- 1 Clinical profile and effects of ductal size on anthropometry of children with Patent
- 2 ductus arteriosus (PDA)

- 4 Abstract
- 5 Background
- 6 It is not known at which size of PDA do severe malnutrition ensue in children, neither is it
- 7 known, the effect of ductal size on anthropometry of children with PDA.
- 8 **Objectives**
- 9 This study was aimed to determine if ductal size had any effect on anthropometry of
- 10 children with PDA and at which size do severe malnutrition ensues in children with PDA.
- 11 **Methods**
- 12 This is an observational cross-sectional study on children who presented with PDA over a
- 13 five-year period in three tertiary institutions.
- 14 **Results**
- 15 Although there was a negative correlation between the size of PDA and the weight of
- patients, the correlation was not significant (Pearson correlation coefficient = -0.1, p = 0.7).
- 17 There was a negative correlation between the size of PDA and patient's height/length, but
- the correlation was still not significant (correlation coefficient = -0.1, p = 0.5).
- 19 The association of size of PDA with the severity of malnutrition, showed greater proportion
- 20 35.3% (6/17) of wasting and stunting in patients who had large PDA compared with fewer
- 21 proportion, 26.1% (6/23) in those whose PDA sizes were 3-6mm and 33.3% (10/30) among
- 22 those with tiny PDA <3mm (χ 2 = 10.21, p = 0.8). Although there is a positive correlation
- between ductal size and nutritional status of patients, severe malnutrition ensues from ductal
- size of 3.2mm.

- 25 The majority of children with PDA presents with severe forms of malnutrition (wasting and
- stunting). Severe malnutrition ensues when ductal size is 3.2mm. The size of PDA has no
- 27 effect on weight and height of children with PDA.
- 28 **Keywords:** children; PDA; ductal size; anthropometry

29 Key Messages

- 30 1. Majority of children with PDA present with severe forms of malnutrition.
- 2. Symptoms of severe malnutrition ensues when the ductal size is 3.2mm.
- 32 3. There is no gender difference in the severity of malnutrition among children with PDA

Introduction

- Patent ductus arteriosus (PDA), occurs when there is a persistent communication between the
- descending a orta and the left pulmonary artery. [1] This is usually due to the failure of the
- 36 closure of the ductus arteriosus. [1]
- 37 PDA could also coexist with other congenital heart anomaly or could even occur as a ductal
- dependent lesson as in TGA with an intact septum and critical pulmonary stenosis. [2,3]
- 39 The reported prevalence of PDA in term neonates is 1 in 2,000 births, accounting for 5%–
- 40 10% of all congenital heart disease. [4] These prevalence rates are higher in preterm neonates
- 41 with values ranging from 20%–60%. [5] The increased prevalence in the preterm infant is
- 42 probably due to the lack of normal closure mechanisms from immaturity. [5]
- 43 Previous documentation revealed malnutrition as a very common issue in congenital heart
- 44 disease and even worse in PDA. [6-9] No known study in this locality ha considered any link
- between anthropometry and the size of PDA among children. Studies abound on the
- 46 nutritional status of children with congenital heart disease, but very few focused on assessing

47	the effects of anthropometry on the size of PDA. This study is therefore aimed to determine if
48	the size of PDA has any effect on anthropometry (weight, height, z scores). It also determines
49	at which size of PDA does severity of malnutrition begin to ensue.
50	Methods
51	Study design
52	This study was an observational cross-sectional study conducted in three institutions from the
53	year 2016 to 2020. During the study period, echocardiography was done on children with
54	various forms of cardiac disease.
55	
56	Study Area and Study Population
57	Children aged 1 day to 18 years with a congenital heart defect who fulfilled the diagnostic
58	criteria for patent ductus arteriosus from 2016 to 2020 at the University of Nigeria Teaching
59	Hospital were recruited in the study. We defined patent ductus arteriosus PDA as a defect
60	seen between the descending aorta and the left pulmonary artery and with a left to right shunt.
61	Relevant clinical features were also elicited by a thorough history taking and
62	sociodemographic variables were also enumerated.
63	Anthropometric measurements included height in centimetre for age more than 2 years and
64	supine length in centimetres for age below two. Weight was measured by standardized
65	methods and recorded in kilograms. Z scores for weight for age (WAZ), weight for height
66	(WHZ), and height for age (HAZ) were also calculated using the WHO Anthro software. The
67	clinical feature was also elicited.
68	Echocardiographic Measurement of Patent Ductus Ateriousus

