Serum cystatin C levels in preterm newborns in our setting: Correlation with serum creatinine and preterm pathologies

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ABSTRACT

Background: Cystatin C (CysC) is a renal function marker that is not as influenced as creatinine (Cr) by endogenous or exogenous agents, so it is proposed as a marker in preterm infants.

Objectives: To determine serum CysC values in preterm infants during the first week of life, compared to Cr. To analyse alterations caused by prematurity diseases.

Method: The design involved a longitudinal, observational study of prospective cohorts. Groups were based on gestational age (GA): Group A (24–27 weeks), Group B (28–33 weeks), Group C (34–36 weeks). Blood samples were collected at birth, within 48–72 h and after 7 days of life.

Statistics: SPSS v.20 software was used. The statistical methods applied included chi-squared test and ANOVA.

Results: A total of 109 preterm infants were included in the study. CysC levels were 1.54 mg/l (±0.28) at birth, 1.38 mg/l (±0.36) within 48–72 h of life, and 1.50 mg/l (±0.31) after 7 days (p<0.05). Cr levels were 0.64 mg/dl (±0.17) at birth, 0.64 mg/dl (±0.28) within 48–72 h, and 0.56 mg/dl (±0.19) after 7 days (P<.05). CysC values were lower in hypotensive patients and in those with a respiratory disease (P<.05), and no alterations associated with other diseases were observed. There were no differences in Cr levels associated with any disease. Creatinine levels were higher in patients ≤1.500 g (P<.05).

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Conclusions: Serum CysC decreased within 48–72 h of life, and this decline showed significance (\(P < .05\)). The levels increased after 7 days in all 3 GA groups, and there was no difference in CysC levels among the groups. More studies in preterm infants with hypotension and respiratory disease are required. CysC is a better glomerular filtration rate (GFR) marker in ≤1,500 g preterm infants.

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Valores de cistatina C sérica en recién nacidos pretérmino en nuestro medio. Relación con valores de creatinina sérica y patologías de la prematuridad

RESUMEN

Antecedentes: La cistatina C (CisC) es un marcador de función renal no tan influenciado como la creatinina (Cr) por agentes endógenos o exógenos, por lo que se propone como marcador en el pretérmino.

Objetivos: Determinar valores de CisC sérica en pretérminos en la primera semana de vida, comparándola con la Cr. Analizar modificaciones por patologías de la prematuridad.

Método: Estudio longitudinal, observacional, de cohortes prospectivo.

Grupos por edad gestacional (EG): grupo A (24–27 semanas), grupo B (28–33 semanas), grupo C (34–36 semanas). Se recogieron muestras de sangre al nacimiento, a las 48–72 h y a los 7 días.

Estadística: Programa SPSS v.20. Métodos estadísticos utilizados χ² y ANOVA.

Resultados: N = 109 pretérminos. CisC al nacimiento: 1,54 mg/l (± 0,28), a las 48–72 h de vida: 1,38 mg/l (± 0,36), a los 7 días: 1,50 mg/l (± 0,31) (p < 0,05). Cr al nacimiento: 0,64 mg/dl (± 0,17), a las 48–72 h: 0,64 mg/dl (± 0,28), a los 7 días: 0,56 mg/dl (± 0,19) (p < 0,05). Valores de CisC más bajos en pacientes con patología respiratoria e hipotensos (p < 0,05) sin modificación según patologías restantes. No diferencias en valores de Cr según patología. Valores de creatinina más altos en pacientes ≤ 500 g (p < 0,05).

Conclusiones: Descenso de CisC sérica a las 48–72 h de vida, siendo esta caída en el tiempo significativa (p < 0,05), ascenso a los 7 días, en los 3 grupos de EG y sin diferencias en valores de CisC entre los grupos. Se requieren más estudios en pretérminos con patología respiratoria y situaciones de hipotensión. En ≤ 1,500 g la CisC es mejor marcador de filtrado glomerular (FG).

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Introduction

The best renal function assessment index is the estimation of glomerular filtration rate (GFR). Procedures for the measurement of such filtrate are based on the renal clearance of exogenous or endogenous molecules. Creatinine is the most commonly used endogenous marker. Given that its production is proportional to the muscular mass of individuals and that it is not only freely filtered by the glomerulus but also secreted by the proximal tubule and, in addition, serum Creatinine is not sensitive enough for the identification of early stages of renal damage. Serum Creatinine is not the most adequate marker for the paediatric population in general and the neonatal population in particular. As an alternative to creatinine, several biological markers have been suggested, cystatin C (Cys C) being the most commonly analysed.

