Total Plasma Protein in Very Preterm Babies: Prognostic Value and Comparison with Illness Severity Scores

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Abstract

Objective: We aimed to investigate the predictive value for severe adverse outcome of plasma protein measurements on day one of life in very preterm infants and to compare total plasma protein levels with the validated illness severity scores CRIB, CRIB-II, SNAP-II and SNAPPE-II, regarding their predictive ability for severe adverse outcome.

Methods: We analyzed a cohort of infants born at 24–31 weeks gestation, admitted to the tertiary intensive care unit of a university hospital over 10.5 years. The outcome measure was “severe adverse outcome” defined as death before discharge or severe neurological injury on cranial ultrasound. The adjusted odd ratio (aOR) and 95% confidence interval (95% CI) of severe adverse outcome for hypoproteinemia (total plasma protein level <40 g/L) was calculated by univariate and multivariate analyses. Calibration (Hosmer-Lemeshow goodness-of-fit) was performed and the predictive ability for severe adverse outcome was assessed for total plasma protein and compared with CRIB, CRIB-II, SNAP-II and SNAPPE-II, by calculating receiver operating characteristic (ROC) curves and their associated area under the curve (AUC).

Results: 761 infants were studied: 14.4% died and 4.1% survived with severe cerebral ultrasound findings. The aOR of severe adverse outcome for hypoproteinemia was 6.1 (95% CI 3.8–9.9). The rank order for variables, as assessed by AUCs and 95% CIs, in predicting outcome was: total plasma protein [0.849 (0.821–0.873)], SNAPPE-II [0.822 (0.792–0.848)], CRIB [0.821 (0.792–0.848)], SNAP-II [0.810 (0.780–0.837)] and CRIB-II [0.803 (0.772–0.830)]. Total plasma protein predicted severe adverse outcome significantly better than CRIB-II and SNAPPE-II (both p<0.05). Calibration for total plasma protein was very good.

Conclusions: Early hypoproteinemia has prognostic value for severe adverse outcome in very preterm, sick infants. Total plasma protein has a predictive performance comparable with CRIB and SNAPPE-II and greater than other validated severity scores.

Introduction

Advances in perinatal care have resulted in an improvement in the survival rate of very low birth weight infants (VLBWI) as well as in a decrease of the disability rate in VLBW survivors [1,2]. Amidst all the progress made over decades, neonatal outcome has been benefiting from the development and the use of illness severity scores, which have permitted quality of care evaluation, risk adjustment comparisons in benchmarking studies, management and resource implementation. The Clinical Risk Index for Babies (CRIB) [3], its revised version (CRIB-II) [4], the Score for Neonatal Acute Physiology II (SNAP-II) and the SNAP Perinatal Extension II (SNAPPE-II) [5] are the most widely used scores to estimate illness severity and in-hospital mortality risk in the neonatal intensive care unit (NICU).

Some authors have underlined that risk adjustment using these scores is imperfect because additional perinatal factors may significantly influence VLBWI survival [6]. Others have noted the deterioration over time in their predictive performance, due to incremental improvement of care [7]. Actually, the relationship between physiological status and mortality risk may change as preventive interventions, monitoring strategies and new treatment protocols are introduced. Infants with similar illness severity scores may differ for their risk of death [6] and a better understanding of all the perinatal factors influencing mortality remains a meaningful challenge for neonatologists.

Another point at issue is that, even if survival has progressively improved especially after extremely preterm birth, the high rate of disability in survivors is still a concern [8]. So, for the clinician involved in assessing the initial risk of high vulnerable neonatal populations, death is not the only important adverse outcome, and a primary goal is also to quantify the risk of severe brain damage.

Recently we reported, for the first time, that hypoproteinemia (total protein level of less than 40 g/L) on day 1 of life is an independent factor associated with severe adverse outcome (SAO), defined as in-hospital death or severe neurological injury on
cranial ultrasound, in a large sample of critically ill preterm babies [9]. In order to further investigate the clinical interest of this finding, we performed a study with the following objectives: 1) to confirm the prognostic value for SAO of hypoproteinemia in another population of VLBWI; 2) to compare total plasma protein levels alone with composite scores CRIB, CRIB-II, SNAP-II and SNAPPE-II, regarding their predictive ability for SAO in this population.

