUM(+) on the other hand had still not exhibited any change at 90 s; for although it had begun to receive positive contributions from previously nonischemic sites (shaded) it also lost contributions from (a) sites from the originally ischemic area whose average deflection began to decrease (solid) and (b) nonischemic sites overlying the RV which were initially positive but whose magnitudes declined towards zero after the subsequent large occlusion (see Table III).

**Preocclusion TQ-ST segment deflection.** While preocclusion control values for the TQ-ST deflection (TQ-STo) closely approached an isoelectric value for the animal population as a whole (TQ-STo = +0.05±0.10 mV), the values in individual animals and at different electrode sites showed greater variation. This is illustrated in Fig. 7. The SEM for TQ-STo at all sites in a single animal was regularly 0.10 mV while the mean was

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often significantly different from zero (either positive or negative) depending upon the animal studied (dark circles).

Nonspatial influences. The effect of progressive hyperkalemia (3.0 to 8.0 mM) on the evolution of the TQ-ST segment deflection was studied in six animals. Results in a representative animal are shown in Fig. 8. The average ΔTQ-ST recorded from all ischemic and nonischemic regions decreased with steadily increasing plasma potassium levels varying in a linear fashion with the logarithm of the [K+]₀ level ($r = 0.974 \pm 0.013$). The value of [K+]₀ at which no changes in TQ-ST segment are expected was found by extrapolation to be 8.72±0.15 mM ($n = 5$). Changes in whole ventricular dynamics did not change significantly in this study until [K+]₀ exceeded 8.0 mM. By this time the TQ-ST segment changes were nearly nonexistent. To insure that the effect of K⁺ on the TQ-ST deflection was nonspatially mediated, in the sixth animal the polyethylene occluder was brought out of the chest cavity, saline-soaked gauze pads were placed around the heart (to insure proper conductive pathways to the precordium), and the chest was closed. Needle electrodes were then placed under the skin at various points on the right and left precordium. With 11 repeated coronary occlusions at varying [K+]₀ levels the ΔTQ-ST again declined in a logarithmic manner ($r = 0.98$).

![Graph](image)

**Figure 6** Effect of a sudden increase in ischemic area on the TQ-ST segment deflection. Average change in the TQ-ST segment (ΔTQ-ST) was measured at all sites at 30 and 60 s after seven repeated small artery occlusions (one star; solid symbols) in one animal. During occlusions 1, 3, 5, and 7 the ischemia was maintained for 100 s and the ΔTQ-ST at 90 s also measured. During occlusions 2, 4, and 6 the large artery was occluded at 65 s and ΔTQ-ST again measured at 90 s (two stars; clear symbols). Despite the increase in ischemic area as evidenced by the doubling in the number of ischemic sites, the ΔTQ-ST at all ischemic sites in the distribution of the occluded small artery (dark circles) declined in magnitude within the following 25 s. SUM + (●), the sum of all positive ΔTQ-ST at all sites remained unchanged after the large artery occlusion while the sum of the negative ΔTQ-ST, SUM−(■), rapidly declined. Dashed lines indicate ischemic boundary after small and large artery occlusions. *$P < 0.001$ (unpaired t-test). **$P < 0.01$ and ***$P < 0.05$ (paired t test at individual ischemic sites). See text for additional discussion.
FIGURE 7 Deflections from the isoelectric line in the epicardial electrogram recorded before coronary artery occlusion (TQ-ST0). TQ-ST0 values varied among different animals and electrode sites. Histogram constructed from measurements of the TQ-ST deflection at all electrode sites and in all animals used in the compilation of the data in Table II. The main trunk of the anterior descending artery divided right from left ventricular sites, however, neither spatial nor hemodynamic factors were thought to be responsible for these variations. Although the mean of this population is close to zero, the mean of the deflections in individual animals (●) did, in some instances, assume values which were significantly different from zero.

DISCUSSION

Spatial influences and epicardial mapping. After acute coronary artery occlusion in the pig TQ-ST segment deflection changes (ΔTQ-ST) during early ischemia may be recorded from all epicardial regions of the heart. This and earlier studies (12, 35–37) have demonstrated that the relative magnitudes and polarities of these deflections in each animal are determined primarily by (a) the position of the recording electrode with respect to the ischemic tissue boundaries; (b) the area of ischemic involvement; (c) the transmural shape of ischemic involvement; and (d) the thickness of the ventricular wall at the boundary.

