Neuro-motor mechanisms of Pharyngo-esophageal Motility in Dysphagic Infants with Congenital Heart Disease

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Abstract

INTRODUCTION—Aero-digestive morbidities are common in congenital heart disease infants and mechanisms are unclear. We hypothesized that adaptive pharyngo-esophageal motility reflexes are different in surgical congenital heart disease infants (S-CHD) vs. nonsurgical congenital heart disease infants (CHD) and healthy controls.

METHODS—Abrupt pharyngeal provocation was performed with graded water infusions using purpose-built micro-manometry. Data from 12 S-CHD were compared with 10 CHD and 12 controls. 197 water stimulations were examined for the frequency, latency, duration and magnitude of Pharyngo-Upper Esophageal Sphincter contractile response (PUCR), Pharyngeal reflexive swallow (PRS), esophageal body peristalsis and lower esophageal sphincter (LES) relaxation characteristics. Mixed statistical models were applied.

RESULTS—Frequency distribution (%) of PUCR: PRS: None in S-CHD vs. CHD vs. controls respectively were 36:46:17 vs. 9:80:11 vs. 15:61:24 (p < 0.05). Response latency to the final esophageal body waveform (p = 0.01) and the response duration of esophageal body peristalsis (p = 0.04) were prolonged in S-CHD vs. controls but were similar to CHD (p = 0.22). Pharyngeal infusion induced LES relaxation characteristics were similar in all 3 groups.

CONCLUSIONS—Abnormality in the recruitment of PUCR or PRS reflexes and esophageal body peristalsis in S-CHD implicate dysregulation in vagal cholinergic excitatory neuromotor responses.
Introduction

Congenital heart defects are the most common type of birth defects affecting nearly 1% or approximately 40,000 births per year in the USA (1). These infants need parenteral nutrition, innovative enteral feeding strategies or prolonged respiratory support (2-4). The prevalence of feeding disorders in post-surgical infants with congenital heart disease varies from 22% to 50% (5-7). Poor nutritional status resulting from inadequate feeding capabilities leads to an imbalance of energy intake resulting in growth failure. Malnutrition is a major problem, which affects the subsequent stages of cardiovascular surgery (8). Furthermore, the acquisition of feeding skills is further delayed among infants with cyanotic congenital heart disease compared to acyanotic congenital heart disease and foregut dysmotility and oropharyngeal dysphagia neuromotor mechanisms are often implicated but have not been systematically evaluated before (9).

Antecedent conditions for the prototype of the dysphagic infant may include but are not limited to: aero-digestive tract manipulation and surgical trauma in the form of injury to thoracic visceral innervations, post-surgical inflammation, chronic mechanical ventilation, changes in circulation and influence of anesthesia and narcotics. Furthermore, the risk factors for laryngopharyngeal dysfunction and dysphagia in such infants also include preoperative acuity, duration of intubation, types of congenital heart defect, vocal cord injury, growth characteristics at birth and the type and duration of surgical procedures (5,6,10-13). Alternatively, feeding difficulties may follow underlying central neurological sequelae or immature brain development (14,15). Regardless of the etio-pathological mechanisms for the infant with dysphagia, the final level of functional swallowing coordination and safe regulation occurs at the level of pharyngoesophageal level. These observations form the basis for the current study.

Clarification of the underlying patho-physiological mechanisms of pharyngo-esophageal motility reflexes can help us devise better evidence-based management strategies for the prevention and treatment of infant feeding disorders. In the current study, our objective was to test the hypothesis that dysphagic infants with congenital heart disease who underwent cardiac surgery (S-CHD group) have distinct basal and adaptive pharyngo-esophageal motility characteristics compared with that of healthy controls or those infants who did not undergo cardiac surgery (CHD group).

Methods

Participants

Infants with congenital heart disease (n = 22, 10 males) born at gestational age (GA) range 25 – 40 weeks were evaluated between 38-52 weeks post menstrual age (PMA) for dysphagia at the neonatal and infant feeding disorders program, at the Nationwide Children’s Hospital, Columbus, Ohio. Among these, twelve infants underwent cardiac surgeries for severe congenital heart disease for various reasons and 10 infants did not undergo cardiac surgery. To compare data, 12 healthy control infants that had independent oral feeding skills (6 male; born at GA range 24-40 weeks) were studied at PMA range 37-45 weeks. These controls were part of other ongoing research studies, and were feeding and
thriving appropriately during the hospital course and did not have any genetic or birth defects. All subjects were evaluated by us and the primary neonatologist and were deemed appropriate for study procedures. The study observations reported are from retrospectively collected pharyngo-esophageal manometry data recordings, and IRB approvals were obtained at the Nationwide Children’s Hospital Research Institute, Columbus, Ohio. Written, informed consents were obtained from parents before study, and HIPAA compliance was followed.

