**Pregnancy in Sickle Cell-Haemoglobin C (SC) Disease,**

**A Retrospective Study of Birth Size and Maternal Weight Gain**

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**Condensation**

Mothers with SC disease have lower maternal weight gain and lower birth weight infants than women with a normal haemoglobin genotype.

# Abstract

Title **Pregnancy in Sickle Cell-Haemoglobin C (SC) Disease. A Retrospective Study of Birth Size and Maternal Weight Gain**

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# *Objective*: To assess pregnancy and fetal outcomes in Jamaican subjects with sickle cell-haemoglobin C (SC) disease.

*Study Design*: A retrospective chart review over 21 years (1992-2012) of all pregnancies in SC disease and a comparison group matched by gender and date of delivery in mothers with a normal haemoglobin (AA) phenotype at the University Hospital of the West Indies, Jamaica. There were 118 pregnancies in 81 patients with SC disease and 110 pregnancies in 110 in the normal comparison group. Corrections were made for repeat pregnancies from the same mother. Outcome measures included maternal weight at 20, 25, 30, 35 and 38 weeks gestation, maternal pregnancy complications, birth weight, head circumference and crown heel length and were used to analyse possible predictors of birth weight.

*Results*: First antenatal visits occurred later in women with SC disease, who also had lower haemoglobin level and lower systolic blood pressure. The prevalence of pregnancy-induced hypertension, pre-eclampsia, ante-partum or postpartum haemorrhage did not differ between genotypes. Maternal weight gain was significantly lower in SC disease and there was a significantly lower birth weight, head circumference, and gestational age.

*Conclusions*: Pregnancy in SC disease is generally benign but mothers had lower weight gain and lower birth weight babies, the difference persisting after correction for gestational age.

**Word Count**: 211

**Key Words:** Weight gain; Haemoglobin SC Disease; Birth weight

**Introduction**

The term sickle cell disease covers a group of conditions in which pathology results from the presence of sickle haemoglobin (HbS)1. Inheriting the sickle cell gene from both parents results in homozygous sickle cell (SS) disease which is generally severe, whereas double heterozygotes with sickle cell-haemoglobin C (SC) disease are usually mildly affected.

In SS disease, pregnancy was associated with increased bone pain crises, acute chest syndrome, urinary tract infections and maternal mortality, increased fetal loss at every stage, and a low birth weight baby2,3. In the Jamaican Cohort Study4, which followed 150 females with SS disease from birth, 36% of pregnancies ended in spontaneous abortion, and completed pregnancies showed a lower gestational age and birth weight5. Four deaths occurred, two published in an earlier report5 giving a mortality rate of 2.1%but two further deaths occurred later (unpublished observations). Although pregnancy outcome in SS disease is variable and unpredictable, there is often a severe clinical course for both fetus and mother.

Sickle cell-haemoglobin C (SC) disease is the second most common form of sickle cell disease among mothers of West African ancestry and results from the inheritance of HbS and HbC genes. Early reports, certainly influenced by symptomatic selection, suggested that pregnancy in SC patients ran a more severe clinical course than in SS disease6,7 but it is now clear that SC patients have a more benign outcome2,8,9. These reports have tended to focus on maternal performance and complications, and the data on fetal outcome and birth weight are conflicting. To clarify this issue, the present study has addressed the birth outcome and infant size in a retrospective study of patients and an appropriate comparison group over 21 years at a single institution. This group has also provided an opportunity to examine some of the potential determinants of birth weight.

**Materials and Methods**

*Patients* Retrospective chart review of patients with SC disease delivering at the University Hospital of the West Indies (UHWI), Kingston, Jamaica over 21 years (Jan 1, 1992 and Dec 31, 2012) found 118 singleton pregnancies in 81 women (57 single pregnancies, 13 with two, 9 with three and 2 with four pregnancies). A comparison group of singleton pregnancies in 110 females with a normal haemoglobin (AA) phenotype was derived from the same source, matching maternal age within 1 year and delivery date within 1 day. Maternal measurements at the first antenatal visit included weight, height, haemoglobin level, systolic and diastolic blood pressure and serial measurements of weight were performed at 20, 25, 30, 35 and 38 weeks gestation. Indices of birth outcome included gestational age, birth weight, head circumference, crown-heel length and APGAR scores. Placental weight and estimated blood loss were also recorded. Postpartum haemorrhage was defined as blood loss greater than 500 ml in a spontaneous vaginal delivery or exceeding 1000 ml at Caesarean section. The study was approved by the University of the West Indies/UHWI Ethics Committee.