Cys C is a non-glycosylated, low molecular weight (13,343 Da), cationic protein, with 120 amino acids and two disulphide bridges. It has a protective function: inhibition of enzymes involved in protein metabolism, collagen catabolism and cellular matrix degradation and possible involvement in defense mechanisms against viral and bacterial infections. This protein is regularly synthesised by most nucleated cells, with a considerable distribution volume in bodily fluids. At the renal level, it is freely filtered by the glomerulus, due to its low molecular weight and positive charge at physiological pH, and it is reabsorbed and catabolised by proximal tubular cells. Under normal conditions, urinary concentration is very low if there is no tubular damage. It is not affected by muscular mass, nutrition status, size, age, gender, serum proteins, bilirubin or drugs, though it may present variations in cases of thyroid dysfunction, tumours or inflammatory diseases. Therefore, authors such as Filler recommend the use of equations based on Cys C, rather than the Schwartz formula to estimate GFR in children.

GFR is low in foetal and neonatal life. It increases after birth and reaches a maximum of 60 ml/min/1.73 m² during the first
3–5 weeks of life in normal-term and premature newborns (NB). Approximately 7.36% of NB in our setting are preterm and 1.2% weigh less than 1500 g. This population requires a more adequate filtration marker than the current one due to its physiological characteristics: low weight, low body mass index, reduced muscular mass, tendency to early renal failure arising from the prematurity itself, as well as from the added pathology and the use of nephrotoxic drugs.

This study was planned because of the difficulty to identify Cys C reference values in neonates, since these values are defined in a reduced number of preterm newborns and only during the first 3 postnatal days. The conducted study determines said values during the first 7 days after delivery by relating them with gestational age (GA) and serum levels of creatinine.

**Objectives**

- To determine values of serum concentration of Cys C, as a renal function marker, in preterm NB during the first week of life in our population upon conducting a comparative study with serum creatinine.
- To analyze if the serum concentration of Cys C and creatinine is modified due pathologies related to being premature and/or nephrotoxic drugs.

**Material and methods**

This is a longitudinal, observational, prospective, cohort study. It was conducted in the newborn intensive care unit on preterm new born (NB) born in and/or admitted to the Hospital Universitario Virgen Macarena de Sevilla from July 2010 to May 2012. The protocol was approved by the Ethics Committee of our centre. Informed consent was obtained from the neonates’ legal representatives. Preterm newborns were divided into three groups according to the (degree of prematurity) (GA): Group A (24–27 weeks), Group B (28–33 weeks) and Group C (34–36 weeks). The analyzed variables were GA, weight, gender, administration of nephrotoxic drugs: antibiotics (cephotaxime, vancomycin, gentamicin), furosemide and ibuprofen IV at different doses based on weight in kg, GA in weeks and days of postnatal life, according to the Neofax® handbook, 2011 version. There was no pharmacokinetic control of the drugs that could potentially be monitored; there was coexistence of associated respiratory pathologies (transient tachypnea or hyaline membrane of I–IV degree, which increases according to the severity of the radiological pattern), hypoxic ischaemic encephalopathy, acute renal failure (ARF), intraventricular haemorrhage, necrotising enterocolitis, sepsis and severe low blood pressure (blood pressure below the third percentile, according to GA, gender and days of life). Blood, Cys C and creatinine samples were collected at birth, at 48–72 h of life and at 7 days. Blood samples were obtained from arteries, veins or capillaries during the first 7 days after delivery, when routine samples were collected. Serum Cys C was measured through nephelometry (Particle-Enhanced Nephelometric Immuno-Assay) (BNII Siemens) with an ERM-DA471/IFCC60 certification that ensures its standardisation and traceability. Serum creatinine was measured through the Jaffe method using the IDMS (isotope dilution mass spectrometry)-standardised analyser, Cobas 6000 from Roche.

To calculate the intra-assay precision of Cys C and serum creatinine determinations, two different samples were selected: a sample with normal values and a pathological sample, with known concentrations in both cases; and these were entered 21 consecutive times into the corresponding analysers for each technique. As a result, we obtained the intra-assay coefficients of variation (CV) shown below:

- Cystatin: at 1 mg/l, CV: 2.9%; and at 2.1 mg/l, CV: 2.9%.
- Creatinine: at 1.8 mg/l, CV: 2%; and at 8.5 mg/l, CV: 1.2%.