- 69 Though PDA can be seen from many windows, left-sided parasternal otherwise called the
- ductal view is the best option used in this study to obtain a clear image. The ductal size was
- ascertained and measured at the narrowest diameter, which is at the pulmonary end. [10]

73

77

Assessment of size of Patent ductus arteriosus

- Size of patent ductus arteriosus of 1-3mm is taken as a small size PDA in this study.
- 75 Moderate size PDA were those PDAs with a diameter of 4-6mm while the ductal diameter of
- equal to and more than 7mm is classified as large PDA.

Data analysis

- 78 The data were analysed with the IBM SPSS statistics for windows, version 20 (IBM Corp,
- 79 Chicago). Differences in proportions were compared using the chi-square test. The weight
- and height z-scores were calculated using WHO Anthro and Anthro Plus software. The
- nutritional status was based on the WHO classification of weight for age (WAZ), weight for
- height (WHZ), and height for age (HAZ). P-value < 0.05 was regarded as significant.

Result

- There was a total of 758 children with heart anomalies examined within the study period, of
- which 70 children had confirmed diagnosis of PDA. The patients with PDA were made up of
- 45.7% males and 54.3% females. The age distribution of the patients is as in table 1, with a
- predominance of infants. Their mean age was 30.0±39.2 months.
- The patients' mean weight and height were 11.8 ± 10.5 kg and 84.6 ± 29.2 cm respectively. The
- mean weight for males, 11.3 ± 8.3 kg was comparable to that for females, 12.2 ± 12.5 kg (t = -
- 90 0.29, p = 0.8). Also, the mean height/length for males, 84.7 ± 27.3 was comparable to that for

- females, 81.1 ± 35.2 cm (t = 0.41, p = 0.7). Out of 48 children assessed for nutritional status,
- 92 29.2 % were well-nourished, 45.8% (22/48) were both wasted and stunted, 14.6% wasted,
- 93 8.3% stunted while 2.1% were obese.

96

94

Table I: age distribution of the patients

Age group	Frequency	%
Infants	35	50.0
preschool	24	34.3
school age	7	10.0
adolescents	4	5.7
Total	70	100.0

97 infants; 1-12 months, preschool; > 12 months to 5 years, school age; > 5 years to 10 years,

98 adolescents; >10 to 18 years

Table 2: Nutritional status among the males and females

100

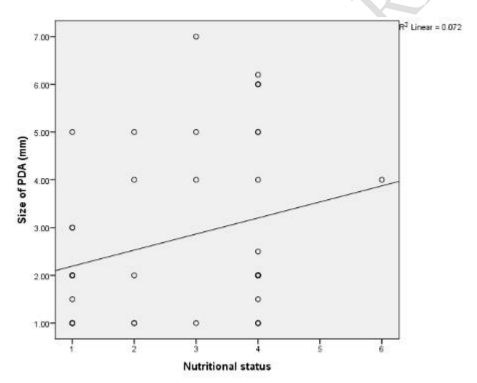
		Nutritional status		Total			
		Normal	Wasted	Stunted	wasted and	Obese (%)	
		(%)	(%)	(%)	stunted (%)		
0	male	5 (20.8)	2 (8.3)	3 (12.5)	13 (54.2)	1 (4.2)	24
Sex	female	9 (37.5)	5 (20.8)	1 (4.2)	9 (37.5)	0 (0)	24
Total		14 (29.2)	7 (14.6)	4 (8.3)	22 (45.8)	1 (2.1)	48

Chi-square = 5.2, p = 0.3. Wasted = Z-score weight-for-age or weight-for-height <2SD, stunted = height/length-for-age <2SD, obese = BMI for age $\ge 2SD$. The calculation was made using WHO "anthro" and "anthroPlus" software.

There was no significant difference in the nutritional status between the males and females as illustrated in table 2.