Spatial factors which affect the TQ-ST segment deflection may be investigated in a physiologic and quantitative fashion by way of solid angle analysis. Predictions obtained from this analysis may then be tested and validated in the intact pig heart, a model which permits production and measurement of a standardized and well-defined area of ischemic involvement.

Boundary location in anteroseptal ischemia. Although it was assumed that negative TQ-ST segment deflections should be recorded from both right and left ventricular nonischemic regions, the finding of positive deflections on the RV came initially as a surprise. This finding indicated that the spherical heart model with an ischemic region of uniform wall thickness from which the theoretical curves were derived (see Appendix) rather incompletely characterized the geometry of the anteroseptal region produced experimentally. A closer look at this geometry suggested that three different boundaries were present: right, left, and septal ventricular boundaries. Each of these boundaries would make its own contribution to the total TQ-ST segment deflection recorded at each electrode site. In Fig. 9, the positions of the boundaries are depicted in a theoretic heart model containing an ischemic region of 18 cm² in area. Also shown are solid angle values calculated for a number of ischemic and nonischemic sites. It is observed that electrodes positioned over the ischemic region subtend positive solid angles, the magnitudes of which are influenced by the electrodes proximity to the different ischemic boundaries and the boundaries' thicknesses. Thus, the solid angle increases as the thicker LV boundary is approached (LV > septum > RV) as did the TQ-ST segment deflections in the experimental model. Electrodes positioned over nonischemic regions, however, sense current flow from different directions and thus the polarity and magnitudes of their solid angles

FIGURE 8 Plot of changes in the TQ-ST segment deflection recorded during multiple occlusions of the anterior descending artery in the pig at steadily increasing plasma potassium (K⁺) levels. The ΔTQ-ST recorded from both ischemic (cyanotic and nonischemic (left ventricle only) sites appeared to vary in a linear fashion with the transmembrane potential (Ek) calculated from the Nernst equation at 37°C, assuming intracellular [K⁺] = 140 mM and with the log of the K⁺ level. Values are means ± SEM. Number of electrodes overlying ischemic and schematic areas were 10 and 7, respectively.
reflect the difference (as opposed to the sum when the electrodes overlie the ischemic region) of contributions from the different boundaries. At nonischemic RV sites, despite the nearby contribution from the thin-walled RV boundary, the more distant, but thicker, septal and LV boundaries subtend the larger solid angles and therefore, as seen experimentally, positive TQ-ST segment deflections may be recorded from this nonischemic region (electrode sites B and C).

Since spatial influences on the TQ-ST deflection depend upon the position of the electrode to the various boundaries, movement of the LV boundary to a more distant location should result in the following. First, the positive voltage at the RV nonischemic sites should eventually become negative if the boundary is moved far enough. Second, the positive potentials of the ischemic regions should decrease in magnitude. Finally, the LV nonischemic sites should become slightly more negative. In the experimental model such a relocation of the LV boundary was accomplished by proximally occluding the LAD artery, thereby creating a larger area of ischemia. The results (Table III and Fig. 6) fulfilled these expectations.

Boundary vs. local influences on the TQ-ST segment deflection. Recognizing that it is the relationship of the electrode to the ischemic boundary which determines the relative magnitude and polarity of the TQ-ST segment deflection, only then can it be appreciated that although sites of TQ-ST segment elevation are usually found to overlie tissue exhibiting lactate production (38), ATP and CPK depletion (29, 38, 39), histologic abnormalities (31, 40), QRS complex alterations (5, 30, 41) and lowered PO\textsubscript{2} (42–44), contractile activity (45), and coronary blood flow (46–48), the frequent lack of quantitative relationships existing between the magnitude of the TQ-ST segment deflection at a particular site and these alternated markers should not be surprising. Although the magnitude of the TQ-ST deflection varies depending upon the electrode’s orientation to the boundary, there is no reason to expect CPK levels, histology, etc., to vary in the same manner. Recent studies in which TQ-ST segment magnitudes at individual sites on the epicardium were compared to later changes (24 h) seen in tissue blood flow (49), histologic abnormalities (40), and CPK levels (49) at the same site, demonstrate the absence of such quantitative relationships.

Electrode location and ischemic shape. Further appraisal of the solid angle concept reveals additional spatial factors such as heart size and ischemic shape which may influence the polarity as well as the magnitude of the TQ-ST deflection. Differences in ischemic shape, first considered by Prinzmetal et al. (50), provide a good example of how solid angle analysis is most useful when employed in a quantitative manner. Some of the most difficult conceptual problems with

![Figure 9](image-url)
The last of the few functionally significant collateral vessels and one expects the boundary to be narrow and well defined (24). However, in the dog, with its richer collateral blood supply (23), evidence suggests that the boundary lacks such definition (60). In this species a wide gradient of transmembrane potentials most likely exists at the periphery of the ischemic region and the boundary or separation between ischemic and nonischemic tissue is wider and much less distinct (18).

