

Original Research Article

Effect of acute placenta inflammatory changes on fetal outcome among parturients in Nigeria.

Key words: acute inflammation, placenta, fetal outcome, Nigeria.

Abstract

Introduction

The placenta unit is significant for the survival of the fetus. Infections from the mother can cause histologically identifiable inflammatory changes in the placenta, which may adversely affect the fetus.

Aim: to identify the inflammatory changes in the placenta and its effect on fetal outcome.

Study design: Cross sectional study

Place and duration of study: Department of Obstetrics and Gynaecology and Department of Anatomical pathology of the University of Port Harcourt Teaching hospital, between 1st September – 31st of December, 2015.

Methods: Histological analysis of 189 placenta tissues of singleton birth parturients was carried out. The sociodemographic characteristics of patients and the fetal outcome was collated and analyzed. The information obtained was processed using the SPSS version 20 software and Epi info software version 7. Results were presented in tables; test of association was done using student's t-test with P value < 0.05 set as significant.

Results: The mean age of patients was 30.9± 4.6 years with age range of 19-48years. Acute placental inflammatory changes of varying grades were noted in 64.6%(122) of placenta with severe inflammatory changes constituting

25 8.5%(16) of all examined placentae. Severely inflamed placenta was associated
26 with birth asphyxia ($P=0.0000033$) and fetal demise ($P=0.0352$)
27 Conclusion. Acute inflammatory changes are common among parturients in Port
28 Harcourt Nigeria. These changes in placenta are associated with birth asphyxia
29 and fetal demise especially when they are severe.

30

31 **Introduction**

32 The placenta is an integral part of the existence of the fetus, it serves as an
33 interface in the transmission of requisite nutrients and other metabolic
34 materials.[1,2] The placenta histology can be altered by intrauterine infections
35 which is usually acquired as a result of ascending infection from the genital tract,
36 gut or hematogenously resulting in chorioamnionitis which complicates about
37 10% of all pregnancies with up to 2% occurring during labour.[3,4]
38 Several organisms have been implicated in the causation of chorioamnionitis
39 inclusive of bacterial and viral organisms. Bacterial organisms are linked to acute
40 inflammation while viral organisms were associated with chronic changes
41 [3,5,6]. Polymicrobial bacterial organisms contribute to these acute
42 inflammatory changes of which ureaplasma urealyticum and gardnella
43 vaginitis are the two commonest organisms.[3]
44 The presence of infectious organisms in the chorioamnion engenders a maternal
45 and fetal inflammatory response characterized by pro-inflammatory and
46 inhibitory cytokines and chemokines in the maternal and fetal compartments;
47 with maternal inflammatory response preceding the fetal response.[7] The
48 placenta changes in the presence of infection can be categorized into acute,
49 subacute or chronic with acute chorioamnionitis with or without fetal

50 inflammatory response, villitis and decidualitis been the most common types of
51 placenta inflammations observed.[8]

52 Acute chorioamnionitis can be diagnosed clinically or histologically. Histological
53 chorioamnionitis is defined by the presence of acute histological changes on
54 examination of the amniotic membrane and chorion of the placenta.[9]

55 Sometimes subclinical infections can be missed but captured by histological
56 examination of placenta and histological diagnosis is the gold standard for
57 evaluation of antenatal inflammatory process that might influence fetal
58 development[10,11]. This modality of diagnosis of infection increases the rate of
59 identification of chorioamnionitis than clinically diagnosed chorioamnionitis
60 confirmed by amniotic fluid culture because of lack of detection of some
61 organisms cultured.[10]

62 Some series have identified that impairment of placental development as a result
63 of infection may have a profound impact in fetal development and pregnancy
64 outcome such as cerebral palsy and high rate of stillbirths.[12,13] It is on these
65 basis that this study seeks to determine the prevalence of acute inflammatory
66 placenta changes and its effect on the fetal outcome among parturients who
67 presented for delivery at the University Teaching Hospital in Port Harcourt,
68 Nigeria

69

70 **Methodology**

71 A cross sectional histological study of 189 placentae of women who had
72 singleton deliveries between 1st September to 30th December 2015 was
73 conducted. The women were recruited into the study as they presented to the
74 labour ward after due counseling on the scope of the study by the investigators.

