Brugada Syndrome and Carbohydrate Metabolism

Abstract

Brugada syndrome is an arrhythmogenic and autosomal dominant disease with incomplete penetrance that may cause syncope and sudden cardiac death in young individuals with a normal heart. In addition, it is the syndrome of right bundle branch block (RBBB) that characterized by ST-segment elevation in precordial ECG leads V1 to V3. There are many factors contributing to the syndrome, including genetic factors, cellular and ionic mechanisms abnormalities, exercise, carbohydrate (CHO) and blood electrolyte. Thai ministry of Public Health reported in 1990 that there was an association between a large meal of glutinous ("sticky") rice or carbohydrates ingested on the night of death in Brugada syndrome victims. Some investigators have reported that a glucose load, glucose alone, glucose combined with insulin infusion, and Thai high glycemic index (HGI) meals influenced ST-segment elevation in Brugada syndrome patients. A reduction in Ikr or Ito or an increase in Ito and/or any other potassium current may change the configuration of the action potential notch that leads to J-point elevation on the ECG. Food ingestion is accompanied by the release of hormones such as insulin, glucagon and gut peptides. A larger meal produced a longer gastric emptying and a higher effect on postprandial profile of the gut hormone. Insulin is an essential hormone that controls the level of BG. Food ingestion activated sympathetic activity indicating by an increased postprandial plasma noradrenaline. Increased postprandial sympathetic nerve activity muscle. In addition, it was known that HGI and carbohydrate load contribute to high CHO oxidation rate. According to the above information, the increasing of digestive related hormones and the CHO oxidation rate induced by having HGI or CHO load may result in abnormal metabolic and autonomic responses in these patients.

Keywords: Brugada syndrome; Electrocardiogram; Electrolyte; Heart rate variability; Glycemic index

Introduction

Brugada syndrome

Brugada syndrome is recognized worldwide, although the prevalence seems much higher in some areas. In Asia, the plenty of cases were reported from Japan, Thailand, Taiwan, China, India, Laos, Vietnam, Singapore and Cambodia. The incidence of the syndrome was nine times higher among men than women and the average age was 45 ± 10.5 years [1]. For Southeast Asian countries, Brugada syndrome was the major cause of sudden death in younger population [2,3]. In Thailand, an epidemiological survey of young men living along the northeastern border of Thailand abutting Laos and Cambodia found that the annual SUDS death rate was 26-38 per 100,000 men (range, 20-49 years old) [4,5].

Brugada syndrome is an arrhythmogenic and autosomal dominant disease with incomplete penetrance that may cause syncope and sudden cardiac death (SCD) in young individuals with a normal heart [6]. In addition, it is the syndrome of right bundle branch block (RBBB) that characterized by ST-segment elevation in precordial ECG leads V1 to V3 [7]. There are many factors contributing to the syndrome, including genetic factors, cellular and ionic mechanisms abnormalities, gender difference, exercise, autonomic nervous system, and blood electrolyte [8]. A genetic mechanism explaining the pathology of the syndrome is beyond this review.

Brugada syndrome and carbohydrate metabolism

Based on Thai ministry of Public Health report in 1990 there was an association between a large meal of glutinous ("sticky") rice or carbohydrates ingested on the night of death in Brugada syndrome victims [9]. This report motivated many investigators to approve that association. In general, glutinous rice (Oryza sativa) is one type of carbohydrate staple food of Asian people. Glycaemic index (GI) value is defined as blood glucose produced after the ingestion of a carbohydrate food [10]. In fact, there are large ranges of GI of rice which are due to variations of rice’s species [11], process conditions [12,13] and differences in amylase and amylopectin content. The GI value of the glutinous rice when compared with glucose is 98 [14] and CHO composition per 100 g of the glutinous rice is 52 g [15].

Some investigators have reported that a glucose load [16], glucose alone [16], glucose combined with insulin infusion [17],
and Thai HGI meals [18] influenced ST-segment elevation in Brugada syndrome patients. It has been known that the epicardial action potential notch is the result of the balance between three different currents: $I_{\text{Na}}$, $I_{\text{CaL}}$ and $I_{\text{CaT}}$ [19]. Theoretically, a reduction in $I_{\text{Na}}$ or $I_{\text{CaL}}$, or an increase in $I_{\text{CaT}}$, and/or any other potassium current may cause the changes of the configuration of the action potential notch [19,20] that leads to J-point elevation on the ECG. In the general concept, glucose induces insulin secretion. From many previous studies, insulin has been shown to cause hyperpolarization of membrane potential by activation of the Na$^+$ and K$^+$ pump [20,21], which activated outward current during the plateau phase of action potential. Therefore, it tends to promote outward current. In addition, activation of the Na$^+$ and K$^+$ pump also could produce a high K$^+$ gradient across the cell membrane with depletion of K$^+$ from the extracellular space. However, the study of Nishizaki and co-workers found serum K$^+$ concentrations were unchanged after the glucose load. They still proposed the possibility that increased K$^+$ gradient could affect the repolarization process differently among the three regions of the ventricular walls (endocardium, epicardium and myocardium) cannot be excluded and might contribute to the ST-segment abnormalities observed in patients with Brugada syndrome [16].

Aulbach and co-workers have reported that insulin stimulates the $I_{\text{CaL}}$ in isolated rat ventricular myocytes in a dose-dependent manner [22]. Therefore, it tends to promote inward current that thought to reduce ST-segment elevation [23,24]. Until now, it seems to be not fully understood how insulin affects the repolarizing currents. However, Nishizaki and co-workers suggested that an overall insulin effect on ST-segment elevation in Brugada syndrome patients might be a balance between the modulation of Na$^+$ and K$^+$ pump current and $I_{\text{CaL}}$, and additional influences on other channels or regional differences of its effects on repolarizing currents. Moreover, the report from previous studies proposed that the glucose load would change the characteristics of other factors, such as the ANS, plasma viscosity, and other ion channels [25] which might influence ST segment to fluctuate occasionally in Brugada syndrome patients. Thereafter, it needed to be explored further.

Due to the Brugada ECG which was characterized by a prominent coved ST-segment elevation displaying a wave amplitude or ST-segment elevation ≥ 2 mm or 0.2 mV at its peak followed by a negative T-wave, with little or no isoelectric separation the earlier study by Nogami and co-workers investigated infusion of glucose or glucose and insulin solution then identified the hypokalemia of membrane potential by activation of the Na$^+$ and insulin secretion. Also, it was known that HGI and CHO load contribute to high CHO oxidation rate [32]. According to the above information, an increasing of digestive related hormones and the CHO oxidation rate induced by having HGI or CHO load may result in abnormal metabolic and autonomic responses in these Brugada Syndrome patients.

Conclusion

This review shows that there are many factors contributing to the syndrome, including genetic factors, cellular and ionic mechanisms abnormalities, gender difference, exercise, autonomic nervous system, and blood electrolyte [8]. In addition, an increasing of digestive related hormones and the CHO oxidation rate induced by having HGI or CHO load may result in abnormal metabolic and autonomic responses in these Brugada Syndrome patients.

References