Manometry Methods

The use of esophageal manometry methods and multimodal provocation techniques in neonates was described by our group (16-19). Briefly, the catheter assembly (Dentsleeve International; Mui Scientific, Mississauga, Ontario, Canada) was connected to the pneumohydraulic micromanometric water perfusion system via the resistors, pressure transducers and amplifiers (solar modules, Solar 2; MMS Medical Instruments, Dover, NH). The esophageal manometry catheter assembly with dual sleeves [recording from Upper Esophageal Sphincter (UES) and Lower Esophageal Sphincter (LES)] and four side ports recording from pharynx, proximal, middle, and distal esophageal loci and a terminal gastric recording port was used. The catheter also includes a dedicated infusion channel to provide esophageal stimulus. The water perfusion rate was 0.02 ml/min per port for esophageal ports, 0.01 ml/min per port for the pharyngeal port, and 0.04 ml/min per port for the sleeves. The catheter was passed nasally in the unsedated supine lying neonate, and all studies were done in the same manner, with the transducers at the level of the subject’s esophagus (midaxillary line). Vital signs were monitored for safety during the manometry study.

Manometric Experimental Protocol

Continuous data acquisition and analysis were performed based on manometric waveform characteristics (16-19). Pharyngeal provocations with sterile water were performed to test the effects of osmosensitive stimulation. During catheter placement and pull through, the UES and LES sleeves were positioned such that they straddled the UES and LES high-pressure zones respectively, and were identified by the presence of a consistent increase in pressure of 5.0 mmHg above the baseline for at least 15 s, in addition to the changes in pressure with respiration. After neonates were allowed to adapt for about 15 min, we evaluated responses to pharyngeal provocation. In our study we gave graded infusion volumes of water (0.1 ml, 0.3 ml and 0.5 ml) into the pharynx with each dose given thrice. However, in non-responders higher volume infusions were not given in some cases due concerns for patient’s safety.

Manometry Data Analysis

The signature manometric waveforms related to UES, LES, and esophageal reflex characteristics were analyzed as defined before (16-20). Briefly, Pharyngeal reflexive swallow (PRS) was defined as a deglutition response within 5 seconds of pharyngeal infusion, which begins with onset of the pharyngeal waveform associated with UES relaxation, propagates into the proximal, middle, and distal esophageal segments, and is accompanied by LES relaxation. On the other hand, Pharyngo-UES-contractile reflex (PUCR) was defined as an increase in UES pressure > 4 mm Hg within 5 seconds of
pharyngeal infusion (18). Response latency was defined as the duration from the onset of pharyngeal stimulus to the onset of PRS or PUCR or peristaltic reflex. As referenced before, resting UES pressure and LES pressure were measured (16-20). Resting pressures were measured as an average of five pressure measurements at the end of expiration observed before the onset of stimulus. Response latency to PUCR was defined as the time taken from the onset of stimulus for an increase in UES pressure of at least 4 mmHg above baseline. Maximum UES contractile pressure was also measured to ascertain the UES contractile reflex magnitude as a difference in onset and peak UES contractile pressures. Specific to the present study, LES resting pressure and relaxation were evaluated as follows: 1) LES resting pressure was evaluated before the onset of the infusion, as a mean of five observations of LES pressure in relation to gastric pressure at end expiration 2) LES response latency is duration from the onset of pharyngeal infusion to onset of LES relaxation i.e. when LES pressure dropped at least 5 mmHg below LES resting pressure. 3) Duration of active LES relaxation was defined as duration from the onset of LES relaxation to the start of LES nadir 4) Nadir was defined as the lowest pressure point reached by the LES, and nadir pressure was the lowest pressure taken during nadir duration 5) Nadir duration was considered as the period during which LES pressure dropped to 5.0 mmHg or less and up to the point LES pressure recovered to 5.0 mmHg All pressures were taken at the end of expiration and in relation to gastric pressure (20).