Furthermore, to calculate the intra-assay precision of Cys C and serum creatinine determinations, two samples with known concentrations (once again, a normal sample and a pathological sample) were selected and determined by duplicate analysis during 10 consecutive days in the corresponding analysers for each technique. As a result, we obtained the inter-assay coefficients of variation (CV) shown below:

- Cystatin: at 1 mg/l, CV 2.6%; and at 2.1 mg/l, CV 2.4%.
- Creatinine: at 1.8 mg/l, CV 3.7%; and at 8.5 mg/l, CV 2.8%.

The statistical analysis was performed using the SPSS v.20 software, and the statistical methods used included the x2 method and the variance analysis of repeated measures on one factor. The p < 0.05 value was considered statistically significant.

**Results**

Total number of studied preterm newborns (n) was 109; 62 (56.9%) were male individuals and 47 (43.1%) female individuals. There were no differences among GA groups based on gender (p = 0.97). Group A had 10 preterm newborns; group B had 50 preterm newborns; and Group C had 49 preterm newborns.

Mean GA of the studied population was 32 weeks, range: 24–36 weeks. For Group A it was 26 weeks; range: 24–27 weeks, for Group B 31 weeks; range: 28–33 weeks and Group C 35 weeks; range: 34–36 weeks.

Mean weight of the studied population was 1767 g, range: 620–3505 g. For Group A it was 911 g; range: 620–1210 g, for Group B 1612 g; range: 845–2435 g and for Group C 2099 g; range: 1260–3505 g.

Table 1 shows the values of serum Cys C and creatinine at birth, at 48–72 h and at the first week of life, as well as the values in each subgroup of the population. Fig. 1a shows the changes in the values of Cys C in each GA group over time. There is a decrease in the serum concentration of Cys C at 48–72 h of life and an increase at 7 days (p < 0.05). This decrease over time occurs in the 3 GA groups, and, though the values of Cys C are higher as the GA of preterm newborns increases, there are no differences among groups (p = 0.07).

Fig. 1b shows the changes in the values of creatinine in each GA group over time. There is a decrease over time, and...
Table 1 – Plasma cystatin C and creatinine values during the first week of life and arranged by gestational age groups.

<table>
<thead>
<tr>
<th>Time</th>
<th>Cystatin C (mg/l)</th>
<th>Creatinine (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Birth Mean ± SD (range)</td>
<td>Birth Mean ± SD (range)</td>
</tr>
<tr>
<td><strong>Group A (n = 10)</strong></td>
<td>1.44±0.28 (0.91–1.81)</td>
<td>0.57±0.19 (0.60–1.00)</td>
</tr>
<tr>
<td><strong>Group B (n = 50)</strong></td>
<td>1.46±0.22 (1.05–1.74)</td>
<td>0.53±0.15 (0.30–1.00)</td>
</tr>
<tr>
<td><strong>Group C (n = 49)</strong></td>
<td>1.41±0.31 (0.94–2.56)</td>
<td>0.62±0.18 (0.30–1.00)</td>
</tr>
</tbody>
</table>

### 48–72 h Mean ± SD (range)

<table>
<thead>
<tr>
<th>Time</th>
<th>Cystatin C (mg/l)</th>
<th>Creatinine (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.49±1.40 (0.91–2.56)</td>
<td>0.57±0.19 (0.60–1.00)</td>
</tr>
<tr>
<td><strong>Group A (n = 10)</strong></td>
<td>1.44±0.44 (0.20–2.71)</td>
<td>0.62±0.18 (0.30–1.00)</td>
</tr>
<tr>
<td><strong>Group B (n = 50)</strong></td>
<td>1.38±1.48 (0.20–2.71)</td>
<td>0.62±0.18 (0.30–1.00)</td>
</tr>
<tr>
<td><strong>Group C (n = 49)</strong></td>
<td>1.50±1.51 (0.76–2.81)</td>
<td>0.68±0.19 (0.30–2.10)</td>
</tr>
</tbody>
</table>

