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2 **Title: Hypoxaemia in Nigerian Children Presenting to the Children**
3 **Emergency Ward (CHEW) of a Tertiary Hospital.**

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5 **Abstract:**

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6 **Aim** of this study is to determine the prevalence of hypoxaemia and predictors of signs of
7 hypoxaemia in children with various disease conditions admitted into the CHEW of a tertiary
8 health facility.

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9 **Study Design:** Descriptive, Cross sectional study

10 **Place and Duration:** Department of Paediatrics (Children Emergency Ward). Study was done
11 from 1st February to 30th April 2015

12 **Methods:** We included 129 children. Seventy two (55.8%) males and 57(44.2%) females and
13 age range 0.08 to 17 years admitted into the CHEW with various disease conditions. Biodata and
14 clinical examination was done in all patients. Oxygen saturation (SpO₂) was determined on
15 admission using pulse oximeter for every sick child admitted. Hypoxaemia is defined as SpO₂
16 less than 90%.

17 **Results:** One hundred and twenty nine children were studied. Ages ranged from 0.08 to 17 years
18 with a mean age of 3.06 ± 3.65 years. Modal age was 4 years. The mean age of 3.34 ± 3.97 years
19 for males was higher than 2.70 ± 3.22 years for females. Thirty one (24%) children had
20 hypoxaemia on admission with 20(64.5%) with respiratory diseases. Infants ($p=0.004$) and
21 children with respiratory disease ($p=0.047$) had a significantly higher prevalence of hypoxaemia
22 among the study group.

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23 Chest in drawing is a common feature but grunting (100%) and wheezing (50%) have the best
24 positive predictive values.

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25 **Conclusion:** Hypoxaemia is prevalent in children who are ill and need emergency care.
26 Respiratory diseases and infants account for a major proportion of hypoxaemic children seen in
27 emergency wards. Chest in drawing is a common feature from different studies; presence of
28 grunting was highly predictive in this study.

29 **Keywords:** Children, Hypoxaemia, Emergency care

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38 **Introduction**

39 Hypoxaemia is defined as reduced oxygen content of blood specifically in arterial blood or the
40 reduced percentage of saturation of haemoglobin with oxygen. It is an under recognized
41 complication of most severe illnesses in neonates and children in developing countries and a
42 common predictor of death. [1, 2, 3]

43 Hypoxaemia can be determined by measuring the level of oxygen in a sample of arterial blood or
44 by determination of oxygen saturation in the blood using the pulse oximeter. Hypoxaemia can be

45 defined as arterial oxygen concentration of less than 75 mmHg or blood oxygen saturation of less
46 than 90%. [1, 3] Hypoxaemia is a common manifestation of severe illnesses in children and a
47 major contributor to mortality. Several clinical signs and symptoms have been found to predict
48 hypoxaemia in sick children with or without acute lower respiratory tract infection, this include
49 inability to feed, fast breathing, grunting, lower chest wall in drawing, nodding and
50 convulsion.[4,5]

51 Hypoxaemia is known to correlate well with disease severity and occurs mainly in diseases that
52 impair ventilation, gaseous exchange or increase oxygen demand in the body. [6] A disease of
53 the respiratory tract such as pneumonia which accounts for more than 2million deaths in children
54 worldwide is commonly complicated by hypoxaemia.[6]The prevalence of hypoxaemia in
55 children varies with disease condition and severity of illness. Prevalence of hypoxaemia in ill
56 children range from 11 to 52% and can be as high as 73 % in children with acute lower
57 respiratory tract illnesses.[3,5] In a study on hypoxaemia as a measure of disease severity in
58 young hospitalized Nigerian children with pneumonia, 41.5% had hypoxaemia with hypoxaemic
59 children 48 times more likely to die.[2]

60 The blood gas analysis is the gold standard for detecting hypoxemia. Other methods include
61 pulse oximetry and less objectively clinical signs.[3] The use of the pulse oximeter is a reliable,
62 safe, non-invasive and reproducible tool which compares well with the results from the blood gas
63 analysis.[7,8] Its use ensures early detection and commencement of efficient treatment of
64 hypoxaemia in sick children in resource limited setting.

65 There is a persisting high rate of childhood mortality in Nigeria from various disease conditions.
66 The objective of this study is to determine the prevalence of hypoxaemia and predictors of signs

67 of hypoxaemia in children with various disease conditions admitted into the CHEW of a tertiary
68 health facility.

69 **Methods**

70 The study was a descriptive cross sectional study conducted at the Paediatric Emergency
71 Department of the University of Port Harcourt Teaching Hospital, Nigeria. The study comprised
72 of all children presenting to the Paediatric Emergency Ward, from 1st February to 30th April 2015
73 whose parents gave consent. The University of Port Harcourt Teaching Hospital is a tertiary
74 health care facility in Port Harcourt, Rivers State in Southern Nigeria. It is the largest health care
75 facility in the State and offers health care to people living in the State and its environs.