Figure 1: Graph of PDA size and nutritional status

Although there is a positive correlation between ductal size and nutritional status of patients, the effect size was small as shown in the figure 1, with ETA square of 0.072



1, well nourished; 2, wasted; 3, stunted; 4, wasted and stunted; 5, overweight; 6, obese

The graph shows that severe malnutrition ensues when ductal size is 3.2mm.

The frequency of some clinical features varied among these patients with PDA as illustrated in table 3. The commonest feature was fast breathing, observed in 68.9% of the patients assessed for the clinical feature, followed by pulmonary hypertension in 51.4%.

Table 3: Frequency of different clinical features among patients with PDA

Clinical feature	Frequency (n/N)	% (n/N) X 100
Cough	0/39	0.0
Fast breathing	40/58	68.9
Failure to gain weight	22/46	47.8
Easy fatigability	30/67	44.8
Pulmonary hypertension	36/70	51.4

N = number of patients with complete data for the assessed feature, n = actual number of

patients with the symptoms.

The majority of the patients (42.9%) had tiny PDA while 32.8% and 24.3% had small and large PDA respectively. Analysis of the size of PDA with nutritional status indicates that

35.3% (6/17) of patients with large PDA are wasted and stunted compared with 26.1 (6/23)

and 33.3% (10/30) of those with small and tiny PDA respectively ($\chi 2 = 10.21$, p = 0.8).

Although there was a negative correlation between weight and the size of PDA, the correlation was not significant (Pearson correlation coefficient = -0.1, p = 0.7). There was a negative correlation between the height/length and size of PDA, but the correlation was still not significant (correlation coefficient = -0.1, p = 0.5).

Discussion

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

This study was aimed to determine if size of PDA had any effect on anthropometry. The study showed no effect of size of PDA on weight and height on the children with PDA. We noted that severe malnutrition begins in children whose PDA size is 3.2mm and above. Increased metabolic stress from cardiac failure, high fat-free mass to fat mass ratio, prolonged hypoxia, metabolic acidosis, and worsening sympathetic system activity could explain this finding. [11-17] Other reason for children with large size PDA presenting with wasting and stunting could be due to elevated pulmonary artery pressure in children with large PDA. The pulmonary pressure is caused by pulmonary over-circulation and pulmonary vascular disease, either in combination or alone could create a nidus for chest infections. [15] This could further worsen malnutrition. Accentuated pulmonary hypertension, poor intake due to anorexia, easy fatigability, uncoordinated breast sucking, neurological dysfunction, easy satiety, and fast breathing all get accentuated in children with large size PDA. [16] The commonest symptom seen in this study were fast breathing and this was seen mostly in children with large PDA. This could be caused by pulmonary hypertension which is seen in over 50% of the children. Some studies have also documented PDA-associated symptoms as been triggered by mesenteric, cerebral hypo-perfusion, renal, and pulmonary oedema secondary to pulmonary hypertension seen in over 50% of those with persistent PDA. [18-21] Abhijeet et al [22] also noted breathlessness and history of recurrent respiratory tract

159 infections as the commonest symptoms in his series and noted that these symptoms are see in majority of children who had large PDA 160

The prevalence of severe malnutrition in children with Patent Ductus Arteriosus noted in this study is high, this is seen mostly among the under-fives. This prevalence is higher compared with prevalence values seen in children without any congenital heart disease. Chinawa et al have documented that children with congenital heart disease who are less than five years old are prone to malnutrition when compared to those who had no congenital heart disease. This could be explained by increase metabolic demands seen at this age, late surgical intervention, progressive hypoxemia and progressive, pulmonary hypertension which is usually seen in children less than five years old who had congenital heart disease and who had no intervention. [23-29] Other forms of malnutrition seen in this study included stunting, wasting, or both. Mechanisms for malnutrition are multifaceted. These include associated chromosomal

anomalies or genetic syndromes, feeding difficulties, poor absorption from congestive cardiac failure (CCF). Besides, increased caloric demand altered respiratory, and neuro-humoral dysfunction with attendant chronic hypoxia with impaired cellular metabolism have all been implicated in malnutrition in children with PDA. [30] Malnutrition in children with PDA is a

known cause of frequent hospitalization, pulmonary hypertension, and death. [31,32]

177

178

179

180

181

182

161

162

163

164

165

166

167

168

169

170

171

172

173

174

175

176

Limitations

It is known that the echocardiography has many limitations in the measurement of the size of the duct. In older children and young adults, the lung limits the visualization of the duct and hence difficulty in measuring the size.