The influence of boundary width on the distribution of solid angle values at ischemic and nonischemic sites is shown in Fig. 11. Although different boundary widths do not alter the precordial distribution, changes in the epicardial distribution in the vicinity of the boundary are seen. When the boundary is wide, solid angle values now decrease, instead of increasing, as electrodes are moved from the center to the periphery of the ischemic region.

One would expect this difference in the solid angle distribution from the center to the periphery of the ischemic region to be reflected by differences in the TQ-ST segment voltages in species with (e.g., pig and baboon) and without (e.g., dog) a well-defined boundary. Studies in the dog have shown that the TQ-ST segment deflection is greater in the center of the ischemic region than at the periphery (61). Studies in the baboon, on the other hand, show only a rapid transition in polarity as electrodes are moved from the ischemic to the nonischemic region (15). The lack of uniform wall thickness of the anteroseptal ischemic region and the size of the electrodes did not permit us to investigate the possibility that TQ-ST segment deflections might actually increase as the boundary is approached in the pig. The suggestion that the boundary is both diffuse and dependent upon collateral blood flow in the dog raises a number of interesting possibilities. First, precise localization of the boundaries of the ischemic region is less likely to be accomplished in the dog than in the pig. Second, one would expect the width of this boundary to depend upon collateral vessel patency and perfusion pressure as suggested by Bayley (18). Finally, one would also expect that the area of ischemic involvement might be modified in the dog by interventions which either alter the level of collateral blood supply to this wide boundary or the oxygen demands of the tissue at the boundary. In comparison, in the pig such modification might not be possible. Here the amount of tissue becoming ischemic closely approaches that normally perfused by the occluded artery (Table 1). Modification of eventual infarct size in humans may similarly depend upon available collateral flow.

**Spatial influences and precordial mapping.** Although solid angle analysis was carried out for both precordial and epicardial locations, experimental veri-
verification was limited to the latter. In this instance the polarity and magnitude of the TQ-ST segment at each site could be associated with the position the electrode maintained with respect to the ischemic region. Similar verification at precordial sites though desirable will be difficult to accomplish. Not only does the distance from the electrode to the heart’s surface vary depending upon the electrodes’ position on the precordial, but whether changes in ischemic area and shape, heart size, etc., decrease or increase the magnitude of the TQ-ST deflection depends in turn upon this distance. This matter has received little attention despite its significant influence on the magnitude of the TQ-ST deflection. For instance, a change in the precordial-to-heart distance of only 1 cm (i.e., from 5 to 6 cm) may reduce the magnitude by over 40%. Similar variations in this distance may occur in a variety of physiologic (posture, pregnancy, etc.) and pathologic (pulmonary congestion, pneumothorax, cardiac dilatation, etc.) conditions.

TQ-ST segment data obtained at the precordium which does suggest general agreement with solid angle analysis include observations that: (a) the magnitude of the precordial TQ-ST segment deflection increases with increases in ischemic area (12, 62); and (b) maximal precordial TQ-ST segment deflections are recorded over the center of the ischemic region, decreasing at more peripheral locations (5, 63).

Nonspatial influences on the TQ-ST segment deflection. According to the solid angle theorem (Fig. 1), the TQ-ST segment deflection is also a function of the difference in transmembrane voltage existing between the normal (\(V_{mn}\)) and ischemic (\(V_m\)) cells during both diastole (TQ segment) and systole (ST segment). During early ischemia this difference is believed to arise primarily as a consequence of anoxia and potassium leakage out of the ischemic cells, and its accumulation in the surrounding extracellular space (13, 32, 64, 65). If the difference in transmembrane voltage between the normal and ischemic cells widens, the TQ-ST deflection magnitude increases. If, for any reason, these differences disappear, then so must the TQ-ST deflection.