Methods

Ethics statement
This study was approved by the institutional medical research ethics committee (Comité de Protection des Personnes Sud-Ouest et Outre Mer III, authorization number 2012/36).

According to French legislation, written parental consent was not needed for this study.

Design and study population
The study design was an observational cohort analysis of all the infants born between 24 and 31 weeks of gestational age (GA) and admitted within 12 hours of life to the tertiary NICU of Saint Pierre University Hospital (Reunion Island, France) during a 10.5 year period (1 January 2001 to 30 June 2011). Patients were excluded if they died within the first 12 hours after birth, if clinical data were incomplete, if plasma protein value on the first day of life was not available or if any of the items for calculating the CRIB, CRIB-II, SNAP-II and SNAPPE-II were missing.

Data collection
Clinical data were drawn from the unit perinatal database, which has prospectively recorded demographic, gestational and perinatal variables of all mother-infant pairs since 2001. This recording of mother-infant clinical information was approved by the National Committee for data protection (Commission Nationale de l’Informatique et des Libertés, registration number 1620660). Information was collected at the time of delivery and at the time of infant hospital discharge and regularly audited by appropriately trained staff. For the purpose of this study, records have been validated and have been used anonymously. Available information in this data set included maternal age, parity and gravidity, pre-existing clinical and gynecological diseases, whether singleton or multiple pregnancy, obstetric history and illness, antenatal steroid administration, labor and delivery complications, mode of delivery (vaginal/caesarean section), outborn (transfer after birth, admitted into NICU within 12 hours of life), gender, GA, birth weight (BW), cord blood lactate concentration, 1 and 5 minute Apgar score, neonatal morbidities and neonatal death.

Total protein values, CRIB, CRIB-II, SNAP-II and SNAPPE-II
Data on total protein values, CRIB, CRIB-II, SNAP-II and SNAPPE-II were abstracted from the medical charts.

According to the unit guidelines for assessing fluid and electrolyte status in VLBWI, babies born less than 32 weeks of GA and hospitalized in our NICU had a blood sample at around 12 hours of life. The plasma total protein measurement provided by this blood sample was used for the purpose of this study.

Hypoproteinemia was defined as a total protein level of less than 40 g/L [9,10].

CRIB was calculated as already described [3], using the six variables: BW, GA, presence of congenital malformations, maximum base excess in the first 12 hours of life, and minimum and maximum appropriate fraction of inspired oxygen – Fo2– (FiO2 min) and (FiO2 max) in the first 12 hours. CRIB-II was calculated using the variables BW, GA, temperature and base excess on admission [4]. Following the original paper from Richardson et al [5], the six item score (urine output, lowest mean blood pressure, worst PaO2/FiO2 ratio, lowest pH, occurrence of seizures, lowest temperature) was used for SNAP-II. To this, BW, small for GA, Apgar at 5 minutes were added for calculating the SNAPPE-II [5]. The data collection window for these two scores was the first 12 hours after birth.

Outcome measure
The outcome measure was SAO. This was defined as death or severe (grade 3 or 4) intraventricular hemorrhage or cystic periventricular leukomalacia occurring before infants’ discharge from hospital. Intraventricular hemorrhage was graded at cerebral ultrasound according to Papile et al [11]. Cystic periventricular leukomalacia was defined by cranial ultrasound as an area of increased echogenicity of the periventricular white matter in acute phase which subsequently evolved into cystic lesion [12]. Cranial ultrasounds were routinely performed during the infant hospital stay by experienced examiners according to the following protocol: day 1, 7, 10, 13 and then at least every 2 weeks or more often as clinically indicated, until discharge.

Statistical analysis
We assessed the prognostic value for SAO of total plasma protein in two sets of analyses.

In the first analysis, we estimated and compared the incidence of SAO in hypoproteinemiac patients and in patients with normal protein values. The crude odd ratio (OR) and 95% confidence interval (95% CI) of SAO for hypoproteinemia were calculated. We then used backward stepwise logistic regression to obtain the adjusted OR for potential confounding variables. Univariate analysis was first performed and variables significant at a p value <0.10 were entered in the multivariate model. For this analysis continuous variables were categorized by cut-off values chosen as having the highest Youden index [13].