Conclusion

183	The majority of children with PDA presents with severe forms of malnutrition (wasting and
184	stunting). Severe malnutrition ensues when ductal size is 3.2mm. The size of PDA has no
185	effect on weight and height of children with PDA.
186	Declaration
187	Ethics approval and consent to participate: Ethical approval was obtained from the Ethics
188	and Research committee of the University of Nigeria Teaching hospital Enugu
189	Verbal informed consent was obtained from parents or guardians (care-givers) of the subjects
190	and controls.
191	.Consent for publication:
192	Not applicable
193	Availability of data and materials: data supporting the findings of this study are available
194	from the corresponding author (JMC) on request.
195	Competing Interest: We declare that we have no competing interests.
196	Funding: This study was not funded by any organization. The authors bore all the expenses
197	that accrued from the study.
198	Author contributions statement
199	JMC was involved in the conception and design of the article BFC was involved in the
200	analysis and interpretation of the data. JMC was involved in the drafting of the paper, while
201	COD, ATC and ACA were involved in critical revision of the article for intellectual content;
202	and the final approval of the version to be published. All authors agree to be accountable for
203	all aspects of the work.
204	
205	
206	COMPETING INTERESTS DISCLAIMER:

207	
208	Authors have declared that no competing interests exist. The products used for this research
209	are commonly and predominantly use products in our area of research and country. There is
210	absolutely no conflict of interest between the authors and producers of the products because
211	we do not intend to use these products as an avenue for any litigation but for the advancement
212	of knowledge. Also, the research was not funded by the producing company rather it was
213	funded by personal efforts of the authors.
214	
215	
216	
217	References
218	1.Kaemmerer H, Meisner H, Hess J et al. Surgical treatment of patent ductus arteriosus: a
219	new historical perspective. Am J Cardiol 2004.; 94:1153-4.
220	2. Hagau N, Culcitchi C. Nutritional support in children with congenital heart disease. Nutr
221	Ther Metab. 2010;28:172-84.
222	3, Condo M, Evans N, Bellu R et al. Echocardiographic assessment of ductal significance:
223	retrospective comparison of two methods. Arch Dis Child Fetal Neonatal Ed. 2012; 97: 35-
224	8.
225	4. Dice JE, Bhatia J. Patent ductus arteriosus: an overview. J Pediatr Pharmacol Ther
226	2007;12:138-46.
227	5. Schneider DJ, Moore JW. Patent ductus arteriosus. Circulation. 2006;114:1873–1882.
228	6. Owens JL, Musa N. Nutrition support after neonatal cardiac surgery. Nutr Clin Pract.
229	2009;24:242-9.

- 7. Tokel K, Azak E, Ayabakan C et al.. Somatic growth after corrective surgery for
- congenital heart disease. Turk J Pediatr 2010; 52:58–67
- 8. Vaidyanathan B, Nair SB, Sundaram KR et al. Malnutrition in children with congenital
- 233 heart disease (CHD) determinants and short-term impact of corrective intervention. Indian
- 234 Pediatr 2008; 45:541–6.10.
- 9. Anant K, Suthep W. Patent ductus arteriosus associated with pulmonary hypertension and
- desaturation. Cardiology Journal 2012; 19: 543-546
- 10. Tschuppert S, Doell C, Arlettaz-Mieth R, Baenziger O, Rousson V, Balmer C, et al. The
- 238 effect of ductal diameter on surgical and medical closure of patent ductus arteriosus in
- preterm neonates: size matters. J Thorac Cardiovasc Surg. (2008) 135:78–82. doi:
- 240 10.1016/j.jtcvs.2007.07.027
- 241 11.Chugh R, Salem MM. Echocardiography for patent ductus arteriosus including closure in
- 242 adults. *Echocardiography* 2015;32:125-139.
- 243 12. Okoromah CA, Ekure EN, Lesi FE et al. Prevalence, profile and predictors of malnutrition
- in children with congenital heart defects: a case-control observational study. Arch Dis
- 245 *Child* 2011;96:354–60.
- 246 13. Forchielli ML, McColl R, Walker WA et al. Children with congenital heart disease: a
- 247 nutrition challenge. *Nutr Rev* 1994; 52:348–53.
- 248 14. Freeman LM, Roubenoff R. The nutrition implications of cardiac cachexia. *Nutr*
- 249 *Rev* 1994;52:340–7.
- 250 15. Niu MC, Mallory GB, Justino H et al. Treatment of severe pulmonary hypertension in the
- setting of the large p atent ductus arteriosus. Pediatrics 2013;131:1643-9
- 252 16.Ranjit P, Jason NJ, Ronak N et al. Effect of patent ductus arteriosus on pulmonary
- vascular disease. Congenital heart disease 2019; 14: 37-41