### 7 days Mean ± SD (range)

<table>
<thead>
<tr>
<th>Time</th>
<th>Cystatin C (mg/l)</th>
<th>Creatinine (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.47±1.47 (0.91–2.56)</td>
<td>0.57±0.19 (0.30–1.00)</td>
</tr>
<tr>
<td><strong>Group A (n = 10)</strong></td>
<td>1.54±1.54 (0.91–2.56)</td>
<td>0.64±0.17 (0.30–1.20)</td>
</tr>
<tr>
<td><strong>Group B (n = 50)</strong></td>
<td>1.38±1.38 (0.20–2.71)</td>
<td>0.64±0.17 (0.30–1.20)</td>
</tr>
<tr>
<td><strong>Group C (n = 49)</strong></td>
<td>1.47±1.47 (0.91–2.56)</td>
<td>0.62±0.14 (0.40–1.20)</td>
</tr>
</tbody>
</table>

### 95% CI

- **Cystatin C**: 1.49–1.60
- **Creatinine**: 0.58–1.00

**Fig. 1** – (a) Cys values arranged by GA group ($p = 0.07$) and postnatal days of life ($p < 0.05$). (b) Creatinine values arranged by GA group ($p = 0.11$) and postnatal days of life ($p < 0.05$).

Serum cystatin C concentrations are lower at 7 days ($p < 0.05$). However, there were no differences among GA groups ($p = 0.11$).

**Fig. 2a** shows how serum Cys C continues to decrease at 48–72 h depending on the presence or absence of respiratory pathology, and how it increases at 7 days ($p < 0.05$). Besides, it shows that the concentration of Cys C is lower in the group with respiratory pathology ($p < 0.05$). **Fig. 2b** shows that serum creatinine decreases over time depending on the presence or absence of respiratory pathology, with lower values at 7 days ($p < 0.05$). However, there are no differences in the values of creatinine among groups based on the presence or absence of respiratory pathology ($p = 0.73$).

**Fig. 2c** shows that serum Cys C continues to decrease at 48–72 h of life and that it increases at 7 days in both weight groups ($p < 0.05$). There are no differences among groups >1500 g compared to ≤1500 g ($p = 0.08$). **Fig. 2d** shows that serum creatinine continues to decrease over time in both weight groups ($p < 0.05$), and creatinine values are higher in the group ≤1500 g ($p < 0.05$).

**Fig. 2e** shows the values of Cys C in relation to blood pressure figures. Serum Cys C values are slightly lower in patients with low blood pressure ($p < 0.05$) during the first 48 h.
compared to patients with normal blood pressure. These values subsequently increase in both groups. Creatinine values increase at 48 h in patients with low blood pressure, and these values decrease at the first week of life \( (p < 0.05) \). These figures are more stable in patients with normal blood pressure \( (p = 0.59) \) (Fig. 2f).

The decrease in serum Cys C at 48–72 h of life and the increase at 7 days also depend on whether nephrotoxic drugs are administered or not, there being no differences in the values of Cys C among groups \( (p = 0.61) \). In relation to creatinine, it continues to decrease over time \( (p < 0.05) \), without there being differences between both groups \( (p = 0.94) \) (Table 2).

No group statistical analysis was not conducted to determine the presence or absence of the following pathologies: hypoxic ischaemic encephalopathy \( (n = 3) \), acute renal failure (ARF) \( (n = 3) \), intraventricular haemorrhage \( (n = 3) \), necrotising
Table 2 – Plasma cystatin C and creatinine values in our population arranged by weight, respiratory pathology, low blood pressure cases and administration of nephrotoxic drugs.

| Time | Weight | Respiratory P. | Nephrotoxic drugs | Blood pressure | Normal blood pressure | Low blood pressure | Blood pressure | Nephrotoxic drugs | Blood pressure | Nephrotoxic drugs |
|------|--------|---------------|-------------------|-----------------|---------------------|-------------------|----------------|-------------------|----------------|----------------|----------------|
| Birth | ≤1500 g | Yes | No | Yes | No | Yes | No | Yes | No | Yes | No |
|      | n=35 | n=12 | n=61 | n=96 | n=12 | n=34 | n=34 | n=34 | n=34 | n=34 | n=34 |
| 48–72 h | Creatinine (mg/dl) Birth Mean ± SD 0.61 ± 0.18 0.63 ± 0.18 0.65 ± 0.16 0.65 ± 0.17 0.60 ± 0.18 0.65 ± 0.17 0.65 ± 0.16 0.65 ± 0.17 0.60 ± 0.18 0.65 ± 0.17 |
| 7 days | Cystatin C (mg/l) Birth Mean ± SD 1.46 ± 0.28 1.41 ± 0.37 1.50 ± 0.29 1.43 ± 0.27 1.48 ± 0.34 1.43 ± 0.27 1.48 ± 0.34 1.43 ± 0.27 1.48 ± 0.34 1.43 ± 0.27 |

Discussion

It is widely known that serum Cys C values are very high in neonates and that these are exclusively related to NB, since serum Cys C does not cross the placental barrier as creatinine, which comes from both mothers and NB.