76 The study was approved by the Ethics and Research Committee of the University of Port
77 Harcourt Teaching Hospital and written informed consent was obtained from all caregivers that
78 participated. A detailed history was taken and physical examination conducted on all children
79 admitted into the Paediatric Emergency Ward and a clinical diagnosis made. Presence or absence
80 of symptoms and signs of respiratory distress were particularly sought for and recorded using a
81 study proforma. Pulse oximetry was done for all patients at presentation using a pulse oximeter
82 (Contec CMS0DL) with appropriate probe size placed on the finger and peripheral capillary
83 oxygen saturation (SpO₂) was recorded while breathing room air. Recordings were taken after
84 stabilization of the pulse oximetry reading for one minute. Hypoxaemia was defined as SpO₂ of
85 less than 90% recorded by pulse oximetry.

86 All subjects were treated by the children emergency ward managing team with appropriate
87 medications and interventions based on their individual diagnoses. They were followed up to
88 monitor the outcome of their admission.

89 Data was analyzed using the Epi info version 7.1.3.3 software. The Chi square and Student's t
90 tests were used to test for statistically significant differences in proportions and means
91 respectively. A p value of less than or equal to 0.05 was considered as statistically significant.

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93 **Results**

94 A total of 129 children participated in the study, Seventy two (55.8%) males and 57(44.2%)
95 females giving a male to female ratio of 1.3:1. Ages ranged from 0.08 to 17 years with a mean
96 age of 3.06 ± 3.65 years. Modal age was 4 years. The mean age of 3.34 ± 3.97 years for males
97 was higher than 2.70 ± 3.22 years for females. The difference was not statistically significant
98 ($t=0.98$, $df=1$, $p=0.327$). The primary diagnosis in 66 (51.2%) of the children was a respiratory
99 disease and otherwise in the remaining 63 (48.9%).

100 The SpO₂ ranged from 54% to 99% with a mean of $91.53 \pm 8.57\%$, mode of 98% and median of
101 95%. Thirty one (24%) of the children had hypoxaemia with 20(64.5%) having primary
102 respiratory disease. The mean SpO₂ for the hypoxaemic children was $78.13 \pm 6.90\%$ while the
103 mean SpO₂ for the non-hypoxaemic children was $95.78 \pm 2.55\%$ ($t=21.27$, $df=1$, $p<0.001$).

104 **Table 1** shows the levels of SpO₂ measured among the patients.

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106 **Table 1: Levels of SpO₂ measured**

SpO ₂ (%)	Frequency (N)	Percent (%)
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≥95	69	53.5
90-94	29	22.5
75-89	27	20.9
<75	4	3.1
Total	129	100.00%

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108 **Table 2** shows the relationship between presence of hypoxaemia and some variables among the
 109 study group. Infants (p=0.004) and children with respiratory disease (p=0.047) had a
 110 significantly higher prevalence of hypoxaemia among the study group.

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112 **Table 2: Relationship between some variables and presence of hypoxaemia**

Variable		Hypoxaemia	No hypoxaemia	Total	p
Age group	Infants	19 (35.8)	34 (64.2)	53 (42.1)	0.004*
	Children	11 (15.1)	62 (84.9)	73 (57.9)	
Gender	Males	18 (25.0)	54 (75.0)	72 (55.8)	0.390
	Females	13 (22.8)	44 (77.2)	57 (44.2)	
Primary diagnosis	Respiratory disease	20 (30.3)	46 (69.7)	66 (51.2)	0.047*
	Non-respiratory disease	11 (17.5)	52 (82.5)	63 (48.8)	

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*significant

114 **Table 3** shows the predictive value of some clinical signs of hypoxaemia. Tachypnoea was the
 115 most sensitive (sensitivity=61%) for hypoxaemia, followed by intercostals recession
 116 (sensitivity=58%) and subcostal recession (sensitivity=55%). Grunting (specificity=100%),
 117 wheezing (specificity=99%) and suprasternal recession (specificity=99%) were the most specific
 118 for hypoxaemia. The best predictors of hypoxaemia were grunting (100%) and wheezing (50%).

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119 **Table 3: Predictive value of some clinical signs of hypoxaemia**

	Hypoxaemic patients (n=31)	Non-hypoxaemic patients (n=98)	p value	Sensitivity	Specificity	Positive predictive value	Negative predictive value
Flaring	13	29	0.107	42	70	31	79
Intercostal recession	18	40	0.05	58	59	31	82
Subcostal recession	17	30	0.009	55	69	36	83
Suprasternal recession	0	1	0.76	0	99	0	76
Grunting	1	0	0.24	3	100	100	77

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Wheezing	1	1	0.424	3	99	50	76
Tachypnoea	19	47	0.102	61	52	29	81

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121 One of the 31 cases with hypoxaemia died giving a case fatality rate of 3.2%.