In the second analysis, discrimination-that is the ability to correctly predict SAO-was assessed for total plasma protein levels and compared with CRIB, CRIB-II, SNAP-II, SNAPPE-II, BW and GA discrimination, by calculating receiver operating characteristic (ROC) curves and their associated area under the curve (AUC), with 95% CI based on the observed values entered on a continuous scale [14]. An AUC value of 0.5 indicates no better than chance ability to discriminate and larger values indicate increasing ability. A value above 0.8 is considered good [15]. Calibration of total plasma protein levels, CRIB, CRIB-II, SNAP-II, SNAPPE-II, BW and GA was investigated using the Hosmer and Lemeshows (HL) goodness-of-fit test, which categorizes the observations into groups according to their predicted risk [15]. The numbers of predicted and observed outcomes within each of these groups are compared. A non-significant p value of the HL indicates an acceptable calibration.

Comparisons between groups were performed using χ²-test or Fisher’s exact test for categorical variables; the ANOVA test was used for parametric variables and the Mann-Whitney U test for non-parametric continuous variables. Comparison of the AUCs was evaluated by the DeLong method [16]. All statistical analyses were carried out using the MedCalc. ver. 12.3.0.0 statistical software package (MedCalc Software Mariakerke, Belgium) and p values <0.05 were considered statistically significant.

Results
From 1 January 2001 to 30 June 2011, 841 neonates born below 32 weeks of gestation were hospitalized in our NICU within the
first 12 hours of life. Three babies died within 12 hours after birth, clinical data were missing or incomplete for 26, and total protein value was not obtainable for 51 infants. The items to calculate CRIB, CRIB-II, SNAP-II and SNAPPE-II were always available, so all the remaining 761 newborns were eligible for the analysis. Infants with missing data were similar in their characteristics and outcomes when compared to those included in the study (data not shown).

SAO was present in 140 patients (18.4% of the study population): 109 patients (14.4%) died and 31 (4.1%) survived with severe cerebral ultrasound findings. Table 1 shows antenatal characteristics of the study population, characteristics at birth and postnatal diseases.

Hypoproteinemia occurred in 24% of all patients. The blood sample was carried out at 14.7±5.3 (mean±SD) hours after birth. The rate of SAO was 52.5% in patients with hypoproteinemia and 7.6% in those with normal protein values (p<0.001; crude OR 13.3; 95% CI 8.7–20.4). The univariate procedure yielded 16 variables for inclusion into the multivariate logistic model: maternal age≥25 years, multiple pregnancy, time-frame, steroid administration, cesarean section delivery, outborn, male gender, gestational age, FiO2 max>40%, FiO2 min>23%, Apgar score≤3 at 1 minute, hypoproteinemia, cord blood lactate>4 mmol/L, congenital malformations, admission temperature≤35.1°C and anemia at birth (data not shown).

Results of the multivariate analysis are reported in table 2.

There was a significant linear association between GA and plasma protein levels, (r² 0.23, p<0.001, data not shown). However at the multivariate analysis both were independent factors strongly associated with SAO. The adjusted OR of SAO for hypoproteinemia after correcting for the confounding variables was 6.1 (95% CI 3.8–9.9).

Figure 1 shows the scattergram illustrating the total plasma protein values in infants with SAO compared to those who survived without severe neurological injury.

ROC curves for total plasma protein levels, illness severity scores and BW are reported in Figure 2.

The rank order for variables, as assessed by AUCs and 95% CIs in predicting SAO, was plasma protein levels, SNAPPE-II, CRIB, SNAP-II and CRIB-II. Plasma protein levels predicted SAO significantly better than CRIB-II and SNAP-II, while no significant difference was seen with CRIB and SNAPPE-II. Calibration for plasma protein levels was very good (p = 0.79 at the HL goodness-of-fit test, result shown in table 3).