75 All patients who gave consent for the study were included. Exclusion criteria
76 included: multi-order pregnancies, women who had evidence of
77 immunosuppression like diabetes mellitus, on steroids or diagnosed with
78 retroviral disease, patients with preeclampsia, previous antenatal infections,
79 intrauterine growth restriction of known cause, patients with PROM and
80 Chorioamnionitis were also excluded. All other patients who presented to the
81 labour ward during the study period were included in the study. The
82 sociodemographic characteristics of patients, which include: age, parity,
83 educational status and booking status; birth weight, presence or absence of birth
84 asphyxia and fetal outcome (dead or alive) were collated in a prestructured
85 spread sheet and analyzed. The placenta immediately after delivery was
86 collected and blood stain removed from it using gentle running water, preserved
87 in 10% formaldehyde solution and transferred to the anatomical pathology
88 laboratory of the teaching hospital for processing. Grossing of the placenta was
89 done and representative sections were taken. Tissues were processed in the
90 automated tissue processor and later stained using the hematoxylin and eosin
91 methods.

92 Two independent pathologists reviewed processed slides and areas of disparity
93 were resolved following a consensus decision of the two pathologists. Histologic
94 grading of acute inflammation was assessed by the density of neutrophil
95 infiltrates and the relative distance migrated from the vessels of origin. Placental
96 inflammatory changes were scored and categorized as: (Nil)-no
97 chorioamnionitis; Grade 1(Mild) :presence of polymorphonuclear leucocytes at
98 the subchorionic plate and lower third of chorion; Grade 2(moderate): At least

99 two separate foci of leucocytes infiltrates in the chorion and Amnion, and Grade
100 3(severe)- extensive leucocyte infiltrates in the chorion/amnion[14].

101 The information obtained was processed using the SPSS version 20 software
102 (SPSS Inc; Chicago USA) and Epi info software version 7. Results were presented
103 in tables, test of association was done using student's t-test with P value < 0.05
104 set as significant.

105 **Results**

106 The mean age of patients was 30.9± 4.6 years with an age range of 19-48 years.
107 69.8%(132) were in the age group 25-34 years and 74.6%(141) were
108 nulliparous patients, 74.6%(141) had tertiary education and 93.7%(177) of
109 examined placenta belonged to booked patients while 6.3%(12) were unbooked.

110 Other sociodemographic variables are as shown in Table 1.

111 Table two showed the relationship of booking status to placenta inflammatory
112 changes. Acute placenta inflammatory changes were observed in 122(64.6%) of
113 all patients studied with 16(8.5%) having severe inflammatory changes. Among
114 the patients, 177(93.7%)were booked while 6.3% were unbooked. In patients
115 with severe inflammatory changes 14(87.5%) were booked while 2(12.5%)
116 were unbooked. Booking status was not significantly associated with placenta
117 inflammatory changes (P=0.08541). Among babies delivered 11(5.8%) had low
118 birth weight while 94.2%(178) were not low birth weight. Among all babies born
119 to mothers with severe placenta inflammatory changes, 2(12.5%) were low birth
120 weight while 14(87.5%) were not low birth weight. Other distributions are as
121 shown in Table 2. Patients with severe placenta inflammatory changes were not
122 significantly associated with low birth weight babies (P=0.15)

Table 3 showed the relationship between fetal outcome and placenta inflammatory changes: 140(74.1%) had no birth asphyxia while 49(25.9%) had some form of birth asphyxia. Among those with severe placenta inflammatory changes 12(75%) had some form of asphyxia while 4(25%) had no birth asphyxia. Placenta inflammatory changes are significantly associated with birth asphyxia ($P = 0.00367$). Severe inflammatory changes are significantly associated with severe birth asphyxia ($P = 0.0000033$).

Among the placenta examined, 186(98.4%) belong to babies that were alive while 3(1.6%) were dead. Among the dead babies 2(66.7%) had severe inflammatory changes while one had moderate inflammatory changes (33.3%)

Fetal demise was not significantly associated with general placenta inflammatory changes ($P = 0.2666$), however severe placenta changes was significantly associated with fetal demise ($P = 0.0352$)

Tables

137

138 Table 1:Sociodemographic characteristics of patients

139

Variables	Frequency	Percentage
Age		
15-24	16	8.5
25-34	132	69.8
35-44	40	21.2
45-54	1	0.5
Parity		
0	141	74.6
1	20	10.6
2	21	11.1
3	5	2.6
4	2	1.1
Educational status		
-Primary	2	1.1
-Secondary	46	24.3
-Tertiary	141	74.6

Booking status		
-booked	177	93.7
-unbooked	12	6.3

Table 2: Relation of placenta inflammatory changes to booking status and birth weight.