122 **Discussion**

123 The prevalence of hypoxaemia of 24% in this study was similar to that of 20.6% in a study in Ife,
 124 Nigeria[9] and 23.8% in Kanpur, India.[10] It is however higher than the 5.8%, 11.9% and 13%
 125 found in studies done in The Gambia,[11] Chandigarh, India[12] and Enugu, Nigeria[13]
 126 respectively, and lower than 41.5% and 73% seen in Ilorin, Nigeria² and Papua New Guinea
 127 respectively.[3] Various factors including altitude, health care settings, diagnoses in subjects, age
 128 group of subjects, cut off values for hypoxaemia could have contributed to the differences seen
 129 among the various studies. Differences may also be due to prevalence of respiratory illnesses
 130 which can be affected by incidence of viral illnesses which may be seasonal.

131 The significantly higher prevalence of hypoxaemia among infants (35.8%) compared to children
 132 above one year in this study is corroborated by similar studies among children in Enugu,[13]
 133 Ife,[9] The Gambia[11] and India.[12] This may be due to the fact that infants have a lower tidal
 134 volume and relative inefficient compensatory mechanisms to improve ventilation. Infants are
 135 also less unable to compensate for ventilation perfusion mismatch in situations of increased dead

136 space.[14] Emodi et al[13] showed an equal occurrence of hypoxia in both genders in their study
137 as was also seen in the present study.

138 Hypoxaemia was significantly higher among patients with a respiratory disease compared to
139 other diseases in this study. This is similar to findings in other studies.[2,3,11,12] While this was
140 the finding in studies [3,11] among ill children with respiratory and non-respiratory illnesses,
141 studies [2,12] among children with respiratory illnesses all revealed a higher prevalence of
142 hypoxaemia in children with severe pneumonia. Pneumonia results in airway obstruction from
143 swelling, abnormal secretions, and cellular debris. Atelectasis, interstitial edema, and ventilation-
144 perfusion mismatch causing significant hypoxemia often accompany airway obstruction.[15]

145 The present study showed tachypnoea and chest wall retractions (intercostals and subcostal) to
146 be the most sensitive clinical features of hypoxaemia, and grunting, wheezing and suprasternal
147 recession to be the most specific clinical features of hypoxaemia. In a study by Rao, et al [10] the
148 sensitivity for hypoxaemia was highest with chest wall retraction which is similar to finding in
149 this study. He also reported flaring of alar nasi, inability to feed as other strong indicators of
150 hypoxaemia with sensitivity of 84% and 81% respectively especially in children with
151 pneumonia. Chest in drawing was also significantly associated with presence of hypoxaemia in
152 the study by Kuti and colleagues in Ile Ife, Nigeria.[9] In this study, grunting and wheezing had
153 the highest positive predictive value in occurrence of hypoxaemia, also a study in Ibadan,
154 Nigeria by Adebola and colleagues [16] showed that flaring and chest in drawing were predictive
155 of hypoxaemia. The use of clinical signs in determination of hypoxaemia in ill children has met
156 diverse views. A systematic review and meta-analysis of prospective diagnostic studies that
157 evaluated the accuracy of individual or combined clinical symptoms and signs in predicting
158 hypoxemia among children aged <5 years with ARI, revealed that cyanosis, inability to feed,

159 head nodding, respiratory rate >70/min and unresponsiveness/impaired arousability had high
160 specificity but low sensitivity.[17] This report was supported by Lodha *et al.*[18] and Dyke *et*
161 *al.*[19] who concluded that clinical symptoms and signs alone or in combination do not have
162 sufficient sensitivity and specificity to predict hypoxemia in children with ALRI. Using clinical
163 symptoms, about 20% of hypoxemic children would be missed, and 17–50% of children given
164 supplemental oxygen would not need it in high sensitivity models based on clinical signs.[3]
165 Pulse oximetry when used correctly provides a reliable bedside standard for detecting
166 hypoxaemia even in developing countries. Pulse oximetry can correctly identify 20–30% more
167 children with hypoxemias than using clinical signs alone and will ensure judicious and efficient
168 use of oxygen therapy in resource limited setting where oxygen is not readily available.[2,20,21]

169 Although pulse oximetry remains the reliable means of determining oxygen saturation in
170 children, using of clinical symptoms may be very helpful in deciding presence of hypoxaemia in
171 ill children where pulse oximeter is not available in resource limited settings.

172 **Conclusion:** Hypoxaemia is prevalent in children who are ill and need emergency care.
173 Respiratory diseases and infants account for a major proportion of hypoxaemic children seen in
174 emergency wards. Chest in drawing is a common feature from different studies; presence of
175 grunting was highly predictive in this study.

176 **Consent:** As per international standard or University standards, the patient written consent has
177 been collected and preserved by authors

178 **Ethical Approval:** As per international standards or University standards, a written ethical
179 approval has been collected and preserved by authors.

180 **Competing Interest:** Authors have declared that no competing interest exist.

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