**Table 1.** Population characteristics (n=761 preterm infants<32 weeks gestation).

| Antenatal variables | Maternal age, mean±SD, y | 27.7±6.8 |
|---------------------|--------------------------|----------|
| Hypertensive Disease of Pregnancy, % | 22.3 |
| Maternal Diabetes, % | 9.7 |
| Maternal Body Mass Index>30 before pregnancy, % | 16.0 |
| Antenatal Steroids, % | 69.6 |
| Singleton Pregnancy, % | 78.8 |
| Population characteristics | Birth Weight, mean±SD, g | 1203±346 |
| Gestational Age, mean±SD, wk | 28.8±1.9 |
| Small for Gestational Age, % | 9.9 |
| Male Gender, % | 50.3 |
| Apgar Score≤3 @ 1 minute, % | 10.1 |
| Caesarean Section, % | 56.4 |
| Outborn, % | 2.0 |
| Congenital malformations, % | 2.7 |
| Cord blood lactate concentration, mean±SD, mmol/L | 4.0±3.5 |
| CRIB, mean±SD | 2.6±3.1 |
| CRIB-II, mean±SD | 6.2±3.4 |
| SNAP-II, mean±SD | 14.7±14.8 |
| SNAPPE-II, mean±SD | 20.7±20.2 |
| Hypoproteinemia, % | 24.0 |
| RDS requiring invasive mechanical ventilation, % | 59.4 |
| Anemia at birth, % | 15.6 |
| Treated hypotension on day 1 of life, % | 22.6 |
| Treated Patent Ductus Arteriosus % | 15.1 |
| Necrotizing Enterocolitis, % | 4.5 |
| Bronchopulmonary Dysplasia @ 36 weeks, % | 4.7 |

(CRIB) Clinical Risk Index for Babies; (CRIB-II) revised Clinical Risk Index for Babies; (SNAP-II) Score for Neonatal Acute Physiology II; (SNAPPE-II) Score for Neonatal Acute Physiology Perinatal Extension II; (RDS) Respiratory Distress Syndrome.

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Table 3 shows the results of AUCs curves, AUCs comparison and calibration for all these variables. Results are compared to predictive accuracy of BW and GA.

The model which analyzed GA and plasma protein simultaneously had an AUC of 0.873 (0.847–0.896) and the calibration at the HL goodness-of-fit test showed p = 0.39.

Discussion

Our study has shown that low plasma protein values in the first day of life have a strong predictive ability for severe outcome in VLBWI. This result is consistent with our previous finding [9], which could be confirmed in another population, differing from the former for perinatal characteristics, rates of impaired outcome, and care practice. Moreover, the result was stable over time in this study cohort.

In the present study, the performance of protein levels was comparable to that of the CRIB and of the SNAPPE-II and greater than that of other validated severity scores, indicating that additional real time indices of neonatal illness, based on pathophysiological concepts, must be taken into account when considering factors influencing mortality and morbidity in vulnerable newborns.

The interest of this approach in predicting outcome is not new and it has been reported elsewhere: recently, two papers [17,18] have highlighted the prognostic value of high early lactate levels, which in one case showed a mortality predictive ability similar to that of CRIB and CRIB-II in VLBWI [18]. One study from De Felice et al. [19], which investigated the predictive accuracy of CRIB and CRIB-II in VLBWI has underlined the need to seek new clinical risk-adjustment markers in high-risk newborns; these authors had previously demonstrated the relationship between physiological markers of early peripheral microcirculatory changes and neonatal illness severity scores in sick babies after birth [20,21].

What our study adds, in relation to previous investigations in this domain, is the reproducible and remarkable prognostic value for the outcome, in very preterm infants, of plasma protein levels measured early in postnatal life. This biological variable has never been considered in establishing risk scores for neonatal disease severity [22], even when these scores were designed for measuring morbidity and mortality risk by mainly taking into account individual and physiological characteristics of the infants, as in the SNAP [23], in the SNAP-II or in the Neonatal Therapeutic Intervention Scoring System [24].

This assertion, far from representing a criticism of the above cited, well validated scores, simply aims to draw attention to a simple and easily available marker of physiological impairment, possibly influencing the in-hospital outcome of the vulnerable preterm baby.

The present study was not designed to investigate the pathophysiological basis of the association ‘plasma protein and in-hospital outcome’ of VLBWI, but we have previously described a significant positive correlation between low colloid oncotic pressure, low total protein levels and hypotension on day 1 of life in newborns with respiratory distress [25], and we hypothesised in one recent paper that low early protein levels may impair maintenance of
intravascular volume and adequate blood flow to vital organs in critically ill premature babies [9].

Obviously, the hypothesis that low plasma protein levels may influence the cardiovascular adaptation and blood flow perfusion to organs in the immediate postnatal period must be rigorously established in future, prospective investigations.