**Statistical Methods**

Subject characteristics, manometric measurements, and outcome variables were compared between the S-CHD, CHD and control groups. Multinomial mixed models and linear mixed models with compound symmetry matrix were used to analyze the repeated measures data. Statistical tests were adjusted for multiple comparisons using the Tukey-Kramer method. These models were fit using PROC GENMOD for categorical responses and PROC MIXED for continuous responses in SAS (SAS v.9.2; SAS Institute, Cary, NC). To evaluate the effect of graded volume on the recruitment of reflexes, we used ANOVA and Chi square and compared between the groups (Figure 3). Descriptive data are reported as least-square means ± SE, percentages, or as range unless stated otherwise.

**Results**

**Demographic and disease characteristics**

Demographic characteristics are summarized in table 1. At evaluation, all the 12 control infants had independent oral feeding skills contrasting 12 S-CHD infants and 10 CHD infants who were dysphagic and nasogastric tube-dependent. Specific symptom profiles in congenital heart disease infants were: poor oral extraction, (n= 4, 18%), aero-digestive symptoms such as bradycardia, oxygen desaturation, coughing and arching during swallowing (n=9, 41%), arching and irritability (n=14, 64%) and feeding aversion or behavioral symptoms (n=2, 9%). 50% of infants in S-CHD group and 40% of infants in CHD group had gastrostomy tube for feeding at discharge in contrast to none in control group (p < 0.05). Table 2 & 3 provides the detail for the individual demographic and disease characteristics of S-CHD and CHD infants respectively.
**Recruitment Frequency of Pharyngo-UES Responses to Pharyngeal Stimuli**

Responses to 74, 67 & 56 pharyngeal water stimulations were analyzable for the aerodigestive reflex characteristics in the S-CHD, CHD and control groups respectively. The cumulative response rate of PUCR and PRS was 81% (54 of 67) in S-CHD vs. 89% (50 out of 56) CHD vs. 72% (53 out of 74) in control ($p = 0.047$). However, further analysis of immediate first responses revealed that the distribution frequency of PUCR, PRS and none were different (Figure 1 and 2). Further analysis revealed that volume-response analysis was different for PUCR (Figure 3).

**Impact of flow rates on type of response**

The various infusion volumes were given abruptly in all cases. The catheters are not rigid as they are made of silicone and offer resistance. Hence the infusion flow rates are different but on further analysis, the infusion flow rates were similar in 3 groups (ANOVA, $p = 0.73$). On further analysis, we found that the distribution of infusion flow rates were similar when PUCR and PRS response were present (ANOVA, $p =0.49$). Using logistic regression, the flow rates were not found to affect the type of response (PUCR or PRS) ($p =0.49$). On further analysis even within the 3 groups the flow rates did not affect the type of response S-CHD ($p = 0.80$), CHD ($p = 0.91$) and control ($p = 0.31$).

**Pharyngeal & Upper Esophageal Sphincter response characteristics**

In S-CHD vs. CHD vs. controls respectively, resting UES pressures (12.3 ± 1.9 vs. 8.7 ± 2.0 vs. 13.5 ± 2.2 mmHg respectively,) and response latency of PUCR and PRS (4.3 ± 0.6 vs. 4.8 ± 0.6 vs. 5.0 ± 0.6 seconds) were similar ($p > 0.05$, Figure 4).

**Esophageal body peristaltic response**

Response latency to esophageal peristalsis was similar in all 3 groups (Figure 4). On the other hand, S-CHD infants showed prolonged duration of esophageal peristalsis compared to controls (Figure 4). For the esophageal peristaltic responses, 49 out of 54 (90.7%) in S-CHD, 46 out of 50 (92%) in CHD and 51 out 53 (96%) in Controls were completely propagated. The remainder of esophageal peristaltic responses failed or incompletely propagated. There was no retrograde esophageal peristaltic propagation seen in any of the groups. Among the esophageal peristalsis that were completely propagated, 83.7% from S-CHD, 78.2% from CHD and 90.1% from Controls propagated in anterograde manner and remainder propagated in synchronous manner ($p = 0.27$).