*Statistical Analysis* Since observations of weight gain required knowledge of the pre-pregnancy weight, analysis was limited to a subset of 88 SC pregnancies where mothers attended antenatal clinics before 16 weeks gestation when weight still reflected pre-pregnancy levels10. Maternal weight gain (overall and rate of weight gain), hospital admissions and the outcome variables were compared between the SC and comparison groups using a regression model. Adjusting for differences within, and between, mothers with more than one pregnancy was performed by a mixed linear random effects model for continuous outcomes and a mixed logistic regression random effects model for binary outcomes. Body mass index (BMI) was calculated as weight (kg)/height (m) squared. Statistical Package for the Social Sciences (SPSS) Version 22 was used.

**Results**

*First Antenatal Clinic Visit* Patients with SC disease presented later at first visit, had lower haemoglobin levels, and lower systolic blood pressure but maternal weight, height, body mass index, age or diastolic blood pressure did not differ (Table 1).

*Maternal Outcome* Prior to delivery-related admissions, there were 44 admissions among SC patients (23 bone pain crisis, 11 pre-eclampsia, 5 pregnancy induced hypertension, 4 urinary tract infection, 2 acute chest syndrome, 2 gestational diabetes), and 15 admissions in controls (5 pregnancy induced hypertension, 4 pre-eclampsia, 3 urinary tract infection, 2 vomiting, one gestational diabetes). There was no maternal mortality in either group. Pre-eclampsia was more common in SC mothers (11/118 [9.3%]) than comparison group (4/110 [3.6%]), although the difference was no longer significant (p=0.098) after correction for repeat pregnancies and there were no significant genotype differences in the prevalence of pregnancy induced hypertension (SC 5/118 [4.2%]; comparison group 5/109 [4.6%]), urinary tract infections (4/118 [3.4%]; 3/110 [2.7%]), mean duration of labour (8.64 hours; 8.02 hours), mode of delivery (Caesarean section 31/118 [26.2%]; 22/110 [20.0%]), or postpartum haemorrhage (6/115 [5.2%]; 7/109 [6.4%]). Maternal weight gain was consistently lower in SC disease throughout pregnancy and the total weight gain in mothers with SC disease completing 38 weeks gestation was 2.42 kg (Table 2) less than in the comparison group (p<0.0001).

*Infant Outcome*  Infants of SC mothers had lower gestational age and birth weight (Table 3), despite similar rates of induction of labour (SC 5; comparison group 6) and of Caesarean Sections. Infants of SC mothers weighed 443g less (95% CI 266-620) after controlling for repeat pregnancies. The difference was reduced to 299g (CI 156-441) after controlling for gestational age but remained highly significant (300g, CI 157-442g) after controlling for both gestational age and induction/operative deliveries (p<0.001). Focusing on the 51 pregnancies derived from the Cohort Study, the birth weight was 414g lower (CI 177-651, p=0.001) than the AA controls, after correction for individuals contributing more than one pregnancy. Low birth weight babies (<2500g) were more frequent in SC pregnancies (SC 22.9%, AA 5.5 %, p<0.0001) and head circumference and placental weight were also significantly lower than controls. Apgar scores of 7 at 1minute (SC 21/110 [18.8%], comparison group 17/110 [15.5%]) and 5 minutes (SC 7/118 [5.9%] versus 4/110 [3.6%]) did not differ between the groups, 2 0.6 and 0.2 respectively.