It is worth noting here, that one of the factors associated with impaired outcome in VLBWI is the failure in cardiovascular adaptation after birth [26–28]. As we know, this variable is not among the items integrated in risk scores based on fixed covariates, such as the CRIB-II, and it is measured by parameters such as hypotension and urine output in scores conceived to capture the mortality risk also linked to physiological variables (SNAPP and SNAP-II). Now, hypotension is a rough and delayed sign of impaired cardiovascular adaptation and in addition, more subtle disturbances in cardiovascular function display significant relationships with clinical illness severity [29,30].

In our view, the search for new risk-indicators in sick preterm infants should take into account markers which could influence transitional circulation.

The limitations of our analysis include that, due to the study design, the collection window for protein value was variable around the first 12 hours after birth and extending up to 20 hours of life, so that both obstetrical and neonatal care measures during the early postnatal period may have influenced this parameter. Similarly, some data were recorded afterwards and this may have reduced the reliability and accuracy of the collected information. Finally, this report does not allow the conclusion that modifying plasma protein levels is necessary to improve outcome in VLBWI.

Nonetheless, our result supports the new finding that protein values represent a marker of abnormal physiological state which is strongly predictive of impaired outcome in preterm babies.

**Conclusion**

The emphasis of this paper was to encourage neonatologists to pay close attention to plasma protein levels during postnatal transition in preterm babies, as these have a prognostic value for...
adverse in-hospital outcome of very premature, sick infants. Further investigations are needed to determine whether any benefit can be obtained by adding protein measurements to other validated illness scores for newborns. Moreover, our finding raises the interest in addressing the above issues regarding new markers of physiological derangement in early postnatal life, in future, prospective studies on VLBW, thus providing additional insight into factors influencing mortality and morbidity for this vulnerable population.

References

1. Bode MM, D’Eugenio DB, Forsyth N, Coleman J, Gross CR et al (2009) Outcome of extreme prematurity: a prospective comparison of 2 regional cohorts born 20 years apart. Pediatrics 124: 466–474.

2. van Haastert IC, Groenendaal F, Uiterwaal CS, Termote JU, van der Heide-Jalving M et al (2011) Decreasing incidence and severity of cerebral palsy in prematurely born children. J Pediatr 159: 86–91.

3. International Neonatal Network (1995) The CRIB (clinical risk index for babies) score: a tool for assessing initial neonatal risk and comparing performance of neonatal intensive care units. Lancet 346: 193–196.

4. Parry G, Tucker J, Tarnow-Mordi WO (2005) CRIB-II: an update of the clinical risk index for babies score. Lancet 366: 1789–1791.

5. Richardson DK, Coccoran JD, Escobar GJ, Lee SK (2000) SNAP-II and SNAPPE-II: Simplified newborn illness severity and mortality risk scores. J Pediatr 138: 92–100.

6. Gugliardi L, Cavazza A, Brunelli A, Battaglioli M, Merazzi D et al (2004) Assessing mortality risk in very low birthweight infants: a comparison of CRIB, CRIB-II, and SNAPPE-II. Arch Dis Child Fetal Neonatal Ed 89: F419–F422.

7. Bührer C, Metz B, Ohlade N (2000) CRIB, CRIB-II, birth weight or gestational age to assess mortality risk in very low birth weight infants? Acta Paediatr 97: 999–1005.

8. Wood NS, Marlow N, Costeloe K, Gibson AT, Wilkinson AR (2000) Neurologic and developmental disability after extremely preterm birth. EPICare Study Group. N Engl J Med 343: 576–584.

9. Iacobelli S, Bonnante F, Lacoutière C, Ferdynus C, Cottenet J, et al (2012) Hypoproteinemia on the first day of life and adverse outcome in very preterm babies admitted to the neonatal intensive care unit. J Perinatol 32: 520–524.

10. Reading RF, Ellis R, Fleweto A (1998) Plasma albumin and total protein in preterm babies from birth to eight weeks. Early Hum Dev 57: 37–42.

11. Papile LA, Burstein J, Burstein K, Koffee H (1978) Incidence and evolution of subependymal and intraventricular hemorrhage: a study of infants with birth weights less than 1,500 gm. J Pediatr 92: 529–534.