	Placenta inflammatory changes				Total n(%)
	Mild	Moderate	Severe	0 Nil	
Booking status					
-booked	58	40	14	65	177(93.7)
-unbooked	4	4	2	2	12 (6.3)
Total	62	44	16	67	189(100)
Fetal weight (kg)					
1.5-2.4	3	2	2	4	11(5.8)
2.5-3.4	37	30	8	42	117(61.9)
3.5-4.4	22	11	6	20	59(31.2)
4.5-5.4	-	1	0	1	2(1.1)
Total	62	44	16	67	189(100)

Table 3: Placenta histology and fetal outcome

	Placenta inflammatory changes				
	Nil	Mild	Moderate	Severe	Total n(%)
Birth Asphyxia					
Nil	58	48	30	4	140(74.1)
Mild	6	11	5	1	23(12.2)
Moderate	3	3	8	7	21(11.1)
Severe	0	0	1	4	5(2.6)
Total	67	62	44	16	189(100)
Fetal outcome					
Alive	67	62	43	14	186(98.4)
Dead	0	0	1	2	3(1.6)
Total	67	62	44	16	189

153

154

155 **Discussion**

156 The average age of parturients was similar to what was observed in other areas in
157 Nigeria [15,16], which **was** the peak of the sexual and reproductive ages of the
158 patients studied. This study also noted that a large proportion of women with
159 inflammatory placenta changes **were** nulliparous which **was** in conformity with
160 what was observed by Baker et al, who identified that acute placenta infections
161 decrease with increasing parity [17]. The reason as postulated by Lagadari et al
162 [18] is the development of protective layer of macrophages between decidua and
163 trophoblastic layers as parity increased as demonstrated in rat models.

164 Acute placenta inflammatory changes of varying degree were observed in more
165 than half of all patients' placentae examined, with similar distribution of placenta
166 changes observed by Rhone et al [19], where about 50% of placenta studied by
167 his group identified changes consistent with inflammatory changes. Variations in
168 the prevalence rate of acute inflammatory changes are related to the differences
169 in tissue sampling techniques and diagnostic criteria.

170 **Acute placenta inflammatory changes** have a direct correlation to clinical
171 chorioamnionitis, which is linked to poor fetal outcome [20]. These observations
172 brings to the front burner the need to screen parturients for possible organisms
173 that are linked to acute infections such as bacterial vaginosis which is a known
174 etiological factor for chorioamnionitis and preterm membrane rupture.

175 The unskilled supervised deliveries (unbooked) have been associated with
176 increased risk of chorioamnionitis and puerperal sepsis due to some unhygienic
177 birth practices [21]. In this group of patients, placenta inflammatory changes

178 was not associated with the booking status of the paturients; thus there may be
179 some other confounding factors which are not related to the booking status of
180 the patient that need to be unearthed.

181 Daniele et al and Gracia[20, 22] demonstrated the association between fetal
182 placenta inflammation and poor neonatal growth, which is as result of distortion
183 of placenta function. This observation **did not** agree with the authors' findings,
184 which showed **no relationship between low birth weight and the** presence of
185 placenta inflammatory changes..

186 It was observed that, as in this study, severe placenta inflammatory changes are
187 associated with poor fetal outcome such as asphyxia and even stillbirths [23,24].
188 The mechanism by which this is made possible is via placenta damage with loss
189 of function, preterm labour, release of inflammatory mediators, which result in
190 fetal organ damage and transplacental infection [5]. The fetal inflammatory
191 response syndrome which is related to placenta infections had also be linked
192 with the development of cerebral palsy and the development of neurological
193 deficits in the babies that survive the infectious onslaught [25].

194 **Conclusion,**

195 Based on the above it is pertinent to know that placental inflammatory changes
196 being the hallmark of fetal infection is associated with poor fetal outcome in Port
197 Harcourt Nigeria. Hence it is imperative that steps be instituted to reduce the
198 risk of placenta infection among paturients by creating protocols for screening of
199 bacterial pathogens, reduce the factors that increase risk and possibly
200 prophylaxis therapy of all at risk patients. Also a protocol of histological
201 examination of placentae of stillborns and babies with severe birth asphyxia is

202 also recommended to eliminate the long-term complications of infections related
203 morbidities.

204 Conflict of interest: Nil

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209 Ethical clearance was obtained from the ethics committee of the University of
210 Port Harcourt Teaching hospital before the commencement of the study.

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