**Lower esophageal sphincter relaxation characteristics**

Resting LES pressures were similar in S-CHD vs. CHD vs. controls (17.1 ± 1.8 vs. 18.8 ± 1.9 vs. 19.3 ± 2.1 mmHg respectively). LES relaxation frequency and type, LES nadir pressure, Latency to LES relaxation, LES relaxation duration, LES nadir duration were all similar ($p > 0.05$, Table 4).
DISCUSSION

Oro-pharyngeal dysphagia and feeding difficulties are common in infants with congenital heart disease, and mechanisms remain unclear. This is the first study that attempted to clarify the effects of pharyngeal provocation on the upper and lower esophageal sphincter functions in addition to peristaltic reflexes, in a prototype of infants with congenital heart disease. In the current study, we demonstrated significant pharyngo-esophageal dysmotility mechanisms in the infants with congenital heart disease especially among those that underwent cardiac surgery. Importantly, peristaltic reflexes, UES contractile responses were aberrant while LES relaxation responses were preserved. Also, our data suggest increased efferent cholinergic excitatory activity as the basis for: (1) the increased frequency of pharyngo-UES contractile responses and (2) decreased cholinergic inhibitory activity resulting in prolonged esophageal body response in congenital heart disease infants that underwent cardiac surgery. However the response latencies of studied reflexes were similar in these infants compared to controls, signifying normal reaction to evoke a motor response. Interestingly, we observed that the cholinergic inhibitory responses at the LES level are unaffected, thus implying that nitric oxide or vasoactive intestinal polypeptide mediated LES relaxation is intact.

S-CHD infants show longer duration of esophageal peristalsis and longer duration from stimulus onset to terminal esophageal response compared to controls. This suggests that cholinergic excitatory activity at the level of esophageal smooth muscle is prolonged implicating slower clearance, which is further supported by delay in restoring esophageal quiescence. Alternatively, these findings also imply prolonged afferent sensitivity resulting in prolonged efferent motor outputs at the esophageal level.

Aerodigestive reflex responses are important in the clearance of aero-digestive tract during deglutition in addition to maintaining airway safety. Using a novel experimental design and provocative interrogation of pharyngo-esophageal motility reflexes, we determined the sensory-motor characteristics. Prevalence of feeding difficulties in infants with CHD is high especially in those undergoing cardiac surgeries with incidences varying from 22-50% (5-7). It is likely that the effects of associated patho-biologic factors such as inflammation, surgical and visceral trauma, circulatory changes, chronic ventilation or airway disease, may have altered sensory-motor characteristics of the studied reflexes (5,6,11) (21-23). Inflammation modifies the myoelectric and peristaltic functions of viscera (24,25). Also, it is well known various neural pathways are affected due to thoracic surgeries and various factors such as intubations or hypothermia have been attributed (26,27). These factors potentially alter pharyngo-esophageal reflexes responsible for pharyngo-esophageal clearance leading to aerodigestive problems (28).

The selection of subjects in the current study is by natural referral, and may represent the severe forms of dysphagia. Nevertheless we made pilot observations such that there are differences between the surgical CHD and non-surgical CHD. Indeed, the severity of CHD may vary between subjects, and both of these groups are representative of heterogeneity with varying severity, surgical approaches and variable duration of post-operative care.
Further studies are needed to clarify differences in mechanisms in specific congenital heart defects before and after major interventions.

The implications of this study are several. It is well recognized that infants with differing congenital heart conditions have difficulty transitioning from tube to oral feeding. This is a first attempt at identifying potential causes for such issues. It is commonly speculated that surgical manipulation and perioperative course in the congenital heart disease infants interferes with central and enteric nervous system communication with potential for significant aerodigestive maladaptation affecting safety and function. Our study supports this concept. Pharyngo-UES contractile reflex is the frequent response to pharyngeal infusion in adults in contrast to neonates in whom PRS is common response (18,29). Though delineation of exact pathways is difficult in human subjects, earlier study in cats has shown that glossopharyngeal nerve acts as afferent while vagal nerve acts as efferent for this reflex (30). In our study we have shown that response latency of Pharynx, UES and esophagus being similar in all 3 groups points to relative sparing of afferent pathways. At the same time prolonged esophageal peristaltic motor activity point to involvement of vagal efferent motor neural pathways. Further studies in this domain will provide a better understanding of the etio-pathological mechanisms and open doors for the development of preventive and interventional strategies to assist infants with congenital heart conditions become more successful at oral feeding. This would be an important area of clinical research. Minimally invasive surgery and lesser manipulation of esophagus during the surgery may reduce local injury and or inflammation around the neural pathways; thus potentially helping in prevention of dysphagia. Early oro-motor therapy may help in redevelopment of neuronal pathways to Upper esophageal sphincter thereby ameliorating dysphagia. Development of mechanisms of neuroplasticity, sucking-swallowing-peristalsis rhythms, and appropriate oral sensory processing may help in regeneration and restoration of neuromotor functions.