The number of potential interventions and mechanisms by which the TQ-ST segment deflection may be modified in a nonspatial manner is large and diverse. These would include factors which directly modify the transmembrane potentials of either the normal or ischemic cells during diastole or systole (e.g., electrolytes, temperature, catecholamines, heart rate, metabolic inhibitors, and anti-arrhythmic therapy). Also included are agents which alter the transmem-

**FIGURE 11** Influence of boundary thickness (T) on the polarity and magnitude of the solid angle computed at precordial and epicardial locations. Area of ischemic involvement equaled 13.8 cm² (\(P = 0.50\) radians). At the precordium, whether the boundary is well defined (\(T = 0\)) or diffuse (\(T = 15\) mm), the solid angle values are essentially unchanged. At the epicardium, differences in solid angle values for the well-defined and diffuse boundaries occur only in the region of the boundary. Although solid angle values in the ischemic region increase in magnitude as a distinct boundary is approached, they decrease when the boundary is wide and diffuse. See text for additional discussion.
brane voltage of the ischemic tissue by changing the rate and degree of potassium leakage out of these cells. In the presence of ischemia-inhibited Na\(^+\)-K\(^+\) ATPase pump, chronotropic (66), inotropic (catecholamines [30, 67], ouabain [68], calcium [69]), and metabolic (hyperthermia [70], acidosis [71]) factors serve to accelerate potassium efflux. This facilitates potassium accumulation in the extracellular space of the ischemic tissue and causes the diastolic resting potential to decline and the action potential duration to shorten. Factors which slow potassium loss (e.g., hypothermia, potassium solutions, and myocardial depressants) may help to restore the transmembrane potential of the ischemic cells back to more normal values.

The possibility, therefore, that a change in the ischemic TQ-ST segment deflection elicited by a given intervention can be reliably and exclusively attributed to a change in the spatial geometry of the ischemic region (ischemic area and shape) and not to a nonspatial change in the transmembrane potential difference (ΔV\(_m\)) between the two tissues would seem to be unlikely. For example, one indication that propranolol may benefit patients with angina pectoris is the observation that it decreases the magnitude of the ischemic TQ-ST segment deflection (29, 72). Hemodynamically, propranolol decreases heart rate and the force and velocity of contraction (73). In this way it may reduce demands of the ischemic tissue, and may, as shown in the dog (74), help to decrease ischemic area. As shown earlier (Figs. 3–5), at the precordial a decrease in ischemic area will decrease the solid angle and the TQ-ST deflection magnitude (12, 62). There are other, nonspatial ways, however, in which propranolol may decrease the deflection. As suggested by the work of Sarnoff et al. (75), Parker et al. (66), and Polwmeni and Vassalle (76), decreases in oxygen consumption will also decrease the rate of potassium leakage and accumulation in the ischemic region. This in turn will limit the development of ΔV\(_m\) and the TQ-ST deflection magnitude will decline. Specific electrophysiologic actions of propranolol may also affect ΔV\(_m\). Kupersmith et al. (77) and Wittig and Williams (17) found that propranolol returns the action potential duration of ischemic cells back towards normal. The systolic ΔV\(_m\); therefore, is reduced, resulting in a decline in the ST and total TQ-ST segment deflection. This effect of propranolol may be related to its ability to antagonize catecholamine effects on calcium currents across the myocardial membrane (78). The mechanism by which verapamil, a specific calcium current blocker, reduces the ischemic TQ-ST segment may be similar (79, 80).

Potassium ion infusion was chosen for this study because it has minimal effects on oxygen consumption and contractile activity of the heart while having significant electrophysiologic actions. Potassium can (a) directly alter the transmembrane potential (depolarizes cells during diastole and shortens their action potential duration during systole [81]); (b) influence the rate of potassium leakage and accumulation in the extracellular space (stimulates uptake by the cell of potassium ion [82]); and (c) increase potassium conductance (83). The precise mechanism by which potassium infusion reduced the magnitude of the TQ-ST segment deflections in this study cannot be ascertained without making simultaneous measurements of transmembrane potentials before and during coronary artery occlusion in the normal and ischemic regions. The possibility that some of the changes in the TQ-ST segment may be due to hyperkalemic changes in conduction velocity (30) should also be investigated. The finding that the decline in the TQ-ST deflection appeared to vary in a linear fashion with the logarithm of K\(^+\) value is interesting, however, for both E\(_K\) and the action potential duration (which correspond to the TQ and ST segments of the electrogram) both vary with the log of K\(^+\). The latter discovery was made by examining data from Surawicz’ original study (84) and then plotting action potential duration versus the log of K\(^+\); the resultant curve was remarkably linear over the range of K\(^+\) values from 2.0 to 10.00 mM (r = 0.992). The finding that changes in the TQ-ST segment were similar at either nonischemic or ischemic regions on the epicardium and precordium suggests that potassium exerts a primarily nonspatial influence on the TQ-ST segment and thus is not dependent upon either electrode location or ischemic geometry. This nonspatial influence of potassium may help to account for any previously observed absences of TQ-ST deflections in the ECG from the acute myocardial infarction patient who is also in renal failure (85), as well as suggest an alternate mechanism by which glucose-insulin-potassium improves the ECG picture of patients with an acute myocardial infarction (86).