12. deVries LS, Eken P, Dhowowitz LMS (1992) The spectrum of leukomalacia using cranial ultrasound. Behav Brain Res 49: 1–6.

13. Fluss R, Faraggi D, Reiner B (2005) Estimation of the Youden Index and its associated cutoff point. Biom J 47: 458–72.

14. Hosmer DW, Lemeshow S (1989) Applied logistic regression. In: J Wiley and Sons editors, New York. pp 135–175.

15. DeLong ER, DeLong DM, Clarke-Pearson DL (1988) Comparing the areas under two or more correlated receiver operating characteristic curves: a nonparametric approach. Biometrics 44: 837–845.

16. DeLong ER, DeLong DM, Clarke-Pearson DL (1988) The area under the receiver operating characteristic (ROC) curve. Radiology 143: 29–36.

17. Hussain F, Gilhman K, Gray PH (2009) Does lactate level in the first 12 hours of life predict mortality in extremely premature infants? J Paediatr Child Health 45: 263–267.

18. Phillips LA, Dewhurst CJ, Yoxall CW (2011) The prognostic value of initial blood lactate concentration measurements in very low birthweight infants and their use in development of a new disease severity scoring system. Arch Dis Child Fetal Neonatal Ed 96: F275–280.

19. De Felice C, Del Vecchio A, Latini G (2005) Evaluating illness severity for very low birth weight infants: CRIB or CRIB-II? J Matern Fetal Neonatal Med 17: 257–260.

20. De Felice C, Toi P, Parrini S, Del Vecchio A, Bagossi F et al (2005) Histologic chorioamnionitis and severity of illness in very low birth weight newborns. Pediatr Crit Care Med 6: 298–302.

21. Dorling JS, Field DJ, Manktelow B (2005) Neonatal disease severity scoring systems. Arch Dis Child Fetal Neonatal Ed 90: F11–F16.

22. De Felice C, Vacca P, Del Vecchio A, Criscuolo M, Lozapone A et al (2004) Early postnatal skin colour changes in term newborns with subclinical histological chorioamnionitis. Eur J Pediatr 163: 550–554.

23. Dorling JS, Field DJ, Manktelow B (2005) Neonatal disease severity scoring systems. Arch Dis Child Fetal Neonatal Ed 90: F11–F16.

24. Richardson DK, Gray JE, McCormick MC, Workman K, Goldmann DA (1993) Score for Neonatal Acute Physiology: a physiological severity index for neonatal intensive-care. Pediatrics 91: 617–623.

25. Gray JE, Richardson DK, McCormick MC, Workman K, Goldmann DA (1992) Neonatal Therapeutic Intervention Scoring System: a therapy-based severity-of-illness index. Pediatrics 90: 561–567.

26. Zimmermann B, Françoise M, Germain GF, Lallemand C, Gouyou-BB (1997) Colloidal oncotic pressure in neonatal respiratory distress syndrome. Arch Pediatr 4: 952–958.

27. Micali-Aless VM, de Vries LS, Welsh AG (1987) Mean arterial blood pressure and neonatal cerebral lesions. Arch Dis Child 62: 1066–1069.

28. Fanaroff JM, Wilson-Costello DE, Newman NS, Montpetite MM, Fanaroff AA (2006) Treated hypotension is associated with neonatal morbidity and bearing loss in extremely low birth weight infants. Pediatrics 117: 1131–1135.

29. Pollicer A, del Carmen Bravo M, Madero R, Salas S, Quero J et al (2009) Early systemic hypotension and vasopressor support in low birth weight infants: impact on neurodevelopment. Pediatrics 123: 1369–1376.

30. Sokoloff M, Borzagge M, Serti I (2010) Hemodynamic monitoring in neonates: advances and challenges. J Perinatol 30: 830–845.

31. Stark MJ, Clifton VL, Wright IMR (2008) Microvascular flow, clinical illness severity and cardiovascular function in the preterm infant. Arch Dis Child Fetal Neonatal Ed 93: F271–F274.

Author Contributions

Conceived and designed the experiments: SI FB CQ. Performed the experiments: SI FB CQ. Analyzed the data: SI FB CQ. Contributed reagents/materials/analysis tools: FB CQ PYR CB. Wrote the paper: SI. Reviewed and critically revised the manuscript drafts: CB JBG.