This study has limitation in that the patient population was heterogeneous with variety of different types of congenital heart disease and underwent different cardiac surgeries. In addition, temporal changes in pharyngo-esophageal motility due to maturational changes can happen with time, and we need further studies to elucidate the maturational or modifiable changes. Limited specific CHD subsets did not permit us to perform sub group analysis. In patients who did not respond to lower volume of infusions, higher volume infusions were not given in some cases due to concerns for patient’s safety. In such cases threshold volume may be generally greater thus supporting extreme hyposensitivity. Generally, we see the responses to 0.3 ml in 90% of infusions (18).

Feeding difficulties in children at risk for undergoing cardiac surgeries further lead to inadequate nutritional intake, resulting in poor growth. Also, a strong association exists between decrease in weight for age and mortality during the first year of life and during the first months following surgery for correction of congenital cardiac disease (31). The neuro-motor dysfunction and immaturity of cerebral development that exists in infants with congenital heart disease also places them at risk for aerodigestive problems (14,15). With this in mind, multiple studies, surgeries, and medications may put them at additional risk given the lack of proven therapies for esophageal or aerodigestive malfunctions. Procedures such as gastrostomy or fundoplication, although frequently performed, do not account for
underlying dysmotility. Given the fact that LES characteristics were normal in infants with congenital heart disease, fundoplication should be performed with caution since increasing LES tone further may lead to worsening of dysphagia in these infants as well as increasing chances of anterograde aspiration in dysphagic infants. In the current study, the dysmotility mechanisms were indicative of proximal aero-digestive maladaptation. Furthermore, infants with CHD who got gastric fundoplication have been found to be associated with significantly increased inter stage mortality than infants with CHD who did not get this procedure (32).

In summary, based on the current study, UES hyperactivity may be an early therapeutic target. This is the first study to highlight these findings. Differences as well as longitudinal changes in UES and LES functions in infants with congenital heart diseases will need further investigations.

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Figure 1. Frequency distribution of PUCR & PRS as primary response to pharyngeal provocation

□ PUCR ■ PRS □ No response. * $p < 0.001$, † $p = 0.02$, § $p < 0.001$. PUCR occurred more frequently in S-CHD vs. CHD ($p = 0.02$) and Controls ($p < 0.001$). PUCR: Pharyngo-UES contractile response, PRS: Pharyngeal reflexive swallows, NR: No response.
Figure 2. Pharyngo-Esophageal manometry: Effect on UES of 0.3 ml pharyngeal water infusion in A. Control, B. S-CHD & C. CHD
In response to pharyngeal water infusion, UES contraction is seen in S-CHD vs. UES relaxation in Control and CHD.
Figure 3. Distribution of PUCR at different stimulus volumes

Overall, stimulus-response relationships were evident with increased recruitment of PUCR frequency with increasing volumes (S-CHD vs. CHD, \( p = 0.06 \), ANOVA; and S-CHD vs. Control, \( p = 0.06 \), ANOVA). Specifically, PUCR frequency is similar in all 3 groups at 0.1 ml. In contrast, at 0.3 ml stimulus, PUCR frequency is significantly higher in S-CHD vs. CHD (\( p = 0.002 \)) and S-CHD vs. controls (\( p = 0.01 \)). However, with 0.5 ml infusions, increased recruitment of PUCR frequency were noted though not statistically significant (S-CHD vs. CHD, \( p = 0.16 \), and S-CHD vs. controls, \( p = 0.06 \)); this may be due to limited number of 0.5 ml infusions administered.
Figure 4. Evaluation of relationship between stimulus and peristaltic response

Control □ CHD ■ S-CHD * p = 0.04, † p = 0.01. Response latency was similar in all 3 groups for PUCR or PRS and onset of esophageal waveform. However duration of esophageal peristalsis and duration from stimulus onset to terminal esophageal waveform was significantly longer in S-CHD vs. control (p < 0.05).
Table 1
Demographic and Outcome characteristics

| Characteristics     | S-CHD N=12 | CHD N=10 | Controls N=12 |
|---------------------|------------|----------|---------------|
| Birth GA, weeks     | 32.6 ± 1.1 | 32.9 ± 1.6 | 32.8 ± 1.6   |
| PMA at study, weeks | 46.8 ± 1.4* † | 42.5±0.7* | 40.6±0.7†    |
| Birth weight, Kg    | 1.64 ± 0.2 | 2 ± 0.4   | 2.3 ± 0.4    |
| Weight at study, Kg | 3.44 ± 0.75| 3.3 ± 0.64| 3.86 ± 1.06  |
| Male, n (%)         | 7(58)      | 3(25)     | 6(50)         |

Values are Mean ± SE or as stated.