*Possible Determinants of Lower Birth Weight* In mothers with SC disease, birth weight was not influenced by admissions for bone pain crisis (p=0.755), acute chest syndrome (p=0.25), urinary tract infection (p=0.68), or pregnancy induced hypertension (p=0.42). Pre-eclampsia reduced the birth weight in both patients and comparison group but the difference was greater in SC disease (mean 979g, 95% C.I. 606 to 1353, p<0.001) than in the comparison group (734g, 158 to 1309, p=0.01) although only the difference in SC disease remained significant (542g, 229 to 855, p=0.001) after correction for gestational age. Haemoglobin levels at first antenatal clinic attendance did not influence birth weight, an identical birth weight of 2.91 kg occurring in 74 SC mothers with haemoglobin levels >10g/dl and in the 34 SC mothers with levels <10g/dl.

**Comments** *Main Findings:* Women with SC disease began antenatal attendance later than the comparison group, had lower total weight gain and their offspring had lower birth weight and lower placental weight even after correction for gestational age.

*Interpretation:* The later registration for antenatal care in SC disease, previously noted elsewhere11,12 is an enigma, although some of the delay in Jamaica may be artefactual since the University of the West Indies allows later registration of perceived ‘high-risk’ patients.

Consistent with previous observations2,9, pregnancy was generally well tolerated in SC disease with similar frequencies of pre-eclampsia, pregnancy induced hypertension, urinary tract infections, mode of delivery, and blood loss to the comparison group. Previous observations on fetal growth and birth weight in mothers with SC disease have reached conflicting conclusions and are difficult to evaluate without appropriate controls2,3,12,13, stratifying the results by genotype of sickle cell disease13, or have the weakness of combining results from multiple centres with different treatment regimens14. Some studies report intra-uterine growth retardation (IUGR) among SC mothers12,14, or an increase in low births weights (<2500g) without presenting birth weight figures15.In Jamaica, an earlier study of 21 pregnancies among 8 SC women found birth weight reduced compared to ‘normal values’ in the same hospital16, but data derived from the Jamaican Cohort Study9, found no difference in birth weight (SC 2,980g, AA 3.030g) in 95 pregnancies in 43 SC patients compared with matched AA controls followed from birth. These data appear to conflict with the lower birth observed in 51 deliveries in 32 SC mothers from the same Cohort in the present report and the difference remains unexplained although the earlier report9 included all deliveries at multiple institutions compared with a single hospital in the present study.

The lower birth weight in babies of SC mothers remained highly significant after correction for gestational age and possible contributing factors include a lower total haemoglobin or hospital admissions for bone pain crisis, acute chest syndrome and urinary tract infections. Low first trimester total haemoglobin levels (8.0-9.9g/dl) were associated with lower birth weight in a study in China17 but although one third of SC patients in the present study had haemoglobin levels below 10g/dl, there was no relationship with birth weight. This observation is consistent with the lowered oxygen affinity of HbS implying that haemoglobin levels do not reflect the oxygen carrying capacity. Hospital admissions were more frequent in SC disease but although a reduced birth weight was associated with total admissions, those for the three common specific sickle cell related pathologies, bone pain crisis, acute chest syndrome or urinary tract infection failed to show any relationship with the lowered birth weight. A lower birth weight was associated with pre-eclampsia in both cases and the comparison group but the difference remained significant only in SC disease after correction for gestational age.

*Strengths and Limitations:* A strength of this study is the relatively large number of pregnancies studied at a single institution, with complete follow-up and no policy for routine transfusion support. Limitations include its retrospective nature and the 21 year interval over which therapeutic approaches could have changed but there was no evidence of this.

**Conclusion**

Pregnancy outcome in SC disease is generally benign but maternal weight gain and birth weight is lower than in controls even after correction for the lower gestational age.

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**Disclosure of Interest:** None of the authors has any interest to disclose or any conflict of interest

**Author’s Contribution:** All authors played major roles in contributing to this work.MT, ISM, RMC and GS contributed to the conception of the study and the planning of the methodology of the study. The collection of the data was performed by MT and ISM while the analysis of the study was performed by CO . All authors contributed to the writing of the manuscript and accept responsibility for the manuscript.

**Details of Ethics Approval**

### The University of the West Indies/UHWI Ethics Committee which is the institutional ethics committee responsible for human experimentation gave ethical approval the 4th February, 2013 and the reference number is 107, 2004/2005.

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