Preocclusion TQ-ST segment deflections. The cause of control (preocclusion) TQ-ST deflections above or below zero in a normal animal or man is not known (87, 88). In this study heart rate was kept constant (89) and electrode artifact, if present, would presumably account only for positive deflections. Since neither spatial (right vs. left ventricular sites) nor hemodynamic correlation apparently accounted for this variation, it is assumed that slight changes in the temperature at the epicardial surface, because of the open-chest preparation, might be responsible (90, 91). The conventional view, therefore, that the normal TQ-ST segment is isoelectric, is not supported by this study.

Solid angle theory and the experimental model. The solid angle theorem employed in this study is a
mathematical expression which relates the potential recorded at an electrode site to the flow of currents in homogeneous volume conductors of infinite extent. Its conception can be traced at least as far back as Helmholtz in 1853 (92) and was utilized extensively in the formulation of modern electrocardiographic theory by Wilson and Bayley (93) and McFee and Johnston (94). It requires that the recording electrode and excitable tissue both be located in a homogeneous conductor of infinite extent. However, in both clinical (precordium) and experimental (epicardium) situations the boundaries are finite and the substances surrounding the heart (skin, bones, lung, blood) of somewhat varying conductivity. These factors, however, may tend more to alter the absolute magnitude of the electrocardiographically recorded potentials than their relative distribution (18, 95). For example, Bayley showed that the potential at a point on a volume conductor of arbitrary shape (as in the case of the thorax) is approximately twice the value obtained at the same point were the volume conductor infinite in extent (18). A review of the advantages and limitations of solid angle analysis as related to ECG interpretation has recently been written (19).

Summary and conclusions. The TQ-ST segment deflection arises due to differences in transmembrane potential at the boundary between normal and ischemic tissue during diastole and systole. Nonspatial factors influence the magnitude of this potential gradient, while spatial factors alter the relative position of the boundary to the electrode site. Spatial factors including electrode location, ischemic area and shape, heart size, and wall thickness can be quantitatively analyzed with the solid angle theorem and then experimentally verified in the intact porcine heart, a model which permits production and measurement of a well-defined area of ischemic involvement. Nonspatial influences are of equal importance, though less easily analyzed in a theoretical fashion. One such influence, the elevation of plasma K⁺ levels, results in a decrease in the magnitude of the TQ-ST deflections recorded from both ischemic and nonischemic portions of the heart after coronary artery occlusion. The majority of pharmacologic agents previously shown to influence the TQ-ST segment deflection during ischemia probably do so via both spatial and nonspatial mechanisms.

APPENDIX

Spherical coordinate transform. The spherical coordinates (R, ψ, θ) of a point P in space are illustrated in Fig. 12 (left). Rectangular coordinates (X, Y, Z) are related to spherical coordinates by the equations:

\[ X = R \sin\psi \cos\theta; \quad Y = R \sin\psi \sin\theta; \quad Z = R \cos\psi. \]

The distance \( \overline{AB} \) between two points in space \( A(X_1, Y_1, Z_1) \) and \( B(X_2, Y_2, Z_2) \) can be obtained by the equation:

\[ \overline{AB} = \sqrt{(X_1 - X_2)^2 + (Y_1 - Y_2)^2 + (Z_1 - Z_2)^2}. \]

Solid angle computations. In Fig. 12 (right) a small segment of ventricular wall is diagrammed illustrating the normal and ischemic (shaded) myocardial tissue with the boundary between them. The solid angle constructed at the electrode site N by the ischemic boundary may be computed from:

\[ \Omega = \int_0^{\pi} \left[ \cos\alpha - \cos\psi \right] \: d\theta, \]

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where $\alpha$ and $\gamma$ are obtained from the Cosine Law or:

$$\alpha = \cos^{-1}\left(\frac{KL^2 - KN^2 - LN^2}{2 \cdot KN \cdot LN}\right)$$

and

$$\gamma = \cos^{-1}\left(\frac{LM^2 - MN^2 - LN^2}{2 \cdot MN \cdot LN}\right).$$

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