- 254 17. Farber HW, Loscalzo J. Mechanism of disease: Pulmonary hypertension. Discov Med,
- 255 2005; 5: 80–87.
- 256 18.Lemmers PM, Benders MJ, D'Ascenzo R et al. Patent Ductus Arteriosus and Brain
- 257 Volume. Pediatrics. 2016;137:20153090.
- 258 19. Clyman RI, Couto J, Murphy GM. Patent ductus arteriosus: are current neonatal treatment
- options better or worse than no treatment at all? Semin Perinatol. 2012;36:123–9.
- 260 20. Reese J, Laughon MM. The Patent Ductus Arteriosus Problem: Infants Who Still Need
- 261 Treatment. J Pediatr. 2015;167:954–6.
- 262 21. Koch J, Hensley G, Roy L et al. Prevalence of spontaneous closure of the ductus
- arteriosus in neonates at a birth weight of 1000 grams or less. Pediatrics. 2006;117:1113-
- 264 1121.
- 265 22. Abhijeet M, Dashetwar A, Manoharrao D et al. A study of clinical profile and
- 266 management of patients with Patent ductus arteriosus. Journal of Evidence Based Medicine
- 267 and Healthcare 2016; 3:4533-4539
- 268 23. Monteiro FM, Araujo TL, Lopes MO et al. Nutritional status of children with congenital
- heart disease. Rev Latino-Am Enfermagem. (2012) 20:1024–32.
- 271 24. Schuurmans FM, Pulles-Heintzberger CFM, Gerver WJM et al. Long-term growth of
- 272 children with congenital heart disease: a retrospective study. *Acta Pædiatr.* (1998) 87:1250–
- 273 5.

- 274 25. Varan B, Tokel K, Yilmaz G. Malnutrition and growth failure in cyanotic and acyanotic
- congenital heart disease with and without pulmonary hypertension. *Arch Dis Child.* (1999)
- 276 81:49–52.

- 277 26. Vaidyanathan B, Nair SB, Sundaram KR et al. Malnutrition in children with congenital
- heart disease (CHD) determinants and short term impact of corrective intervention. *Indian*
- 279 *Pediatr.* (2008) 45:541–6.
- 280 27. Vaidyanathan B, Radhakrishnan R, Sarala DA et al. What determines nutritional recovery in
- 281 malnourished children after correction of congenital heart defects? *Pediatrics*. (2009)
- 282 124:294–9.
- 283 28. Daymont C, Neal A, Prosnitz A et al. Growth in children with congenital heart
- 284 disease. *Pediatrics*. (2013) 131:236–42.
- 285 29. Kyle UG, Shekerdemian LS, Coss-Bu JA. Growth failure and nutrition consideration in
- 286 chronic childhood wasting diseases. *Nutr Clin Pract.* (2015) 30:227–38.
- 287 30. Okoromah CA, Ekure EN, Lesi Feet al. Prevalence, profile and predictors of malnutrition
- in children with congenital heart defects: a case-control observational study. Arch Di s
- 289 *Child* 2011;96(4):354–60.
- 290 31. Forchielli ML, McColl R, Walker WA et al Children with congenital heart disease: a
- 291 nutrition challenge. *Nutr Rev* 1994; 52:348–53.
- 292 32. Freeman LM, Roubenoff R. The nutrition implications of cardiac cachexia. Nutr
- 293 *Rev* 1994;52:340–7.

295

296

297

298

299

300