* \( p = 0.003 \),
† \( p < 0.001 \).

Other characteristics were similar in all 3 groups \( p > 0.05 \).
### Table 2

**Characteristics of S-CHD infants**

| Patient ID. | Cardiac defect                        | Cardiac Surgery                                      | GA (wks) | PMA at Surgery (wks) | PMA at study (wks) |
|-------------|---------------------------------------|------------------------------------------------------|----------|----------------------|--------------------|
| 1           | Critical aortic valve stenosis        | Damus-Kaye-Stansel, Blalock-Taussig shunt            | 29       | 39.9                 | 48.9               |
| 2           | Hypoplastic right heart syndrome      | Right modified Blalock-Taussig shunt                 | 34       | 43.6                 | 51.1               |
| 3           | Coarctation of aorta, VSD             | Aortic arch augmentation, VSD closure                 | 33       | 39.4                 | 42.1               |
| 4           | Interrupted aortic arch               | Aortic arch repair and augmentation                  | 29       | 29.3                 | 45.6               |
| 5           | Hypoplastic left heart, VSD, ASD, PDA | Coarctation repair, Pulmonary vavloplasty            | 39       | 41.4                 | 47.7               |
| 6           | Dilated cardiomyopathy               | Heart transplant                                     | 40       | 47.1                 | 52.7               |
| 7           | Transposition of Great arteries       | Great arteries switch                                | 27.7     | 40.4                 | 48.1               |
| 8           | ASD, VSD, PDA                        | ASD & VSD patch, PDA ligation                       | 33       | 44.3                 | 49                 |
| 9           | Coarctation of Aorta, PDA             | Coarctation repair, PDA ligation                     | 31       | 37.0                 | 40.0               |
| 10          | ASD                                   | ASD patch                                            | 31       | 47.4                 | 49                 |
| 11          | Complete heart block Tetralogy of Fallot | Cardiac Pacemaker implantation                   | 32       | 32.1                 | 41.9               |
| 12          | Tetralogy of Fallot                   | Right ventricle outflow tract patch repair, VSD patch repair and a ASD and PDA closed | 33       | 42                   | 45.7               |

ASD - Atrial septal defect, VSD - Ventricular septal defect, PDA - Patent ductus arteriosus.
Table 3

Characteristics of CHD infants

| Patient ID | Cardiac defect                        | GA in weeks | PMA at study in weeks |
|------------|---------------------------------------|-------------|-----------------------|
| 1          | VSD, ASD                              | 35          | 45.7                  |
| 2          | VSD                                   | 27          | 39.9                  |
| 3          | VSD                                   | 36          | 45.0                  |
| 4          | VSD                                   | 39          | 41.3                  |
| 5          | Pulmonary valve stenosis, ASD         | 25          | 38.5                  |
| 6          | VSD                                   | 36          | 42.6                  |
| 7          | Complete AV canal defect              | 38          | 45.3                  |
| 8          | ASD                                   | 35          | 40.9                  |
| 9          | Pulmonary valve stenosis              | 30          | 41.1                  |
| 10         | L-transposition of great arteries, ASD| 28          | 40.71                 |

ASD - Atrial septal defect, VSD - Ventricular septal defect
## Table 4
Characteristics of LES relaxation reflex

| Characteristics                              | S-CHD | CHD   | Controls |
|----------------------------------------------|-------|-------|----------|
| LES Relaxation, frequency, %                 | 79%   | 82%   | 72%      |
| Response Latency to LES Relaxation, sec      | 4.6 ± 0.6 | 5.1 ± 0.6 | 5.6 ± 0.7 |
| Duration of active LES Relaxation, sec       | 2.3 ± 0.5 | 3.0 ± 0.6 | 2.9 ± 0.5 |
| Duration of complete LES Relaxation Nadir, sec | 22.4 ± 4.9 | 15.0 ± 5.3 | 13.0 ± 2.1 |
| Magnitude of LES Relaxation, Nadir pressure, mmHg | -1.5 ± 0.8 | -0.8 ± 0.9 | -1.4 ± 1.0 |

Values are Least Square Mean ± SE or as stated otherwise

LES relaxation characteristics were similar in all 3 groups ($p > 0.05$).