Upon determination of GFR, Cys C provides better sensitivity (97%) and specificity (96%) than creatinine (83% and 87%, respectively), which is supported by recent studies where Cys C reference figures are determined in both preterm NB and term NB from different populations. The objective of this study was not to estimate GFR in our population, but rather serum concentration of Cys C in preterm newborns; thus, the GFR rate is not mentioned in the study. However, it is worth mentioning the reference method used for the measurement of GFR in these patients. There are different estimation equations based on creatinine, Cys C and both endogenous markers. These equations arise from a newborn population with specific characteristics (in relation to age and chronic renal disease stage) and from certain procedures for the measurement and calibration of several endogenous markers, which differ from equations used for the adult population. The most commonly used estimation equation based on creatinine is the Schwartz equation. It was originally published in 1976. It is called the original Schwartz equation. In 2009, the original equation was updated and it is currently known as the updated or modified Schwartz equation. The creatinine measurement procedure in the lab in this equation is an enzymatic procedure with IDMS traceability (standardised creatinine). It is the first paediatric estimation equation with standardised creatinine based on the size of the patient, 2009 updated Schwartz equation (ml/min/1.73 m² \[ \frac{K \times \text{size}}{\text{pCr}} \]), where K is 0.413, with size in cm and plasma creatinine (pCr) in mg/dl.

Equations based on serum Cys C are simpler than equations based on creatinine, since the former do not include anthropometric data: Filler, 2003 (ml/min/1.73 m² \[ 91.62 \times \text{CysC} \cdot 1.123 \times \text{cystatin C: mg/l} \]).

The results of our study show that Cys C values progressively decrease at 48–72h of life, this decrease being statistically significant over time \( p<0.05 \), followed by an increase at 7 days. This decrease over time occurs in the 3 GA groups, and, though the values of Cys C are higher as the GA of preterm newborns increases, there are no statistically significant differences among GA groups. However, Je-Hyan found that serum levels of Cys C tend to decrease as GA increases, as from the third postpartum day. However, we have found certain concordance with studies that show lower values of serum Cys C in bigger preterm newborns, which means that serum Cys C might be affected by renal immaturity that favours excessive urinary loss in more immature newborns, compared to term newborns. Interestingly, according to some authors, newborns with low birth weight have higher serum levels of Cys C during childhood. The specific meaning of high serum levels of Cys C during the first days of life is
In the values of Cys C as a predictor of independent risk for ARF.

There are several limitations in this study. Firstly, it is difficult to determine which subjects are “healthy”, although we tried to classify patients according to the clinical severity of their pathologies associated with prematurity. Subjects may have different degrees of disease severity, may be subject to different mechanic breathing modalities and may have a certain degree of subclinical renal failure, which is inherent to their immaturity. Secondly, we could not standardise levels in the first group, because this is for a smaller and statistically less valuable group.

Conclusions

Serum Cys C decreases at 48–72 h of life, this decrease being statistically significant over time (p<0.05), followed by an increase at 7 days. This decrease over time occurs in the 3GA groups, and, though the values of Cys C are higher as the GA of preterm newborns increases, there are no statistically significant differences in Cys C values among GA groups. These changes are probably due to the maturation of renal function.

Statistically significant differences in serum creatinine values are evidenced when groups are divided according to weight ≤1500 g or >1500 g (p<0.05), and creatinine levels are higher in patients with lower weight, although this difference is not present with serum Cys C. Thus, Cys C is a better GFR marker in very preterm newborns, since its values are less variable for the determination of the renal function in neonates and it is independent of body mass.

Lower Cys C values are found in cases of haemodynamic instability as well as in preterm newborns with associated respiratory pathology. However, more studies are needed for the identification of the exact measures of Cys C, creatinine and GFR in preterm newborns, taking into account that slight increases in serum creatinine probably indicate renal lesion, especially in this vulnerable population. Renal damage during the neonatal period and its subsequent follow-up should be assessed.

Conflicts of interest

The authors have no conflicts of interest to declare.

References


